Dynamic Model of Paclitaxel Biosynthesis Suggests That the Key Enzyme Is Taxadiene 5alpha-Hydroxylase in *Taxuschinensis* Cell Suspension Culture

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Abstract—*Taxus* suspension cell cultures are a sustainable source of paclitaxel. In this study, transcript profiles of *T. chinensis* cells were analysed systematically by RT-qPCR to investigate the transcriptional dynamics of *Taxus* cell during cultures. Correlation between the expression levels of genes involved in paclitaxel biosynthesis enzymes and the specific synthesis rate of paclitaxel in suspension culture of *Taxuschinensis* had been studied by a statistic model. The model suggested that, taxadiene 5-alpha-hydroxylase, with the highest correlation coefficient, was the rate-limiting enzyme in paclitaxel biosynthetic pathway in *Taxuschinensis*.

Keywords—paclitaxel biosynthesis genes; relative expression; rate-limiting step

I INTRODUCTION

Paclitaxel, isolated from the bark of several *Taxus* species, is diterpene alkaloid with remarkable anticancer properties, specially effective against breast cancer and non-small cell lung cancer [1]. The most promising and environmentally friendly way for the sustainable production of paclitaxel is provided by plant cell cultures. Though more than 20 molecular cloning of cDNA sequences encoding these enzymes have been studied [2], the genes that control the bottleneck steps of paclitaxel synthesis are still unknown. Taxadiene synthase (TS), catalysing geranylgeranyl pyrophosphate to generate taxane skeleton, was thought as the most promising candidate, only because it is responsible for the branch point biochemical reaction and its cyclization activity is very low [3].

This study systematically detected the expression of more than 20 genes involved in paclitaxel biosynthesis in *Taxuschinensis* suspension cultures. A model had been constructed to study the correlation between gene transcript profiling with paclitaxel synthesis kinetics in order to show light on the committed steps of paclitaxel biosynthetic.

II MATERIAL AND METHODS

A. Cell Culture and Paclitaxel Measurement

The *Taxuschinensis* cell line was provided by Shanghai Jiaotong University. Cell cultures were grown as Miao [4] described. Paclitaxel was measured by HPLC as Yu [5] described.

B. Total RNA Extraction and cDNA Preparation

Cell samples were frozen with liquid nitrogen and stored at -80°C. Total RNA was extracted with Tiangen RNA extraction kit. The first-strand cDNA synthesis was carried with Prime Script® RT-PCR Real Time kit provided by Takara.

Primers were designed using primer premier 5. Tiangen Super Real PreMix Plus (SYBR Green) kit was used for RT-qPCR. The β-actin gene of *Taxus cuspidate* was used as a housekeeping control. The experiments were repeated three times. Data was analyzed using 2-ΔΔCT method.

III RESULTS AND DISCUSSION

A. Transcript Profiling

1) The upstream pathway genes: The upstream pathway is composed of two alternative compartmental pathways: MVA pathway in cytosol and MEP/DXP pathway in plastid.

![Gene Expression Patterns](image-url)
The expression of all upstream genes rapidly and slightly increased during the first 1.5 days due to cellular microenvironment disturbance in the early vaccination. And then the expression decreased slightly and the lowest decrease in transcriptional level was observed on day 8 or day 12. The gene expression went up sharply after day 12 when nutrients have been depleted. (Figure 1)

The expression of genes involved in MVA pathway, such as HMGR and HMGS genes, were always less than the control (Figure 1a), while the expression of five genes in DXP pathway increased sharply after day 8 or 12 and up to 2-5 times more than the control. It was consistent with the theory that terpene precursor was mainly from the DXP pathway. [6].

2) The downstream genes:
Following the formation of the special taxane skeleton, 18 sequential biochemical reactions, being responsible for the modification to the nucleus mainly by hydroxylation and acylation, constitute the downstream pathway.

(a) TS expression patterns, (b) T2H and T5H expression patterns
(c) T13H, T7H and NADPH expression patterns, (d) T10H and T14H expression patterns
(e)DBT/NBT, BAPT and TAT expression patterns (f)BAPT, PAM and TBT expression patterns

FIGURE II. EXPRESSION PATTERNS OF THE DOWNSTREAM GENES.

The highest increase in transcriptional level of TS was observed in late culture (Figure 2a). The expression patterns of cytochrome P450 oxygenases genes activating taxane skeleton fluctuated violently. Similarly with other genes, the expression of hydroxylase genes increased slightly in the first 2 days of culture due to cellular microenvironment disturbance (Figure 2b-d). After the lag phase, the expression of hydroxylase genes decreased rapidly after day 2. In the late culture, their expression went up again as the second peak because paclitaxel biosynthesis rebloomed after the inhibition to the secondary metabolism reduced with the depletion of nutrients.

Comparing with the violent wave of cytochrome P450 oxygenases genes, the expression of acyltransferase genes in late paclitaxel biosynthesis pathway varied little in the cell culture. The expression increased significantly in early culture but no second expression peak was observed in late culture (Figure 2c-f).

B. Dynamic Model Analysis of Paclitaxel Biosynthesis

3) Dynamic model of paclitaxel biosynthesis:
Paclitaxel specific synthesis rate was obtained:

\[ r = \frac{1}{x} \frac{dp}{dt} \]

where x stands for cell volume, p is for paclitaxel content, t is short for time.

As been shown in Figure 3, the first 12 days was accumulation phase for paclitaxel synthesis and then the paclitaxel content began to decrease. The highest paclitaxel biosynthetic activity was observed around day 8 according to paclitaxel specific synthesis rate trends. Paclitaxel biosynthetic activity of cells improved gradually from vaccination to day 8 and then decreased rapidly after day 8 down to 0 in day 12 which meant paclitaxel synthesis rate and degradation rate were equal in that day. And after day 12, paclitaxel degradation rate exceeded paclitaxel synthesis.

4) The committed step model based on the correlation analysis between paclitaxel biosynthesis rate and the transcription expression of enzyme: Paclitaxel complex biosynthetic pathway can be simplified to a series of reaction dynamics system and the slowest rate-limiting steps determine the paclitaxel synthesis rate of Taxus cells. Assuming that expression level determines transcriptional level and decides the catalytic activity of enzymes, there would be positive correlation between the transcription level of the rate-limiting enzymes and paclitaxel special synthesis rate. On the other hand the correlation coefficient of non-limiting enzyme is
little. Therefore, the committed step could be deduced based on the statistics correlation between the paclitaxel synthesis and the transcriptional of paclitaxel synthesis.

Comparing with the upstream, there was obviously positive correlation between transcriptional level of the downstream enzymes and the paclitaxel synthesis rate ($r = 0.33$). The possible reason was that the topological distance of these enzymes for paclitaxel was shorter and less byproducts other than paclitaxel existed so that more products are converted to paclitaxel. It suggested that strongly expressing the downstream genes, especially T5H, would be more beneficial for improving paclitaxel biosynthesis than strongly expressing the upstream genes.

IV CONCLUSION

The correlation coefficient of T5H, was the highest which suggest that T5H was the rate-limiting enzyme of paclitaxel biosynthesis. Cytochrome P450 oxygenases, activating the skeleton, were slow and control the biosynthesis of paclitaxel. Improving the expression of hydrogenases, especially T5H, would give more opportunities to manipulate paclitaxel production.

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