Screening and identification of desulfurization microorganisms

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Abstract: In recent years, biological desulfurization technology has attracted more and more attention from domestic and foreign researchers due to its mild reaction conditions, high efficiency, and environmental protection. In this study, bacteria were collected from oil depots, sewage treatment plants and sewage, and using a screening medium with DBT as the only sulfur source. Three strains that can degrade DBT were isolated, numbered 1, 2, and 3. Through the identification of morphological characteristics, physiological and biochemical identification and molecular identification, it is preliminarily determined that the bacteria No. 1 belongs to the genus Kinetococcus, the bacteria No. 2 is Micrococcus mutans, the bacteria No. 3 is Micrococcus luteus, and the desulfurization capacity of the three bacteria is No. 3. Bacteria > Bacteria No.1 > Bacteria No.2.

1 Introduction

During the processing and utilization of coal and petroleum, harmful substances such as sulfur oxides will be discharged into the surrounding environment in various ways, which results in air pollution, acid rain, greenhouse effect, and destruction of the ozone layer[1]. The pre-combustion biological desulfurization of coal and oil is carried out at room temperature and pressure. It is a potential clean technology with many advantages of safety and environmental protection, low investment, low consumption and high efficiency at this stage[2,3]. However, due to the instability of microbial desulfurization gene expression, there are problems such as easy variation during the growth of desulfurization microorganisms and complicated desulfurization process mechanism. At present, using microbial desulfurization is still in the laboratory research stage. Screening stable and efficient desulfurization microorganisms by effective experimental methods is the focus of current research[4,5].

Coal and petroleum contain a large amount of extremely difficult-to-degrade and heterocyclic form organic sulfur. Dibenzothiophene (DBT) is one of the representative species. Using DBT as a model compound to carry out research on organic sulfur removing have been achieved. In this study, samples were taken from oil depots, sewage treatment plants, and sewage. The screening medium with DBT as the sole sulfur source was used to screen desulfurization microorganisms, and the morphological, physiological and biochemical characteristics were identified. Molecular identification of 16SrDNA was conducted. Finally, three microorganisms with desulfurization ability were screened out for future research.

2 Materials and methods

2.1 Materials and reagents

2.1.1 Medium

(1) Screening medium: K₂HPO₄·3H₂O 4g, NaH₂PO₄·2H₂O 4g, NH₄Cl 2g, MgCl₂·6H₂O 0.2g, CaCl₂·2H₂O 0.001g, FeCl₃·6H₂O 0.001g, glycerol 2g, distilled water 1000mL, DBT concentration changes with the progress of the experiment.

(2) Beef extract peptone medium: beef extract 5.0g, peptone 10.0g, NaCl 5.0g, water 1000mL, pH 7.2~7.4.

(3) Sugar fermentation medium: peptone 2g, glucose 10g, K₂HPO₄ 0.2g, NaCl 5.0g, bromothymol blue 1% aqueous solution 3mL, distilled water 1000mL, pH 7.0~7.2.

(4) Nitrogen-free medium (Axube medium): glucose 10.0g, potassium dihydrogen phosphate 0.2g, magnesium sulfate 0.2g, sodium chloride 0.2g, calcium sulfate 0.1g, calcium carbonate 5.0g, distilled water 1000mL, pH 7.0.

(5) Glucose peptone medium (methyl red experiment): Peptone 5.0g, glucose 5.0g, NaCl 5.0g, distilled water 1000mL, pH 7.0~7.2.

(6) Experimental medium for producing hydrogen sulfide: peptone 10.0g, NaCl 5.0g, beef extract 10.0g, cysteine 0.5g, distilled water 1000mL, pH 7.0~7.4.

(7) Gelatin liquefaction medium: peptone 5.0g, gelatin 100~150g, distilled water 1000mL, pH 7.2~7.4.
2.1.2 Experimental method

(1) Sample collection and processing
Three samples collection locations were selected: the oil-soaked soil in the Huangdao Oil Depot, the sewage from the sewage treatment plant, and the sewage at the school gate. The three kinds of bacteria obtained were named number 1, 2, and 3.

The inoculation amount of enriched culture bacteria was shown in table 1.

Table 1: Inoculation of enriched culture bacteria

| Sample bacteria | Medium amount /ml | Inoculation amount |
|-----------------|-------------------|--------------------|
| number 1        | 210               | 10mL               |
| number 2        | 210               | 10mL               |
| number 3        | 210               | 10mL               |

(2) Desulfurization bacteria screening
The solid medium with DBT concentration of 1mmol/L, 2mmol/L, 5mmol/L, 6mmol/L, 7mmol/L, 8mmol/L and 9mmol/L were prepared respectively. The bacterial culture of No.1, No.2 and No.3 were diluted to 10^-6 and coated to the medium of the corresponding concentration of DBT concentration. The bacteria were cultured at 30℃, 270 r/min on a shaker for 3 days. The culture results of three strains growth in DBT medium with different concentrations were observed and counted.

(3) Identification of desulfurization microorganisms
1) Cell morphology observation
   Cell morphology identification was conducted by microscope observation.

2) Molecular biology identification
   The genetic attributes were determined by 16SrDNA.

3 Results

3.1 The growth of bacteria at different DBT concentrations
When screening bacteria with different concentration of DBT, the growth of different bacteria were shown in table 2 below.

Table 2: The growth of bacteria No.1, No.2 and No.3

| DBT concentration (mmol/L) | Colony Number of No.1 bacteria | Colony Number of No.2 bacteria | Colony Number of No.3 bacteria |
|---------------------------|--------------------------------|--------------------------------|--------------------------------|
| 1                         | Uncountable                    | Uncountable                    | Uncountable                    |
| 2                         | Uncountable                    | Uncountable                    | Uncountable                    |
| 5                         | 100                            | 100                            | Uncountable                    |
| 6                         | 80                             | 60                             | 100                            |
| 7                         | 30                             | 0                              | 50                             |
| 8                         | 4                              | 0                              | 30                             |
| 9                         | 0                              | 0                              | 12                             |

According to the growth of the bacteria at different DBT concentrations, it could be concluded that bacteria No.1 could tolerance a maximum DBT concentration of 8 mmol/L. Bacteria No.2 could grow in a screening medium with a DBT concentration of 6 mmol/L. Bacteria No.3 could grow at the selection medium with the highest DBT concentration of 9 mmol/L. It was preliminarily judged that the desulfurization capacity of the three bacteria was No.3 > No.1 > No.2.

3.2 Morphological characteristics and physiological and biochemical identification of desulphurization bacteria

3.2.1 Observation of colony morphology
The colony morphology of three bacteria were shown in table 3 and fig. 1.

Table 3: The colony morphology of three bacteria

| Bacteria   | Colony morphology                                                                 |
|------------|-----------------------------------------------------------------------------------|
| Bacteria No. 1 | Round, milky white, opaque, protruding surface, moist, easy to pick up, diameter 0.3cm~0.8cm |
| Bacteria No. 2 | Round, milky white, opaque, protruding surface, moist, easy to pick up, diameter 0.3cm~0.8cm |
| Bacteria No. 3 | Round, milky white, opaque, protruding surface, moist, easy to pick up, diameter 0.3cm~0.8cm |
3.2.2 The physiological and biochemical identification

The results of physiological and biochemical identification of three bacteria were shown in Table 4.

Table 4: Physiological and biochemical identification results

| Bacteria | Gram staining | Aerobic test | Methyl red test | Starch hydrolysis test | Gelatin liquefaction test | Sugar fermentation test | Hydrogen sulfide production test |
|----------|---------------|--------------|-----------------|------------------------|--------------------------|-------------------------|---------------------------------|
| No.1     | Purple        | Bacteria grow at the puncture site | Yellow          | Transparent circle, Φ 1.6cm | Gelatin is liquefied     | No acid, no bubbles         | Hydrogen sulfide production    |
| No.2     | Purple        | Bacteria grow at the puncture site | Yellow          | Transparent circle, Φ 1.6cm | Gelatin is liquefied     | Few bubbles               | Hydrogen sulfide production    |
| No.3     | Purple        | Sterile growth at the puncture site, bacteria growth on the surface of the medium | Yellow          | No transparent circle | Gelatin is not liquefied | Few bubbles               | Hydrogen sulfide production    |

3.3 Molecular biology identification

The molecular biology identification were performed by 16SrDNA identification, and the PCR amplification results of random primers were shown below. The gel image of S119 PCR amplification system was as follows.

![Gel image of S119 PCR amplification system](image1)

The RAPD phylogenetic tree of S119 primer was obtained by cluster analysis with NTSYS-pc software as shown in Fig. 3.

![RAPD phylogenetic tree of S119 primer](image2)

4 Discussion

As shown in the above figures, the genetic relationship between No.1 and No.3 bacteria was relatively close.

4.1 Identification results and analysis of No. 1 bacteria

The characteristics of the screened strain No.1 were as follows: the colonies were round, opaque, and protruding on the surface. The early colonies were milky white, and the front and back sides had the same color. Observed by staining, the cells were round, spherical, single or stacked in pairs or multiple irregularly, with an average size of 0.5 - 1.0 μm. Observed by gram staining, the
bacterial cells were purple and positive. During the puncture culture, bacteria grew inside and on the surface of the medium, which were facultative aerobic bacteria. It had the ability to fix nitrogen and could hydrolyze starch but could not decompose cellulose. Fermentation of glucose did not produce acid or gas. Producing catalase. Hydrolyzing gelatin. Producing hydrogen sulfide.

Refer to "Berger's Bacteria Identification Manual", No.1 bacteria belonged to part fourteen (Gram-positive cocci), part a (aerobic and facultative anaerobic cocci), family I (micrococcus), *kinesococcus* genus.

4.2 Identification results and analysis of No.2 bacteria

The characteristics of the No. 2 strain were as follows: the colonies were round, opaque, and the surface was protruding. The early colonies were milky white, and the later colonies became yellow, moist, easy to provoke, and the front and back were the same color. Observed by staining, the cells were round, spherical, single or stacked in pairs or multiple irregularly, with an average size of 0.5-1.0 μm. Observed by gram staining, the bacterial cells were purple and positive. During puncture culture, bacteria grew inside the medium, which was a facultative aerobic bacteria. No nitrogen fixation ability. It could hydrolyze starch but could not decompose cellulose. Fermentation of glucose did not produce acid and gas. Producing catalase, and could hydrolyze gelatin. Producing hydrogen sulfide.

Refer to "Berger's Bacterial Identification Manual", No.2 bacteria belonged to the fourteenth part (Gram-positive coccus), part a (atrophic and facultative anaerobic coccus), family I (micrococcaceae), genus I (*micrococcus*), *micrococcus mutans*.

4.3 Identification results and analysis of bacteria No. 3

The characteristics of the No. 3 strain were as follows: the colonies were round, opaque, and protruding on the surface. The early colonies were milky white, and the later colonies became yellow, moist, easy to provoke, and the front and back sides had the same color. Observed by staining, the cells were round, spherical, single or stacked in pairs or multiple irregularly, with an average size of 0.5 - 1.0 μm. Observed by gram staining, the bacterial cells were purple and positive. During the puncture culture, there was no spreading growth in the punctured place, which was aerobic bacteria. It could not fix nitrogen, hydrolyze starch, decompose cellulose and use glucose as a carbon source. It could produce catalase, protease that could decompose gelatin and hydrogen sulfide. It did not produce acid and gas with glucose as the carbon source.

Refer to "Berger's Bacteria Identification Manual", No. 3 bacteria belonged to the fourteenth part (gram-positive coccus), part a (atrophic and facultative anaerobic coccus), family I (micrococcaceae), genus I (*micrococcus*), *micrococcus luteus*.

5 Conclusion

Using DBT as the only sulfur source, three bacteria named bacteria No.1, No.2 and No.3 were screened out. By bacteria morphology observation, physiological and biochemical characteristics analyze and molecular identification experiments, it was preliminarily determined that No.1 bacteria belonged to the genus *Kinetococcus*, No.2 was *Micrococcus mutans*, and No.3 was *Micrococcus luteus*. The desulfurization capacity of the three strains was preliminarily determined to be No.3 bacteria> No. 1 bacteria > No.2 bacteria.

References

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