Transcriptome comparison between pluripotent and non-pluripotent calli derived from mature rice seeds

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In vitro plant regeneration involves a two-step practice of callus formation and de novo organogenesis. During callus formation, cellular competence for tissue regeneration is acquired, but it is elusive what molecular processes and genetic factors are involved in establishing cellular pluripotency. To explore the mechanisms underlying pluripotency acquisition during callus formation in monocot plants, we performed a transcriptomic analysis on the pluripotent and non-pluripotent rice calli using RNA-seq. We obtained a dataset of differentially expressed genes (DEGs), which accounts for molecular processes underpinning pluripotency acquisition and maintenance. Core regulators establishing root stem cell niche were implicated in pluripotency acquisition in rice callus, as observed in Arabidopsis. In addition, KEGG analysis showed that photosynthetic process and sugar and amino acid metabolism were substantially suppressed in pluripotent calli, whereas lipid and antioxidant metabolism were overrepresented in up-regulated DEGs. We also constructed a putative coexpression network related to cellular pluripotency in rice and proposed potential candidates conferring pluripotency in rice callus. Overall, our transcriptome-based analysis can be a powerful resource for the elucidation of the molecular mechanisms establishing cellular pluripotency in rice callus.

Callus is a pluripotent cell mass, which can be produced from a single differentiated somatic cell1-2. Pluripotent callus can undergo de novo organ formation or embryogenesis, giving rise to a new organ or even an entire plant2. A balance of two phytohormones, auxin and cytokinin, underlies a two-step in vitro tissue culture3: incubation of tissue explants on auxin-rich callus-inducing medium (CIM) activates cell proliferation to facilitate callus formation4, whereas de novo shoot regeneration can be initiated by incubation on the cytokinin-rich shoot-inducing medium (SIM)5,6.

A particular emphasis has been placed on the callus formation process, because active cell proliferation drives pluripotency acquisition, which is a fundamental basis of plant regeneration7. Molecular mechanism underlying callus formation is starting to emerge. Accumulating evidence has shown that the CIM-derived callus resembles root primordium, regardless of origin of tissue explants7-9. In Arabidopsis, callus formation is initiated from the founder cell, such as pericycle (or pericycle-like) cells8. The founder cell undergoes asymmetric cell division and facilitates the acquisition of root primordium identity with the activation of genes including WUSCHEL-RELATED HOMEOBOX 11 (WOX11) and LATERAL ORGAN BOUNDARIES DOMAINS (LBDs)9,10. After acquiring root primordium characteristics, callus cells establish a regeneration competence via expression of root stem cell regulators, including PLT1PLT2, SHORT-ROOT (SHR), SCARECROW (SCR), WOX5, WOX7, and WOX147,8,11-13. Consistently, plt1plt2, scr, and wox5 wox7 wox14 mutants exhibit impaired de novo shoot regeneration possibly due to the failure of pluripotency acquisition7,8,11,12. It has been proposed that PLT and WOX-induced root stem cell identity is likely suspected as cellular nature of pluripotency14-16.

Several upstream regulators of the pluripotency factors have been identified. The PLT3, PLT5, and PLT7 genes, which are induced upon incubation on CIM11, promote the acquisition of pluripotency through the induction of PLT1 and PLT211. Additionally, the root stem cell regulators are further coordinated by epigenetic modification. The HISTONE ACETYLTRANSFERASE OF THE GNAT/MYST SUPERFAMILY 1 (HAG1)/GENERAL CONTROL NONREPRESSED 5 (GCN5) protein binds to the promoters of the WOX5, WOX14, SCR, PLT1, and

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PLT2 genes and deposits H3 acetylation (H3ac) to activate gene expression during callus formation\textsuperscript{12,17}. The hag1 mutant exhibits reduced plant regeneration with low expression of root stem cell genes\textsuperscript{12}.

Although several molecular factors controlling cellular pluripotency during plant regeneration have been identified, the cellular feature of pluripotent callus cells has remained to be fully elucidated. Furthermore, a molecular signaling network involved in pluripotency acquisition has been investigated mostly in a model dicot plant \textit{Arabidopsis}, and it is elusive whether the regulatory modules are conserved in monocot plants.

Cellular reprogramming is associated with changes in transcript accumulation\textsuperscript{12,18–21}. Quantification of transcriptome can thus provide global insights into pluripotency acquisition during callus formation. In this investigation, we performed next-generation transcriptome sequencing to extend our understanding of the molecular mechanism controlling callus formation and pluripotency acquisition in rice. A comprehensive transcriptomic comparison between pluripotent and non-pluripotent calli, allowing to identify genes with potential roles in pluripotency establishment. In addition, based on the bioinformatics approach, we also proposed potentially relevant biological processes for pluripotency acquisition, including antioxidant metabolism. Overall, our findings suggest that cellular pluripotency is delicately regulated by readjusted hormonal and metabolic balance in plants.

Results

Transcriptome analysis identifies DEGs in pluripotent rice callus. While several molecular players involved in pluripotency acquisition have been identified in \textit{Arabidopsis}, we wanted to know whether the gene networks are conserved in monocot rice plants. In addition, since we were able to distinguish pluripotent and non-pluripotent calli which were derived from mature rice seeds, we further tried to elucidate genetic candidates that regulate pluripotency acquisition in rice. Emerging calli could be sorted into pluripotent and non-pluripotent calli at 28 days after CIM incubation (DAC), based on their phenotypes (Fig. 1). Pluripotent calli with a globular-shaped compact cluster of yellowish cells had cellular competence for tissue regeneration, whereas non-pluripotent calli with semitransparent dark yellow color had a limited ability for de novo shoot regeneration (Fig. 1). The separation was reliable and reproducible, and the pluripotent and non-pluripotent calli divided at 28 DAC showed distinguishable regenerative potential during subsequent incubation on SIM (Supplementary Fig. S1 and Supplementary Table S1).

We collected pluripotent and non-pluripotent calli at 28 DAC and extracted mRNAs to perform Illumina sequencing. Using the Illumina Hiseq-2000, we obtained 76,653,249 and 75,047,011 reads from pluripotent and non-pluripotent calli, respectively. Among these reads, 59,747,844 and 57,835,842 reads accounting for 77.9% and 77.1% of the total reads, respectively, were properly paired and mapped to the reference genome of japonica (MSU v.7.0; \texttt{https://phytozome.jgi.doe.gov/pz/portal.html}) (Table 1). Although transcriptome of the two different types of calli was considerably similar (Fig. 2a), principal component analysis (PCA) showed that the different callus types were also significantly separated along the first principal component, which explains 99.8% of the variability (Fig. 2b).

Based on the adopted cut-off (fold-change > 2 and \(p\) value < 0.05) (Fig. 2c), 60 up-regulated genes (Supplementary Table S2) and 184 down-regulated genes (Supplementary Table S3) in pluripotent callus relative
Table 1. Summary statistics of RNA-seq reads.

|                      | Pluripotent callus Replicate 1 | Pluripotent callus Replicate 2 | Pluripotent callus Total | Non-pluripotent callus Replicate 1 | Non-pluripotent callus Replicate 2 | Non-pluripotent callus Total |
|----------------------|--------------------------------|--------------------------------|--------------------------|-----------------------------------|-----------------------------------|-----------------------------|
| No. of total reads   | 34,957,214                     | 41,696,035                     | 76,653,249               | 38,386,915                        | 36,660,096                        | 75,047,011                  |
| No. of mapped reads  | 34,957,214                     | 41,696,035                     | 76,653,249               | 38,386,915                        | 36,660,096                        | 75,047,011                  |
| No. of properly paired and mapped reads | 27,753,148                  | 31,994,696                     | 59,747,844               | 29,488,630                        | 28,347,212                        | 57,835,842                  |
| Properly paired and mapped reads (%) | 79.4%                         | 76.7%                          | 77.9%                    | 76.8%                             | 77.3%                             | 77.1%                       |

Figure 2. Comparison of global transcriptome between pluripotent and non-pluripotent calli. (a) Scatter plot comparing gene expression pattern in pluripotent and non-pluripotent calli. (b) PCA analysis of transcript abundance for two different types of callus mass. X- and y-axis indicate first and second principal components (PCs), respectively. Percent variance explained by each PC was presented along with axis labels. (c) Volcano plot showing significantly up-/down-regulated genes. X- and y-axis represent log_{2} transformed fold-change and log-transformed p value, respectively. (d) Heatmap showing expression profile of all DEGs. Hierarchical clustering was applied to cluster gene expression pattern in pluripotent and non-pluripotent calli. DEGs were distinguished by color map located in left side of dendrogram. Blue and purple bars represent up- and down-regulated DEGs, respectively.
to non-pluripotent callus were collected, which were validated by RT-qPCR analysis for randomly selected 23 DEGs (Supplementary Fig. S2 and Supplementary Table S4). Since a limited cell population may be related to pluripotency, observation of the moderate number of DEGs was reasonable. We performed a visual inspection of the hierarchical clustering results to identify major subgroups and clusters (Fig. 2d). Collectively, we successfully distinguished pluripotent and non-pluripotent calli to analyze transcriptome possibly related to cellular pluripotency.

KEGG analysis reveals metabolic processes enriched in DEGs. Changes in gene expression underlie differences in biological and metabolic functions between pluripotent and non-pluripotent calli. Gene ontology (GO) analysis of DEGs did not enrich a certain category. We thus alternatively explored the Kyoto Encyclopedia of Genes and Genomes (KEGG) terms that were enriched for up- and down-regulated genes in pluripotent calli. Within the group of up-regulated genes, we identified overrepresented KEGG categories for metabolic pathways (map01100), biosynthesis of secondary metabolites (map01110), fatty acid elongation (map00062), α-linolenic acid metabolism (map00592), and ascorbate metabolism (map00053) (Fig. 3a). This might reflect that pluripotent calli possibly promotes accumulation of secondary metabolites, such as ascorbic acid and antioxidants, as well as primary metabolites including fatty acids and lipids, compared with non-pluripotent calli (Fig. 3a), consistent with the fact that reactive oxygen species (ROS) and lipid molecules are closely associated with pluripotency acquisition in callus22.

In the case of the down-regulated genes, KEGG terms related to metabolic pathways (map01100), diterpenoid biosynthesis (map00904), phenylpropanoid biosynthesis (map00940), carbohydrate metabolic processes (map01200 and map00500), and hormone metabolism and signaling (map04075 and map04010) were enriched (Fig. 3b). The down regulation of these gene sets may be related to global alterations in plant metabolism and hormone signaling. Furthermore, consistent with the fact that callus tissues resemble to root primordium7,8, photosynthesis-related genes (map00195) were also included in this group.

Primary and secondary metabolism and hormone signaling are related to pluripotency acquisition. We next wanted to know molecular factors underlying pluripotency establishment in rice callus. To this end, we first checked whether genes known to regulate cellular pluripotency were included in DEGs of pluripotent rice calli. As a result, the PLT1 gene encoding an auxin-responsive transcription factor responsible for establishing root stem cell niche was up-regulated in pluripotent rice calli (Fig. 4a; Supplementary Table S2)11,15. Furthermore, a HSF transcription factor was also included (Fig. 4a), consistent with a possible role of HSFs in plant regeneration23. Other key transcription factors for various developmental processes were found in DEGs of pluripotent calli (Fig. 4a), implying that they might act as potential upstream regulators of pluripotency acquisition.
We expanded our analysis and examined more genes implicated in diverse aspects of primary and secondary metabolism in plants. Genes related to photosynthesis and primary metabolism for sugar, lipid, and amino acids were particularly enriched in our DEG lists (Fig. 4b–f), and they showed dynamic changes in pluripotent callus relative to non-pluripotent callus. Phenylpropanoid and glucosinolate metabolism genes were also differentially expressed (Fig. 4g,h), and most of which were down-regulated in pluripotent calli. Notably, a majority of antioxidant genes were up-regulated in pluripotent calli (Fig. 4i), as supported by KEGG analysis. In addition, several genes involved in developmental processes and stress responses were also changed in pluripotent calli (Fig. 4j,k), indicating that plant metabolic and developmental processes are massively reprogrammed during pluripotency establishment.

Callus formation and pluripotency acquisition are strongly associated with hormone metabolism and signaling. To further inspect the relevance of hormone signaling in pluripotency acquisition in rice, a list of genes involved in hormone metabolism were investigated in DEGs. Although auxin and cytokinin signaling are known to play crucial functions in plant regeneration,24 the genes related to auxin and cytokinin metabolism were negligibly influenced in pluripotent rice calli (see below). Instead, metabolic genes of gibberellic acid (GA) and abscisic acid (ABA) were particularly changed (Fig. 4l,m), suggesting that pluripotency acquisition in a limited cell population of callus might be conferred by intimate interactions between GA and ABA signaling.

Both pluripotent and non-pluripotent calli were generated by incubation on auxin-rich CIM, and they showed a comparable cell proliferation activity (Fig. 1). Since we compared both proliferating calli, genes involved in cell proliferation and cell cycle progression were negligibly influenced in pluripotent calli compared to non-pluripotent calli. It may also explain the observation that auxin and cytokinin signaling, which primarily affect cell division and proliferation, were not significantly changed. Since a limited cell population have distinguishable cellular features between pluripotent and non-pluripotent calli, the number of DEGs was small, and biological processes typically required for plant regeneration were limitedly enriched in the DEGs. Nonetheless, our analysis is still relevant in that the results enable to gain a new insight into the pluripotency acquisition in rice callus.

Antioxidant activity is increased in pluripotent calli. Our RNA-seq analysis showed that many metabolic processes are regulated during callus formation, among which increased antioxidant activity in pluripotent callus was repeatedly suggested (Figs. 3a, 4i). In support, several previous studies have also shown that pluripotent stem cells contain higher levels of antioxidants in the maintenance of stemness.25–32 We thus put our focus more on the relevance of antioxidant activity in pluripotency acquisition in rice callus.

We asked whether antioxidant levels are indeed increased in pluripotent calli. To this end, we measured levels of polyphenols, which includes several types of antioxidants such as phenolic acid, hydrolysable tannins and flavonoids with robust ROS-scavenging activity, in pluripotent and non-pluripotent callus extracts. Polyphenol measurement revealed that total polyphenol levels of pluripotent calli were 1.2 times higher than those of non-pluripotent calli independently of callus incubation period (Table 2).
Table 2. Polyphenol content in pluripotent and non-pluripotent calli. Total content of polyphenolic compounds was determined according to Prussian Blue method (Hagerman and Butler55). Quercetin was used as a standard. Three biological replicates were averaged, and the data are represented as the mean ± standard deviation. DW, dry weight.

| Callus incubation period on CIM (days) | Pluripotent callus (mg/g DW) | Non-pluripotent callus (mg/g DW) |
|----------------------------------------|-----------------------------|---------------------------------|
| 28                                     | 48.3 ± 2.5                  | 36.7 ± 1.6                      |
| 56                                     | 62.8 ± 3.1                  | 49.8 ± 2.6                      |

Table 3. Antioxidant activity in pluripotent and non-pluripotent calli extracts. Radical scavenging activity was determined using 2,2-diphenyl-1-picrylhydrazyl (DPPH). Butylated hydroxytoluene (BHT), 6-hydroxy-2,5,7,8-tetramethylchroman-2-carboxylic acid (Trolox), and ascorbic acid were used as positive controls. The EC50 value represents the concentration of antioxidant required to decrease the initial DPPH concentration by 50%. Three biological replicates were averaged, and the data are represented as the mean ± standard deviation.

| Callus incubation period on CIM (days) | EC50 (mg/mL)       |
|----------------------------------------|-------------------|
|                                        | Pluripotent callus | Non-pluripotent callus |
| 28                                     | 4.0 ± 0.8         | 3.4 ± 0.4               |
| 56                                     | 1.1 ± 0.07        | 0.09 ± 0.003            |
| BHT                                    | 0.073 ± 0.012     |                      |
| Trolox                                 | 0.006 ± 0.0009    |                      |
| Ascorbic acid                          | 0.123 ± 0.003     |                      |

The radical scavenging activity was further determined by the 2,2-diphenyl-1-picrylhydrazyl (DPPH) test. As a result, antioxidant activity was distinguishable between pluripotent and non-pluripotent calli. Pluripotent callus extract (EC50 = 1.1) had 12 times greater antioxidant activity compared to extracts from non-pluripotent (EC50 = 0.09) at 8 weeks after callus incubation (Table 3). These results indicate that enhanced antioxidant activity is critical for pluripotency maintenance in callus.

**Network analysis identifies novel genes involved in pluripotency establishment in callus.** Our dataset indicates that 244 genes were differentially expressed between pluripotent and non-pluripotent rice calli. The robustness of our transcriptome prompts us to query the expanded gene set for pluripotency acquisition. To this end, we decided to perform gene regulatory network (GRN) analysis to identify more candidate genes, whose expression is linked to core regulators of pluripotency acquisition. Based on a publicly-available GRN (RiceFREN; http://https://ricefrend.dna.affrc.go.jp/), we first extracted the sub-network using QUISCENT CENTER SPECIFIC HOMEOBOX (QHB/OsWOX3) as the seed gene, which is known as a core regulator of pluripotency establishment16,34,35. In the subnetwork, PLT1 and PLT4, the key players of plant regeneration, co-expressed with QHB (Supplementary Fig. S3). We also investigated the DEG genes in this sub-network containing QHB and PLTs and found that LOC_Os05g35070 was included as a potential regulator of pluripotency acquisition (Supplementary Fig. S3).

To obtain additional candidates regulating cellular pluripotency, we reciprocally extracted subnetworks using our DEGs. All DEGs in pluripotent callus were used as queries and a sub-network for each DEG was extracted. Among the input DEGs, only three genes, LOC_Os04g32300, LOC_Os06g08340, and LOC_Os03g42420, formed GRNs coexpressed with QHB and/or PLTs (Fig. 5a). The three subnetworks obtained could be merged (Fig. 5a; Supplementary Table S5), constructing a putative GRN related to pluripotency acquisition. Interestingly, the GRN basically contained core regulators of plant regeneration, such as PLT1, PLT4, QHB, WOX9, ESR1, and CUC3, suggesting that at least the three DEGs are potentially important for pluripotency acquisition in rice callus. From the putative pluripotency GRN, the top 10 hub genes were predicted based on the lowest average path length and higher connectivity (Fig. 5b): CHR38, PLT1, ANT, CUC3, WOX9, HB33, OSH6, OBP2, PLT4, and RFL/LFY. Among them, the potential role of CHR38, ANT, HB33, OSH6, and RFL/LFY have not yet been investigated for plant regeneration to date, and thus they can be considered as novel potential regulators of pluripotency acquisition.

To validate our prediction, we chose a candidate gene and analyzed the potential role in plant regeneration using Arabidopsis model system. The OSH6 gene was of particular interest, because it is expressed at a confined pluripotent region of rice embryonic shoot apical meristem (SAM) will be specified36,37. In addition, OSH6 is also highly expressed in callus tissues (Supplementary Figs. S1 and S5). RT-qPCR analysis enabled to measure sensitive gene expression changes, compared with RNA-seq, and revealed that transcript accumulation of OSH6 was significantly increased in pluripotent calli relative to non-pluripotent calli (Supplementary Fig. S4). Hence, we retrieved the Arabidopsis homolog of rice OSH6 gene, KNOTTED1-LIKE HOMEOBOX GENE 6 (AtKNAT6, At1g23380), based on phylogenetic analysis (Supplementary Fig. S6). To convince the AtKNAT6 gene is the functional homolog of rice OSH6 gene, we compared tissue-specific expression of Class I KNOTTED1-LIKE
**HOMEOBOX (KNOX)** genes (*KNAT2* and *KNAT6*) closely related to the *OSH6* in the sub-clade (Supplementary Fig. S6) and confirmed that only *AtKNAT6*, but not *KNAT2*, is highly expressed in callus tissues similar to *OSH6* (Supplementary Fig. S5), suggesting that the *AtKNAT6* gene is a conserved homolog of rice *OSH6* gene possibly in the control of pluripotency acquisition during callus formation. We next obtained a *knat6* loss-of-function mutant (GK-478F03) and conducted in vitro tissue culture. The *Arabidopsis knat6* loss-of-function mutant (GK-478F03) was used. Leaf explant-derived calli incubated for 7 days on callus-inducing medium (CIM) were transferred to shoot-inducing medium (SIM). Photographs were taken at 21 DAS (days after incubation on SIM) (c). For quantification of shoot regeneration capacity, more than 45 calli at 21 DAS were used to count the number of regenerated shoots per callus. Three biological replicates were averaged (d). Bars indicate the standard error of the mean. Statistical significance is indicated by asterisk marks (ANOVA, **P** < 0.01).

Figure 5. Putative pluripotency gene regulatory network (GRN) in rice. (a) Merged putative pluripotency GRN linking three DEGs and putative pluripotency regulators. Red and blue nodes indicate down- and up-regulated DEGs, respectively. Green nodes represent putative pluripotency regulators. Blue and orange edges indicate two different sub-networks identified by query DEGs, LOC_Os03g42420 and LOC_Os04g32300/LOC_Os06g08340, respectively. (b) Analysis of putative pluripotency GRN. X- and y-axis indicate average shortest path length and degree of each node (gene), respectively. (c, d) De novo shoot regeneration efficiency. The *Arabidopsis knat6* loss-of-function mutant (GK-478F03) was used. Leaf explant-derived calli incubated for 7 days on callus-inducing medium (CIM) were transferred to shoot-inducing medium (SIM). Photographs were taken at 21 DAS (days after incubation on SIM) (c). For quantification of shoot regeneration capacity, more than 45 calli at 21 DAS were used to count the number of regenerated shoots per callus. Three biological replicates were averaged (d). Bars indicate the standard error of the mean. Statistical significance is indicated by asterisk marks (ANOVA, **P** < 0.01).
Discussion

In vitro plant regeneration is driven by the exogenous application of phytohormones, auxin and cytokinin, and involves a two-step process: callus formation and de novo shoot regeneration. In Arabidopsis, the hormone-induced callus is analogous to lateral root primordium. The callus tissue further establishes root stem cell niche, through the expression of root stem cell regulators, including PLTs and WOXs. Since PLT and WOX-induced root stem cell identity is a cellular nature of pluripotency, they facilitate to form competence for tissue regeneration in callus. In support, only after expression of the pluripotency factors in callus, PLTs and WOXs, de novo shoot organogenesis can be initiated on SIM. In this study, we compared the transcriptome of pluripotent and non-pluripotent calli to understand molecular networks involved in pluripotent acquisition in rice. Notably, molecular processes underlying pluripotent callus formation in rice are likely similar to those of dicot plants. PLTs were up-regulated in pluripotent callus relative to non-pluripotent callus in rice. Additionally, based on a putative GRN related to cellular pluripotency in rice (Fig. 5a), WOXs, CUCs and ESR1, which are known to be associated with cellular pluripotency in Arabidopsis, were also involved in pluripotency acquisition in rice callus, indicating the similarity of hormone-induced callus of rice and Arabidopsis. Together, PLTs and WOXs are major determinants of cellular pluripotency across diverse plant species and can be used as key marker genes for pluripotency acquisition.

Several metabolic processes are associated with pluripotency establishment in rice callus. Photosynthetic processes significantly suppressed in pluripotent calli. Since callus resembles root primordium, shoot developmental processes are essentially impaired in callus. Consequently, primary metabolism was also globally influenced. In particular, sugar, amino acid, and nitrogen metabolism were suppressed during callus formation and pluripotency acquisition. We suspect that carbon and energy metabolism are substantially reprogrammed to optimize cellular pluripotency.

Notably, genes related to fatty acid metabolism and ROS homeostasis are enriched in upregulated DEGs of pluripotent calli. Several lipid species, such as prostaglandin E2, linoleic acid, and albumin-associated lipids, are demonstrated as key metabolites for pluripotency establishment and maintenance in mammalian stem cells. In addition, ROS acts either as signaling molecules or detrimental agents inducing cell death. ROS levels are delicately balanced in stem cell niche in plants, which is responsible for pluripotency acquisition. Consistently, our study also showed that increased antioxidant activity in pluripotent calli is important for acquiring competence for de novo organogenesis. Overall, acquisition of cellular pluripotency is an active process that accompanies reprogramming of primary and secondary metabolism.

Since the pluripotent and non-pluripotent calli used in our study were incubated for same period of time on CIM, a limited cell population distinguishes callus samples with different capacity of cellular pluripotency. Consistently, a small number of DEGs were obtained, and several key factors involved in auxin and cytokinin signaling and cell proliferation were unidentified from the DEGs. To understand biological relevance of 244 DEGs of pluripotent callus, coexpression network analysis was employed. Using a public database available for extracting coexpression network in rice, we could propose a putative GRN involved in cellular pluripotency (Fig. 5a). Several DEGs formed a GRN with core regulators of plant regeneration, such as PLTs, WOXs, and CUCs, and interestingly, they could be integrated into a larger coexpression network. Based on the results, several potential regulators can be suggested. The first group includes the DEGs that construct a gene networks with PLTs and WOXs. LOC_Os03g42420, LOC_Os04g32300, LOC_Os05g35070, and LOC_Os06g08340. The second group includes genes, which constitute a hub with high connectivity in the putative pluripotency GRN, but have not studied to date: CHR38, ANT, HB33, OSH6, and RFL. To validate our suggestion, we chose a candidate, OSH6, and confirmed its role in de novo shoot regeneration (Fig. 5c,d). The rice OSH6 gene possibly regulates both cytokinin production and auxin responses, linking cytokinin and auxin signaling in the control of pluripotency acquisition, although the detailed molecular mechanism should be elucidated in the future. Further, the remaining 8 potential candidates would also be associated with pluripotency acquisition and should be studied in the future to gain a comprehensive view of cellular pluripotency in rice.

Altogether, we propose a hypothesis how plant callus acquires pluripotency in rice. Based on the DEGs in pluripotent callus, we proposed molecular processes as well as putative GRNs involved in pluripotency acquisition. Our study provides candidates for evaluating the involvement of genes in pluripotency acquisition. It also provides a resource for comparative transcriptome analysis of plant regeneration in other species.

Methods

**Callus induction, subculture, and regeneration.** Mature seeds of Dongjin (Oryza sativa L., ssp. japonica) were soaked with 70% ethanol for 1 min, surface sterilized in 2.5% sodium hypochlorite supplemented with 2 drops of Tween20 for 20 min, and rinsed with sterile distilled water five times. Callus was induced on callus-inducing medium (CIM) supplemented with 2 mg/L 2,4-D, 0.3 mg/L-1 casein, 0.5 mg/L-1 L-proline, 0.5 mg/L-1 L-Glutamine, 3% sucrose, and 0.3% gelrite and incubated for 4 weeks at 28 °C under continuous dark condition. Pluripotent and non-pluripotent calli were selected at 4 weeks after incubation on CIM and sub-cultured on shoot-inducing medium (SIM) containing 2 mg/L-1 Kinetin, 0.2 mg/L-1 NAA, 3% sucrose, and 0.8% agar for de novo shoot regeneration.

**RNA isolation and cDNA synthesis.** Approximately 0.1 g (fresh weight) of pluripotent and non-pluripotent calli at 28 DAC were harvested respectively, and immediately frozen in liquid nitrogen for RNA isolation. Total RNAs were isolated using the RNeasy plant mini kit (Qiagen). Genomic DNA was eliminated by DNase I (Invitrogen, Carlsbad, CA, USA) treatment, as recommended by the manufacturer. The purity of RNA samples was examined by use of the Agilent 2100 Bioanalyzer (Agilent Technologies). cDNA was synthesized from
300 ng RNA using the PrimerScript II First Strand cDNA Synthesis Kit MIX (Takara Bio, Japan) with oligo (dT) primers in a final volume of 20 μL according to the manufacturer’s instructions.

**Primer design and RT-qPCR conditions.** Gene-specific primers for qRT-PCR analysis were designed using the primer 3.0 online tool (http://bioinformatics.psb.ugent.be/webtools/primer3/) according to the sequences of reference genes and a target gene. Primers were synthesized by the Bioneer Genomics Institute (Daejeon, Korea) with the following parameters: Tm values ranging from 50 to 65 °C, GC percent of 45–55%, primer lengths of 17–25 bp, and product lengths of 50–150 bp. RT-qPCR was carried out with the SYBR Green PCR Master Mix system (Bioneer, Daejeon, Korea) on an Applied Biosystems 7500/7500 Fast Real-time PCR System (ABI, CA, USA). The PCR amplification system and program were performed as described previously. Three biological replicates were analyzed. The RefFinder online software was applied to further estimate the stability of reference genes. Relative gene expression levels were calculated using the 2−ΔΔCt method. All data were recorded as mean ± standard deviation and statistical significance was determined by SPSS (version 9.0 for Windows 98, SPSS Inc.). Primers used in RT-qPCR analysis were listed in Supplementary Table S6.

**Illumina sequencing and data analysis.** For RNA-seq library construction, 1 μg of total RNA extracted from pluripotent calli and non-pluripotent calli incubated on CIM for 28 days. Since callus samples are heterogeneous and have significant variation in gene expression, we collected large amounts of samples (> 300 calli) for single biological replicate. Because of the sampling burden, we performed biological duplicates.

The total RNA was purified by DNase and mRNA purification kit. Then the purified mRNA samples fragmented in 94 °C for 8 min. Fragmented mRNAs were converted into cDNA using random primer. The cDNA samples were ligated by adapters using TruSeq RNA kit (Illumina, CA, USA) and amplified by PCR. Libraries were qualified using an Agilent 2100 Bioanalyzer (Agilent Technologies, CA, USA). To conduct paired end sequencing, libraries were subjected to Illumina HiSeq2000 platform (Illumina, CA, USA) for two biological replicates.

To identify high confident DEGs, we mapped raw RNA-seq reads and quantified transcript abundance considering distribution of mate-pair inner distance. To this end, briefly, we preliminarily mapped the raw read sequences against primary reference transcriptome sequences of rice (MSU v.7.0; https://phytozome.jgi.doe.gov/pz/portal.html) using Bowtie2 aligner. The mean and standard deviation of mate-pair inner distance were estimated for each sequencing library (Supplementary Table S7). Then, the raw reads were re-aligned against the rice reference genome (MSU v.7.0) considering mean and standard deviation of mate inner distance, and splice junction using Tophat2 and Bowtie2. Transcriptome abundance was calculated using Cufflinks and Cuffquant along with application of –frag-bias-correct and –multi-read-correct options for precise quantification. Statistical significance and fold change of each transcriptome was analyzed using Cuffdiff. The differential expression analysis of two samples was performed using criteria, including the absolute value of log2 fold change ≥ 1 and P < 0.05, to ensure the significance of gene expression difference.

**KEGG pathway enrichment analysis.** To conduct KEGG pathway enrichment analysis, the number of genes in a specific pathway was counted for gene group of interest and whole reference genome, respectively. The statistical significance was calculated based on the binomial test comparing observed ratio of genes for specific pathway to expected ratio.

**Antioxidant activity analysis.** All samples were homogenized with a mortar, dissolved in three volumes of methanol, and centrifuged at 12,000g for 10 min at room temperature. The supernatant was stored at ~ 80 °C until use for the analysis. The total content of polyphenolic compounds was measured according to the Prussian blue method with several modifications. As a standard, Quercetin (Sigma Chemical Co., St. Louis, MO) was used.

Radical scavenging activity was measured according to a spectrophotometric method using an ethanol solution of DPPH. As positive controls, Trolox (6-hydroxy-2,5,7,8-tetramethylchroman-2-carboxylic acid), BHT (butylated hydroxytoluene), and ascorbic acid were used. The EC50 values, defined as the amount of antioxidant necessary to decrease the initial DPPH concentration by 50%, were calculated. Three biological replicates were averaged, and statistical significance was determined by SPSS (version 9.0 for Windows 98, SPSS Inc.).

**Analysis of GRN for DEGs.** To gain more insights into the acquisition of pluripotency from the DEGs, we searched GRNs carrying DEGs using RiceFREND database containing co-expression gene network which was constructed based on gene expression in various developmental stage and phytohormone treatments in rice. All DEGs were queried in the database. The results were manually inspected whether the DEGs form a sub-network with key pluripotency regulators, such as QHB, PLT3, and CUC3. The sub-networks link DEG with key molecular component of pluripotency were downloaded from the database and merged and visualized using Cytoscape. Degree and average shortest path length of genes in merged network were analyzed using NetworkKAnalyzer embedded in Cytoscape.

**Phylogenetic and tissue-specific expression pattern analyses.** To identify Arabidopsis orthologous gene of rice OSH6 gene, we constructed phylogenetic tree using all KNOTTED1-LIKE HOMEobox (KNOX) genes in the Arabidopsis and rice genomes. The KNOX protein sequences from primary transcripts were retrieved and aligned using MUSCLE algorithm with default parameters. An unrooted phylogenetic tree was constructed using MEGA X software based on the neighbor-joining algorithm with complete deletion for gaps
and missing data treatment, the Jones-Taylor-Thornton (JTT) model for substitution model, and 1000 bootstrap replications for phylogeny test.

In addition, to compare the tissue specific expression pattern of rice and Arabidopsis Class I KNOX genes, transcript accumulation values of the selected genes were downloaded from Rice Expression Database (http://expression.icrri.org/index) and Arabidopsis EFP Browser (http://bar.utoronto.ca/efp/cgi-bin/efpWeb.cgi) to visualize the data.

De novo shoot regeneration. Wild-type (Arabidopsis Col-0 ecotype) and knat6 mutant (GK-478F03) seedlings were grown on Murashige and Skoog (MS) media at 22–23 °C under long-day condition (16-h light/8-h dark cycle) with fluorescent light (150 μmol photons/m²s). Leaf explants of third and fourth leaves from 2-week-old seedling were incubated on callus-inducing medium (CIM, B5 medium supplemented with 0.5 μg/ml 2,4-dichlorophenoxyacetic acid [2,4-D] and 0.05 μg/ml kinetin) for callus induction. The plates were incubated under 22 °C and continuous dark conditions for 2 weeks. Calli incubated on CIM for 7 days were transferred to shoot-inducing medium (SIM, B5 medium supplemented with 0.9 μmol/l 3-indoleacetic acid, 2.5 μmol/l 2-isopentenyladenine). The plates were incubated at 25 °C and continuous light condition up to 3 weeks.

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Author contributions

S.S., H.K.K., and H.J.L. performed the experiments and analysed the data. S.H.B and H.L.L. carried out plant regeneration analysis. Y.J.J. and P.J.S. conceived and designed the study. P.J.S. wrote the manuscript.

Competing interests

The authors declare no competing interests.
