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Removal of hydrogen sulfide using crushed oyster shell from pore water to remediate organically enriched coastal marine sediments

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Abstract

Hydrogen sulfide is highly toxic and fatal to benthic organisms as well as causing depletion of dissolved oxygen and generating blue tide in eutrophic coastal seas. The purposes of this study are to reveal adsorption characteristics of hydrogen sulfide onto crushed oyster shell, and to evaluate removal efficiency of hydrogen sulfide from pore water in organically enriched sediments using container experiment in order to develop a coastal sediment amendment. The crushed oyster shell was mainly composed of CaCO$_3$ with calcite and CaO crystal phase. The batch experiment showed removal kinetics of hydrogen sulfide can be expressed as the first order equation and Langmuir plot fitted well in describing the adsorption behavior with the adsorption maximum at 12 mg-S g$^{-1}$. The container experiments suggested the oyster shell adsorbs hydrogen sulfide in pore water effectively and reduces oxygen consumption in the overlying water. Furthermore, oxidation-reduction potential of the sediment was higher with addition of crushed oyster shell than the control without oyster shell. Thus, it is concluded that crushed oyster shell can be an effective amendment to remediate organically enriched sediments in eutrophic coastal seas.

Key words
adsorption, enclosed water body, hydrogen sulfide, sediment, ORP
1. Introduction

Hydrogen sulfide is a byproduct of sulfate reduction that is highly toxic to benthic organisms in enclosed or semi-enclosed coastal seas. Sulfate reduction is followed by oxygen depletion due to decomposition of organic matter and the resulting hydrogen sulfide sometimes generates blue tide when it is oxidized at the sea surface. Sediments of coastal area that are highly influenced by human activity contain a high much amount of organic matter, which can be a major cause of the formation of hydrogen sulfide.

In Japan, 200 G g of oysters were produced in 2007 and more than half come from Hiroshima prefecture (Ministry of Agriculture, Forestry and Fisheries of Japan, 2007). The intensive oyster culture in Hiroshima Bay causes accumulation of organic matter on the sediment, resulting into high hydrogen sulfide levels. Oyster meat themselves is consumed and a significant amount of oyster shell is discarded. Some oyster shells are reused as soil conditioners and feeds for chicken, while others are left piling up in the field producing unpleasant fishy smell for a long period. Therefore, new applications utilizing these discarded oyster shells are expected to contribute towards promoting recycling consciousness within the society.

While several studies have been carried out for other applications of oyster shell as filtering medium (Park and Polprasert, 2008), catalyst (Nakatani et al., 2009), construction material (Yoon et al., 2003) and soil conditioner (Lee et al., 2008ab), there is few information on the application of crushed oyster shells for the removal of hydrogen sulfide from pore water in order to remediate coastal marine sediments. Since the sediments within the oyster culture grounds are rich in organic matter, using crushed oyster shells to remove the hydrogen sulfide from the sediment will be consistent with the objectives of recycling and environmental restoration.
The purposes of the this study are (1) to determine some adsorption characteristics of hydrogen sulfide onto crushed oyster shells, and (2) to evaluate the removal efficiency of hydrogen sulfide from pore water in organically enriched sediments.

2. Materials and methods

2.1. Characterization of crushed oyster shells

The crushed oyster shells called "Matsu" used in this study were provided by Maruei Corporation, Hiroshima, Japan. They had been left soaking in seawater for 1~2 years to remove any remnants of oyster meat and dried in the field for 3~4 months to remove moisture and salts. The oyster shells were then crushed into coarse pieces and dried in an oven at 400 °C until the water contents falls below 6~7%. Finally, the coarse pieces were further crushed into 7~12 mm pieces. They are sold as feeds for chicken in Japan. It is not difficult to obtain a lot of crushed oyster shells with 7~12 mm pieces to remediate coastal sediment because some mass production plants for the materials are already in operation. The previous study on the material applied for soil conditioner showed that the more crushed or smaller the pieces are, the higher the soil pH increase due to calcium dissolution (Yokota, 1981). Such rapid pH change is not suitable for target benthic ecosystems. Fine grain oyster shells are therefore not suitable. Furthermore, very fine grain oyster shells become viscid and muddy when they are mixed together with coastal sediments. This is why crushed oyster shells with 7~12 mm pieces were selected.

The material was ground into powder using an agate mortar, and the powder X-ray diffraction patterns (XRD) were recorded by an XRD instrument (Rigaku-RINT1100K, Rigaku) using Cu Ka radiation at 20 kV, 20
mA. Organic carbon, carbonate and nitrogen were also determined with a CHN analyzer (MT-5, Yanaco) after removing carbonate by acidifying with HCl (Yamamuro and Kayanne, 1995).

The specific surface area of the crushed oyster shell was determined by the Brunauer-Emmett-Teller method (nitrogen gas adsorption) using a Micromeritics adsorption equipment (Micromeritics Gemini 2370, Shimazu).

Scanning electron microscopic image was taken using a SEM (ABT-150F, TOPCON) at an accelerating voltage of 10 kV.

Wet degradation was carried out to determine the elements comprising the material. A 0.2 g powder sample was degraded at 100 °C on a hot plate with the addition of 5 mL ultra pure nitric acid in a Teflon beaker. After wet degradation, the sample was filtered through a 0.45 μm membrane filter (HA, Millipore). The elements in the filtrate were determined by ICP-AES (Optima3000, Perkin Elmer).

2.2. Adsorption experiment

2.2.1. Removal kinetics

Adsorption kinetics experiment was carried out in a 100 mL BOD bottle containing 50 mL of the hydrogen sulfide solution within the concentration range of 20~800 mg·S·L⁻¹. The hydrogen sulfide solution was prepared as follows: aliquot of Na₂S·9H₂O (Nacalai Tesque) was dissolved in 500 mL of a deaerated 3 % NaCl solution purged with N₂ gas. Thereafter, the pH of the solution was adjusted to 8.2 with 0.1 mol·L⁻¹ HCl. 50 mL of hydrogen sulfide solution was slowly dispensed into the bottle and 1.0 g of the crushed oyster shell was added to the solution. The gas phase in the bottle was replaced with N₂ gas by blowing N₂ gas and capping the bottle tightly. The bottle was agitated moderately at 100 rpm at 25°C in a water bath and time
course of hydrogen sulfide concentration was monitored with a detection tube (200SB, Komyo Rikagaku Kougyo). A blank test in the absence of the crushed oyster shells were also conducted following the same procedure to compensate for hydrogen sulfide loss due to oxidation and volatilization, etc. These same settings were prepared in triplicates.

The adsorption kinetics of hydrogen sulfide onto the crushed oyster shell was calculated on the basis of equation (1).

\[
q_t = \frac{(C_0 - C_t) - (C_{b0} - C_{bt})}{W} \times V
\]  

(1)

where \(q_t\) is the adsorbed hydrogen sulfide (mg-S g\(^{-1}\) of the crushed oyster shell) at time \(t\), \(V\) is the volume of solution (L), \(C_t\) is the crushed oyster shell bottle (mg-S L\(^{-1}\)) at time \(t\), \(C_0\) is the initial concentration. \(C_{bt}\) and \(C_{b0}\) are the concentrations of hydrogen sulfide in the blank (mg-S L\(^{-1}\)) at time \(t\) and the initial concentration, respectively. The \(W\) is the amount of the crushed oyster shell used (g).

2.2.2. Adsorption isotherm

Hydrogen sulfide solution with a concentration range of 5–800 mg-S L\(^{-1}\) was prepared following the same procedure described above. 50 mL of hydrogen sulfide solution was slowly dispensed into a 100 mL BOD bottle and 1.0 g of the crushed oyster shells were added to the solution with the head space in the bottle substituted with N\(_2\) gas and the bottle was closed tightly. The bottle was agitated moderately at 100 rpm at 25°C in a water bath until achieving equilibrium (2~10 d) at which time hydrogen sulfide concentration was measured with a detection tube (200SB, Komyo Rikagaku Kougyo). The blank test in the absence of crushed oyster shells was also conducted.
following the same procedure to compensate for hydrogen sulfide loss during
the oxidation and volatilization. These same settings were prepared in
triplicates.

The amount of hydrogen sulfide adsorbed onto the crushed oyster shell was
calculated using equation (2).

\[ q_e = \frac{(C_0 - C_e) - (C_{be} - C_{be})}{W} V \] (2)

where \( q_e \) is the hydrogen sulfide adsorbed (mg·S g\(^{-1}\) of the crushed oyster
shell) at equilibrium, \( V \) is the volume of solution (L), \( C_e \) is the concentration
of hydrogen sulfide in the crushed oyster shell bottle (mg·S L\(^{-1}\)) at
equilibrium and \( C_0 \) the initial concentration, \( C_{be} \) is the concentration of
hydrogen sulfide in the blank at equilibrium (mg·S L\(^{-1}\)) and \( C_{b0} \) the initial
concentration, while \( W \) is the weight of the crushed oyster shell (g).

2.3. X-ray absorption fine structure (XAFS) of crushed oyster shell

The hydrogen sulfide adsorbed sample was prepared as follows: A 500
mg·S L\(^{-1}\) hydrogen sulfide solution was prepared dissolving 1.9 mg of Na\(_2\)S·
9H\(_2\)O (Nacalai Tesque) in 500 mL of deaerated Milli-Q water purged with N\(_2\)
gas, thereafter pH of the solution was adjusted to 8.2 with 0.1 mol L\(^{-1}\)HCl.
The 50 mL of 500 mg·S L\(^{-1}\) hydrogen sulfide solution was slowly dispensed
into a 100 mL BOD bottle and 1.0 g of the crushed oyster shell was added to
the solution and head space in the bottle was substituted for N\(_2\) gas and
stopper of the bottle was closed tightly. Then, the bottle was agitated
moderately at 100 rpm at 25°C in a water bath for 5 d. After the adsorption of
hydrogen sulfide onto the crushed oyster shell, the sample was air-dried for 2
XAFS analysis was conducted using the BL11 of Hiroshima Synchrotron Research Center (Hayakawa et al., 2008). The synchrotron radiation from a bending magnet was monochromatized with a Si(111) double-crystal monochromator. The sample chamber was filled with He gas, and XAFS spectra were measured both by the X-ray fluorescence yield (XFY) mode and conversion electron yield (CEY) mode simultaneously. Energy of the incident X-rays was calibrated with the S K-edge XAFS spectrum of CuSO$_4$ obtained with the CEY mode, and the main peak corresponding to the SO$_4^{2-}$ was set to be 2.4816 keV according to the previous report (Backnaes et al., 2008). Pieces of relatively flat crushed oyster shell were measured without further treatment to conserve the surface condition of them. Each sample was mounted on a double-stick tape (NW-K15 ;Nichiban) placed in the central hole (15 mm in diameter) of a copper plate. The angle between the incident x-rays and the sample surface was 20 deg, and the X-ray fluorescence was detected from the direction normal to the incident beam in the plane of electron orbit of the storage ring. XAFS spectra were compared between crushed oyster shells with and without hydrogen sulfide treatment.

2.4. Container experiments

2.4.1 Experimental settings

The container experiment was designed for simulating enclosed water bodies and shown in Fig. 1. The crushed oyster shell and sediments (described below) were mixed in round-shaped black polyethylene containers (ϕ: 550 mm, h: 420 mm), and sand-filtered natural seawater was supplied and allowed to overflow at an exchange rate of 0.7 d$^{-1}$, which is the average exchange rate of seawater between the upper and the lower layers of the
water column in the northern part of Hiroshima Bay. These containers were placed in a water bath (1000 L FRP container) to prevent rapid water temperature change during the experimental period. The light intensity was adjusted to 50~120 \( \mu \text{mol m}^{-2} \text{s}^{-1} \) for simulating shallow coastal area conditions using loosely-woven nylon black sheets (cheese cloth) placed over the containers.

The sediment was collected from the Ohzu Inlet located at the head of Hiroshima Bay where municipal wastewater discharging is significant. Debris and pebbles were removed from the sediments before use. In the experimental container, 50 L of sediment and 50 L of the crushed oyster shell were mixed, while 50 L of sediment without the shells was used as a control. These settings were prepared in triplicates.

2.4.2 Overlying seawater analyses

Water temperature, salinity and pH were measured by a multiple electrode (U-10, Horiba). Dissolved oxygen concentration was determined by titration (Winkler method; APHA, 1989).

2.4.3 Sediment and hydrogen sulfide in pore water analysis

Sediment was collected with a container (\( \phi: 77 \text{ mm}, h: 47 \text{ mm} \)), and pH (PRN-40, Fujiwara) and ORP (RM-12P, TOA DK) were measured immediately thereafter. The container was then sealed and kept in a cool, dark place and transported to the laboratory. The sediment was centrifuged at 3,500 rpm for 10 min to separate pore water from the sediment and concentration of hydrogen sulfide in the pore water was determined with a detection tube (200SB, Komyo Rikagaku Kougyo).
3. Results and discussion

3.1 Characterization of the crushed oyster shell

The chemical composition of the crushed oyster shell used in this study is shown in Table 1. The material was mainly composed of calcium carbonate. The concentrations of trace elements in this material were less than those in previous reports (Zn: 0.069~0.084 mg g⁻¹, K: 0.08 mg g⁻¹, Mn: 0.099~0.12 mg g⁻¹, Fe: 0.07~0.39 mg g⁻¹, Yokota, 1981, Yamada et al., 2003, Marine Blue 21, 2003). The specific surface area of the crushed oyster shell used in this study was 0.25 m² g⁻¹, smaller than those of previous reports (1.75~9.59 m² g⁻¹, Yoon et al., 2003; Park and Polprasert, 2008). The powder X-ray diffraction patterns showed the main crystal phase was calcite and CaO referring to a powder diffraction file (ICDD: 47-1743, JCPDS:28-775) (Fig. 3).

Oyster shells have two microstructural types within their shell structure: sheet phase layer and bulky phase layer (Yoon et al., 2003, Marine Blue 21, 2003). The sheet phase layers are composed of leaf shaped, hard, laminated layers. On the other hand, bulky phase layers are brittle and have a lot of pores with lattice-shaped structures.

The crushed oyster shells are classified into two products based on particle size. The coarse fraction which is mainly composed of sheet phase layer is used for feeds for chicken. The fine grain fraction which consisted of bulky phase layer is used for soil conditioners. The trace element contents in the fine grain fraction are higher than those of coarse fractions (Yokota, 1981).

As shown in SEM image (Fig. 2), the crushed oyster shell with coarse grain used in this study is mainly composed of sheet phase layers.

The crushed oyster shell used in this study was coarse grain and its surface mainly consisted of sheet layer structures of calcite crystals and not bulky layers that are highly porous. Thus, the concentrations of trace elements in
the crushed oyster shell and its specific surface area used in this study were less than previous reports. The calcite (CaCO₃) to CaO transformation is verified at 500~600 °C and CaCO₃ decomposition actively occurs at a temperature of around 810~850 °C (Balmain et al., 1999, Wilburn et al., 1991, Lee et al., 2008ab). Thus, the crushed oyster shell used in this study contains CaO initially, not derived from the drying process because drying temperature (400 °C) is lower than the transformation and decomposition temperature of CaCO₃.

3.2 Adsorption experiment

3.2.1 Removal kinetics

Removal kinetics of hydrogen sulfide under different initial concentrations is shown in Fig. 4. The concentrations of hydrogen sulfide decreased dramatically through time and its decreasing rate was expressed as first order equation described by equation (3). The first order rate constants and correlation coefficients between observed values and calculated values by the equation (3) are shown in Table 2.

\[
[C_t] = [C_0]e^{-kt} \quad (3)
\]

Where, \([C_t]\) : concentration of hydrogen sulfide at time \(t\) (mg-S L⁻¹), \([C_0]\) : initial concentration of hydrogen sulfide (mg-S L⁻¹), \(t\) : time (d), and \(k\) : first order rate constant (d⁻¹).

3.2.2 Adsorption isotherm

Adsorption isotherm for hydrogen sulfide is shown in Fig. 5A. Here, three types of adsorption models, namely Langmuir model, Freundlich model and
Henry model, described by equations (4), (5) and (6), respectively, were applied to the hydrogen sulfide adsorption.

\[
\frac{1}{q_e} = \frac{1}{ab} \cdot \frac{1}{C_e} + \frac{1}{a} \quad \text{(4)}
\]

\[
\log q_e = \log K_f + \frac{1}{n} \log C_e \quad \text{(5)}
\]

\[
q_e = HC_e \quad \text{(6)}
\]

Where, \( q_e \): the hydrogen sulfide adsorbed (mg-S g\(^{-1}\)) at equilibrium, \( C_e \): the concentration of hydrogen sulfide (mg-S L\(^{-1}\)) at equilibrium, \( a \): maximum adsorption capacity (mg-S g\(^{-1}\)), \( b \): adsorption coefficient related to bonding energy (L mg\(^{-1}\)), \( H \): Henry constant (L g\(^{-1}\)), \( K_f \), \( n \): Freundlich constant.

Both Freundlich model and Henry model were not suitable for hydrogen sulfide adsorption onto the crushed oyster shell because their correlation coefficients were low: 0.828 for Freundlich model and 0.803 for Henry model. On the other hand, Langmuir model fitted well (Fig. 5B). However, the adsorption equation was separated into two concentration ranges at equilibrium: less than 4.9 mg-S L\(^{-1}\) and more than 4.9 mg-S L\(^{-1}\). The several parameters for hydrogen sulfide adsorption expressed as Langmuir model are shown Fig. 5B. The maximum adsorption capacity was 12 mg-S g\(^{-1}\). In this paper, the maximum adsorption capacity was compared with those of adsorbents for gaseous H\(_2\)S because there is no report on maximum adsorption capacities of hydrogen sulfide in water. The maximum adsorption capacity was 2.3-71, 0.53-12.0, 14-1530 mg-S g\(^{-1}\) for activated carbons, montmorillonites and an activated carbon catalyst (Guo et al., 2007, Xiao et al., 2008, Nguyen et al. 2005, Bashkova et al., 2007). The adsorption capacity of the crushed oyster shell was comparable to momorillonites and some
activated carbons.

3.3. X-ray absorption fine structure (XAFS) of crushed oyster shell

Sulfur K-edge XAFS spectra of crushed oyster shell were compared between those with and without hydrogen sulfide adsorbed (Fig. 6). The spectra were obtained with the XFY mode, and they were normalized with the XFY at the 2.490 keV. After the H$_2$S adsorption a new peak appeared at 2.472 keV while the main peak corresponding to the sulfate (2.4816 keV) decreased. It was concluded that hydrogen sulfide was adsorbed onto the crushed oyster shell, and they were in the form of S$^2-$ (Fleet et al., 2005). The decrease of the sulfate peak may be attributed to the coverage of the adsorbed hydrogen sulfide. Since the outer surface of the crushed oyster shells is covered with hydrogen sulfide after adsorption treatment, the sulfate X-ray fluorescence signal seen after hydrogen sulfide adsorption becomes weaker than that prior to adsorption.

3.4 Container experiments

The water temperature, salinity and pH of overlying water changed in the range of 23.6~29.4°C, 2.85~3.13%, 8.0~8.4, respectively, and did not show significant differences between the experiments and the control throughout the experimental periods. The concentrations of DO in the overlying water of the experimental containers were higher than that of the control after Day 25 with a statistical difference (p<0.05) on Day 35 (Fig. 7). The pH of the sediment did not show a significant difference between the experiments and the control during the experimental periods, changing in the range of 6.7~7.8. The ORP of the sediment was more positive (-348~252 mV) in the experiments than the control (-378~349 mV), showing a
statistical difference (p<0.01) just after the addition of the materials. The concentration of hydrogen sulfide in the pore water was lower than the detection limit (<0.1 mg·S L⁻¹) during the experimental period (Fig. 8). On the other hand, the hydrogen sulfide concentration of the control increased to 7.6 mg·S L⁻¹.

From the results described above, the oyster shells apparently adsorbed hydrogen sulfide from pore water and formed S⁰, S₂O₃²⁻, SO₃²⁻ by oxidation of hydrogen sulfide as mentioned by Hayes and Taylor (2006). As a result, the DO concentration of overlying water and ORP of the sediment in the experiments were higher than those of the control (Fig. 7).

4. Conclusions

The purposes of the present study include investigating adsorption characteristics of hydrogen sulfide onto crushed oyster shell, and to evaluate the removal efficiency of hydrogen sulfide from pore water in organically enriched sediments using container experiments. The results were as follows:

(1) The crushed oyster shell used in this study was mainly composed of calcium carbonate with calcite and CaO crystal phase. The specific surface area was 0.25 m²·g⁻¹.

(2) Removal kinetics of hydrogen sulfide was expressed as a first order equation. Langmuir model was fitted well to the hydrogen sulfide adsorption kinetics with an adsorption maximum at 12 mg·S g⁻¹.

(3) The container experiments showed that the oyster shell adsorbed hydrogen sulfide in pore water effectively and reduced oxygen consumption required for oxidation of hydrogen sulfide. As a result, the DO concentration of overlying water and ORP of the sediment in the experiments were higher.
than those in the control.

Thus, it is concluded that the application of crushed oyster shells is effective in reducing hydrogen sulfide in the pore water of organically enriched sediments, which in turn may prevent DO depletion in the overlying water and decrease in ORP of sediments. The crushed oyster shell is one of the effective amendments to remediate organically enriched sediments in eutrophic coastal seas.

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liming materials. Waste Manage. 28, 2702-2708.


Table 1 Chemical composition of the crushed oyster shell used in this study (mg g⁻¹)

Table 2 Removal rate constant for hydrogen sulfide expressed as first order equation

Fig. 1. Schematic drawing of the container experimental setting

Fig. 2 SEM image of the crushed oyster shell

Fig. 3 X-ray diffraction pattern of the crushed oyster shell

Fig. 4 Removal kinetics of hydrogen sulfide under different initial concentrations with crushed oyster shell

Fig. 5 Adsorption isotherm (A) and Langmuir plots (B) for hydrogen sulfide onto the crushed oyster shell

Fig. 6 Sulfur XAFS spectrum of crushed oyster shell with and without hydrogen sulfide adsorbed by X ray fluorescence yield method

Fig. 7 Change in DO concentrations of overlying water

Fig. 8 Change in hydrogen sulfide concentrations in pore water
Fig. 1. Waste FRP container (1000 L) Overflow seawater circulation for water temperature control Sand filtered seawater supply Sediment (50 L) Round-shaped black polyethylene container (Experimental container) Light intensity: 50-120 μmol m⁻² s⁻¹ Shading sheet
Fig. 2
Fig. 3
Fig. 4

The graph shows the relationship between time (d) and \( \ln \left( \frac{C_t}{C_0} \right) \) for different concentrations of sulfur (mg-S L\(^{-1}\)). The concentrations tested are 20 mg-S L\(^{-1}\), 100 mg-S L\(^{-1}\), 150 mg-S L\(^{-1}\), 200 mg-S L\(^{-1}\), 400 mg-S L\(^{-1}\), and 800 mg-S L\(^{-1}\). The data points and lines indicate a decrease in concentration over time, with higher concentrations resulting in a steeper decrease.
Fig. 5
Fig. 6

Normalized intensity (a. u.)

--- Initial
--- H$_2$S adsorbed

X-ray energy (keV)

S$^{2-}$
Day 0 1 02 03 04 05 06 07 08 09 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43 44 45 46 47 48 49 50

Oyster shell

DO (mg L$^{-1}$)

0.0 3.0 6.0 9.0 12.0

Control

Oyster shell

Fig. 7
Fig. 8

The graph shows the concentration of H$_2$S-S (mg-S L$^{-1}$) over a period of 50 days. Two treatments are compared: Control and Oyster shell. The graph indicates that the Oyster shell treatment has a lower H$_2$S-S concentration compared to the Control throughout the observation period.
Table 1

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