mRNA Differential Display of Tea Leaves under Polyethylene Glycol Stress

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Abstract
Tea (Camellia sinensis L.) is one of the world's three largest beverages which play an important role in agricultural production of China, India, Sri Lanka and other countries. Gene expression of two years old Fuding-dabaicha was analyzed using DDRT-PCR under PEG6000 stress. Through the analysis and screening of experimental conditions and primers combination, a number of different fragments were obtained. By submitting to GeneBank and BLAST comparison, we identified three different fragments while only one has pigment-related gene fragment.

Keywords: Two years old Fuding tea tree, DDRT-PCR, PEG6000 stress

1. Introduction
Drought is one of the main environmental factors which limit the world's crop production. Global arid and semi-arid regions account for about 36% of the total land area and 43% of arable land. In China, about half of the land is arid and semi-arid regions. Drought occupies the first in all abiotic stress to crop yields loss, second only to the loss of biological stress caused by pets (Shanfu Chen & Qingyao Shu,1999,p555-560).

Studies of plants drought resistance started earlier and ranged from the morphological analysis (Hervé Cochard & S.Tete Barigah, Marc Kleinhentz, 2008, p976-982), physiological index (P.Songsri, S.Jogloy & T. Kesmala, 2008, p2245-2253; Carlos German Muñoz-Pereza, Henry Terán, Richard G. Allen, & James L. Wright, 2008, p2111-2120), to the research that focus on the molecular level studies now (Hong hong Hu, Mingqiu Dai & Jialing Yao. 2006, p12987-12992; Yuemin Huang, Benzé Xiao & Lizhong Xiong, 2007, p73-85; Benzé Xiao, Xichen & Chengbin Xiang, 2009, p73-83). With a detailed study on the mechanism of drought gradually down to the molecular level, a lot of drought related genes were found but most of them focus on Arabidopsis, rice, corn and so on.

Tea prefers moisture and largely depends on water. However, some tea-growing areas of precipitation are not enough and this leads to restrictions on tea production. In addition, as global warming, water shortage will become more serious.

The current researches about tea drought resistance mainly in physiological filed and molecular aspects studies are little. In this paper, using mRNA different display technique, gene expression under PEG6000 stress environment was studied in tea.

2. Materials and Methods
2.1 Plant material, stress induction and RNA isolation
Two years old tea (Camellia sinensis cv. Fuding-dabaicha) was selected. It was washed and then cultured in 1/2 Hoagland nutrient solution 3 days based on the same size. After that, it was put in 20% polyethylene glycol (PEG6000) solution to make a drought environment. Stress treatment time was as follows: 1d (P1), 2d (P2), 3d (P3). The control was cultured in the same volume of distilled water (CK) and repeated three times.

Total RNA was isolated by RNA isolation kit following manufacture's protocol (RNAout Column Plant RNAout, TIANDZ, Beijing).Then 4μL total RNA extraction liquid was taken to separated 1.5% (w/v) 10×TAE by 2% agarose gel electrophoresis at 80mv for 30 min.
2.2 Differential display

Three anchored primers (Table 1) were used and cDNA first strand synthesis was performed as following manufacture's protocol (Revert Aid™ First Strand cDNA Synthesis Kit #K1621, Fermentas, Canada). Eighteen random primers (Table 1) were used for differential display PCR and it was performed in 25μL reaction mixture, using four dilutions of the first strand cDNA. Each reaction mixture contained 1μL of each anchored primer and random primers, 2μL cDNA, 12.5μL Master-mix, 14μL ddH₂O. The reaction was performed using a PCR Instrument (Bio-Rad, USA) programmed to 95°C for 5 min to initial denaturation followed by 35 cycles of 45s of denaturation at 95°C, 45s annealing at 45°C, extension at 72°C with a final extension period of 10 min at 72°C. The PCR products (4μL) were separated 1.5% (w/v) 10×TBE by agarose gel electrophoresis at 120mv for 1h. (Primers in table 1 all referenced Ashok K. Jain, 2001,p59-67, and Lina Sha, Fengling FU & Wanzhen Li,2006,p365-370)

2.3 Sequencing and comparison

Selected interested segments, recycling by kit (E.Z.N.A.™ Gel Extraction Kit (50) D2500-1, OMEGA, USA).Connected segments to pMD19-T Simple Vector:T-vector 1μL,segment 4μL,solutionⅠ5μL/link 16°C overnight.Transformed into competent cells of E.coli DH5α (100μL E.coli, 10μL plasmid), after culture chosen positive clones and further with the M13 primers colony PCR identification: 12.5μL Mix, 2.5μL primer M13(±),10μL ddH2O, total 25μL.Finally, chosen positive clones to puncture cultured 16-24h in 37°C and send to Invitrogen biotechnology Co., Ltd sequencing. The results were submitted to the Gene Bank for BLAST alignment.

3. Results

3.1 Changes of plant shape

Compared to control, there was clear droughty trait on the treatment plant. Like Fig 1.

3.2 RNA isolation

Agarose gel electrophoresis results showed that 18s and 28s of total RNA were clear and it meant that total RNA extracted could be used in reverse transcription reaction. Like Fig 2.

3.3 PCR products

Electrophoresis results showed that the PCR product of control was the most abundant, with the longer of processing time, the less abundant of treatments. The different combination of AP and RP had significant differences in bands. Due to space limitation, only a picture was displayed. Fig 3.

A band about 550bp (in white box in figure) was chosen and sent to sequencing. Then BLAST in GeneBank.

3.4 Alignment analysis

By submitting to NCBI and comparing fragments, we found that it had 75% homology with *Camellia sinensis* clone Sajin tea leaf mutant color tag S31.B15 genomic sequence (Accession: DQ443473.1) and 72% with Vitis vinifera contig VV78X260735.4, whole genome shotgun sequence (Accession: AM461193.2). Compared with DNAMAN software, some results were shown as follows:
4. Discussion

DD-PCR was discovered by two scientists at Harvard Medical School P.Liang and A.D Pardee since 1992, it had aroused wide interest for benefits because this technology is simple and quick and can analysis simultaneously two or more samples. It has been successfully applied in a variety of plant resistance research, for example, studies about identification of drought-responsive transcripts in peanut (Ashok K. Jain, Sheikh Mehboob Basha, C. & Corley Holbrook.2001, p59-67) which got several mRNA transcripts that were up-regulated or down-regulated following water stress. Researches about differential gene expression profiles analysis of tea plant induced Tea Looper (Ectropic oblique) attack using DDRT-PCR (Chaoling Wei, Xiangfeng gao & Aihua Ye, 2007, p133-140) found many differential expression fragments and some of them were firstly found in molecular mechanism studies of plant-insect interaction. But this technique has a high false-positive defect, while improving the annealing temperature is a major improvement measure. In this experiment, exploring three annealing temperature of 42°C, 45°C and 50°C, we found 45°C was the best temperature for the better reproducibility results(date not shown). However, some bands maybe disappear due to higher anning temperature which leads to difficulty in primer pair. Therefore, further studies are needed.

In this paper, gene expression has significant differences under drought condition: many down-regulated fragments even disappeared while some of them almost remained unchanged and a part of them up-regulated. By sequencing, we got a nucleic acid band may be close to pigment. Generally, pigment plays an important role in the maintenance of leaf color, flower or other parts of plant, but deeper study found that some pigments had other physiological functions such as secondary metabolite. Anthocyanin is a widely researched pigment and it was found to have many physiological effects. Anthocyanin is a water-solubility plant pigment, existing in 27 genus and 72 species of angiosperms, determining flowers, fruits and seeds’ color (ANNAMARYJU D S,1997, p671-674). There are higher content in plants grape, hawthorn, pine needle, ginggko, peanuts, tea and so on. It belongs to flavonoid of phenolic compounds and has function of anti-oxidant, anti-mutation, strengthening the immune system, etc. A variety of regulatory genes adjust anthocyanin synthesis, in which plant MYB protein was the main factor, which one of the four main stress resistance related transcription factors (Yunrong Zhao, Shilei Wang, 2008, p3095-3097, Zhiru Xu, Chunlei Li, 2008, p597-604). Studis found that numbers of cloning new genes MYB transcription factors are drought-tolerance, high temperature, low temperature effects. For example, AtMYB2 (Hoeren FU, Dolferus R & Wu Y, et al, 1998, p479-490) and AtMYB60 (Cominelli E, Galbiati M & Vavasseur A, 2005, p196-2000) in Arabidopsis were proved to participate in the process of plant drought stress. Over expression osmyb4 gene in Oryza (Vannini C, Locatelli F & Bracale M, 2004, p115-127) could significantly increase GM crops to drought, high salt and UV radiation tolerance. Some studies believe that with the severity of water stress, anthocyanin content was growing and phenylalanine lyase (PAL) activity was the main factor, whiche one of the four main stress resistance related transcription factors (Yunrong Zhao, Arti Rani & Asosii Paul, 2009, 837-846) and successfully cloned this ge ne. So we can speculate that the fragment in this experiment not only associate with tea leaf color change but also have relationship with drought resistance.

Aknowlegement

When completing this paper, Dr.Li Cheng-lei gave much help and I was very grateful.

References


Plant, 2(1):73-83.


Table 1. The Primers’ composition and nature

<table>
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<tr>
<th>Primer type</th>
<th>Primer code</th>
<th>Base composition</th>
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<td>AP1</td>
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Figure 1. The traits’ difference between control and treatment in 3 days

Figure 2. Agaros electrophoresis of the total RNA(From right to left:ck,treatment 1,2,3 days)

Figure 3. The PCR product of the same primer and different drought days