Epistatic interactions of major effect drought QTLs with genetic background loci determine grain yield of rice under drought stress

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Epistatic interactions of QTLs with the genetic background and QTL-QTL interaction plays an important role in the phenotypic performance of introgression lines developed through genomic-assisted breeding (GAB). In this context, NIL pairs developed with various drought QTL (qDTY) combinations in the genetic background of IR64, TDK1-Sub1 and Savitri backgrounds were utilized to study the interactions. Multi-season phenotyping of NIL pairs harboring similar qDTY combinations provided contrasting performance for grain yield under drought (RS) (classified as high and low yielding NILs) but nearly similar performance under non-stress (NS) conditions. Genome wide genotyping data revealed a total of 16, 5 and 6 digenic interactions were detected under RS conditions in low yielding NILs of IR64, TDK1-Sub1 and Savitri respectively while no significant interaction was found in high yielding NILs under RS and NS conditions in any of the genetic backgrounds used in this study. It is evident from this study that existence of epistatic interactions between QTLs with genetic background, QTL-QTL interaction and interactions among background markers loci itself on different chromosomes influences the expression of a complex trait such as grain yield under drought. The generated information will be useful in all the GAB program of across the crops for precise breeding.

Drought is one of the most severe climatic threat to rainfed rice production affecting on more than 23 million hectares of South and Southeast Asia1. A further increase in drought intensity has anticipated due to ongoing global climatic changes. Development of climate-adapted crop varieties is an urgent need to counter the effect of more than 50% yield losses due to abiotic stresses2-4. Widely grown high yielding rice varieties are highly sensitive to drought. The recent advances in molecular marker and genomics technologies have played a major role in selection of drought-tolerant traits. Rice drought breeding programme at IRRI has successfully identified and validated some of the major and consistent effect grain yield QTLs under reproductive stage drought (qDTYs) such as qDTY1.1,2 qDTY2.1,6 qDTY2.2,7 qDTY3.1,6 qDTY3.2,4 qDTY6.1,8 and qDTY12.1,9 using molecular markers. A considerable effort has been made to improve drought susceptible mega varieties of rice through marker-assisted introgression of qDTYs10-13. Assembling multiple alleles/QTLs/genes in a genetic background using marker-assisted introgression may be an appropriate approach to achieve the expected phenotypic variance in enhancing rice grain yield under drought stress12,14, submergence tolerance15, salt tolerance16, cold tolerance17, resistance to blast18-20, bacterial blight21-23, brown planthopper24,25. The introgression of major effect QTLs/genes and its combinations do not always result in desirable improvement for the targeted trait(s). In some cases, the effect of identified genetic loci in a QTL/gene pyramiding program is not sufficient to fully explain the desired phenotypic variation due to various interactions occurring for a complex quantitative trait. The possible interactions between

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the genetic make-up, epistasis, pleiotropy and linkage among/between the introgressed loci, with the genetic background and environment could have negative/positive effects on the expression of the traits that should never be underestimated. QTL pyramiding approach has enormous potential to improve our understanding of interactions among QTLs and also help to develop new strategies to increase the efficiency of marker-assisted selection program. In this context, development of NILs using marker assisted selection (MAS) is a way forward to verify the effects of one or few major effect QTLs as well as increase the recovery of recurrent parent genome at a faster rate. Rice NILs for complex traits such as drought, blast and brown planthopper were developed and evaluated in different genetic backgrounds of popular rice varieties.

NILs harboring QTLs for abiotic/biotic stresses in the elite backgrounds with superior performances are useful material for immediate release to farmer’s field. NILs are also useful source to study the stable epistatic interactions between the introgressed donor segment and the rest of the recurrent parent genome which gives an important clue about differences in performance of the same set of NILs under variable conditions. The expressions of QTLs for complex traits are strongly affected by the genetic backgrounds. The estimation of genetic backgrounds or recurrent parent genome (RPG) recovery with different types of markers and number of markers also affect the estimation of RPG recovery as a good number of markers with good coverage estimates better information. Better estimation of RPG recovery at early generation of selection can be useful to eliminate linkage drag which can cause unexpected traits in the MAS breeding products because of neighboring genes of the target QTLs. Also, the favorable allele at one locus in a particular background may become unfavorable in another genetic background due to the epistatic interaction. Role of epistasis phenomena in plant breeding has been discussed for a long time, but the extent of the expression level of quantitative traits is due to this complex genetic phenomena are poorly understood. Epistatic or non-allelic interaction is a genetic factor and may change the magnitude and direction of phenotypic expression of QTLs by suppressing or enhancing the loci effect associated with a trait to the other loci of genetic background. The epistatic effect is a deviation from the sum of independent effects of QTLs therefore biased estimation of QTL effects is possible in genetic models assuming no interaction in QTL mapping studies. The epistatic interaction of QTLs and background or modifying loci has been recently reported for quantitative traits in various crops.

Genomic interactions play an important role in deciding the performance of MAS lines for grain yield under drought and various significant QTL x QTL interactions in pyramided lines have been reported for drought-related traits. Most of these studies highlighted capture of such interactions for increased performance of QTLs pyramided lines under drought. But none of these studies reported reasons for a lower yield of NILs with same combinations of pyramided QTLs in the same or different recipient backgrounds. Keeping in this view, the present study is an effort (i) to evaluate the performance of drought NILs of IR64, TDK1-Sub1 and Savitri backgrounds under reproductive stage moderate and severe drought stress and non-stress conditions (ii) to identify promising drought tolerant NILs with higher recovery of recipient genome (iii) to study epistatic interaction of major effect grain yield drought QTLs and with background marker loci.

Results
Phenotypic performance of drought NILs possessing qDTY under RS and NS conditions.

Significant phenotypic differences under RS and NS were observed among the parents and NILs in IR64, TDK1-Sub1 and Savitri backgrounds. The results of ANOVA, means, LSD, heritability of IR64 NILs with a qDTY and qDTY combinations under RS (SS and MS) and NS are summarized in Table 1. Among IR64 NILs, the average yield reduction was found 87% under SS and 74% under MS as compared to NS conditions. Trial mean of NILs in IR64 background for GY was 690 kg ha\(^{-1}\) under SS in 2015 DS while it was 1358 kg ha\(^{-1}\) under MS in 2017 DS. Broad-sense heritability (H) of GY under RS ranged from 0.78 to 0.89 in the years of 2015 and 2017 DS. The NIL pairs harboring same qDTY or qDTY combinations with on par background recovery, provided significantly different grain yield (GY) under similar level of drought stress were classified into high and low yielding NILs (Table 1). For instance, nine IR64 NILs possessing qDTY12, QTL yielded 25 to 3375 kg ha\(^{-1}\) under SS situation, whereas no such difference was observed under NS situation (4490 to 6470 kg ha\(^{-1}\)). Another four IR64 NILs possessing qDTY12 yielded 480 kg ha\(^{-1}\) to 2495 kg ha\(^{-1}\) under MS. In NS conditions, trial mean for GY was 5360 kg ha\(^{-1}\) and ranged from 4375 kg ha\(^{-1}\) to 6958 kg ha\(^{-1}\). Nine NILs with same combination of qDTY12 yielded a large variation for grain yield under drought and ranged from 89 to 2141 kg ha\(^{-1}\). The huge variation in grain yield under drought even having same QTL combinations was due to some hidden interactions and analysis of such interaction were explored and discussed in this study. Similarly three NILs with qDTY12 yielded 572 to 1455 kg ha\(^{-1}\); four NILs with qDTY12 yielded 369 to 936 kg ha\(^{-1}\), two NILs with qDTY12 yielded 289 to 611 kg ha\(^{-1}\) and two NILs with qDTY12 yielded 414 to 1031 kg ha\(^{-1}\). The mean plant height (PH) of IR64 NILs under RS was higher than the recurrent background IR64 while PH of NILs under NS was very similar to that of IR64 (Table 1). Days to flowering (DTF) were affected significantly with severity of stress and flowering delay of 10 days under SS and 7 days under MS was observed.

Mean performances of measured traits of TDK1-Sub1 NILs, heritability and LSD is presented in Table 2. Grain yield under severe stress (GYSS) ranged from 0 kg ha\(^{-1}\) to 479 kg ha\(^{-1}\) with trial mean of 327 kg ha\(^{-1}\). The recurrent parent TDK1-Sub1 yielded 14.4 kg ha\(^{-1}\) while the donor parent IR55419-04 performed well with GY of 881 kg ha\(^{-1}\) under SS situation. GY ranged from 0 kg ha\(^{-1}\) to 479 kg ha\(^{-1}\) with range of 300 kg ha\(^{-1}\) to 1555 kg ha\(^{-1}\). The average yield reduction was more than 90% under SS while 84% less grain yield was recorded under MS among TDK1-Sub1 NILs. Heritability for GY was recorded 0.89 to 0.91 in SS and 0.58 in NS. The pyramided three NILs with qDTY1, qDTY2, combination yielded 0 to 479 kg ha\(^{-1}\) under SS situation. However, the performance of both lines was 300 and 1555 kg ha\(^{-1}\) respectively in MS. Similarly, NIL pair consisting of qDTY1, qDTY2, qDTY1, qDTY2, combination yielded 86 kg ha\(^{-1}\) and 479 kg ha\(^{-1}\) under SS and 1085 kg ha\(^{-1}\) to 1555 kg ha\(^{-1}\) under MS. A reduction in PH was observed under SS and MS conditions compared to NS condition.
The mean PH under SS, MS and NS were 64 cm, 69 cm and 92 cm, respectively. Significant differences in DTF with trial mean of 94 days and flowering differences of 7 days between entries was observed under RS. Earliest line flowered in 80 days while most of the lines were of late maturing type and it flowered in 96 days. As observed in the case of IR64 NILs, TDK1-Sub1 NILs was also produced nearly similar grain yield under NS situation.

The same traits were also observed in the Savitri NILs (Table 2). A reduction in mean GY under SS and MS compared with mean GY under NS was 90% and 61%, respectively. The recurrent parent Savitri yielded 48 kg ha$^{-1}$ while the donor parent IR77298-5-6-18 yielded 236 kg ha$^{-1}$ under SS. NIL pair carrying single $qDTY_{12}$ yielded 134 to 517 kg ha$^{-1}$ under SS, whereas four NILs carrying two $qDTY$ ($qDTY_{12} + qDTY_{12}$) yielded 18 to 627 kg ha$^{-1}$. The significant deviation observed in GY suggesting some negative interaction between background markers loci.

Table 1. Mean performance for grain yield (GY), days to flowering (DTF) and plant height (PH) of IR64 NILs under severe stress (SS), moderate stress (MS) and non-stress (NS) conditions. Note: LSD$_{0.05}$: least significant difference at the 5% confidence level; H: Broad sense heritability.
**Mean** of the Savitri NILs was significantly reduced under drought stress conditions ranging from 61 to 85 cm in SS, 58 to 75 cm in MS and from 78 to 102 cm in NS conditions. **PH** was reduced by 17 to 20 cm in drought stress. The mean days to flowering (**DTF**) in Savitri NILs ranged from 72 to 81 days in SS, 76 to 82 days in MS and from 74 to 81 days in NS conditions. **DTF** was severely affected under drought for recipient parent Savitri varied from 103 days in NS while 113 and 96 days was recorded in SS and MS conditions respectively. The heritability (**H**) was high for **DTF** (0.84,0.85,0.92) and medium to high for **PH** (0.72,0.80,0.88) and **GY** (0.74,0.89,0.58) under SS, MS and NS conditions for Savitri NILs (Table 2).

### Promising drought NIL lines with the maximum genetic background coverage of recipient parents.

Recent recurrent genome recovery (**RPG**) were estimated using 111, 107 and 89 polymorphic SSR markers among NILs carrying qDTY5 in IR64, TDK-Sub1 and Savitri backgrounds, respectively. The **RPG** recovery among the NILs of multiple genetic backgrounds ranged from 85 to 93% in the case of IR64 NILs, 86 to 90% in the case of TDK-Sub1 NILs and 80 to 89% in the case of Savitri NILs. Most of the IR64 NIL lines showed more than 90% **RPG** recovery followed by TDK-Sub1 and least recovered lines belong to Savitri background. Graphical representation of the selected pyramided NILs achieving higher genome recovery of respective backgrounds is depicted in Supplementary Fig. S1 (IR64 NILs with qDTY12, qDTY2), Supplementary Fig. S2 (TDK1-Sub1 NILs with qDTY12, qDTY3) and Supplementary Fig. S3 (Savitri NILs with qDTY1, qDTY2). The selected promising NILs capturing higher recurrent genome background and yielded well under SS, MS and NS conditions is presented in Table 3. These superior NILs with excellent drought tolerance could be used a drought tolerant line/variety to be released after multi-location evaluation in national and provincial coordinated trials in target countries.

### QTL–QTL and QTL-background interaction in drought NILs.

A total of 16 digenic interactions between marker loci were detected in low yielding NILs of IR64 background under SS and MS situation. Out of 16 digenic interactions, we identified 6 interactions between drought grain yield QTLs (qDTY2, qDTY3, and qDTY12) and background markers loci while 9 interactions were found among background markers loci of different chromosome. A significant negative QTL-QTL interaction was also found between introgressed qDTY1 (RM543-RM212) located on chromosome 1 with qDTY3 (RM263-RM573) identified on chromosome 2. In context of QTL and background marker loci interactions, qDTY2, (RM154-RM236) interacted negatively with two genetic loci (RM7-RM32) on chromosome 3 and chromosome 4 (RM142-RM119), while qDTY3 (RM263-RM573) was interacted negatively with genetic loci (RM108-RM215) on chromosome 9 and chromosome 10 (RM271-RM269) under SS situation (Table 4, Fig. 1a). Similarly, qDTY12 interacted negatively with...
Table 3. Selected promising drought NILs identified with maximum recurrent parent genome recovery. Note: RPG: Recurrent parent genome recovery.

| Background | Promising NILs | Parentage | Foreground | RPG recovery % | Severe stress | Moderate stress | Non-stress | GV (kg ha⁻¹) |
|------------|---------------|-----------|------------|----------------|--------------|----------------|-----------|-------------|
| IR64       | IR 102784-2-42-16-1-1-2 | IR 99621-181/IR 99620-158//IRRI 149 | qDTY_{12}, qDTY_{12} | 93 | 1426 | 1860 | 5430 |
| IR64       | IR 102793:1-11-229-3-1-1 | IR 99622-312/IR 99620-158//IRRI 149 | qDTY_{12}, qDTY_{12} | 92 | 656 | 1900 | 5618 |
| IR64       | IR 102793:1-11-192-1-1-4 | IR 99621-14/IR 99619-361//IRRI 149 | qDTY_{12}, qDTY_{12} | 91 | 1386 | 1110 | 4748 |
| IR64       | IR 102784-62-66-1-1-2 | IR 99621-181/IR 99620-158//IRRI 149 | qDTY_{12}, qDTY_{12} | 91 | 611 | 845 | 5255 |
| IR64       | IR 102796-14-140-1-1-1 | IR 99622-306/IR 99620-228//IRRI 149 | qDTY_{12}, qDTY_{12} | 90 | 788 | 2740 | 4928 |
| IR64       | IR 102796-14-124-1-1-3 | IR 99626-306/IR 99620-228//IRRI 149 | qDTY_{12}, qDTY_{12} | 90 | 996 | 2235 | 5220 |
| IR64       | IR 102782-70-112-3-1-1 | IR 99621-14/IR 99619-361//IRRI 149 | qDTY_{12} | 90 | 3375 | 2210 | 5143 |
| IR64       | IR 102793:1-11-64-1-1-2 | IR 99621-181/IR 99619-361//IRRI 149 | qDTY_{12}, qDTY_{12} | 89 | 2002 | 2750 | 6145 |
| IR64       | IR 102793:1-11-66-3-1-1 | IR 99622-312/IR 99620-158//IRRI 149 | qDTY_{12}, qDTY_{12} | 89 | 2141 | 1455 | 4375 |
| TDK1-Sub1  | IR 102777:5-83-1-2-7 | IR 90266-B-542-1/IR07289 | qDTY_{12}, qDTY_{12} | 90 | 316 | 465 | 5590 |
| TDK1-Sub1  | IR 102777-6-86-2-2-11 | IR 90266-B-542-1/IR07289 | qDTY_{12}, qDTY_{12} | 89 | 479 | 1555 | 4966 |
| Savitri    | IR 106523-21-28-1-2-B | IR 102773-1061/IR 90252-B-548-2-B/IR 90250-B-475-1-B | qDTY_{12} | 89 | 517 | 2565 | 7098 |

Discussion

Epistasis is a genetic phenomenon of interaction which may enhance or reduce the expression (depending on degree and direction) of interacting loci underlying QTLs associated with the complex trait. QTL mapping and interaction studies in recent years in different crops suggested the possible presence of epistasis such as (i) interactions between introgressed QTLs (2) interactions between introgressed QTLs and 'background' (modifying) loci and (3) interactions between complementary background loci affecting overall expressions of trait. To further understand this interaction (QTL-QTL and QTL to background), we have selected range of rice drought NILs developed in different genetic backgrounds.

Several large and consistent effect QTLs for grain yield under drought (qDTY_{12}, qDTY_{12}, qDTY_{12}, qDTY_{12}) were identified and validated in various rice genetic backgrounds using the QTL associated markers. Exploiting the available molecular markers for introgression of consistent drought grain yield QTLs (qDTYs)
Table 4. Epistatic interaction of major effect loci of grain yield under drought with background loci of IR64, TDK1-Sub1 and Savitri NILs. Note: GYMS: Grain yield under moderate stress; GYSS: Grain yield under severe stress; *Chromosome ID at the first scanning position; †Scanning position in cm of the first flanking marker pair; ‡Chromosome ID at the second scanning position; ‡‡Scanning position in cm of the second flanking marker pair; *LOD score caused by epistatic effects; PVE(%): Phenotypic variation explained by epistatic effects; ††LOD score caused by additive effects; **PVE(%): Phenotypic variation by the additive effects; †‡Estimated additive effect of position 1; †††Estimated additive effect of position 2; †‡‡Additive by additive epistatic effect at the two scanning positions.

| NILs Background | Trait Name | Chromosome | Position 1 | Chromosome | Position 2 | LOD ep1 (%) | PVE_epi (%) | LOD add2 (%) | PVE_add (%) | Add1 (%) | Add2 (%) | Add1 by Add2 (%) |
|----------------|------------|------------|------------|------------|------------|-------------|-------------|-------------|-------------|-----------|-----------|-----------------|
| IR64           | GYSS 3     | 146        | 2          | 4.21       | 11.78      | 2.88        | 4.01        | 124.65      | 458.93      | -298.92   |          |                  |
| IR64           | GYSS 1     | 141        | 9          | 86.7       | RM108-MR215| 3.33        | 4.73        | 2.50        | 2.60        | 603.78    | -297.59   |                  |
| IR64           | GYSS 3     | 144        | 11         | 8          | RM286-MR332| 3.60        | 4.66        | 2.68        | 2.47        | -363.62   | -797.04   |                  |
| IR64           | GYSS 2     | 124.7      | 3          | 86.7       | RM108-MR215| 3.82        | 4.43        | 2.79        | 2.77        | -396.48   | -930.95   |                  |
| IR64           | GYSS 4     | 115        | 9          | 86.7       | RM108-MR215| 3.75        | 4.63        | 2.99        | 2.60        | 552.35    | -997.21   |                  |
| IR64           | GYSS 2     | 9.8        | 3          | 65         | RM7-RM232  | 3.26        | 3.92        | 2.94        | 2.65        | -426.60   | -966.28   |                  |
| IR64           | GYSS 2     | 9.8        | 4          | 75         | RM142-RM119| 3.29        | 4.66        | 2.68        | 2.47        | -363.62   | -797.04   |                  |
| IR64           | GYSS 2     | 84.8       | 10         | 21.3       | RM222-RM331| 3.67        | 3.54        | 2.84        | 2.58        | 434.01    | -942.80   |                  |
| IR64           | GYSS 2     | 84.8       | 9          | 86.7       | RM108-MR215| 3.75        | 4.63        | 2.99        | 2.60        | 552.35    | -997.21   |                  |
| IR64           | GYSS 2     | 124.7      | 10         | 66.3       | RM271-RM269| 3.52        | 4.62        | 2.74        | 2.58        | -408.88   | -999.2   |                  |
| IR64           | GYSS 3     | 65         | 10         | 86.7       | RM108-MR215| 3.81        | 4.63        | 2.94        | 2.59        | 554.08    | -999.2   |                  |
| IR64           | GYSS 3     | 65         | 10         | 36.3       | RM311-RM271| 3.52        | 4.43        | 2.76        | 2.58        | 504.44    | -991.78   |                  |
| IR64           | GYSS 4     | 75         | 11         | 86.7       | RM108-MR215| 3.60        | 4.63        | 2.92        | 2.59        | 553.86    | -992.10   |                  |
| IR64           | GYSS 8     | 90         | 19         | 94.7       | RM108-MR215| 3.27        | 4.64        | 2.92        | 2.58        | 558.65    | -982.32   |                  |
| IR64           | GYSS 10    | 16.3       | 11         | 71.3       | RM269-RM228| 3.39        | 3.47        | 2.68        | 2.50        | -501.44   | 328.77    |                  |
| TDK1-Sub1      | GYSS 2     | 91.9       | 6          | 10         | RM108-MR215| 3.06        | 14.09       | 13.30       | 4.09        | -73.18    | 72.46     | -72.46         |
| TDK1-Sub1      | GYSS 3     | 150        | 6          | 10         | RM108-MR215| 3.08        | 12.40       | 11.16       | 5.80        | -22.97    | 50.06     | -26.05         |
| TDK1-Sub1      | GYSS 5     | 36.7       | 6          | 10         | RM108-MR215| 3.60        | 17.90       | 13.89       | 3.93        | -122.05   | 113.11    | -113.11        |
| TDK1-Sub1      | GYSS 6     | 10         | 11         | 100        | RM275-RM380| 2.80        | 14.88       | 10.87       | 4.98        | 45.55     | -42.51    | -44.08         |
| Savitri        | GYMS 4     | 8.5        | 6          | 65         | RM136-RM275| 3.43        | 10.56       | 2.41        | 2.60        | -211.62   | -382.62   |                  |
| Savitri        | GTMS 10    | 15         | 6          | 85         | RM136-RM275| 3.43        | 12.76       | 2.58        | 4.22        | -211.62   | -382.62   |                  |
| Savitri        | GYSS 2     | 49.8       | 3          | 182        | RM55-RM570 | 4.31        | 10.78       | 2.80        | 5.01        | 10.66     | -58.66    | -238.33        |
| Savitri        | GYSS 3     | 67         | 3          | 187        | RM55-RM570 | 4.10        | 12.76       | 2.61        | 7.94        | -58.66    | 10.66     | 238.33         |
| Savitri        | GYSS 3     | 167        | 7          | 73.2       | RM10-MR47  | 6.33        | 14.31       | 2.44        | 2.12        | -49.38    | 144.08    | -94.60         |
| Savitri        | GYSS 3     | 182        | 11         | 0          | RM48-RM332 | 4.21        | 11.78       | 2.88        | 4.01        | -58.66    | 10.66     | -238.33        |
moderate to high heritability for grain yield under drought stress for QTL identification. In present study, 600 genome-wide SSR markers were used for estimating the similarity between backgrounds and their derived NILs. Promising NILs have captured 89–93% recovery of respective backgrounds (Table 3) while 44 NILs used in this study possessed 85–93% genome recovery. Phenotypic selection combined with marker-assisted background selection approach after qDTY introgression was followed, that could be one of the reasons not to achieve the expected recovery in some of the NILs. Similar concern was discussed earlier by Dixit et al. in his study of marker-assisted breeding to combine submergence and drought tolerance in rice. However, in future drought NILs advancement, donor segment can be further minimized by using the recombinant selection markers and generation of whole genome profiling data through SNP-chips of respective DTY QTLs. A similar approach was also explained by Jena et al. in context to rice NILs developed for brown planthopper (BPH) resistance having low recovery (82–89.5%) of recurrent parent genome IR24 even after three back crosses.

Role of digenic interactions between DTY QTLs affecting the performance of one QTL over the other drought QTL has been well documented in various QTL mapping and marker-assisted drought breeding programme. Positive interaction of qDTY2.2 with qDTY3.2 and qDTY5.2 have been reported by Dixit et al. which increases grain yield under drought significantly while Shamsudin et al. identified positive digenic interactions of qDTY2.2 and qDTY3.1, with qDTY3.2 enhancing overall expression of drought-related traits. Interaction of qDTY3.1 and qDTY2.2 reduces the flowering duration and subsequently increases GY under stress. A positive epistatic interaction of qDTY4.1 and qDTY5.2 loci with qDTY7.1 have been reported in the pyramided line of Samba Mahsuri enhancing the grain yield under drought. Apart from these known qDTY’s QTL x QTL interactions, there are many unknown genomic interactions exist with background loci called as background noise which should be identified for effective deployment of drought QTLs in MAS breeding program.

Interaction between the target QTL/gene with the genetic background has been discussed earlier for various complex traits in rice. However, there have been limited attempts to identify particular background markers loci interacting with introgressed QTLs and influencing the overall expression of target trait. There is an urgent need to identify such epistatic gene interactions which complicate the genotype-phenotype relationship of complex traits such as drought. In present study, we have detected negative interactions between drought QTLs

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Cyclic illustrations of epistatic QTLs of IR64 NILs for grain yield under various levels of drought stress. The dotted lines indicate marker pairs interacting significantly on same or different chromosomes with their corresponding LOD value due to epistatic effect. (a) Grain yield under severe stress (GYSS): chromosome 1 (qDTY1.2 RM543-RM212) showed epistatic negative interaction with qDTY2.3 (RM263-RM573) on chromosome 2 while qDTY2.2 (RM154-RM236) and qDTY2.3 (RM263-RM573) showed epistatic interaction with background markers loci on chromosomes 3, 4, 9 and 10 (b) Grain yield under moderate stress (GYMS): chromosome 3 (qDTY3.2 RM232-RM22, 10 cM) showed epistatic interaction with background loci of chromosomes 4 and 11 (c,d) GYMS and GYSS of high yielding NILs: no significant interactions was found (e) Grain yield under non-stress (GYNS): no interaction.
and \( q_{DTY3.2} \) with IR64 background markers loci located on chromosomes (2, 3, 4, 10 and 11), among background markers loci located on different chromosomes and between drought grain yield QTL \( q_{DTY2.3} \) with \( q_{DTY2.3} \), which might be one of the reason of poor performance of low yielding NILs carrying DTY QTLs (Table 1). The influence of epistatic drought QTL interactions explained PVE% ranging from 3.47% to 25.46% which could be very crucial in determining the grain yield under drought. Five digenic interactions were found in low yielding NILs of TDK1-Sub1 background, where \( q_{DTY6.1} \) (RM587-RM510) had shown negative interaction with background marker located on five different chromosomes. Savitri background NILs had shown 6 epistatic interactions between background markers loci in low yielding lines. Many of such epistatic interactions are background specific. However, in our study, the high effect of many of the \( q_{DTY} \) QTLs interaction with the genetic background marker loci indicate that these are biological interactions and not just statistical. The QTLs \( q_{DTY2.2} \), \( q_{DTY2.3} \), \( q_{DTY3.2} \) and \( q_{DTY6.1} \) have shown interactions with the background markers on the same chromosome in three other genetic backgrounds (data unpublished). Some earlier studies\(^{32,52}\) highlighted the importance of epistatic QTL effect over the additive QTLs are influencing the expression of introgressed trait in crops like Arabidopsis and rice. Thomson et al.\(^{53}\) reported a loss of Saltol QTL effect in developed NILs and suggested an interaction between Saltol and other background loci. Similarly, Babu et al.\(^{54}\) observed the difference in level of seedling stage salt tolerance among Saltol introgressed advanced lines and assumed the existence of a possible interaction between Saltol QTL and genetic background but these reports did not report on the particular background markers loci interacting negatively with Saltol QTL, resulting in variable expression of Saltol introgressed lines. Revealing such epistatic interactions in present study, between introgressed QTL and background markers loci sheds more light in understanding the differences in phenotypic expression of MAS introgressed/pyramided lines for quantitative traits that can be useful in future MAS programs.

### Conclusions

Performance of NILs carrying same major effect drought grain yield QTLs/QTLs combinations are strongly affected by interactions with the genetic background loci. This is the first report on in-depth analysis of loss of introgressed QTL effects in SS, MS due to negative interactions between drought QTLs and background markers loci, drought QTL-QTL interaction and among interacting background markers loci on various chromosomes in SS, MS in low yielding NILs but not under NS. For both low as well as high yielding NILs, no interaction was detected under NS. Absence of such negative interactions in high yielding NILs under MS, SS indicate future MAS program to carefully select against such interactions to increase grain yield under drought. Promising NILs in background of popular varieties (IR64, TDK1-Sub1 and Savitri) with absence of negative interactions and
performing well in drought stress and non-stress conditions might be a very useful lines for use in future MAS/variety to be released for drought-prone ecosystems.

**Methods**

**Plant materials.** A total of 44 drought NILs derived from three backgrounds (33 NILs of IR64, 5 NILs of TDK1-Sub1 and 6 NILs of Savitri) were used in this study (Supplementary Table S2). We initiated the process of NILs development with a total of 130 lines (70 lines from IR64, 32 lines from TDK1-Sub1 and 28 lines from Savitri) developed through marker-assisted breeding for the three genetic backgrounds for the present study. However, before including all the lines we attempted foreground selection with QTLs linked markers as well as background marker selection of all the lines with 600 genome wide background SSR markers and those lines which follows the minimum criteria of RPG recovery (>80%) were taken for the selection of NILs for further evaluation. As a result, finally we have selected a total of 44 drought NILs derived from three backgrounds (33 NILs of IR64, 5 NILs of TDK1-Sub1 and 6 NILs of Savitri) were used in this study. A total of eight sets of IR64 NILs (20 high yielding and 13 low yielding) possessing different QTL and QTL combinations was developed through crossing of IR64 with different drought tolerant donors contributing different grain yield QTLs. Four different drought tolerant donors were utilized for development of the selected NILs, for instance IR77298-14-1-2-17 (qDTY2, and qDTY4), IR 86918-B-315 (qDTY1, and qDTY1), Vandana (qDTY2, and qDTY3) and Way Rarem (qDTY12) were crossed with IR64 and through marker assisted backcross breeding the above mentioned NILs were developed. The pyramided version of NILs was developed through intercrossing of NILs of different gene combinations. Two sets of TDK1-Sub1 NILs (2 high yielding and 3 low yielding NILs) possessing different QTL combinations were utilized in the present study. The pyramided version of NILs were developed from the cross between IR55419-04/2*TDK1. After confirming the presence of three QTLs (qDTY1, qDTY4, and qDTY6) in pyramided line, it was utilized for backcrossing with submergence tolerant parent TDK1-Sub1 to develop NILs in TDK1-Sub1 background. Similarly, two sets of Savitri NILs (3 high yielding and 3 low yielding NILs) were developed from intercrossing of two BC1 population (IR 77298-5-6-18/2*Savitri IR 74371-46-1-1/2*Savitri) to develop lines carrying qDTY7 and qDTY12. QTLs in Savitri background were all of the NILs used in the present study in all the three background were re-confirmed with earlier reported markers for the presence of QTL and QTL combinations.

![Figure 3. Cyclic illustrations of epistatic QTLs of Savitri NILs for grain yield under various levels of drought stress (a) GYSS: chromosome 2 (RM71-RM290, 50 cM) and chromosome 3 (RM7-RM251 67 cM, RM319-RM55 167 cM and RM55-RM570, 182 cM) have shown epistatic interaction with background loci of chromosomes 3, 7 and 11 (b) GYMS: chromosome 4(RM337-RM335) and chromosome 10(RM244-RM239) was interacted negatively with a common genetic loci (RM136-RM275) on chromosome 6 (c,d) GYMS and GYSS of high yielding NILs: no significant interactions was found (e) GYNS: no interaction.](https://www.nature.com/scientificreports)
Evaluation of NILs under reproductive-stage drought stress and non-stress conditions. To study the grain yield variability of NILs pyramided with same QTL and QTL combinations, a total of 33 NILs derived from IR64, 5 NILs from TDK1-Sub1 and 6 NILs from Savitri backgrounds were evaluated during the dry seasons (DS) of 2015 and 2017 in a transplanted lowland ecosystem under reproductive-stage drought stress (RS) and non-stress (NS) conditions at the Ziegler Experiment Station of IRRI, Los Baños, Laguna, Philippines (14°30’N longitude 121°15’E latitude). Seedlings were raised in a wet-bed nursery for 21 days before transplanting into the main experimental field. The field lay out was generated using the statistical software Plant Breeding Tools (PBTools v1.4) developed at IRRI. The experimental design was alpha lattice in 2 replications in two-row plots of 5 m length with row to row and plant to plant spacing of 20 × 20 cm. Continuous supply of irrigation at 5 cm standing water was provided till the crop maturity in non-stress experiment. Fertilizer nitrogen, phosphorus, and potassium (NPK) were applied at recommended dose of 120:30:30 kg ha⁻¹.

Reproductive stage drought stress experiment was conducted as described previously. In brief for RS experiment normal irrigation was maintained in the field up to 30 days after transplanting, after which the water was drained out to initiate stress and continued till the harvesting. Water table depth was measured using PVC pipe with a plugged hole at the bottom and inserted in soil to 1 m depth and 15 cm above the soil surface in experimental stress field at regular interval. Life-saving irrigation was provided at level of severe stress when all the susceptible checks showed severe leaf rolling with minimum probability to recover upon watering and water table remained below 1 meter. Life-saving irrigation provided through flash flooding was drained out after 24 hrs to initiate further cycle of stress.

The stress imposed was classified from severe to moderate based on the relative yield reduction to non-stress (NS) experiments explained earlier by . Experiments with yield reduction of <70% were characterized as severe stress (SS) and experiments with yield reduction of 31–70% were characterized as moderate stress (MS) as earlier reported by Kumar et al.

Observations recorded. Data on days to 50% flowering (DTF), plant height (PH), and grain yield (GY) were recorded in RS and NS experiments. DTF was observed as the number of days from seeding to the 50% the plants in a plot flower. The PH of three random plants from each NILs was measured before maturity from the soil surface to the tip of the main tiller and then averaged for mean analysis. Grains from individual plot were harvested at physiological maturity, oven-dried to a moisture content of 14% before weighing. The grain yield in grams was further converted in kg ha⁻¹ before entry means analysis in PBTools.

NILs genotyping. The presence of introgressed drought QTLs and QTLs combinations in IR64 (qDTY 1.1, qDTY 1.2, qDTY 1.2, qDTY 1.3, qDTY 1.4, and qDTY 1.12), TDK1-Sub1 (qDTY 1.1, qDTY 1.2, and qDTY 1.12) and Savitri backgrounds (qDTY 1.2 and qDTY 1.12) were confirmed in respective NILs using the earlier reported peak and flanking markers of respective grain yield drought QTLs. Fresh and young leaves from 25 days old transplanted seedlings were collected in plot-wise bulk from each NIL- and their respective parents. Genomic DNA was extracted from collected leaves using a modified CTAB protocol developed by Murray and Thompson, dissolved in 200 μl of TE (Tris-EDTA) buffer and stored at −20°C. PCR amplification with the molecular markers was performed on a thermal cycler (G-Storm G51, UK) at Genotyping service laboratory (GSL), IRRI, Philippines. Each PCR reaction mixture of 15 μl consisted of 10 ng of rice genomic DNA, 1× PCR buffer, 100 μM dNTPs, 100 μM oligonucleotide primers and 1 unit of Taq polymerase. PCR products were resolved casting high-resolution 8% (w/v) polyacrylamide gel electrophoresis (PAGE) (CBS scientific, model MGV-202–33) and resolved in a 1x TBE buffer at 90 volts for 1–2 h, depending on the PCR product sizes. The separated DNA fragments after electrophoresis were stained with SYBER Safe™ and visualized under UV trans-illuminator (AlphaImager™ System).

To study the genomic reconstitution of NILs with the similar QTL or QTLs combinations but showing variation in grain yield under drought, background genotyping was carried out on the selected NILs and parents. A total of 600 SSR markers equally distributed on all 12 chromosomes were selected from the Gramene database (http://www.gramene.org/) for identification of polymorphic markers between parents, which were further utilized for background selection in order to estimate recurrent parent genome (RPG) recovery.

Statistical analysis, recurrent parent genome recovery and epistatic interaction analysis. The phenotypic data collected from drought stress and non-stress experiments were statistically analyzed for the computation of trial means and standard error of difference (SED) using PBTools v1.4. Least significant difference (LSD) ‘at the 5% and 1% significant levels’ were used to compare the means of the test entries and to infer the significant differences of the traits studied between parents and each NILs. Linear mixed model for analysis of variance was calculated with following equation:

\[ Y_{ijk} = \mu + G_i + R_j + BK(R_j) + e_{ijk} \]

where, \( Y_{ijk} \) is measurement recorded in plot, \( \mu \) is overall mean, \( G_i \) is effect of \( i^{th} \) genotype, \( R_j \) is the effect of the \( j^{th} \) replicate, \( BK(R_j) \) is the block effect of \( j^{th} \) replicate and \( e_{ijk} \) is the error. The genotypes were considered as fixed and the replicates and block effects were random for estimating the entry means. The recurrent parent genome (RPG) recovery percentage of drought NILs of various background were estimated with the following formula:

\[ \text{Recurrent parent genome (RPG) recovery %} = \frac{2(B) + (H)}{2N} \times 100 \]
Where, \( B \) = SSR marker loci homozygous for genetic background; \( H \) = no. of marker loci still in heterozygous; \( N \) = total no. of polymorphic SSR markers used for background estimation.

Graphical representation of background genome was constructed using software Graphical Genotype (GGT v 2.0) [1]. In epistasis mapping interactions between markers loci of all the NILs including high and low yielding NILs background wise were analyzed considering all the conditions (SS, MS and NS) in model using QTL IciMapping ver. 4.0.1. In epistasis mapping method, a two-stage stepwise regression strategy was adopted to identify the most significant markers and marker-pair followed by two-dimensional scanning to identify significant digenic epistasis using adjusted phenotypic values based on best fitted multiple regression model [46]. In the first stage, the significant markers and marker pairs explaining epistatic and additive variations were selected in model and then in next stage the stepwise regression with stricter probability levels were applied to the residuals from the first stage to select significant marker pairs and estimate their effects in model [46]. The stricter probability level was applied in the second stage to avoid over-fitting due to the large number of regression variables. Extensive simulations have shown epistatic QTLs and QTL epistatic networks can be identified efficiently by ICIM mapping even in case two interacting QTL have some additive effects [46]. The threshold LOD value was determined by a permutation test involving 1000 runs at a significance level of \( P = 0.01 \) to detect significant digenic interactions between marker loci. In mapping parameters, window size and walk speed used for the genome scan was 10 cM and 1 cM respectively.

Data Availability
The data sets supporting the results of this article are included within the article.

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Author Contributions
S.Y. was involved in genotyping, observations, experimental data analysis and drafting the manuscript; N.S. was involved interpretation of data and writing the manuscript; R.R.M. helped in genotyping; S.D., S.K.S., P.S.P., N.P.M., S.P.D., R.B.Y., V.K.S., P.S., R.K.V. were involved in drafting and correction of the manuscript R.K.V. and V.K.S. was also involved with conceptualization of the result from present study on other crops. A.K. was involved in the design of the experiment and in the critical revision of the manuscript. All authors approved the final version of the manuscript.

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