Agrobacterium-Mediated Genetic Transformation of Wild Oryza Species Using Immature Embryos

Sae Shimizu-Sato, Katsutoshi Tsuda, Misuzu Nosaka-Takahashi, Toshiya Suzuki, Seijiro Ono, Kim Nhung Ta, Yuri Yoshida, Ken-Ichi Nonomura and Yutaka Sato *

Abstract

Genetic transformation is one of the most important technologies for revealing or modulating gene function. It is used widely in both functional genomics and molecular breeding of rice. Demands on its use in wild Oryza species is increasing because of their high genetic diversity. Given the difficulties in genetic crosses between distantly related species, genetic transformation offers a way to alter or transfer genetic traits in wild rice accessions. However, transformation of wild Oryza accessions by conventional methods using calli induced from scutellum tissue of embryos in mature seeds often fails. Here, we report methods using immature embryos for the genetic transformation of a broad range of Oryza species. First, we investigated the ability of callus induction and regeneration from immature embryos of 192 accessions in 20 species under several culture conditions. We regenerated plants from immature embryos of 90 accessions in 16 species. Next, we optimized the conditions of Agrobacterium infection using a vector carrying the GFP gene driven by the maize ubiquitin promoter. GFP signals were observed in 51 accessions in 11 species. We analyzed the growth and seed set of transgenic plants of O. barthii, O. glumaepatula, O. rufipogon, and O. brachyantha. The plants grew to maturity and set seeds normally. Southern blot analyses using DNA from T0 plants showed that all GFP plants were derived from independent transformation events. We confirmed that the T-DNAs were transmitted to the next generation through the segregation of GFP signals in the T1 generation. These results show that many Oryza species can be transformed by using modified immature-embryo methods. This will accelerate the use of wild Oryza accessions in molecular genetic analyses and molecular breeding.

Keywords: Agrobacterium-mediated transformation, Immature embryo, Oryza, Wild accession

Background

Rice (Oryza sativa L.) is one of the most important crops in the world. Breeding new rice cultivars with strong tolerance to environmental changes caused by global warming is an urgent goal. Because wild Oryza species have high genetic diversity and thus have better chances of being adapted to fluctuating environ-
ments, their use in basic science and breeding is increasing (Morishima and Oka 1970; Vaughan 1994). Wild Oryza accessions have been studied to analyze their drought tolerance (Zhang et al. 2006), shattering (Ishii et al. 2013), awn formation (Ikemoto et al. 2017), zinc accumulation (Ishikawa et al. 2017), salt tolerance (Yichie et al. 2018), yield (Fan et al. 2019), and disease resistance (Kariya et al. 2019). These studies identified genes regulating potential agriculturally important traits. The accumulation of whole genome sequences of wild Oryza species (Oryzabase,
Gramene, http://www.gramene.org) provides opportunities in both basic and applied studies (Stein et al. 2018; Zhao et al. 2018; Shenton et al. 2020).

Genetic transformation is one of the most important technologies for revealing or modulating gene function, and is often used in functional genomics and molecular breeding. Several methods for transferring genes into rice (Shimamoto et al. 1989) have been developed, including particle bombardment (Toriyama et al. 1988), polyethylene glycol-mediated (Zhang and Wu 1988), and Agrobacterium-mediated methods (Potrykus 1991; Hiei et al. 1994). Through the improvement of Agrobacterium-mediated methods, transformation efficiency has become high, and transgenes with single or low copy numbers or large DNA segments with defined ends can be integrated into host genomes (Hiei et al. 1994). Agrobacterium-mediated transformation in rice uses two main types of starting materials: calli induced from scutellum tissue

| Type of medium          | nN6C, CCMC                  | nN6C | CCMC | No callus induced on either |
|------------------------|-----------------------------|------|------|----------------------------|
| O. barthii, AA         | W1473, W1583                | W1643| W1647| W1605, W1643               |
| O. glumaepatula, AA    | W1189, W1191, W2165, W2184, W2203| W1167| W1169, W1477, W2145, W2160, W2201| W1702, W1702, W2149, W2173, W2192|
| O. longistaminata, AA  | W1573                       |      |      |                            |
| O. meridionalis, AA    | W1635, W2100, W2112         | W1297| W1300, W1631, W1638| W2071, W2077, W2080, W2081, W2105|
| O. rufipogon, AA       | W0107, W0593, W0625, W1230, W1239, W1294, W1551, W1685, W1852, W1865, W1965, W2014, W2265| W0629, W0630, W1235, W1739, W1807, W1866, W1944, W1962, W2003, W2051| W2079, W2103, W0102, W0106, W0137, W0610, W0621, W1114, W1666, W1669, W1681, W1690, W1718, W1921, W1945, W2078, W2109, W2263|
| O. punctata (2x), BB   | W1515, W1582, W1586, W1593 | W1514, W1577, W1590, W1592 | W1526 |
| O. eichingeri, CC      | W1315                       | W1291|      | W0002, W0614, W1301, W1830, W1930 |
| O. officinalis, CC     | W1315                       |      |      | W0665, W0566, W1131, W1200, W1252, W1302, W1308, W1361, W1814 |
| O. rhizomatis, CC      | W1323                       | W1805, W1812 | W1808 |
| O. minuta, BBCC        | W1323                       | W0016, W1213 | W0051, W1318, W1319, W1329, W1331, W1336, W1342 |
| O. punctata (4x), BBCC | W1474(B)                   |      |      | W0045, W1328 |
| O. alta, CCDD          |                             | W0017, W0018 | W1147 |
| O. grandiglumis, CCDD  | W1483                       | W1480(B), W2220 | W0613, W1195, W1476 |
| O. latifolia, CCDD     | W1177, W1197                | W0047, W1539 | W0048, W0542, W1166, W1168, W1181, W1184 |
| O. australiensis, EE   | W0008, W1639                | W1632, W2082, W2084 | W1296, W1628, W1630, W2086, W2104 |
| O. brachyantha, FF     | W1401                       | W0654 | W0656, W1407(B), W1706, W1711 |
| O. graminata, GG       | W0003                       | W0067(B) | W0004, W0615 |
| O. meyeriana, GG       | W2068                       |      | W1348 |
| O. longiglumis, HHJJ   | W1227                       |      |      | W1218, W1219, W1220, W1222, W1223, W1224, W1228, W1229 |
| O. ridleyi, HHJJ       |                             |      |      | W0001, W2033, W2035 |
| Sub-subtotal           | 40                          | 43   | 55   | 54 |
| Subtotal               | 138                         |      |      |   |
| Total                  | 192                         |      |      |   |
| Species, genome type | Accession | A and B | A alone | B alone | C | D |
|----------------------|-----------|---------|---------|---------|---|---|
| *O. barthii*, AA     | W0747     | 0/8     | 4/16    |         |   |   |
|                      | W1467     | 5/8     | 0/8     |         |   |   |
|                      | W1473     | 4/16    |         |         |   |   |
|                      | W1583     | 6/16    |         |         |   |   |
|                      | W1588     | 0/8     | 3/8     |         |   |   |
|                      | W1643     | 5/8     | 0/8     |         |   |   |
|                      | W1702     | 0/8     | 6/8     |         |   |   |
| *O. glumaepatula*, AA| W1183     | 0/8     | 6/8     |         |   |   |
|                      | W1189     | 5/16    |         |         |   |   |
|                      | W1191     | 10/16   |         |         |   |   |
|                      | W1477     | 0/8     | 0/8     | 4/8     | 0/8 |
|                      | W2140     | 0/8     | 6/8     |         |   |   |
|                      | W2145     | 0/8     | 0/8     | 2/8     | 0/7 |
|                      | W2149     | 0/8     | 8/16    |         |   |   |
|                      | W2160     | 0/8     | 0/8     | 3/8     | 0/8 |
|                      | W2165     | 13/16   |         |         |   |   |
|                      | W2173     | 0/8     | 10/16   |         |   |   |
|                      | W2184     | 12/16   |         |         |   |   |
|                      | W2192     | 0/8     | 5/8     |         |   |   |
|                      | W2201     | 0/7     | 0/8     | 2/8     | 0/7 |
|                      | W2203     | 9/16    |         |         |   |   |
| *O. longistaminata*, AA | W1573   | 2/40    |         |         |   |   |
| *O. meridionalis*, AA | W1300    | 2/16    | 0/8     |         |   |   |
|                      | W1625     | 0/8     | 3/24    |         |   |   |
|                      | W1631     | 2/12    | 0/8     |         |   |   |
|                      | W1635     | 3/16    |         |         |   |   |
|                      | W2071     | 0/8     | 1/16    |         |   |   |
|                      | W2080     | 0/8     | 1/16    |         |   |   |
|                      | W2105     | 0/8     | 2/16    |         |   |   |
| *O. rufipogon*, AA  | W0102     | 0/8     | 4/8     |         |   |   |
|                      | W0106     | 0/8     | 3/8     |         |   |   |
|                      | W0107     | 12/16   |         |         |   |   |
|                      | W0137     | 0/8     | 7/16    |         |   |   |
|                      | W0593     | 10/16   |         |         |   |   |
|                      | W0610     | 0/8     | 6/16    |         |   |   |
|                      | W0621     | 0/7     | 2/8     |         |   |   |
|                      | W0625     | 7/16    |         |         |   |   |
|                      | W0629     | 3/8     | 0/8     |         |   |   |
|                      | W1230     | 2/16    |         |         |   |   |
|                      | W1239     | 11/16   |         |         |   |   |
|                      | W1551     | 13/16   |         |         |   |   |
|                      | W1666     | 0/8     | 5/9     |         |   |   |
|                      | W1669     | 0/8     | 6/8     |         |   |   |
|                      | W1681     | 0/8     | 4/8     |         |   |   |
| Species, genome type | Accession | A and B | A alone | B alone | C | D |
|----------------------|-----------|---------|---------|---------|---|---|
| W1685                | 12/16     |         |         |         |   |   |
| W1690                | 0/8       | 4/8     |         |         |   |   |
| W1718                | 0/8       | 4/8     |         |         |   |   |
| W1739                | 3/8       | 0/7     |         |         |   |   |
| W1852                | 5/16      |         |         |         |   |   |
| W1865                | 12/16     |         |         |         |   |   |
| W1944                | 3/8       | 0/8     |         |         |   |   |
| W1945                | 0/8       | 4/8     |         |         |   |   |
| W1962                | 3/8       | 0/8     |         |         |   |   |
| W2014                | 11/16     |         |         |         |   |   |
| W2051                | 4/8       | 0/8     |         |         |   |   |
| W2263                | 0/8       | 5/9     |         |         |   |   |
| W2265                | 13/16     |         |         |         |   |   |
| O. punctata (2x), BB | W1582     | 5/16    |         |         |   |   |
|                     | W1586     | 4/16    |         |         |   |   |
|                     | W1590     | 3/8     | 0/8     |         |   |   |
|                     | W1592     | 2/8     | 0/8     |         |   |   |
|                     | W1593     | 4/16    |         |         |   |   |
| O. eichingeri, CC    | W1525     | 2/24    |         | 0/16    |   |   |
| O. officinalis, CC   | W0614     | 0/8     |         | 2/15    |   |   |
|                     | W1301     | 0/8     |         | 3/16    |   |   |
|                     | W1830     | 0/7     |         | 1/16    |   |   |
|                     | W1930     | 0/8     |         | 2/15    |   |   |
| O. rhizomatis, CC    | W1808     | 0/8     |         | 3/16    |   |   |
|                     | W1812     | 4/16    |         | 0/16    |   |   |
| O. minuta, BBCC      | W0051     | 0/8     |         | 3/16    |   |   |
|                     | W1213     | 2/8     |         | 0/8     |   |   |
|                     | W1318     | 0/8     | 0/8     | 0/8     | 1/8 |
|                     | W1323     | 3/15    |         |         |   |   |
|                     | W1331     | 0/8     | 0/8     | 0/8     | 1/8 |
| O. punctata (4x), BBCC | W1474(8) | 2/16    |         |         |   |   |
|                     | W1564     | 0/16    |         | 2/16    |   |   |
| O. alta, CCDD        | W0017     | 2/16    |         | 0/12    |   |   |
| O. grandiglumis, CCDD| W0613     | 0/8     |         | 4/8     |   |   |
|                     | W1195     | 0/8     |         | 3/16    |   |   |
|                     | W1476     | 0/8     |         | 4/16    |   |   |
|                     | W1480(8)  | 3/16    |         | 0/8     |   |   |
|                     | W1483     | 6/16    |         |         |   |   |
|                     | W2220     | 4/16    |         | 0/8     |   |   |
| O. latifolia, CCDD   | W1177     | 2/16    |         |         |   |   |
|                     | W1539     | 1/24    |         | 0/16    |   |   |
| O. brachyantha, FF   | W0056     | 0/8     |         | 5/8     |   |   |
|                     | W1407(8)  | 0/8     |         | 4/8     |   |   |
|                     | W1706     | 0/8     |         | 3/8     |   |   |
of embryos in mature seeds, and immature embryos at approximately 10 days after fertilization. The former method is more convenient because we can collect and stock enough mature seeds to run experiments at any time. The latter method can be applied to a much wider range of cultivated rice (Hiei and Komari 2008) and is applicable to various crop species, including maize (Ishida et al. 1996), wheat (Ishida et al. 2005), and sorghum (Sato-Izawa et al. 2018). Thus, the immature embryo method may be applicable to wild *Oryza* species, for which the mature embryo method is often unsuccessful.

*Oryza* comprises two cultivated species (*O. sativa* and *O. glaberrima*) and 21 wild species (Vaughan 1989, 1994) with 9 genome types—AA, BB, CC, BBCC, CCDD, EE, FF, GG, and HHJJ—according to cytogenetic observations of interspecific F_1 hybrids and total genomic DNA hybridization (Khush et al. 2001). The two cultivated species have the AA genome. The introduction of agronomically useful genes from wild accessions into cultivated species is highly valuable. However, the introduction of those genes from wild accessions with the AA genome into cultivated species by genetic crosses followed by repeated backcrosses is time consuming and labor intensive. In addition, genes linked to genes for reproductive barriers or for undesirable characters are difficult to introduce alone by crossing. The introduction of

| Species, genome type | No. accessions tested | No. accessions with callus induction^a | No. accessions with regeneration^b | Percentage of regenerated accessions^c |
|----------------------|-----------------------|----------------------------------------|------------------------------------|----------------------------------------|
| O. barthii, AA       | 10                    | 8 (80%)                                | 7 (88%)                            | 70%                                   |
| O. glumaepatula, AA  | 18                    | 17 (94%)                               | 14 (82%)                           | 78%                                   |
| O. longistaminata, AA| 1                     | 1 (100%)                               | 1 (100%)                           | 100%                                  |
| O. meridionalis, AA  | 15                    | 13 (87%)                               | 7 (54%)                            | 47%                                   |
| O. rufipogon, AA     | 44                    | 39 (89%)                               | 28 (72%)                           | 64%                                   |
| O. punctata (2x), BB | 8                     | 8 (100%)                               | 5 (63%)                            | 63%                                   |
| O. eichingeri, CC    | 2                     | 1 (50%)                                | 1 (100%)                           | 50%                                   |
| O. officinalis, CC   | 16                    | 7 (44%)                                | 4 (57%)                            | 25%                                   |
| O. rhizomatix, CC    | 3                     | 3 (100%)                               | 2 (67%)                            | 67%                                   |
| O. minuta, BBCC      | 12                    | 10 (83%)                               | 5 (50%)                            | 42%                                   |
| O. punctata (4x), BBCC | 8               | 4 (50%)                                | 2 (50%)                            | 25%                                   |
| O. alta, CCDD        | 3                     | 2 (67%)                                | 1 (50%)                            | 33%                                   |
| O. grandiglumis, CCDD| 7                     | 6 (86%)                                | 6 (100%)                           | 86%                                   |
| O. latifolia, CCDD   | 10                    | 4 (40%)                                | 2 (50%)                            | 20%                                   |
| O. australiensis, EE | 10                    | 5 (50%)                                | 0 (0%)                             | 0%                                    |
| O. brachyantha, FF   | 7                     | 6 (86%)                                | 4 (67%)                            | 57%                                   |
| O. granulata, GG     | 4                     | 2 (50%)                                | 0 (0%)                             | 0%                                    |
| O. meyeriana, GG     | 2                     | 1 (50%)                                | 0 (0%)                             | 0%                                    |
| O. longiglumis, HHJJ | 9                     | 1 (11%)                                | 1 (100%)                           | 11%                                   |
| O. ridleyi, HHJJ     | 3                     | 0 (0%)                                 | 0 (0%)                             | 0%                                    |
| Total                | 192                   | 138 (72%)                              | 90 (65%)                           | 47%                                   |

^a100 × number of accessions with induced calli / total number of tested accessions
^b100 × number of regenerated accessions / number of accessions with induced calli
^c100 × number of regenerated accessions / total number of tested accessions
genes from wild accessions with other genomes by genetic crosses is even harder. Usually, crosses between different genome types do not produce stable offspring. In some cases, aneuploid plants can be obtained but their chromosome composition is unstable and not healthy. Genetic transformation offers promise in the transfer of genes from wild accessions, especially with genomes other than AA, and allows the functional analysis of genes in wild *Oryza* accessions.

The Japanese National Institute of Genetics (NIG) holds in Oryzabase and distributes more than 1700 accessions of wild *Oryza* species. Here, we used the immature embryo method for the genetic transformation of wild rice accessions from a broad range of species with various media and *Agrobacterium* infection conditions and show that many *Oryza* species can be transformed by the immature embryo method.

### Materials and Methods

#### Plant Materials

Seeds of wild *Oryza* species were supplied by NIG. Plants were grown under natural light in a greenhouse and transplanted into a paddy field in June in Mishima, Japan. At 1 month after transplanting, day-length was controlled 12.5 h light / 11.5 h dark to induce flowering. *Oryza sativa* ‘Nipponbare’ was grown as a control in similar conditions but under natural day-length. At 8 to 10 days after flowering, immature embryos were isolated.

**Plant Regeneration from Immature Embryos**

Regeneration tests using immature embryos were conducted according to Hiei and Komari (2008). All culture media are detailed in Supplemental Table 1. Ovaries were sterilized in 70% ethanol (v/v) for 1 min, followed by 15 min in 50% (v/v) sodium hypochlorite solution with shaking, and washed 5 times in sterile distilled water. Hiei and Komari (2008) used two types of callus-inducing media, nN6C and CCMC, to induce calli from *japonica* and *indica* cultivars, respectively. The immature embryos were isolated under a stereomicroscope on a clean bench, placed on nN6C or CCMC medium containing 25 mg/L meropenem, and cultured at 32 °C for 7 days. Then they were transferred to the same fresh media and cultured for another 7 days under the same conditions. Next, we tested regeneration of shoots and roots in four combinations of media, designated A to D (Supplemental Table 2). Calli formed on the immature embryos were transferred to either regeneration medium (N6R) or pre-regeneration medium (NBPRC). On N6R, calli were cultured at 32 °C for 2 weeks, and regenerated plantlets were transferred to rooting medium (N6F). On NBPRC, calli were cultured at 32 °C for 1 week, and were then transferred to regeneration medium (RNM). After 2 weeks on RNM, regenerated plantlets were transferred to rooting medium (MSI). Plants which formed roots on N6F or MSI were eventually transferred to soil in pots and grown to maturity in the greenhouse.

### Table 4 Efficiency of *Agrobacterium* infection of calli induced from immature embryos of 16 wild *Oryza* species

| Strain         | Centrifugation | LBA4404  | EHA105  |
|----------------|---------------|----------|---------|
|               |               | 1100×g   | 20,000×g| 1100×g  | 20,000×g|
| 1              | *O. barthii*, W1467, AA | 1/9      | 2/8     | 0/8     | 1/8    |
| 2              | *O. glumeapetula*, W2184, AA | 3/8      | 3/9     | 1/8     | 2/8    |
| 3              | *O. longistaminata*, W1573, AA | 0/8      | 0/8     | 0/8     | 1/8    |
| 4              | *O. meridionalis*, W2080, AA | 0/9      | 1/8     | 0/8     | 0/8    |
| 5              | *O. rufipogon*, W1551, AA | 3/8      | 2/8     | 0/8     | 1/8    |
| 6              | *O. punctata* 2x, W1582, BB | 2/8      | 0/8     | 0/8     | 0/8    |
| 7              | *O. eichingeri*, W1525, CC | 0/8      | 0/8     | 1/8     | 0/8    |
| 8              | *O. officinalis*, W1301, CC | 0/16     | 0/16    | 0/17    | 0/16   |
| 9              | *O. rhizomatis*, W1808, CC | 0/16     | 0/15    | 0/16    | 0/17   |
| 10             | *O. minuta*, W0051, BBCC | 0/8      | 0/8     | 1/8     | 0/8    |
| 11             | *O. punctata* 4x, W1564, BBCC | 0/8      | 0/8     | 1/8     | 0/8    |
| 12             | *O. alta*, W0017, CCDD | 1/8      | 1/8     | 0/8     | 0/8    |
| 13             | *O. grandiglumis*, W2220, CCDD | 0/16     | 0/18    | 0/16    | 0/18   |
| 14             | *O. laitifolia*, W1177, CCDD | 0/24     | 0/25    | 0/24    | 0/24   |
| 15             | *O. brachyantha*, W1711, FF | 2/9      | 1/8     | 0/8     | 1/8    |
| 16             | *O. longiglumis*, W1227, HHJJ | 0/24     | 0/24    | 0/25    | 0/24   |
| 17             | *O. sativa*, Nipponbare, AA | 88/90    | 76/79   | 77/83   | 81/85  |

The number of GFP positive calli / total number of tested immature embryos.
Binary Vectors and Transformation of Agrobacterium tumefaciens

The binary plasmid vector (pPUG1–1) used in this study contains a hygromycin phosphotransferase (hpt) gene under the control of the cauliflower mosaic virus 35S promoter, and a green fluorescent protein (GFP) gene under the control of the maize ubiquitin promoter. The vector was transferred into Agrobacterium tumefaciens strains LAB4404 and EHA105 by electroporation in an E. coli pulser (Bio-Rad).

Agrobacterium-Mediated Transformation

Agrobacterium-mediated transformation was conducted according to Hiei and Komari (2008). All media included 25 mg/L meropenem instead of cefotaxime and carbenicillin. Strain LBA4404 or EHA105 harboring pPUG1–1 was grown on LB medium supplemented with 50 mg/L hygromycin at 28 °C for 3 days. The immature embryos were held in a water bath at 43 °C for 30 min, and then cooled in water. They were then centrifuged in an angle rotor at either 1100× or 20,000×g for 10 min at 22 °C. They were placed on co-cultivation medium containing 100 μM acetosyringone with the scutellum side up, and 5 μL of Agrobacterium suspension was dropped on them. After incubation for 7 days at 25 °C in the dark, plants were regenerated as described in the previous section, with the addition of hygromycin into all media for selection.

Observation of GFP Fluorescence

GFP fluorescence was observed under a fluorescence microscope with a GFP filter (SZX16, Olympus).

Extraction of Rice Genomic DNA and Southern Blot Analysis

Total genomic DNA was extracted from ca. 1 g of fresh leaf of each transgenic line with a DNeasy Plant Maxi Kit (Qiagen) according to the manufacturer’s protocol. PCR analysis to detect the full-length hpt gene was performed using GoTaq DNA polymerase (Promega) with the primer set 5′-ATGAAAAAGCCTGAACCTCACC CG-3′ / 5′-CTATTCCTTTGCCCTCGGACGAGT-3′. Southern blot analysis was performed with an ECL Direct Nucleic Acid Labelling and Detection System (GE Healthcare) according to the manufacturer’s protocol. Genomic DNA (5 μg) was digested by HindIII (Takara) and separated by electrophoresis. For the preparation of hpt probe, the full-length hpt gene was amplified with the primers used for PCR analysis above. Southern blot hybridization signals were detected on an Image Quant LAS 4000 Mini imager (GE Healthcare).

Results and Discussion

Optimization of Culture Media for Regeneration of Plantlets from Immature Embryos from Wild Oryza Accessions

First we tested the response of immature embryos to callus induction. From 192 accessions in 20 wild Oryza species, 138 accessions in 19 species formed callus on

![Fig. 1 Agrobacterium-mediated transformation of Oryza brachyantha W1711. a At 10 days after infection, GFP signals were observed in immature embryos. Bar = 1 mm. b Callus was cultured for 2 weeks on selection medium. Part of the callus showed a GFP signal. Bar = 2 mm. c After 2 weeks' culture on regeneration media, shoots with strong GFP signals were formed. Bar = 3 mm. d Regenerated plants on rooting medium. Bar = 2 cm. e, f Leaves through bright field and GFP filters: left, transgenic; right, non-transgenic. Bar = 1 cm. g Transgenic plants growing to maturity without morphological abnormalities. Bar = 5 cm. h, i T1 seeds of a transgenic plant through bright field and GFP filters, T1 seeds with and without GFP signal were segregated. Bar = 4 mm]
| Species, genome type | Strain     | LBA4404 1100xg | LBA4404 20,000xg | EHA105 1100xg | EHA105 20,000xg | Infection efficiencya |
|----------------------|------------|----------------|------------------|----------------|------------------|-----------------------|
| O. barthii, AA       | W1467      | 1/9            | 2/8              | 0/8            | 1/8              | 3/6 (50%)             |
|                      | W1473      | 0/8            | 0/8              |                |                  |                       |
|                      | W1583      |                |                  | 1/8            |                  |                       |
|                      | W1588      | 0/8            |                  | 0/8            |                  |                       |
|                      | W1643      | 0/8            |                  | 0/8            |                  |                       |
|                      | W1702      | 2/9            |                  |                |                  |                       |
| O. glumaepatula, AA  | W1183      | 1/8            |                  |                |                  | 13/14 (93%)           |
|                      | W1189      | 0/8            |                  | 0/8            |                  |                       |
|                      | W1191      |                |                  | 2/8            |                  |                       |
|                      | W1477      | 1/8            |                  | 3/16           |                  |                       |
|                      | W2140      | 1/8            |                  | 1/8            |                  |                       |
|                      | W2145      | 1/8            |                  |                |                  |                       |
|                      | W2149      | 2/8            |                  |                |                  |                       |
|                      | W2160      | 2/8            |                  | 3/16           |                  |                       |
|                      | W2165      | 1/8            |                  |                |                  |                       |
|                      | W2173      | 3/16           |                  | 2/16           |                  |                       |
|                      | W2184      | 3/8            |                  | 3/9            | 1/8              | 2/8                   |
|                      | W2192      | 2/8            |                  |                |                  |                       |
|                      | W2201      | 1/8            |                  | 1/8            |                  |                       |
|                      | W2203      | 1/8            |                  | 2/8            |                  |                       |
| O. longistaminata, AA| W1573      | 0/8            |                  | 0/8            | 0/8              | 1/8                   | 1/1 (100%)            |
|                      | W1300      | 0/9            |                  | 0/8            |                  |                       | 1/7 (14%)             |
| O. meridionalis, AA  | W1625      | 0/8            |                  | 0/8            |                  |                       |
|                      | W1631      | 0/8            |                  | 0/8            |                  |                       |
|                      | W1635      | 0/8            |                  | 0/7            |                  |                       |
|                      | W2071      | 0/8            |                  | 0/8            |                  |                       |
|                      | W2080      | 0/9            |                  | 1/8            | 0/8              | 0/8                   |
|                      | W2105      | 0/8            |                  | 0/8            |                  |                       |
| O. rufipogon, AA     | W0102      | 1/8            |                  |                | 21/26 (81%)      |                       |
|                      | W0106      | 1/8            |                  |                |                  |                       |
|                      | W0107      | 2/8            |                  | 2/16           |                  |                       |
|                      | W0137      | 0/8            |                  | 0/8            |                  |                       |
|                      | W0593      | 1/8            |                  |                |                  |                       |
|                      | W0610      | 1/8            |                  | 1/8            |                  |                       |
|                      | W0621      | 1/8            |                  | 1/8            |                  |                       |
|                      | W0625      | 1/9            |                  | 1/8            |                  |                       |
|                      | W0629      | 1/8            |                  | 1/8            |                  |                       |
|                      | W1230      | 0/8            |                  | 0/8            |                  |                       |
|                      | W1239      | 2/8            |                  |                |                  |                       |
|                      | W1551      | 3/8            |                  | 2/8            | 0/8              | 1/8                   |
|                      | W1666      | 1/8            |                  |                |                  |                       |
|                      | W1669      | 2/8            |                  |                |                  |                       |
|                      | W1681      | 1/8            |                  |                |                  |                       |
either or both media (Table 1). Using these 138 accessions, we tested the regeneration of shoots and roots in conditions A and B, and then using accessions which did not respond to either, in conditions C and D. Twenty-eight accessions regenerated from immature embryo calli in conditions A and B, 18 only in condition A, and 38 only in condition B (Table 2). Of 54 accessions which did not respond to A or B, 4 regenerated under C and 2 under D (Table 2).

Overall, among the 192 accessions tested, 90 formed calli from immature embryos and regenerated in tissue culture under one or more of the conditions tested. These accessions include AA, BB, CC, BBCC, CCDD, FF, and HHJJ genome species. Thus, we have identified...
conditions of tissue culture and the recovery of plantlets suitable for a wide range of *Oryza* species except those with EE and GG genomes. Condition B covered the widest range of wild *Oryza* accessions, followed by condition A. Very few of the accessions that did not respond to condition A or B responded to condition C or D. In particular, 4 accessions of *O. glumaepatula* (W1477, W2145, W2160, and W2201) responded to condition C, while other accessions of the same species responded to A or B. Thus, each species may have a genotype-dependent optimal condition for the regeneration of plantlets from immature embryos.

In general, species with the AA genome had a high regeneration success rate, notably *O. barthii, O. glumaepatula, O. meridionalis*, and *O. rufipogon* (Table 3). Although the success rate in *O. longistaminata* was 100%, we tested only one accession. As *O. longistaminata* generally propagates vegetatively through rhizomes rather than sexually, the immature embryo method is not particularly applicable for its transformation. We could not regenerate plantlets from immature embryos of *O. australiensis, O. granulata, O. meyeriana*, or *O. ridleyi*. Of 10 *O. australiensis* (EE) accessions tested, 5 formed calli and green spots but did not form shoots. Of 4 accessions of *O. granulata* and 2 of *O. meyeriana* (both GG) tested, 2 and 1 accessions, respectively, produced calli but did not form green spots or regenerate shoots and roots. It is well known that the plant hormone cytokinin plays an important role in shoot regeneration from calli. Further consideration of medium conditions, such as the concentration of cytokinin, may be necessary to induce regeneration from immature embryos of wild *Oryza* species with EE and GG genomes.

**Optimization of Agrobacterium Infection of Immature Embryo Calli from Wild Oryza Accessions**

Next, we compared the conditions of *Agrobacterium* infection of calli. Heat treatment and centrifugation of calli before *Agrobacterium* infection can increase the transformation efficiency in cultivated rice, maize, wheat, and sorghum (Hiei et al. 2006; Ishida et al. 2014; Sato-Izawa et al. 2018). Transformation efficiency is also affected by *Agrobacterium* strains (Hiei and Komari 2008). Therefore, we tested heat treatment, two conditions of centrifugation, and two *Agrobacterium* strains (Table 4). To monitor infection, we used a vector carrying the GFP driven by the ubiquitin promoter. Following infection, we counted the number of immature embryos with GFP-positive spots at 10 days (Fig. 1a). Using calli induced on immature embryos of *Nipponbare*, we showed that neither the speed of centrifugation nor the *Agrobacterium* strain affected the efficiency of infection (Table 4), possibly owing to very high infection efficiency in all conditions tested. Next, we infected calli of 16 wild *Oryza* accessions, one per species in which regeneration was achieved, and measured infection (Table 4). As wild accessions tend to flower gradually in a panicle, it is difficult to collect many ovaries at 8 to 10 day after pollination (DAP) at the same time, so we used 8 to 25 immature embryos of one accession of each species. GFP fluorescence showed that 11 out of 16 accessions were infected in one or more conditions tested (Table 4). This rate is considerably lower than that of *Nipponbare*. *Nipponbare* panicles flower in a brief period, so one can collect many immature embryos at once at the ideal stage for *Agrobacterium* infection. This difference could be a reason for the high infection ratio in *Nipponbare*. Another possibility is that the developmental window of immature embryos which can accept *Agrobacterium* infection may be broader in *Nipponbare* than in wild accessions.

Next, we tested *Agrobacterium* infection in 60 more wild accessions (Table 5), mainly under conditions in which an accession of the same species was infected (Table 4). Of the 76 accessions in 16 species, infection was successful in 51 accessions in 11 species. In most species (*O. barthii, O. glumaepatula, O. longistaminata, O. rufipogon, O. punctata* (2x), *O. eichingeri, O. minuta, O. punctata* (4x), *O. alta*, and *O. brachyantha*), infection and GFP fluorescence had a relatively high success rate (≥50%). However, in *O. meridionalis*, only 1 among 7 accessions showed a GFP signal. This result suggests that this species may have a mechanism that suppresses *Agrobacterium* infection in these conditions. Infection was not successful in *O. officinalis, O. rhizomatis, O. grandiglumis, O. latifolia*, or *O. longiglumis*. However, only 1 or 2 accessions were used for these species, so it is not clear whether this is a general tendency in each species. Further consideration of the conditions of infection, such as cold shock treatment, may be necessary (Sato-Izawa et al. 2018).
Generation of Transgenic Plants and Inheritance of Transgene in Wild Oryza Accessions

Next, we tried to generate transgenic plants expressing GFP by introducing the same vector used in the previous analysis into *O. barthii* (2 accessions), *O. glumaepatula* (8), *O. rufipogon* (15), and *O. brachyantha* (1) (Table 6). The transformation of *O. brachyantha* W1711 is shown as an example (Fig. 1a–i). A GFP signal was observed at 10 days after infection (Fig. 1a). Plants were selected on callus-inducing...
medium with antibiotics (Fig. 1b) and regenerated (Fig. 1c). Regenerated shoots were transferred to rooting medium (Fig. 1d) and then to pots with soil (Fig. 1g) and were grown to maturity. The regenerated T0 plants expressed GFP in leaves (Fig. 1e, f) and roots (Fig. 2). We confirmed the introduction of the transgene in regenerated T0 plants of *O. barthii* W1467, *O. glumaepatula* W2184, *O. rufipogon* W1551, and *O. brachyantha* W1711 by PCR (Fig. 3a–d) and found that the transformation events were all independent, with 1 to 4 T-DNA insertion loci, on the basis of the number of bands in Southern blot analysis (Fig. 3e–h).

We obtained five or more regenerated plants from all accessions tested (Table 6). All grew to maturity without any discernable morphological abnormalities. All plants were fertile and set T1 seeds by self-pollination. We confirmed the transmission and segregation of the transgene in T1 seeds of *O. barthii* W1467, *O. glumaepatula* W2184, *O. rufipogon* W1551, and *O. brachyantha* W1711 by PCR (Fig. 3a–d) and found that the transformation events were all independent, with 1 to 4 T-DNA insertion loci, on the basis of the number of bands in Southern blot analysis (Fig. 3e–h).

We generated transgenic plants from *O. rufipogon* accessions W0106 and W1681 by the immature embryo method but not by the conventional scutellum callus method (data not shown). This difference supports the superiority of the immature embryo method for the transformation of wild *Oryza* accessions. This method opens the door to genome editing, accelerating the study of wild *Oryza* genetic resources for molecular genetic analysis and future use in molecular breeding.

**Supplementary information**

Supplementary information accompanies this paper at https://doi.org/10.1186/s12284-020-00394-4.

**Additional file 1:** Supplemental Table 1. Composition of media (1 L) used in this study. Supplemental Table 2. Work flow of test of regeneration from callus derived from immature embryos of wild *Oryza* species.

**Abbreviations**

hpt: hygromycin phosphotransferase; GFP: green fluorescent protein; DAP: day after pollination.

**Acknowledgements**

We thank Dr. Hiei Y and members of the Plant Innovation Center of Japan Tobacco for their technical assistance and tips for immature embryo culture. The wild *Oryza* accessions used in this study were distributed from NIG, which is supported by the National Bioresearch Project, AMED, Japan.

**Authors’ Contributions**

SSS, KT, MNT, TS, SO, KNT, and YY performed the experiments. KN and YS conceived and designed the study. All authors read and approved the final manuscript.

**Funding**

This work was supported by funding from the National Bioresearch Project’s Fundamental Technologies Upgrade Program to YS from AMED, Japan, and supported by a grant from RDA, Korea (PJ01321803).

**Availability of Data and Materials**

All datasets are available from the corresponding author on reasonable request.

**Ethics Approval and Consent to Participate**

Not applicable.

---

**Table 7** Inheritance and segregation of the transgenes

| Species               | Line number | GFP gene in T1 seeds | Segregation ratio (χ², P) |
|----------------------|-------------|----------------------|--------------------------|
|                      |             | Positive  | Negative  |                        |
| *O. barthii*, W1467 | L1          | 56       | 4         | 15:1 (0.02, 0.89)      |
|                      | L2          | 23       | 7         | 3:1 (0.04, 0.83)       |
| *O. glumaepatula*, W2184 | L1      | 57       | 3         | 15:1 (0.16, 0.69)      |
|                      | L4          | 22       | 8         | 3:1 (0.04, 0.83)       |
| *O. rufipogon*, W1551 | L1         | 56       | 4         | 15:1 (0.02, 0.89)      |
|                      | L4          | 22       | 8         | 3:1 (0.04, 0.83)       |
| *O. brachyantha*, W1711 | L1    | 57       | 3         | 15:1 (0.16, 0.69)      |
|                      | L2          | 23       | 7         | 3:1 (0.04, 0.83)       |

*aLine number is the same as in Fig. 3*
References

Fan F, Long W, Liu M, Yuan H, Pan G, Li N, Li S (2019) Quantitative trait locus mapping of the combining ability for yield-related traits in wild rice (*Oryza longistaminata*). J Agric Food Chem 14:8766–8772

Hiei Y, Ishida Y, Kasaoka K, Komari T (2006) Improved frequency of transformation in rice and maize by treatment of immature embryos with centrifugation and heat prior to infection with *Agrobacterium tumefaciens*. Plant Cell Tissue Organ Cult 87:233–243

Hiei Y, Komari T (2008) *Agrobacterium*-mediated transformation of rice using immature embryos or calli induced from mature seed. Nature Prot 3:824–834

Hiei Y, Ohta S, Komari T, Kumashiro T (1994) Efficient transformation of rice (*Oryza sativa* L.) mediated by *Agrobacterium* and sequence analysis of the boundaries of the T-DNA. Plant J 6:271–282

Ikemoto M, Otsuka M, Thanh PT, Phan PDT, Ishikawa R, Ishii T (2017) Gene interaction at seed-awning loci in the genetic background of wild rice. Genes Gen Syst 1:21–26

Ishida Y, Hiei Y, Komari T (2005) Advances in wheat genetics: from genome to field. Springer, Tokyo

Ishida Y, Saito H, Ohta S, Hiei Y, Komari T, Kumashiro T (1996) High efficiency transformation of maize (*Zea mays* L.) mediated by *Agrobacterium tumefaciens*. Nat Biotechnol 14:745–750

Ishida Y, Tsunashima M, Hiei Y, Komari T (2014) Wheat (*Triticum aestivum* L.) transformation using immature embryos. In: Wang K (ed) *Agrobacterium* protocols, vol 1. Springer, New York, pp 189–198

Ishi T, Numaguchi K, Miura K, Yoshida K, Thanh PT, Htun TM, Yamasaki M, Komeda N, Matsumoto T, Terauchi R, Ishikawa R, Ashikari M (2013) *OsLG1* regulates a closed panicle trait in domesticated rice. Nat Genet 45:462–465

Ishikawa R, Ivata M, Taniko K, Monden G, Miyazaki N, Ono C, Tsujimura Y, Ishida S, Ma JF, Ishii T (2017) Detection of quantitative trait loci controlling grain zinc concentration using Australian wild rice, *Oryza meridionalis*, a potential genetic resource for biofortification of rice. PLoS One 12(10):e0187224

Kariya K, Murata K, Kubo T, Ube N, Ueno K, Yabuta Y, Teraishi M, Okumoto Y, Mori N, Ishihara A (2019) Variation of diterpenoid phytolalexin oryzalexin a production in cultivated and wild rice. Phytochemistry 166:12057

Kush GS, Brar DS, Hardy B (2001) Rice genetics IV. IRRI, Science Pub Inc, NH

Morishima H, Oka H (1970) A survey of genetic variations in the populations of wild *Oryza* species and their cultivated relatives. Japanese J Genet 45:371–385

Potrykus I (1991) Gene transfer to plants: assessment of published approaches and results. Annu Rev Plant Physiol Plant Mol Biol 42:205–225

Sato-Izawa K, Tokue K, Ezura H (2018) Development of a stable *Agrobacterium*-mediated transformation protocol for *Sorghum bicolor* cv430. Plant Biotecnol (Tokyo) 25:181–185

Shenton M, Kobayashi M, Terashima S, Ohyanagi H, Copetti D, Hernandez-Hernandez T, Zhang J, Ohmido N, Fujita M, Toyoda A, Iwata M, Ishikawa R, Iwata M, Teraishi M, Okumoto Y, Mori N, Ishihara A (2019) Variation of diterpenoid phytolalexin oryzalexin a production in cultivated and wild rice. Phytochemistry 166:12057

Shimamoto K, Uchimiya H, Hinata K (1988) Transgenic rice plants after direct gene transfer into protoplasts. Bioresearch Technol 6:1072–1074

Vaughan DA (1989) The genus *Oryza*. Current status of taxonomy. IRRI res pap Ser

Vaughan DA (1994) Wild relatives of rice: genetic resources hand book. IRRI, Philippines

Zheng W, Wu R (1988) Efficient regeneration of transgenic plants from rice protoplasts and correctly regulated expression of the foreign gene in the plants. Theor App Genet 76:835–840

Zhang X, Zhou S, Fu Y, Su Z, Wang X, Sun C (2006) Identification of a drought tolerant introgression line derived from Dongxiang common wild rice (*O. rufipogon* Griff.). Plant Mol Biol 62:247–259

Zhou Q, Feng Q, Lu H, Li Y, Wang A, Tian Q, Zhan Q, Lu Y, Zhang L, Huang T, Wang Y, Fan D, Zhao Y, Qang Z, Zhou C, Chen J, Zhu C, Li W, Weng Q, Xu Q, Wang Z, Wei X, Han B, Huang X (2018) Pan-genome analysis highlights the extent of genomic variation in cultivated and wild rice. Nat Genet 50:278–284

Publisher's Note

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.