Marker assisted development and characterization of non-GM herbicide tolerant Near Isogenic Lines of a mega Basmati rice variety, “Pusa Basmati 1121”

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Abstract

Background

Direct-seeded rice (DSR) is a potential technology for sustainable rice farming as it saves water and labor. However, higher incidence of weed under DSR limits productivity. Therefore, there is a need to develop herbicide tolerant (HT) rice varieties.

Results

We used marker assisted backcross breeding (MABB) to transfer a mutant allele of Acetohydroxy acid synthase (AHAS) gene, which confers tolerance to imidazolinone group of herbicides from the donor parent (DP) “Robin” into the genetic background of an elite popular Basmati rice variety Pusa Basmati 1121 (PB 1121). Foreground selection was done using the AHAS gene linked Simple Sequence Repeat (SSR) marker RM6844 and background selection was performed using 112 genome-wide SSR markers polymorphic between PB 1121 and Robin. Phenotypic selection for agronomic, Basmati grain and cooking quality traits in each generation was carried out to improve the recovery of recurrent parent phenome (RPP). Finally, a set of 12 BC$_4$F$_4$ near isogenic lines (NILs), with recurrent parent genome (RPG) recovery ranging from 98.66 to 99.55% were developed and evaluated. PB 1121-HT NILs namely 1979-14-7-33-99-10, 1979-14-7-33-99-15 and 1979-14-7-33-99-66 were found superior to PB 1121 in yield with comparable grain and cooking quality traits and herbicide tolerance similar to Robin.

Conclusion

Overall, the present study reports successful development of non-GM HT NILs in the genetic background of popular Basmati rice variety PB 1121 by introgression of mutated AHAS allele. This is the first report of the development of HT rice in India. Superior NILs are being evaluated in the national Basmati trials, the release of which is likely to provide a viable option for the adoption of DSR technology in Basmati rice cultivation.

Background

In South-East Asian countries, where rice is primarily cultivated by transplanting, DSR holds great promise as it is a potential water and labor saving technology, which offers the possibility of saving huge amount of water and labour needed for puddling and transplanting. However, DSR cultivation suffers from high weed infestation. Weeds pose a serious threat by competing with the crop for water, nutrients and light, thereby hampering productivity (Powles and Yu 2010). In India, upto 30% of the total cost of rice cultivation goes in controlling weeds (Rao et al. 2015). It has been demonstrated that with effective weed management, the yields from DSR is widely comparable to transplanted rice (Pathak et al. 2011). The use of herbicides is the most effective and economical option to control weeds (Anderson et al. 1989).
However, the herbicides that are safe and effective at minimal doses should be used to ensure environmental safety. Imidazolinone group of herbicides (imazapyr, imazapic, imazethapyr, imazamox, imazamethabenz and imazaquin etc.), control weeds by inhibiting the enzyme acetohydroxyacid synthase (AHAS), also called acetolactate synthase (ALS). AHAS is a critical enzyme for the biosynthesis of branched-chain amino acids namely, leucine, isoleucine and valine in plants. These herbicides, also have low mammalian toxicity due to lack of AHAS gene (Tan et al. 2006; Piao et al. 2017). Due to the above benefits, the herbicides of imidazolinone group are most widely used for weed control in crops like soybean, groundnut etc., which possess natural tolerance to these herbicides. However, the crops like rice, maize, wheat, oilseed rape and sunflower etc. are highly sensitive to imidazolinones. Several variants of AHAS genes conferring imidazolinone tolerance have been developed through mutagenesis and selection in different crops and commercialized as Clearfield® crops since 1992. Imidazolinone herbicides control a broad spectrum of grass and broadleaf weeds in imidazolinone-tolerant crops, including weeds that are closely related to the crop itself and some key parasitic weeds (Tan et al. 2005).

Here, we report the transfer of mutant allele of AHAS gene conferring herbicide tolerance from Robin into a mega Basmati rice variety, PB 1121, through MABB and assessment of PB 1121 NILs for agronomic, grain and cooking quality performance in addition to herbicide tolerance.

Material And Methods

Plant materials

In this study, we used the herbicide tolerant rice mutant Robin as donor and a mega-Basmati rice variety, PB 1121 as the recurrent parent (RP) for the transfer of HT trait. The mutant Robin was developed from an upland rice variety Nagina 22 (N22) through induced mutagenesis using ethyl methane sulfonate (EMS) (Shoba et al. 2017). PB 1121, developed by our group at ICAR-Indian Agricultural Research Institute (IARI), New Delhi, is a Basmati rice variety par excellence in grain and cooking quality with an exceptionally high cooked kernel length (22 mm) and elongation ratio of > 2.5. This variety is currently grown on approximately 1.2 million ha area (60 percent of the Basmati area in India) and contributes annually 3.41 billion USD to foreign exchange earnings through the export of Basmati rice (Singh et al. 2018).

MABB Strategy For Development Of HT Lines

PB 1121 and mutant Robin were first evaluated for their tolerance to the herbicide “Imazethapyr” at the dose of 2.5 ml/litre. The F1 seeds were produced by crossing PB 1121 as female and Robin as male and the hybridity of the F1 plants was tested using the SSR marker RM6844 linked with AHAS gene, The F1s were designated as Pusa 1979 and single F1 was backcrossed with PB 1121 to generate BC1F1 seeds. One BC1F1 plant with the highest recovery of RPG and recurrent parent phenome (RPP) was backcrossed to PB 1121 to generate the BC2F1 seeds. A similar strategy was employed till BC4F1 generation, wherein in
each of the generations, the plant heterozygous for the mutant AHAS allele along with maximum recovery for RPG and RPP were identified. The superior BC$_4$F$_1$ plants were advanced to BC$_4$F$_2$ generation and plants homozygous for the mutant AHAS gene were identified. Further, the selected BC$_4$F$_2$ plants were advanced to BC$_4$F$_4$ generation via pedigree-based phenotypic selection (Fig. 1).

**Molecular Analysis**

Total genomic DNA from leaf tissues was extracted using Cetyl Trimethyl Ammonium Bromide (CTAB) method (Doyle 1991). The PCR reaction of a total 10 µl volume was set up which included, 25–30 ng of template DNA, 5 pmol each of the forward and reverse primers and 2X red dye PCR master mix (Genei Laboratories Pvt. Ltd., Bangalore). The program of PCR amplification consisted of initial denaturation at 95 °C for 5 min; 35 cycles of denaturation at 95 °C for 40 s, annealing at 58 °C for 40 s, extension at 72 °C for 1 min; and a final extension at 72 °C for 10 min. The amplified products were resolved on 3.5% Metaphor™ Agarose gel mixed with 0.1 mg/ml ethidium bromide. The amplicons were visualized on ultraviolet trans-illuminator (Gel Doc™ + Imager, Bio- Rad Laboratories Inc., U.S.A).

**Foreground And Background Selection**

Foreground selection for the identification of plants carrying the mutant AHAS allele was carried out using SSR marker RM6844 linked with AHAS gene at a distance of 1.2 cM in chromosome 2 (Shoba et al. 2017), primer details are given in Table S1. For background selection, the primer sequence of genome-wide SSR markers was fetched from the rice marker database of Gramene (http://www.gramene.org). A total of 856 SSR markers were used to identify polymorphic markers between the parents PB 1121 and Robin, for use in background selection (Table S2). During the background selection in backcross generations, the homozygous and heterozygous plants for PB 1121 allele at each marker loci were counted separately. A reductionist strategy was considered for this, markers that were found to be homozygous for PB 1121 allele in a given generation were not included in subsequent generations for background selection. RPG recovery was estimated using the formula:

$$\text{RPG recovery (％)} = \frac{\text{Number of markers homozygous for RP alleles} + (0.5 \times \text{Number of markers heterozygous})}{\text{Total number of polymorphic markers used for background selection}} \times 100$$

The RPG recovery was visualized using Graphical GenoTypes (GGT) Version 2.0 software (Van Berloo 1999). Based on molecular marker analysis, similarity of NILs to PB 1121 was computed using Jaccard’s coefficient of similarity for generating a dendrogram following an unweighted pair group method with arithmetic mean (UPGMA). Further, for cluster analysis, NTSYS-PC-2.02f (Rohlf 1998) was used.

**Molecular Screening For Aroma Gene**
The NILs and the parents were also screened for the presence of badh2 gene, responsible for aroma in Basmati rice, using gene based marker ‘nkssbadh2’ (Amarawathi et al. 2008). Primer details are given in Table S3.

**Screening For Imazethapyr Tolerance**

Twelve HT-NILs of PB 1121 along with PB 1121 and Robin were planted in a randomized complete block design (RCBD) with three replications and sprayed with herbicide, Imazethapyr (commercially available as Pursuit™) @ concentration of 2.5 ml/liter, after 10 days of transplanting. Another set of same experimental material was grown side by side and used as unsprayed control with manual weeding. Visual observation of herbicide tolerance of the HT-NILs was made on 15 days after spray as per the standard protocol in rice (Shoba et al. 2017).

**Evaluation Of Agro-morphological, Grain And Cooking Quality Parameters**

Agro-morphological evaluation of the HT-NILs and parents was done in RCBD with three replications following recommended agronomic practices. Data on the agro-morphological traits viz, days to 50% flowering (DFF), plant height (PH), number of productive tillers per plant (NPT), panicle length (PL), spikelet fertility (SF %), thousand grain weight (TGW) were recorded on five plants taken at random from the two middle rows of each plot under both sprayed and unsprayed conditions. The plot yield was recorded in kilogram/hectare (kg/ha) from each replication. The data on grain and cooking quality traits such as hulling percentage (HUL%), milling percentage (MIL%), head rice recovery (HRR%), kernel length before cooking (KLBC), kernel breadth before cooking (KBBC), kernel length after cooking (KLAC), kernel breadth after cooking (KBAC), kernel elongation ratio (ER), alkali spreading value (ASV) (Little et al., 1958) and aroma (Sood and Siddiq 1978) was recorded.

The statistical analysis of agro-morphological data was carried out using CropStat 7.2 (IRRI CropStat, 2014). Student’s t-test was performed for statistical significance differences for yield between unsprayed and herbicide sprayed condition at P ≤ 0.05.

**Results**

**Development of PB 1121-NILs with herbicide tolerance**

The recurrent parent PB 1121 and donor Robin were validated for their tolerance to herbicide, Imazethapyr. PB 1121 was highly susceptible and exhibited complete mortality within 15–20 days of herbicide spray, while the donor Robin was highly tolerant to Imazethapyr (Fig. 2). MABB was adopted to transfer the HT trait into PB 1121 from Robin. F1 plants obtained from the cross, PB 1121 and Robin were confirmed for their hybridity using AHAS gene linked SSR marker, RM6844. A true F1 plant was
backcrossed to PB 1121 and 262 BC1F1 plants were generated. Of these, 128 plants were heterozygous for RM6844 with RPG recovery ranging from 70.98–80.35%. A plant, Pusa 1979-14 with the highest RPG (80.35%) and with a higher level of phenotypic similarity to PB 1121 was backcrossed and 188 BC2F1 plants were generated, of which 94 plants were heterozygous for foreground marker, RM6844 and the RPG recovery ranged from 81.25 to 92.85%. The superior BC2F1 plant was further backcrossed and 152 BC3F1 seeds were produced, out of which 73 plants were found to be heterozygous for the foreground marker and these plants were subjected to phenotypic evaluation for agro-morphological traits and background selection was done using 16 SSR markers remained heterozygous in the BC2F1 generation. The RPG recovery in the BC3F1 generation ranged from 93.30 to 95.08%. One superior BC3F1 plant, with maximum RPG (95.08%) and relatively superior recovery for RPP was and backcrossed to generate 122 BC4F1 plants. A total of 58 plants were found to be heterozygous for RM6844 with RPG recovery range from 94.64 to 98.21%. A plant with maximum RPG recovery (98.21%) was selfed to produce BC4F2 population. Out of 384 BC4F2 plants, 88 plants were found to be homozygous for AHAS gene linked SSR marker RM6844.

The mutant AHAS homozygous plants were evaluated for grain and cooking quality traits and 40 superior plants were identified. These 40 progenies were further evaluated for yield and quality traits. Based on the family bulk yield and quality traits, twelve families were selected for further detailed evaluation. The RPG recovery of these families ranged from 98.66 to 99.55% with an average 99.10%, some residual donor segments were observed in chromosome 8 and 11, whereas complete recovery was achieved in all other chromosomes (Fig. 3). All the PB 1121 HT-NILs clustered together with PB 1121 with an average similarity index (SI) of 0.9756. Among the HT-NILs, Pusa 1979-14-7-33-99-15 and Pusa 1979-14-7-33-99-66 showed maximum similarity with SI of 0.982 (Fig. 4).

Finally, selected 12 BC4F4 families were evaluated for Imazethapyr tolerance, agronomic, grain and cooking quality characters in the sprayed and unsprayed condition in a replicated trial.

**Evaluation of PB 1121- HT NILs for tolerance to herbicide Imazethapyr**

The PB 1121 HT-NILs along with parents, PB 1121 and Robin were screened for tolerance to herbicide Imazethapyr. All the NILs homozygous for the mutant AHAS allele exhibited complete tolerance to Imazethapyr. The level of herbicide tolerance in NILs was comparable to DP Robin. While, the RP PB 1121 possessing the wild type AHAS gene did not survive after herbicide spray and showed complete mortality within 15 to 20 days of the Imazethapyr spray (Fig. 5).

**Evaluation of PB 1121 HT-NILs for agronomic, grain and cooking quality**

The mean performance of PB 1121 HT-NILs for yield, yield components and phenological traits, when grown under herbicide sprayed and unsprayed conditions were recorded. All the data from the sprayed condition and only yield data for unsprayed conditions are presented in Table 2. Under herbicide sprayed conditions, PB 1121 showed complete mortality within 15–20 days of spray, while the NILs showed
normal growth and development (Fig. 6). Therefore, the yield and quality traits of NILs under sprayed condition were compared with the data of PB 1121 from unsprayed conditions. It was observed that NILs were at par with PB 1121 in yield, grain and cooking quality traits. NILs Pusa 1979-14-7-33-99-66 and Pusa 1979-14-7-33-99-143 were slightly taller in stature and the NILs Pusa 1979-14-7-33-99-15, Pusa 1979-14-7-33-99-10 and Pusa 1979-14-7-33-99-66 were significantly superior in yield as compared to PB 1121 (Fig. 7 & Table 2) which could be attributed to its higher spikelet fertility. There was no significant difference in the performance of NILs when compared under sprayed and unsprayed conditions (based on Student t-test) while PB 1121 was completely dead under sprayed condition.

Table 1
Number of plants generated and recurrent parent genome recovery in the backcross generations during marker aided introgression of mutant AHAS allele in PB 1121.

| Generation | No. of plants generated | No. of plants carrying mutant AHAS allele in heterozygous/homozygous state | No. of selected plants/families | Genome recovery (%) | Mean Observed RPG Recovery (%) | Expected Average Recovery (%) |
|------------|-------------------------|--------------------------------------------------------------------------------|----------------------------------|---------------------|-------------------------------|-----------------------------|
| F₁         | 15                      | 15#                                                                              | 1                                | *                   |                               |                             |
| BC₁F₁      | 262                     | 128#                                                                             | 1                                | 70.98–80.35         | 75.66                         | 75                          |
| BC₂F₁      | 188                     | 94#                                                                              | 1                                | 81.25–92.85         | 87.05                         | 87.5                        |
| BC₃F₁      | 152                     | 73#                                                                              | 1                                | 93.30–95.08         | 94.19                         | 93.8                        |
| BC₄F₁      | 122                     | 58#                                                                              | 1                                | 94.64–98.21         | 96.42                         | 96.9                        |
| BC₄F₂      | 384                     | 88Δ                                                                              | 40                               | *                   | -                             | -                           |
| BC₄F₃      | 40                      | 40Δ                                                                              | 12                               | *                   | -                             | -                           |
| BC₄F₄      | 12                      | 12Δ                                                                              | 12                               | 98.66–99.55         | 99.10                         | -                           |

*Not estimated; #Plants with mutant AHAS allele in heterozygous state; ΔPlants with mutant AHAS allele in homozygous condition.
Table 2
Agronomic performance of PB 1121 HT-NILs in comparison with the recurrent parent, PB 1121.

| Genotype                  | DFF | PH   | NPT  | PL    | SF%  | TGW  | YLD (kg/ha) | %RPG |
|--------------------------|-----|------|------|-------|------|------|-------------|------|
| Pusa 1979-14-7-33-99-8   | 106.0 | 123.15 | 19.30 | 27.89 | 88.20 | 28.87 | 6701 ± 274.40 | 98.66 |
| Pusa 1979-14-7-33-99-10  | 109.0* | 122.45 | 20.60* | 28.40 | 92.70* | 28.73 | 7129 ± 322.95 | 99.11 |
| Pusa 1979-14-7-33-99-15  | 106.0 | 131.53 | 20.20* | 28.52 | 92.87* | 28.93 | 7310 ± 382.05 | 99.55 |
| Pusa 1979-14-7-33-99-34  | 107.0 | 126.25 | 19.70* | 29.28 | 87.53 | 27.99 | 6673 ± 497.09 | 99.11 |
| Pusa 1979-14-7-33-99-56  | 106.5 | 128.85 | 17.20 | 27.56 | 87.75 | 28.88 | 6836 ± 359.89 | 99.11 |
| Pusa 1979-14-7-33-99-66  | 106.5 | 136.85 | 19.10* | 29.29 | 92.74* | 28.32 | 7248 ± 237.14 | 99.55 |
| Pusa 1979-14-7-33-99-87  | 105.5 | 127.95 | 15.70 | 28.79 | 71.48 | 26.43 | 4716 ± 379.94 | 98.66 |

Data on DFF, PH, NPT, PL, SF%, TGW presented is from Imazethpyr sprayed plots while the yield data (YLD) is presented for both unsprayed and herbicide sprayed plots. *Significant at 5%; DFF: days to 50% flowering; PH: Plant height in cm; NPT: Number of productive tillers; PL: Panicle length in cm; SF: Spikelet fertility percentage; TGW: Thousand grain weight in grams (g), YLD: Plot yield in kg/ha; USP: Unsprayed condition; SP: herbicide (Imazethapyr) Sprayed condition; RPG: Recurrent Parent Genome; CD : Critical difference. #Data on DFF, PH, NPT, PL, SF%, TGW for PB 1121 is from unsprayed plots for comparison, as PB 1121 did not survive under Imazethapyr spray.
| Genotype                  | DFF  | PH   | NPT  | PL   | SF%  | TGW  | YLD (kg/ha) USP | YLD (kg/ha) SP | %RPG |
|--------------------------|------|------|------|------|------|------|----------------|----------------|-------|
| Pusa 1979-14-7-33-99-101 | 105.5| 125.10| 13.80| 27.58| 86.64| 27.32| 5799 ± 368.33 | 5866 ± 231.13 | 98.66 |
| Pusa 1979-14-7-33-99-143 | 106.0| 134.35|15.50 | 28.56| 84.80| 28.30| 5832 ± 586.79 | 5559 ± 265.96 | 98.66 |
| Pusa 1979-14-7-33-99-166 | 107.0| 133.47|17.30 | 27.73| 89.11| 28.54| 6910 ± 426.37 | 7017 ± 522.41 | 99.11 |
| Pusa 1979-14-7-33-99-182 | 107.5| 124.85|16.90 | 27.33| 87.11| 27.92| 6502 ± 346.16 | 6506 ± 316.17 | 98.66 |
| Pusa 1979-14-7-33-99-226 | 106.0| 126.00|16.80 | 27.48| 88.14| 27.90| 6599 ± 368.68 | 6547 ± 282.89 | 99.11 |
| PB 1121#                 | 106.0| 125.58|16.06 | 29.81| 86.10| 28.92| 6250 ± 369.38 | -              | -     |
| CD (0.05)                | 2.90 | 7.39 | 2.76 | 1.31 | 5.35 | 1.29 | 868.26         | 828.96         | -     |

Data on DFF, PH, NPT, PL, SF%, TGW presented is from Imazethapyr sprayed plots while the yield data (YLD) is presented for both unsprayed and herbicide sprayed plots. *Significant at 5%; DFF: days to 50% flowering; PH: Plant height in cm; NPT: Number of productive tillers; PL: Panicle length in cm; SF: Spikelet fertility percentage; TGW: Thousand grain weight in grams (g); YLD: Plot yield in kg/ha; USP: Unsprayed condition; SP: herbicide (Imazethapyr) Sprayed condition; RPG: Recurrent Parent Genome; CD: Critical difference. #Data on DFF, PH, NPT, PL, SF%, TGW for PB 1121 is from unsprayed plots for comparison, as PB 1121 did not survive under Imazethapyr spray.

The mean performance of grain and cooking quality traits of the PB 1121 HT-NILs evaluated under sprayed condition is presented in Table 3. All the NILs possessed extra-long slender grain type with a very strong aroma, and ASV of 7.0, which was similar as that of PB 1121. Hulling percent (%), milling (%), and head rice recovery (%), KLBC, KBBC, KBAC were comparable to PB 1121. Three NILs namely, Pusa 1979-14-7-33-99-15, Pusa 1979-14-7-33-99-10 and Pusa 1979-14-7-33-99-143 were found significantly superior for KLAC and elongation ratio as compared to PB 1121 (Table 3 & Fig. 8)
Table 3
Grain quality traits of PB 1121 HT-NILs in comparison to PB 1121.

| Genotype          | HUL  | MIL  | HRR  | KLBC | KBBC | KLAC | KBAC | E/R | ASV | AROMA |
|-------------------|------|------|------|------|------|------|------|-----|-----|-------|
| Pusa 1979-14-7-33-99-8 | 77.77 | 67.46 | 55.96 | 8.77 | 1.61 | 18.54 | 2.28 | 2.11 | 7   | 3     |
| Pusa 1979-14-7-33-99-10 | 79.77 | 70.40 | 58.40 | 8.56 | 1.69 | 19.66 | 2.36 | 2.29*| 7   | 3     |
| Pusa 1979-14-7-33-99-15 | 78.92 | 68.33 | 54.83 | 8.71 | 1.68 | 19.69 | 2.29 | 2.26*| 7   | 3     |
| Pusa 1979-14-7-33-99-34 | 76.70 | 66.73 | 55.13 | 8.83 | 1.67 | 19.35 | 2.32 | 2.18 | 7   | 3     |
| Pusa 1979-14-7-33-99-56 | 78.66 | 67.90 | 53.80 | 8.81 | 1.64 | 19.10 | 2.31 | 2.16 | 7   | 3     |
| Pusa 1979-14-7-33-99-66 | 78.23 | 68.33 | 56.05 | 8.72 | 1.63 | 18.97 | 2.48*| 2.18 | 7   | 3     |
| Pusa 1979-14-7-33-99-87 | 77.50 | 69.23 | 55.36 | 8.62 | 1.69 | 18.46 | 2.31 | 2.14 | 7   | 3     |

*Significant at 5%; HUL: hulling recovery in percentage; MIL: milling recovery in percentage; HRR: head rice recovery in percentage; KLBC: kernel length before cooking in mm; KBBC: kernel breadth before cooking in mm; KLAC: kernel length after cooking in mm; KBAC: kernel breadth after cooking in mm; E/R: kernel elongation ratio; ASV: alkali spreading value; AROMA: aroma score from panel test; #PB 1121 showed complete mortality after herbicide spray, the data of PB 1121 taken from the unsprayed area is presented for the purpose of comparison.
| Genotype       | HUL   | MIL   | HRR   | KLBC  | KBBC  | KLAC  | KBAC  | E/R | ASV | AROMA |
|---------------|-------|-------|-------|-------|-------|-------|-------|-----|-----|-------|
| Pusa 1979-14-7-33-99-101 | 81.12 | 70.25 | 57.15 | 8.82  | 1.67  | 19.32 | 2.33  | 2.19| 7   | 3     |
| Pusa 1979-14-7-33-99-143  | 78.60 | 69.65 | 56.67 | 8.97  | 1.74  | 20.29 | 2.32  | 2.26| 7   | 3     |
| Pusa 1979-14-7-33-99-166  | 77.07 | 67.89 | 54.09 | 8.75  | 1.67  | 18.88 | 2.31  | 2.16| 7   | 3     |
| Pusa 1979-14-7-33-99-182  | 77.76 | 66.64 | 54.95 | 8.73  | 1.64  | 19.32 | 2.31  | 2.21| 7   | 3     |
| Pusa 1979-14-7-33-99-226  | 77.45 | 67.63 | 55.42 | 8.68  | 1.69  | 19.04 | 2.35  | 2.19| 7   | 3     |
| PB 1121 #                | 78.15 | 69.70 | 55.80 | 8.76  | 1.68  | 18.82 | 2.30  | 2.15| 7   | 3     |
| CD (0.05)                | 3.41  | 3.99  | 3.27  | 0.31  | 0.09  | 0.78  | 0.08  | 0.08| -   | -     |

*Significant at 5%; HUL: hulling recovery in percentage; MIL: milling recovery in percentage; HRR: head rice recovery in percentage; KLBC: kernel length before cooking in mm; KBBC: kernel breadth before cooking in mm; KLAC: kernel length after cooking in mm; KBAC: kernel breadth after cooking in mm; E/R: kernel elongation ratio; ASV: alkali spreading value; AROMA: aroma score from panel test; #PB 1121 showed complete mortality after herbicide spray, the data of PB 1121 taken from the unsprayed area is presented for the purpose of comparison.

Based on molecular analysis of badh2 locus, all the HT-NILs were confirmed to have badh2 allele with 8 bp deletion which is identical to PB 1121 (Fig. 9). In sensory evaluation, the HT-NILs were found to be strongly scented with an aroma score of 3.
Discussion

Basmati rice of India attracts consumers worldwide due to its peerless grain and cooking quality characteristics and appealing aroma. Among the Basmati rice varieties developed in India, PB 1121 is considered as the hallmark of Basmati owing to its exceptional KLAC (20 to 25 mm) and high volume expansion upon cooking (> 4 times) (Singh et al. 2018). Basmati rice is protected under Geographical Indications (GI) for cultivation in seven states of north-western India where rice is primarily cultivated under transplanted conditions. However, due to the limited availability of labor on time, high transplanting cost and depleting water table, there is a need to shift rice cultivation to DSR. Although DSR is a potential technology, the response of farmers for its adoption has been lukewarm owing to heavy weed infestation under DSR. Herbicide tolerant rice varieties provide a potential alternative for effective weed management under DSR.

MABB has several advantages over conventional breeding and has been successfully deployed for improving elite rice varieties for resistance to biotic stresses such as bacterial blight and blast (Joseph et al. 2004; Gopalakrishnan et al. 2008; Singh et al. 2012; Singh V.K. et al. 2012; Singh V.K. et al. 2013; Khanna et al. 2015; Ellur et al. 2016a, 2016b), and abiotic stresses such as salinity tolerance (Babu et al. 2017; Singh et al. 2018a) which it has brought a paradigm shift in Basmati rice breeding (Singh et al. 2011).

In this study, we report the transfer of herbicide tolerance trait from Robin into the genetic background of PB 1121 through MABB. Robin is an EMS induced mutant of an upland drought tolerant rice variety N22, which possess tall plant stature, short grains and lacks aroma (Shoba et al. 2017). Using poor grain quality DP to transfer HT trait into the genetic background of high quality Basmati rice variety PB 1121, is a challenging task especially for the recovery of exquisite grain and cooking quality of this Basmati rice variety in the improved HT-NILs. Foreground selection together with background and phenotypic selection led to precise transfer of mutant \textit{AHAS} gene for herbicide tolerance as well as accelerated the RPG recovery to an extent of 98.66 to 99.55% with four backcross generations. The complete recovery of RPG in mutant \textit{AHAS} gene carrier chromosome as well as rest of chromosome was achieved except chromosomes 8 and 11, which retained some residual donor fragment (Fig. 3). Using the SSR markers for background analysis generally target the non-coding and heterochromatic regions and hence could not quantify the recovery of functional part of the genome. Therefore, background selection was coupled with phenotypic selection for RPP which helped in speeding the process of reconstruction of RPP as it entails the indirect selection for functionally expressed part of the genome (Ellur et al. 2016a).

PB 1121 HT-NILs were at par with PB 1121 for agro-morphological, grain and cooking quality traits, while exhibiting herbicide tolerance similar to Robin. This was possible due to stringent phenotypic selection for agro-morphological traits, grain and cooking quality traits for RP PB 1121 phenotype carried out in each of the backcross generations. The significance of phenotypic selection with background selection in the development of NILs with maximum RPG and RPP recovery has also been earlier demonstrated (Ellur et al. 2016b; Babu et al. 2017). Three NILs, Pusa 1979-14-7-33-99-15, Pusa 1979-14-7-33-99-10 and Pusa
1979-14-7-33-99-66 were significantly superior in yield, grain and cooking quality as compared to PB 1121 indicating the effectiveness of augmenting rigorous phenotypic selection for the cooking quality traits along with background selection.

Pleasant aroma is an essential trait of Basmati rice, which is primarily governed by a recessive gene badh2 located on chromosome 8. The accumulation of 2-acetyl-1-pyrroline (2-AP) in aromatic rice is explained by the loss of function mutations in the badh2 gene (Bradbury et al. 2005; Chen et al. 2008). All the PB 1121 HT-NILs and the recurrent parent PB 1121 were found to carry an 8 bp deletion corresponding to the aromatic allele for the production of 2-AP (Fig. 9).

AHAS is an enzyme which catalyzes two reactions for the synthesis of branched chain amino acids namely, valine, leucine, and isoleucine (Singh and Shaner 1995). Condensation of two pyruvate molecules forms 2-acetolactate which leads to the formation of valine and leucine. While for isoleucine biosynthesis, 2-acetoxyhydroxybutyrate is synthesized from pyruvate and 2-ketobutyrate (Zhou et al. 2007). AHAS-inhibiting herbicides block the substrate access channel in the AHAS enzyme by binding to it and elicit deficiency of branched-chain amino acids (Garcia et al. 2017). The resulting decrease of protein synthesis slows down cell division thus leading to growth difficulties in plants (Yu et al. 2014). However, the mutations within the AHAS gene results in altered AHAS enzyme which confers resistance to the AHAS inhibiting herbicides (Tranel et al. 2002; Duggleby et al. 2000; Christoffers et al. 2006). The entire PB 1121 HT-NILs carrying mutated AHAS gene displayed tolerance to Imazethapyr herbicide when applied @ 2.5 ml/liter with no significant difference in the yield performance, grain and cooking quality traits when compared to HT-NILs under unsprayed conditions. This indicates the effectiveness of mutant AHAS gene to overcome the adverse effect of herbicide Imazethapyr.

In all, the present study reports successful development of non-GM HT NILs in the genetic background of popular Basmati rice variety PB 1121 by introgression of mutated AHAS allele using MABB program. This is the first report of the development of HT rice in India. The PB 1121 HT-NILs are being evaluated in the National Basmati Trials for their subsequent release as commercial varieties. These NILs will help the farmers in adopting DSR in Basmati rice production which would help in economizing the rice production.

**Abbreviations**

DSR: Direct-seeded rice; HT: Herbicide tolerant; MABB: Marker assisted backcross breeding; AHAS: Acetohydroxy acid synthase; PB: Pusa Basmati; DP: Donor parent; RP: Recipient parent; NIL: Near-isogenic line; RPP: Recurrent parent phenome; RPG: Recurrent parent genome; SSR: Simple sequence repeat; ALS: Acetolactate synthase; EMS: Ethyl methane sulfonate; 2-AP: 2-acetyl-1-pyrroline; CD: Critical difference.

**Declarations**
Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Availability of data and material

All relevant data are provided as Tables within the paper and in the Supporting Information files.

Competing interests

The authors declare that they have no competing interests.

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Authors Contributions

The study is part of the Ph.D. thesis research of the first author. N.G. performed the experiments and prepared the manuscript; N.G., A.K.Y. conducted herbicide phenotypic screening; A.K.Y., R.K.E. did the statistical analysis; G.K.S., P.K.B., H.B, K.K.V., C.V., A.M.V.S. and N.M. guided field experiments; A.K., R.K.E. reviewed the manuscript; N.K.S., T.M., supervised the project; A.K.S.; G.K.S. conceptualized and supervised the project, edited and finalized the manuscript. All the authors have read and approved the final manuscript.

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Figures
Figure 1

Marker aided backcross breeding strategy used for transferring the herbicide tolerance trait into PB 1121.
Figure 2
Phenotypic evaluation of donor parent, “Robin” and recurrent parent, “PB 1121” for Imazethapyr herbicide tolerance @ concentration of 2.5 ml/liter, (a) Unsprayed, (b) Sprayed.

Figure 3
Graphical representation of PB 1121 HT-NILs carrying AHAS gene showing the extent of recurrent parent genome (RPG) recovery. CH: chromosome.
Figure 4

Dendrogram showing the genetic relationship among different PB 1121 HT-NILs in BC4F4 generation and their parents, PB 1121 and Robin based on 112 polymorphic SSR markers
Figure 5

Phenotypic screening of PB 1121-HT NILs along with Recurrent Parent (RP), PB 1121 and Donor Parent, Robin for tolerance to the herbicide, Imazethapyr, (a) application of Imazethapyr @ 2.5 ml/liter after 10
days of transplanting, (b) 20 days after herbicide spray showing that complete death of RP, (c) 60 days after herbicide spray, (d) 90 days after herbicide spray.

Figure 6

Field view of the PB 1121 HT-NILs at (a) booting and (b) maturity.
Figure 7

Comparison of yield (kg/ha) between PB 1121 HT-NILs and their parents in unsprayed and herbicide sprayed condition; #PB 1121 seedling did not survive Imazethapyr spray; *Significantly superior to the recurrent parent, PB 1121.
Figure 8

Grain and cooking quality of selected PB 1121 HT-NILs and their parents, PB 1121 and Robin.

Figure 9

A representative image showing the amplification profile of the marker, “nksbad2” based on 8bp deletion in badh2 in the PB 1121 HT-NILs along with recurrent parent (RP), PB 1121 and the non-aromatic donor parent (DP) Robin. 1-12: PB 1121 HT-NILs, Pusa 1979-14-7-33-99-8, Pusa 1979-14-7-33-99-10, Pusa 1979-14-7-33-99-15, Pusa 1979-14-7-33-99-34, Pusa 1979-14-7-33-99-56, Pusa 1979-14-7-33-99-66, Pusa 1979-14-7-33-99-87, Pusa 1979-14-7-33-99-101, Pusa 1979-14-7-33-99-143, Pusa 1979-14-7-33-99-166, Pusa 1979-14-7-33-99-182 and Pusa 1979-14-7-33-99-226
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