Comparison of isozyme transformation in maize as a result of insertion of the chitinase gene

Comparación de la transformación de isoenzimas en maíz como resultado de la inserción del gen de la quitinasa

Yan PM¹, HF Zhang¹, Q Wang¹, XY Yan¹, Y Sun²

Abstract. Isozymes of peroxidase (POD), catalase (CAT), esterase (EST) and superoxide dismutase (SOD) were analyzed on transgenic maize (with external chitinase gene) and its parent by vertical polyacrylamide gel electrophoresis (PAGE). This study was made using shoots at the fourth leaf stage. Results showed that: POD and EST were detected in 6 bands. POD-2 and POD-3 were present at the bud and seedling stages. POD-1, POD-4, POD-5 and POD-6 were only present at the seedling stage. POD-6 expressed stronger in the transgenic maize with chitinase than in its parent. EST-2 was present only at the bud stage, and its expression in transgenic maize was stronger than that in its parent. EST-5 only existed at the seedling stage. EST-4 did not exist in the parent maize seedlings and EST-1, EST-3 and EST-6 were present in buds of the parent corn. All data showed that the expression of isozymes in transgenic and parent maize had obvious differences.

Keywords: Chitinase; Isozyme; Maize; Antioxidant enzyme system.

Abbreviations: POD: peroxidase; CAT: catalase; EST: esterase; SOD: superoxide dismutase; PAGE: polyacrylamide gel electrophoresis; SDS: sodium lauryl sulphate; TEMED: tetra-methylethylene-diamine.

Resumen. Las isoenzimas de la peroxidasa (POD), catalasa (CAT), estearasa (EST) y superóxido dimutasa (SOD) fueron analizadas en maíz transgénico (con el gen externo de la quitinasa) y su progenitor por electroforesis de gel de poliacrilamida (PAGE). Este estudio fue hecho utilizando tallos en el estado fenológico de cuarta hoja. Los resultados mostraron que: POD y EST se detectaron en 6 bandas. POD-2 y POD-3 estuvieron presentes en los estados fenológicos de yema y plántula. POD-1, POD-4, POD-5 y POD-6 sólo estuvieron presentes en el estado de plántula. POD-6 se expresó con más firmeza en el maíz transgénico con quitinasa que en su progenitor. EST-2 estuvo presente sólo en el estado de yema, y su expresión en el maíz transgénico fue más firme que en su progenitor. EST-5 sólo existió en el estado de plántula. EST-4 no existió en las plántulas progenitoras de maíz, y EST-1, EST-3 y EST-6 estuvieron presentes en los estados de yema y plántula. Se detectaron cuatro bandas para CAT. Las bandas fueron más débiles para CAT-3 y CAT-4. Este último se expresó más firmemente en el maíz transgénico que en su progenitor. Se detectaron tres bandas de SOD: SOD-1 y SOD-2 existieron en los estados de yema y plántula, pero SOD-3 no estuvo presente en las yemas del maíz progenitor. Todos los datos mostraron que la expresión de las isoenzimas tuvieron diferencias obvias en maíz transgénico versus su progenitor.

Palabras clave: Quitinasa; Isonenzimas; Maíz; Sistema enzimático antioxidante.

Abreviaturas: POD: peroxidasa; CAT: catalasa; EST: estearasa; SOD: superóxido dismutasa; PAGE: electrophoresis en gel de poliacrilamida; SDS: dodecil sulfato de sodio; TEMED: tetra-metiletilendiamina.

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INTRODUCTION

Consumers pay more attention to the availability of transgenic plants in China as a result of the rapid development and importation of these kinds of plants in this country (Rui et al., 2005; Rui et al., 2006). With the introduction of exogenous genes, old genomes and metabolic balances could be broken in receptor plants. This could result in physiological and/or biochemical changes (e.g., changes in enzyme isozymes and nutrition).

Chitinase exists extensively in animals, plants and microbes. It may constitute an induced defense mechanism in some plants. Its distribution, characteristics, localization, induction and presence in transgenic plants have been investigated (Lan & Chen, 1998). It has been an effective approach to prevent plant, insect and fungi diseases (Ouyang et al., 2001). Maize is the second cereal crop in China, and most maize diseases come from fungi. This makes meaningful to breed transgenic maize with the chitinase gene.

Isolation of isozymes has become an important technique in molecular biology to conduct research on biological evolution, physiology and biochemistry, taxonomy, and genetic science (Rider & Taylor, 1980; Luo et al., 1999). Today, control of the insertion site for exogenous genes is difficult, and little is known about this subject. Insertion of exogenous genes could change content of plant metabolic substances and enzymes, which are the primary cause of transgenic plant safety (Zhao, 1994). In this study, transgenic maize seeds (with the chitinase gene) were used to (1) study changes of isozymes in the antioxidant enzyme system [POD (peroxidase), CAT (catalase), SOD (Superoxide Dismutase) and EST (esterase)] and (2) provide a basis for transgenic plant ecological safety.

MATERIALS AND METHODS

Plant materials. The transgenic and non-transgenic maize seeds were provided by the Agricultural Biotechnology Center of Shanxi, Taiyuan, China. The transgenic maize was achieved by the pollen-mediated approach on maize (Zea mays L.) inbred with line Hai 921. Plasmid DNA of pGL C -RC-1 was mixed with fresh pollen of maize inbreds in sucrose solution (Wang et al., 2001). Plump and smooth seeds were selected for research.

Culture of maize. Plump and healthy seeds were soaked in water during 24 hours, and placed on white porcelain plates. During the first three days, seedlings were cultured in distilled water, and thereafter cultured in 1/2 Hoogland solution (Table 1). Juveniles were transplanted to light loam soil after root length was 2 cm. Samplings were conducted after sprouting and at the four-leaf-stage.

<table>
<thead>
<tr>
<th>Macronutrients</th>
<th>Volume in 1 L culture solution</th>
</tr>
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<tbody>
<tr>
<td>1 mol/L KH₂PO₄</td>
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</tr>
<tr>
<td>1 mol/L KNO₃</td>
<td>5</td>
</tr>
<tr>
<td>1 mol/L Ca(NO₃)₂</td>
<td>5</td>
</tr>
<tr>
<td>1 mol/L MgSO₄</td>
<td>2</td>
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<table>
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<tr>
<th>Micronutrients</th>
<th></th>
</tr>
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<tbody>
<tr>
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</tr>
<tr>
<td>1.81 g/L MnCl₂•4H₂O</td>
<td>1</td>
</tr>
<tr>
<td>0.22 g/L ZnSO₄•7H₂O</td>
<td>1</td>
</tr>
<tr>
<td>0.08 g/L CuSO₄•5H₂O</td>
<td>1</td>
</tr>
<tr>
<td>0.02 g/L H₂MoO₄</td>
<td>1</td>
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</tbody>
</table>

Experimental methods

Discontinuous polyacrylamide gel electrophoresis

The enzyme extracts underwent SDS–PAGE according to Sanchez’ method (Sanchez et al., 2003) on vertical slab gels. The polyacrylamide gel consisted of a (1) 4.5%T/1.2%Cbis stacking gel in 0.125 M Tris-HCl buffer (pH 6.8) with 0.1% (w/v) SDS, and (2) 12.5%T/3.3%Cbis resolving gel in 0.375 M Tris-HCl buffer (pH 8.8) with 0.1% (w/v) SDS.

Thereafter, we mixed 100 µL enzyme solution with 10 µL 0.025% bromophenol blue. Samples of 30-40µL were added into the sample with Syringes. The voltage was regulated to 150V, stabilizing it to 200V after bromophenol blue come into a separation gel. The electrophoresis was ended when the distance from bromophenol blue to bottom was 1 cm (Yang & Zeng, 1984).

Isozyme of POD

Sample preparation. Fresh maize shoots were washed by diluted water and dried using filter paper. One gram of sample was ground in a precooled mortar with 5 ml of 1mol/L phosphate buffer (pH 7.8). Materials were stored at 4 °C in the refrigerator for 2 h. Thereafter, they were centrifuged for 10 min at 10000 rpm. The extracted material contained the enzyme sample, which was stored at –20 °C.

Staining – Acetic acid – benzidine method (Hu & Wan, 1985). The acetic acid - benzidine solution was prepared with 2 g benzidine, 18 ml Acetic acid and 72 ml double distilled water. The staining solution was made using 5 ml Acetic acid - benzidine solution, 2 ml 3% H₂O₂ and 93 ml double distilled water.

The gel was washed three times using double distilled water after electrophoresis. It was then soaked 5–10 min in the staining solution, and photographs were taken after bands were clear.

Isozyme of CAT

Preparation of sample was similar to that of POD. However, concentration of separation gel was 10–11%. Also, 0.5% soluble starch solution was added to the separation gel.
The gel was soaked in the A solution for 15 min at room temperature, washed with double distilled water, and placed into the B solution. The resulting material was fixed in 10% glycerin.

Isozyme of EST
Sample preparation. The operational procedures were the same as for POD.

Staining solution preparation (Rider & Taylor, 1980): 50mg α- naphthylacetate, 50mg β- naphthylacetate and 100mg fast blue RR salt were dissolved in 2 ml acetone solution (V_{acetone}/V_{water}=1:1). The resulting solution was diluted with 150ml 0.1mol/L phosphate buffer (pH 6.5).

The gel was washed with double distilled water after electrophoresis was finished. This material was then put into the staining solution for 1-2 h at 25 °C, soaked with 7% acetic acid solution, and photographs were then taken.

Comparison of CAT isozyme in transgenic maize versus its parent. Results showed that CAT isozyme had four bands in transgenic and non-transgenic corns: CAT-1, CAT-2, CAT-3 and CAT-4. CAT-1 and CAT-3 expressed less than CAT-2 and CAT-4, and they existed only at the seedling stage; transgenic maize expressed more CAT-1 and CAT-3 than non-transgenic maize. CAT-4 was the primary band of CAT isozyme; there was no difference between transgenic maize and its parent on this regard (Fig. 2).

Comparison of EST isozyme between transgenic maize and its parent. EST isozyme of transgenic maize showed six bands: EST-1, EST-2, EST-3, EST-4, EST-5 and EST-6; non-transgenic maize only expressed five bands: EST-1, EST-2, EST-3, EST-5 and EST-6 (Fig. 3). EST-2 only existed at the bud stage, and its expression quantity in transgenic maize was more than that in non-transgenic maize. Transgenic maize expressed EST-4 at the seedling stage, but non-transgenic maize did not.

**Fig. 1.**, POD differentiation in different tissues of corn genotypes. Gel at the left is at the bud stage. Gel at the right is at the seedling stage.
TR: transgenic maize; CK: Control. Every band represents an isozyme.

TR: maíz transgénico; CK: Control. Cada banda representa una isoenzima.
Comparison of SOD isozyme between transgenic maize and its parent. SOD isozyme of transgenic maize had three bands: SOD-1, SOD-2 and SOD-3 (Fig. 4). Isozymes SOD-1 and SOD-2 existed in transgenic maize and its parent at the bud and seedling stages. However, SOD-3 was not detected at the bud stage in non-transgenic maize.

DISCUSSION

It is well known that phenotypic traits are the result of a series of metabolic activities, which are dependent on many enzymes; their activities and concentrations regulate the metabolic reactions. More than 50% enzymes consist of isozymes which catalyze the same reaction, but isozymes have the highest reaction speeds at different conditions, such as substrate concentration, pH value, temperature and distribution in different organs (Chen et al., 2001).

POD, CAT, EST and SOD are four kinds of antioxidant enzymes which are induced under stress conditions. When an external gene is inserted into another genome, the foreign protein of that external gene is perhaps regarded as an impurity by the inherited antioxidant systems. Therefore, it is important to study the isozyme changes in POD, CAT, EST and SOD after inserting the foreign gene. The isozymes of POD, CAT, EST and SOD in transgenic maize were obviously different from those in its parents. The number (number of bands) and concentration (color depth) of isozymes usually increased. Transgenic maize expressed more POD-6, CAT-1 and CAT-3, but less POD-5, than non-transgenic maize. Also, EST-4 and SOD-3 were expressed only in transgenic maize. This indicates that the insertion and expression of the chitinase gene changed the parent map of isozymes.

Genetic engineering could accelerate biological mutation and evolution, but insertion of exogenous genes can also alter the inherited genome. This could result into either deactivation or disorder of some enzymes. Thus, the number (number of bands) and concentration (color depth) of increasing isozymes may enhance the resistance and adaptation to stress (Chen et al., 2001).
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REFERENCES


