Production of domoic acid by laboratory culture of the red alga *Chondria armata*

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**Highlights**

> The red alga *C. armata* was cultured

> Excessive manganese inhibited *C. armata* growth

> Domoic acid content of cultured explants was 4-5 fold that of wild specimens

> *C. armata* produced domoic acid
Abstract

To clarify the production mechanisms and biologic functions of domoic acid (DA) by the red alga *Chondria armata*, we established a laboratory culture of *C. armata*. The alga grew better in modified PES medium (mPES) without trace metals or manganese than in unmodified mPES (seawater + nitrate, phosphate, iron, trace metals, vitamins, and 2-[4-(2-hydroxyethyl)-1-piperazinyl]ethanesulfonic acid), suggesting that *C. armata* is especially hypersensitive to the toxicity of excessive manganese. *C. armata* cultured in N•P•Fe medium (seawater + nitrate, phosphate, and iron) grew best (mean growth rate 828.4%) at a relative nutrient concentration of 50%. Liquid chromatography-mass spectrometry analysis of the algal extracts revealed that the DA content of the cultured explants (2273-3308 ppm) was 4 to 5 fold higher than that of wild specimens. The extract of pooled explants (60 g) was purified by activated charcoal treatment and several types of column chromatography to afford ca. 10 mg DA. The \(^1\)H-nuclear magnetic resonance spectrum of the preparation was indistinguishable from the previously reported spectrum of DA, indicating that *C. armata* itself has an ability to produce DA.

Keywords: Domoic acid (DA); amnesic shellfish poisoning (ASP); red alga; *Chondria armata*; laboratory culture; modified PES medium
1. Introduction

In November 1987, mass food poisoning occurred in Eastern Canada following the ingestion of the mussel *Mytilus edulis*. In addition to general signs, such as abdominal pain, diarrhea, and vomiting, the victims exhibited unique nervous system symptoms including memory loss, and the poisoning was thus called amnesic shellfish poisoning (Teitelbaum et al., 1990). Subsequent studies revealed that domoic acid (DA) was the causative substance of the human intoxication, and that the mussels became toxic by ingesting and accumulating DA originally produced by the diatom *Pseudo-nitzschia pungens* forma *multiseries* (Bates et al., 1989; Wright et al., 1989). DA is an excitatory amino acid primarily isolated as an anthelminthic principal from the red alga *Chondria armata* (Daigo, 1959), which inhabitants of an isolated island in the Kagoshima Prefecture used to take to expel their intestinal worms. DA has extremely high affinity for glutamate receptors in the central nervous system (Zaczek and Coyle, 1982; Debonnel et al., 1989), and a heavy overdose of DA causes dysmnesia by destroying the CA3 region of the hippocampus, which coordinates memory in the cerebrum (Strain and Tasker, 1991). After the occurrence of amnesic shellfish poisoning, several studies were conducted to examine the distribution, growth characteristics, and DA productivity of diatoms, the transfer/accumulation of DA via the food chain to other marine organisms, abnormal behavior and death of animals following DA ingestion, and the mechanism of human intoxication (Perl et al., 1990; Kotaki et al., 1999). The biosynthetic pathway of DA has been studied in diatoms (Ramsey et al., 1998; Thomas et al., 2012). *C. armata*, however, is quite difficult to culture, and, although there are some reports of DA in wild algal specimens and coexisting DA isomers (Noguchi and Arakawa, 1996; Zaman et al., 1997), little information is available on the production mechanisms and physiologic functions of DA.
in the alga. To elucidate this point, we established a laboratory culture of *C. armata*.

### 2. Materials and methods

#### 2.1. Culture materials

Thalli of *C. armata* were collected at Hanasezaki, Kagoshima Prefecture, Japan, in August 2007. They were immediately placed in a bottle containing natural seawater and brought back alive to the laboratory of Nagasaki University. The thalli were preserved in autoclaved seawater at 21°C under photosynthetically active radiation of 80 µmol photons·m⁻²·s⁻¹ provided by cool-white fluorescent lamps with a photoperiod of 14:10 light:dark. The branches of the thalli were rinsed several times with the autoclaved seawater, and 50- to 100-µm long apexes were cut with a needle sharpened into a microscopic blade under a stereoscopic microscope (SZ60; Olympus). These apex explants were used for the following experiments.

#### 2.2. Investigation of culture medium

Modified PES medium (mPES; seawater + nitrate, phosphate, iron, trace metals, vitamins, and 2-[4-(2-hydroxyethyl)-1-piperaziny1]-ethanesulfonic acid) (Provasoli, 1968; Kuwano et al., 1998) prepared with filter-sterilized or autoclaved seawater was used as the primary medium in the present study (Table 1). Various media were prepared by removing components from the primary medium to determine the suitable composition of nutrients for the growth of *C. armata* [experiment (Exp) 1]. In the experiment, an apex was placed in each well of a 24-well plate filled with the prepared media (3-4 wells used for each medium), and
incubated under the conditions described above. Growth was evaluated based on the development of a trichome (Fig. 1, upper panel) and the color of the apex explants. DA production by the cultured thalli was examined in Exp 2. Apex explants grown without other algae, fungi, or protozoa in wells of the 24-well plate were carefully selected under an inverted microscope (CKX41; Olympus), and cultured for 30 days in 1-L flat-bottom flasks containing various media under the conditions described above. Three pieces of the branches were placed in each flask and filter-sterilized air was continuously provided through an inlet at the bottom corner of the flask. The growth rate was calculated as the percentage of the fresh weight of the thalli at the end of the culture to that at the beginning of the culture, and the DA content was determined by the following method.

2.3. Determination of DA content

The explants obtained in Exp 2 (3 explants cultured with each medium were combined), and wild algae (3 lots) were extracted with water. Each extract was passed through an HLC-DISK membrane filter (0.45 µm; Kanto Chemical Co., Inc.), and subjected to liquid chromatography-mass spectrometry (LC-MS) (Japan Food Hygiene Association, 2005). Analysis was performed using an Alliance LC-MS system (Waters) equipped with a ZsprayTM MS 2690 detector. A Mightysil RP-18 GP column (250 x 2.0 mm; Kanto Chemical Co., Inc.) was used with 1% acetic acid in 12% aqueous acetonitrile as the mobile phase. The flow rate was set to 0.2 ml/min, and the column temperature at 35°C. DA was ionized by a positive mode of electrospray ionization (ESI) with a desolvation temperature of 350°C, source block temperature at 120°C, and cone voltage of 30 V, and then monitored through a MassLynxTM NT operating system.
2.4. Nuclear magnetic resonance spectral analysis

From the explants (60 g) cultured in 1-L flat-bottom flasks with N·P·Fe medium (seawater + nitrate, phosphate, and iron; Table 1) for 87 to 157 days (Fig. 1, lower panel), putative DA (CaDA) was extracted with water, and purified by activated charcoal treatment, followed by several types of chromatography using a Bio-Gel P-2 column (3×30 cm; Bio-Rad Laboratories) with 0.03 M acetic acid, a P-P-C column (Nihon Seimitsu), and an ODS column (2×25 cm; Waters) with 1% acetic acid in 5% or 4% aqueous acetonitrile as the mobile phase to afford ca. 10 mg of a chromatographically single component. A part of the component was dissolved in D₂O, and placed in a nuclear magnetic resonance (NMR) tube to measure the ¹H-NMR spectrum (Zaman et al., 1997) with a JEOL JNM-AL400 instrument at 400 MHz.

3. Results

In Exp 1, the apex explants developed few or no trichomes and remained small in size in unmodified mPES. They appeared, however, vigorous and had well-developed trichomes when trace metals or manganese were removed from the medium (Fig. 1, upper panel), suggesting that manganese was harmful to C. armata. The removal of HEPES and the vitamin mix did not affect the results, but as further attempts to remove iron, nitrate, and phosphate were ineffective, some or all of these components were suggested to be necessary for C. armata growth. Based on the above results, medium with added nitrate, phosphate, and iron to seawater (N·P·Fe; Table 1) was used to investigate DA production by C. armata culture in Exp 2. Growth varied according to the concentration of the nutrient mix, and the maximum growth rate was 828.4% at half the original concentration (Fig. 2). Explants from each
medium were pooled, extracted with water, and then submitted to LC-MS analysis, in which all of the extracts produced a peak whose retention time was identical to that of the DA standard in a selected ion chromatogram at m/z 312 (Fig. 3). The DA content calculated from the peak area was 2273-3308 ppm, 4- to 5-fold that in wild specimens (mean 587.5 ppm) (Fig. 4).

The putative DA (CaDA) extracted from the pooled explants (60 g) was purified by activated charcoal treatment and several types of column chromatography to afford ca. 10 mg of a single component. The $^1$H-NMR data of the purified CaDA are provided in Table 2. Both chemical shifts and signal configurations were identical with those of previously reported DA (Wright et al., 1990).

4. Discussion

The results of the present study demonstrated that C. armata could grow in the laboratory. The composition of the growth medium was key to promoting the growth. Although Provasoli’s ES medium (Provasoli, 1968), the original mPES medium, is one of the most common media used for culturing algae in the laboratory, the concentration of manganese (3.6 µM) had detrimental effects on C. armata growth. Manganese is required for a number of essential processes in plants, including oxygen evolution in photosynthesis and detoxification of oxygen-free radicals (Fox and Guerinot, 1998), although damage to terrestrial plants by excess manganese has been reported (Mukhopadhyay and Sharma, 1991), and it is also added to other growth media, such as F medium (Guillard and Ryther 1962) and ASP$_{12}$NTA (Provasoli 1963), at concentrations close to or above 3.6 µM. Therefore, the difficulty of culturing C. armata is primarily due to its specific sensitivity to manganese. The
requirement of manganese for the growth of *C. armata* is likely to be low, and the original concentration of manganese in natural seawater is sufficient for the fundamental needs of *C. armata*.

The addition of iron to the growth medium was necessary to grow *C. armata*. Iron is an essential element for plants and required for photosynthesis and respiratory electron transport, nitrate reduction, chlorophyll synthesis, and detoxification of reactive oxygen species (Sunda and Huntsman, 1995). Iron deficiency easily occurs in the marine environment, especially the open ocean, however, because of its insolvency in oxygenated seawater (Martin et al., 1991). The addition of iron to a site in the Pacific Ocean resulted in the increased productivity of phytoplankton, which supports the notion of an iron limitation (Martin et al., 1994). The iron requirement of coastal phytoplankton species was found to be much higher than that of ocean phytoplankton species in culture experiments (Brand et al. 1983). As *C. armata* grows on rocky shores, its iron requirement level is likely to be similar to that of coastal phytoplankton species.

Although the mean growth rate was highest in the 1/2 N·P·Fe medium (Fig. 2), the effect of nitrate and phosphate concentrations on the growth of *C. armata* remains unknown because the growth rate of each explant varied considerably, even under the same conditions. Although the present study led to the development of a basic technique for the laboratory cultivation of *C. armata*, further studies are needed to improve the culture technique.

The DA content of all of the explants in the present study exceeded 2000 ppm, an amount much higher than that of the wild specimens and corresponding values (201-381 ppm) in the previous study (Noguchi and Arakawa, 1996). The DA content of the explants before beginning the culture was not evaluated, but the detection of DA in laboratory culture
explants that more than doubled in weight at concentrations 4 to 5 times higher than that in
the wild specimens, and the fact that the $^1$H-NMR spectra of the DA extracted, purified, and
isolated from the cultured explants were indistinguishable from the previously reported
spectrum of DA strongly suggest that C. armata itself has the ability to produce DA, although
the involvement of symbiotic bacteria cannot be ruled out. There are some reports on the
productivity or biosynthetic pathway of DA in diatoms, but this, to our knowledge, is the first
study to culture C. armata and indicate its ability to produce DA.

Although some possibilities, including the discharge of residual energy in the cells and
osmoregulation are physiologic and ecologic functions of DA in diatoms (Bates, 1998), no
information is available on its physiologic and ecologic functions in C. armata. Further
studies are in progress to elucidate this point, as well as the biosynthetic pathway of DA in C.
armata.

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Conflict of interest

The authors declare that there are no conflicts of interest.
References


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**Figure captions**

Fig. 1. Explant of *C. armata* cultured in a 24-well plate for 7 days (upper), or in 1-L flat-bottom flasks for 157 days (lower). Arrows indicate trichomes.

Fig. 2. Growth rate of explants cultured with media containing different concentrations of nutrients. N·P·Fe (A) was prepared with autoclaved seawater, and the other media with filter-striilized seawater. Relative concentration of nutrients in N·P·Fe (A), N·P·Fe, 1/2 N·P·Fe, and 1/4 N·P·Fe were 100, 100, 50, and 25%, respectively. Data are shown as individual values (open circles) and the mean of each medium (bars).

Fig. 3. Selected ion chromatograms (m/z 312) of a *C. armata* explant extract (upper) and the DA standard (lower).

Fig. 4. DA content of explants cultured with media containing different concentrations of nutrients (see the legend of Fig. 2), and of wild *C. armata* specimens. Data are shown as the value of pooled explants for each medium (grey columns), and mean (black column) and SD (error bar) of three wild lots.
Fig. 2

Growth rate (%) vs. N·P·Fe

- N·P·Fe (A)
- N·P·Fe
- 1/2 N·P·Fe
- 1/4 N·P·Fe

Growth rate (%) range from 0 to 2000.
Fig. 4

The graph shows the DA content (ppm) for different treatments: N·P·Fe (A), 1/2 N·P·Fe, Wild, 1/4 N·P·Fe, and Cultured. The Cultured group shows the lowest DA content compared to the other treatments.
Table 1. Nutrient composition of mPES and N·P·Fe

<table>
<thead>
<tr>
<th>Medium</th>
<th>Nutrient</th>
<th>Component</th>
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<tr>
<td>mPES</td>
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<td></td>
<td>Na₂glycerophosphate</td>
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<tr>
<td></td>
<td>Fe-EDTA·3H₂O</td>
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<td></td>
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<td>H₃BO₃ (BO₃³⁻)</td>
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</tr>
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<td></td>
<td>Fe-EDTA·3H₂O</td>
<td></td>
<td>8.78</td>
</tr>
</tbody>
</table>

*Media were prepared by adding 2 ml of nutritive salt solution to 100 ml of sterilized natural seawater, and the numerical values here indicate the final concentration. EDTA = ethylenediaminetetraacetic acid, HEPES = 2-[4-(2-hydroxyethyl)-1-piperazinyl]-ethanesulfonic acid.
Table 2. $^1$H-NMR data of CaDA in comparison with DA (Wright et al., 1990)

<table>
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<tr>
<th>Position</th>
<th>DA</th>
<th>CaDA</th>
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<tbody>
<tr>
<td>2</td>
<td>3.98 d</td>
<td>3.94 d</td>
</tr>
<tr>
<td>3</td>
<td>3.05 dddd</td>
<td>3.02 m</td>
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<td>4</td>
<td>3.84 ddd</td>
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</tr>
<tr>
<td>5′-Me</td>
<td>1.27 d</td>
<td>1.23 d</td>
</tr>
</tbody>
</table>

Chemical shifts are expressed in ppm (internal standard, CH$_3$COOD = 2.06 ppm). Letters following the chemical shifts indicate the configuration of signals as follows; s = singlet, d = doublet, q = quartet, m = multiplet.