Interactive Effects between Cytokinin and Ethephon on Shoot Formation in Rhizome Cultures of *Cymbidium kanran* Makino

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Rhizomes of *Cymbidium kanran* were cultured in liquid Murashige and Skoog (MS) media supplemented with zeatin, BA and/or ethephon. Zeatin or BA induced shoot formation from rhizome cultures, but inhibited branching of rhizomes. Ethephon added to the culture medium alleviated the reduction of rhizome branching by cytokinins, but inhibited the cytokinin–induced shoot formation. Both zeatin and BA reduced the ethylene evolution from rhizome cultures. Zeatin reduced the ethephon–induced ethylene evolution, whereas, 0.01–1 μM BA significantly enhanced the ethylene evolution by ethephon.

Introduction

Several investigators have previously reported cytokinin–induced shoot formation in rhizome cultures of *Cymbidium kanran* Makino and closely related species.

Among these studies, we found that MS medium containing lower concentrations of nitrogen salts or the medium containing ethylene inhibitors, such as aminoethoxyvinylglycine (AVG) or silverthiosulfate (STS) promoted shoot formation of *C. kanran* rhizomes. Therefore, it was considered that the rhizome cultures producing lower levels of ethylene can develop shoots from the rhizome branches. In contrast, rhizome formation from shoot culture was enhanced in MS medium containing higher levels of auxin or ethylene, but was inhibited by AVG.

This report describes the interactive effects between cytokinin and ethephon on shoot formation and ethylene evolution from rhizome cultures of *C. kanran* Makino.

Materials and Methods

Approximately 5 mm apical segments of *C. kanran* rhizome prepared by the methods described previously were used as explants. Fifteen rhizome segments were cultured for each treatment. Liquid MS media containing 20 g/l sucrose and zeatin or benzylaminopurine (BA) at 0, 0.01, 0.1, 1, 10 and 10 μM alone, or in combination with 10 μM ethephon were used. Each medium was adjusted to pH 5.5 with 100 mM 2-(N-morpholino)ethanesulfonic acid sodium salts (MES-Na). Culture tubes (25×120 mm) with plastic caps were used as culture vessels. Each contained 10 ml of liquid medium. The cultures were incubated for 8 weeks at 25±1°C without forced aeration under fluorescent light of 25 μE/m²/s for a 16 hour photoperiod.

For measuring the amount of ethylene, approximately 0.5 g of rhizome cultures were collected
after 4 weeks of culture, rinsed 3 times with sterilized distilled water, and transferred into 20 ml syringe vials. One ml of gas sample was collected from each vial after incubating the rhizomes for 24 hours at 25±1°C under fluorescent light of 25 μE/m²s for a 16 hour photoperiod, and the ethylene content was determined by gas chromatography (GC14A, Shimazu-Seisakusho, Kyoto) equipped with flame ionization detector. A stainless column, 100 cm long and 3 mm diameter, containing active alumina (70~80 mesh) was used. The column temperature was kept at 100°C. Rates of ethylene evolution were expressed on the fresh weight basis. At least 3 replications of rhizome cultures were measured for each treatment.

**Results**

Effects of zeatin and ethephon on rhizome growth and shoot formation are shown in Table 1. All of the rhizomes showed branching on MS media containing no phytohormone, whereas addition of zeatin in the culture medium inhibited the branching of rhizome. However, inhibiting effect of zeatin on rhizome branching was alleviated by ethephon and 100% of rhizome branching was obtained in the medium containing 0~0.1 μM zeatin with 10 μM ethephon. Rhizomes could produce rhizome branches in the phytohormone-free medium (Fig. 1-a), whereas a single application of zeatin at 0.01~1 μM and the combination of ethephon with 10 μM zeatin was effective for the induction of the rhizome-like shoots which differentiated leaves from a rhizome-like stem.

![Fig. 1 Effects of cytokinins and ethephon added to liquid MS media on growth of rhizome cultures (8 weeks of culture). Bar=5 mm](image)

(a) Rhizome cultured in hormone free medium.
(b) Protocorm-like shoot formation from a rhizome cultured with 10 μM of zeatin.
(c) Protocorm-like shoot formation from a rhizome cultured with 10 μM of BA.
(d) Rhizome cultured in liquid MS medium supplemented with 0.1 μM BA and 10 μM ethephon.
Table 1. Effect of zeatin and ethephon added to liquid MS media on rhizome branching and shoot formation from rhizome cultures.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Mean number/explant</th>
<th>Formation rate(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Rhizome branch</td>
<td>Rhizome-like shoot</td>
</tr>
<tr>
<td>Zeatin ((\mu M))</td>
<td>Ethephon (10 (\mu M))</td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>4.3(^*1)</td>
</tr>
<tr>
<td>0.01</td>
<td>0</td>
<td>1.6(^{bcd})</td>
</tr>
<tr>
<td>0.1</td>
<td>0</td>
<td>0.7(^{cde})</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>0.5(^{de})</td>
</tr>
<tr>
<td>10</td>
<td>0</td>
<td>0.3(^a)</td>
</tr>
<tr>
<td>0</td>
<td>+</td>
<td>3.6(^a)</td>
</tr>
<tr>
<td>0.01</td>
<td>+</td>
<td>1.8(^{bc})</td>
</tr>
<tr>
<td>0.1</td>
<td>+</td>
<td>1.8(^{bc})</td>
</tr>
<tr>
<td>1</td>
<td>+</td>
<td>2.2(^b)</td>
</tr>
<tr>
<td>10</td>
<td>+</td>
<td>0.9(^{cde})</td>
</tr>
</tbody>
</table>

\(^1\) Mean separation within column by Duncan's multiple range test.
\(^2\) (Number of explants forming rhizome branches/total number of explants) \times 100
\(^3\) (Number of explants forming shoots/total number of explants) \times 100

Data were recorded after 8 weeks of culture.
Table 2. Effect of BA and ethephon added to liquid MS media on rhizome branching and shoot formation from rhizome cultures.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>BA (µM)</th>
<th>Ethephon (10 µM)</th>
<th>Mean number/explant</th>
<th>Formation rate(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Rhizome branch</td>
<td>Rhizome branch*2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Rhizome-like shoot</td>
<td>Rhizome-like*3</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Protocorm-like shoot</td>
<td>Protocorm-like*3</td>
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<tr>
<td>0</td>
<td>–</td>
<td></td>
<td>4.3**1</td>
<td>100</td>
</tr>
<tr>
<td>0.01</td>
<td>–</td>
<td></td>
<td>0.1**1</td>
<td>100</td>
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<tr>
<td>0.1</td>
<td>–</td>
<td></td>
<td>0.3ab</td>
<td>100</td>
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<td></td>
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<td></td>
<td>1.0c</td>
<td>75.0</td>
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<td>0.4bc</td>
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<td></td>
<td>0.7bc</td>
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<td>+</td>
<td></td>
<td>0.1a</td>
<td>33.3</td>
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<td></td>
<td>0.1a</td>
<td>25.0</td>
</tr>
</tbody>
</table>

*1 Mean separation within column by Duncan's multiple range test.  
*2 (Number of explants forming rhizome branches/total number of explants) × 100  
*3 (Number of explants forming shoots/total number of explants) × 100  
Data were recorded after 8 weeks of culture.
When 10 μM ethephon was used alone or in combination with 0.01–1 μM zeatin, it inhibited the formation of rhizome-like shoots. However, in the treatment of ethephon with 10 μM zeatin, the highest rate (66.7%) of rhizome-like shoot formation was obtained. The addition of zeatin at concentrations of 0.1, 1, 10 μM, and the combination of ethephon with these high concentrations of zeatin resulted in the formation of protocorm-like shoots, and the maximum number and formation rate of the shoots were recorded for the treatment of 10 μM zeatin. The inductive effect of the protocorm-like shoot (Fig. 1-b) by 0.1 μM zeatin was completely removed by the addition of 10 μM ethephon to the culture medium.

BA had a similar effect to zeatin on rhizome growth and shoot formation (Table 2). The emergence of the rhizome branch was inhibited by application of BA. Simultaneous addition of 10 μM ethephon significantly removed the inhibitive effect of 0.01–1 μM BA on rhizome branch formation. The formation rate of the rhizome branch was 100% at 0–0.01 μM BA alone and at three combinations of BA less than 1 μM with ethephon.

The rhizome-like shoot formation was increased by the application of BA but reduced by ethephon. The addition of BA at concentrations of 0.01–10 μM resulted in the formation of protocorm-like shoots (Fig. 1-c). The combination of ethephon with BA inhibited the differentiation of protocorm-like shoots from rhizome cultures (Fig. 1-d).

Fig. 2 shows the effect of zeatin and ethephon on ethylene production from C. kanran rhizome cultures. Zeatin suppressed the ethylene evolution from rhizome cultures. This inhibitory effect

![Figure 2](image2.png)

**Fig. 2** Effect of zeatin and ethephon on ethylene evolution from rhizome cultures. Bars indicate SE (n=3)

![Figure 3](image3.png)

**Fig. 3** Effect of BA and ethephon on ethylene evolution from rhizome cultures. Bars indicate SE (n=3)
of zeatin on ethylene evolution was partially overcome by the application of ethephon.

The effect of BA on the ethylene production from rhizome cultures was slightly different from that of zeatin (Fig. 3). The ethylene production was inhibited by low concentrations (0.01～0.1 μM) of BA, but no appreciable effect on ethylene evolution was observed at high concentrations (1～10 μM). The application of 10 μM ethephon with 0.01～1 μM BA increased ethylene evolution twice of the control from rhizome cultures. However, the combination of ethephon with 10 μM BA had no effect on the ethylene evolution.

**Discussion**

This study shows that both zeatin and BA in the culture medium reduce the branching of rhizomes and the addition of ethephon is able to alleviate the inhibition of rhizome branching by these cytokinins.

A single addition of zeatin or BA to the culture medium accelerated the shoot formation from rhizome cultures, although both cytokinins had little effect on longitudinal shoot elongation. The cytokinin-induced shoot formation was apparently inhibited by the ethephon treatment, which, however, promoted proliferation of rhizomes regardless of cytokinin in the culture medium. These results indicate that ethephon in the culture medium may play an important role in regulating cytokinin-induced shoot formation.


References


《和文要約》

カンラン(Cymbidium kanran Makino)のライゾームからの
シュート形成に及ぼすサイトカイニンとエチフォンの相互作用

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カンラン(Cymbidium kanran Makino)のライゾームをゼアチン、BA及びエチフォンを添加した液体MS培地で培養した。ゼアチン及びBA単独添加区でシュート形成が認められたが、ライゾームの分枝は抑制された。エチフォンはサイトカイニンによるライゾームの分枝抑制作用を軽減したが、シュート形成を抑制した。ゼアチン並びにBA単独処理区ではともにライゾームからのエチレン放散が抑制された。エチフォン処理によって誘導されるエチレン生成はゼアチン処理区で抑制されたが、0.01〜1 μM BA処理区では著しく促進された。