Repression of Lignin Synthesis in Rice by C4H and 4CL Using RNAi
Xia Gengshou

Abstract—Cinnamate-4-hydroxylase (C4H) and 4-hydroxyxcinamate CoA ligase (4CL) are two enzymes involved in Phenylpropanoid synthesis pathway from lignin. They catalyze hydroxy-L-phenylalanine into cinnamic acid and coenzyme A esters. This paper constructs RNA interference vectors of C4H and 4CL genes, transforms “Zhonghua 11”, a rice variety into single-gene C4H and 4CL and double-gene C4HL transgenic plants using Agrobacterium mediation. By measuring the lignin content in the straw of mature T₀ generation transgenic rice plant, it shows that lignin content in transgenic plants is lower than that of non-transgenic ones. According to semi-quantitative RT-PCR analysis of T₁ generation, C4H and 4CL gene expressions in transgene-positive plants are significantly lower than those of wild type. C4H transgenic rice plant has lower lignin content but does not affect normal field agronomic traits. The study provides reference for comprehensive utilization of rice straw.

Index Terms—Agrobacterium transformation, lignin, rice, RNA interference.

I. INTRODUCTION

As nature’s highest-yielding organic substance only second to cellulose, lignin is a complex phenolic polymer of four alcohol monomers (Coumadin alcohol, coniferyl alcohol, 5-hydroxy coniferyl alcohol, mustard alcohol) polymerized by over 20 chemical bonds. Due to the constraints of its structure and the current utilization capacity, although lignin possesses a large natural reserve and a growth speed of 50 billion tons per year, so far more than 95% of lignin fails to be effectively utilized. Instead, it is discharged into river as “black liquor” or burned after concentration [1]. People have been exploring a transgenic method to repress lignin biosynthesis, in order to resolve its environmental pollution and make lignin be effectively utilized. The study shows that RNA interference (RNAi) exerts significant inhibitive effect on plant gene expression, and it is a commonly used method to artificially control gene expression. This study constructs RNA hairpin structured (hpRNA) RNAi vector by use of C4H and 4CL genes, and it loads the vector with forward and reverse segments of target gene to transform rice plant by Agrobacterium, so that the plant can transcribe and process siRNA, which can degrade endogenous or homologous genes, and eventually reduce or prevent the expression of these genes. In this way, transgenic rice plants with normal phenotype and low lignin content can be identified and they act as the parent for low-lignin crop breeding.

II. MATERIALS AND METHODS

A. Materials
Japonica rice varieties (Oryza sativa L subsp.japonica): “Nipponbare” and “Zhonghua 11” saved in laboratory. DNA and RNA were extracted from leaves in the seedling stage for detection.

B. The Construction of RNAi Vector for Lignin Biosynthetic Regulatory Genes C4H1, 4CL6 and C4HL
Homologous analysis was carried out on the amino acid sequences coded by C4H, 4CL and C4HL genes listed in GenBank by DNAMAN software, and then specific primers were designed to amplify C4H and 4CL gene fragments so as to construct C4H1, 4CL6 and C4HL RNAi vector.

C. Agrobacterium-Mediated Transformation of Rice Plant
Hiei’s Agrobacterium transformation method [6] was adopted.

D. Detection of T₀ Generation Transgenic Rice Plant
Plant height and number of tillers were measured in field,
and then plants were harvested from the roots, taken back to the laboratory, and dried at room temperature after measuring panicle length and striping spike. After that, grain number per plant, thousand seed weight and straw dry weight were measured. Current commonly-used lignin detection methods were reviewed and detection object and lab conditions were considered so as to determine an appropriate method. First, the Klason method was used, but the operation easily produced losses and affected results. Therefore, after referring to the method of Li Jing et al to measure lignin in ginseng, the classical Klason method and ultraviolet spectrophotometry were combined to constitute the detection method of this study.

E. Detection of T₁ Generation Transgenic Rice Plant

When seedlings grew into about two leaves and one heart, tender leaf was taken and the actin gene acted as internal control (Act-F and Act-R as primer) for RT-PCR (reverse transcription PCR). By this method, the expression of single gene 4CL and C4H and double gene C4HL in transgenic plants were detected.

III. RESULTS

A. Analysis of Agronomic Traits of Transgenic Rice and Contrast of Field Growth Conditions

According to the experimental purposes, 10 numbered single plants (8 C4HL plants and 8 control plants) from T₀ generation were randomly selected, and their straw dry weight, spike length, plant height, number of tillers, number of grains, and thousand seed weight were measured and counted. As in Table I, the three transgenic strains all showed obvious reduction in lignin content, among which C4H strain exhibited most significant lignin decrease, up to 9.35%. Moreover, agronomic traits of this strain were the best compared to the other two strains. Therefore, according to the RNA interference of the two genes, transgenic strains with interference on C4H gene resulted in sharpest lignin decline and minimal impact on agronomic traits.

<table>
<thead>
<tr>
<th>Material</th>
<th>CK</th>
<th>4CL</th>
<th>4CL</th>
<th>C4H</th>
</tr>
</thead>
<tbody>
<tr>
<td>Straw dry weight (g)</td>
<td>33.33</td>
<td>19.34</td>
<td>44.37</td>
<td>32.01</td>
</tr>
<tr>
<td>Spike length (cm)</td>
<td>22.75</td>
<td>24.20</td>
<td>24.38</td>
<td>23.00</td>
</tr>
<tr>
<td>Plant height (cm)</td>
<td>111.5</td>
<td>104.2</td>
<td>110.8</td>
<td>120.2</td>
</tr>
<tr>
<td>Number of tillers</td>
<td>9</td>
<td>7</td>
<td>12</td>
<td>11</td>
</tr>
<tr>
<td>Number of grains</td>
<td>1153</td>
<td>509</td>
<td>1023</td>
<td>1066</td>
</tr>
<tr>
<td>Straw lignin content (%)</td>
<td>14.00</td>
<td>10.17</td>
<td>12.20</td>
<td>9.35</td>
</tr>
</tbody>
</table>

CK MEANS CONTROL RICE

B. Comparative Analysis on Field Growth Conditions of T₀ Generation Transgenic Rice Plant

Fig. 1 is a contrast of photographs of C4H, 4CL, and C4HL transgenic rice plants and CK plant, which were randomly taken from rice harvest. It found out that 4CL rice developed worse than other strains, and its plant was short and small. There was no significant difference between C4H and C4HL transgenic varieties from CK, and their plants of C4H and C4HL were even stronger than CK.

According to the above figure, the roots of all the three genetically modified varieties are significantly more developed than the control plant. In addition, during the subsequent harvesting process, other single plants also exhibited a well-developed root system.

C. Semi-Quantitative RT-PCR on T₁ Transgenic Rice Plant

In order to test the effectiveness of RNA interference, RNA were extracted from T₁ tender shoots for semi-quantitative RT–PCR detection of 4CL, C4H and C4HL gene expression. Results showed that there appeared segregation of character in the T₁ generation and it complied with Mendel's laws; the expression of single-gene 4CL and C4H, and double-gene C4H and 4CL in shoots of positive clone plants all showed decrease (Fig. 2, showing part of the results). Although expressions of the two single-genes and one double-gene in transgenic plant strains were different to each other, they were all lower than the untransformed plants. This proves that the gene silencing effect of constructed RNAi vectors is significant, but it varies from strains.

IV. DISCUSSION

The premise of low-lignin transgenic rice application is that it should not harm the normal growth and development of rice, i.e. there should be no bad and slow growth, lodging
or other results which will affect rice yield. According to the detection of the agronomic traits of T₀ generation, although transgenic 4CL single-gene plants can significantly suppress lignin synthesis and reduce lignin content in the straw, it also inhibits growth seriously. All the indicators are significantly lower than those of non-transgenic varieties of the same strain. Worse still, lodging is prevalent, causing serious production decline. All these would be certain defects during application.

Based on straw lignin detection of T₀ generation and semi-quantitative RT-PCR test on tender shoots of T₁ generation, it proves that the expression of these three genes is suppressed and lignin content in straw decreases. This basically agrees with related literatures on the analysis results of lignin and cellulose content in transgenic plants [7]. The reduction of lignin content in transgenic plants shows that the RNAi vectors into genes interfered normal RNA transcription or expression. However, generally speaking, repression and regulation of lignin gene can significantly reduce lignin content in transgenic plants, and meanwhile obtain C₄H gene which will not affect plant’s normal growth. Accordingly, inhibition of C₄H gene expression would be an ideal way to improve timber or agricultural products, for it can reduce lignin content without harming the normal growth of plants. This study constructs transgenic carriers by RNAi to inhibit key enzymes in lignin synthesis in the rice plant straw, so as to relatively decrease lignin content in rice plant straw, increase cellulose content, and eventually provide high-quality raw materials for rice straw bioavailability.

REFERENCES

Xia Gengshou was born on October 15, 1968, in Jinyun Zhejiang, and he is now an associate professor. He graduated from Ningbo Agricultural Technology College in 1996, got Bachelor’s degree in agronomy of China Agricultural University, got Bachelor’s degree in agronomy of China Agricultural University in 2004. He is mainly engaged in the application of biotechnology and resistance physiology research. He has hosted or participated in more than 10 projects, and published more than 30 academic papers. Prof. Xia is a member of Zhejiang Plant Physiology Association, and Chinese Society for Ecological Economics Education Committee Association.