ISOLATION AND PRELIMINARY CHARACTERIZATION OF A O-NITROBENZALDEHYDE-DEGRADING Alcaligenes SP. ND1

Yu Fang-Bo1*; Guan Li-Bo1; Zhou Shan1*

1Department of Environmental Sciences, School of Environmental Technology, Zhejiang Forestry University, Zhejiang Province, China, 311300

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ABSTRACT

This paper reports the isolation and characterization of a new o-nitrobenzaldehyde (ONBA)-degrading bacterium, Alcaligenes sp. ND1. ND1 degraded almost all ONBA (100 mg L⁻¹) in M9 medium within 36 hours. The key enzyme(s) involved in the initial biodegradation was a constitutively intracellular enzyme(s). This bacterium has great potential utility for bioremediation.

Key words: Alcaligenes sp.; Biodegradation; o-nitrobenzaldehyde.

A large variety of chemicals are commercially produced and newly synthesized each year. Some compounds and byproducts are discharged into the environment during use and manufacture of these chemicals. Their widely use has caused great damages to the environment and to living organisms (2). In China, as an important intermediate for the synthesis of pharmaceuticals, such as nifedipine (Adalat, Procardia), dyes, agrochemicals and other organic compounds, industrial demand of ONBA is great (19). Meanwhile, severely pollution hazard is created because of its toxic and recalcitrant nature. It is toxic either by ingestion, by contact, or by inhalation at low concentrations (https://fscimage.fishersci.com/msds/91095.htm). It is, therefore, necessary to treat ONBA-containing wastewaters before their discharge into the environment, although no specific acceptable limit of ONBA has been decided in treated wastewater.

Microbial metabolism is the main mechanism responsible for degradation of ONBA in the environment, and there is a need to develop remediation with efficient ONBA-degrading microorganisms to eliminate or minimize the contamination. However, before this study only one Pseudomonas sp. strain, ONBA-17, which could utilize ONBA as sole carbon and nitrogen source, has been reported (19). Although, this strain could completely degrade 100 mg L⁻¹ ONBA in 48 h and tolerate up to a higher concentrations (400 mg L⁻¹), and showed potential to be a good candidate for bioremediation, isolation of novel strains is still necessary due to the consideration of new bacterial resources for biodegrading ONBA.

In this work, we have isolated an ONBA-degrading bacterium from activated sludge. The isolate, strain ND1, was Alcaligenes sp., which has not been reported previously to own the ability to degrade ONBA. The characteristics of ONBA degradation by the isolate and the distribution of ONBA-degrading enzymes from strain ND1 were preliminarily studied.

Activated sludge samples were sampled from a municipal wastewater treatment plant in the city of Hangzhou, Zhejiang Province, China. The enrichment mixture was incubated at 25°C and shaken at 150 rpm on a rotary shaker (19). Six subcultures were performed before isolation was done. The final enrichment culture was plated on MSM agar plate containing 15 g agar L⁻¹. After incubation for 3 days at 28°C, single colonies were re-streaked on the fresh MSM agar plates. The isolates were used to inoculate MSM to test their degradation activity. One isolate, designated as ND1, from among those with the best ONBA metabolism was selected for further identification and characterization. The concentration of ONBA was determined with a gas chromatography method as described by Yu et al. (19).

The identification was performed according to Bergey’s Manual of Determinative Bacteriology (9). The 16S rRNA genes of bacteria were amplified with the universal primers 8F and
reverse primer 1492R (18). Multiple alignments of the sequences were performed using the CLUSTAL W program version 1.8 (17). Phylogenetic and molecular evolutionary analyses were conducted using MEGA version 3.0 (10). Tree was constructed by using the neighbor-joining method and dataset was bootstrapped 1000 times (13).

The optimum temperature and pH for ONBA removal was determined according to the method described by Dai et al. (6), but with ONBA as degradation substrate. Determination of MICs for 13 metal ions was performed using the methods described by Yu et al. (19).

Cells from the early stationary phase of growth were centrifuged, washed, and suspended in M9 minimal salts medium (16) without ammonium chloride. After the value of OD_{60\text{nm}} had been adjusted to 1.0, 5% inoculum was inoculated into M9 medium with 100 mg L^{-1} ONBA as sole carbon or nitrogen source. Fructose, 100 mg L^{-1}, was supplemented when ONBA was used as sole nitrogen source. Samples were collected from the cultures at 6-hour intervals. The degradation effect was evaluated according to the GC method.

The localization of degrading enzymes was conducted by the method of osmotic shock (8). The solutions were each transferred (10%, v/v) into phosphate buffer solution (pH 7.0) containing 100 mg ONBA L^{-1} and incubated at 30°C with shaking at 120 rpm. The reaction mixtures at zero time and 10 h were examined against a blank control containing the same reaction mixture, except that ONBA was omitted.

To determine whether the enzyme system responsible for ONBA biodegradation was inducible or constitutive, two batches of cells were grown in M9 medium containing 1% fructose in the presence and absence of 100 mg L^{-1} of ONBA. Total protein was estimated by the method of Lowry et al. (11), using bovine serum albumin as the standard.

The reaction mixture (1.0 mL) contained 50 mmol of phosphate buffer (pH 7.4), 13.24 mmol of ONBA, and cell crude extract. Reactions were performed at 30°C for 12.5 min without shaking, and the residual ONBA was quantified by the GC method. Enzyme activities were expressed as units (micromoles of ONBA decreased per minute) per milligram of protein.

Bacterial colonies could be observed on the isolation and purification medium using ONBA as the sole carbon and energy sources after 3-5 days incubation. All the colonies were identical morphologically, circular, low-convex, about 2 mm in diameter, smooth, shining and entire, from which strain ND1 was isolated. It was a Gram-negative, short-rod or coccobacillary bacterium, 1.2-2.5 μm long and 0.7-1.0 μm in diameter, with peritrichous flagella. The optimum temperature and pH for strain growth were 30°C and 7.6, respectively.

A 1.5 kb 16S rDNA fragment was amplified from the total DNA of the isolate and sequenced. Its GenBank accession number is EU072720. It showed high sequence similarity to Alcaligenes species (especially to A. faecalís) (Fig. 1). According to its morphology, cultural appearance, and physiologic and biochemical characteristics mentioned above, together with the phylogenetic analysis, the strain ND1 was preliminary identified as Alcaligenes sp. Until now, no described ONBA-degrading bacterium has been related to Alcaligenes.

The ability to utilize ONBA as the sole source of carbon seems to be shared by relatively few microorganisms. ONBA is so recalcitrant that large numbers of bacteria cannot use it as a substrate for growth. To date, only Yu et al. (19) reported a pure culture of Psudomonas species, strain ONBA-17, with the ONBA-degrading capability. Alcaligenes species are widespread in nature and can be obtained from various sources, such as water, soil, and living organisms. Some strains of this genus are known to be involved in the biodegradation of quite a few pollutants, such as benzoate and chlorobenzoates, polychlorinated biphenyls, pyridine, phenol, and so on (1,3,4,7,14). This was the first report that Alcaligenes has ONBA-degrading ability. Most Alcaligenes strains have been isolated from wastewaters containing ONBA as good candidate for biotreatment of industrial wastewaters containing ONBA.

As presented in Table 1, strain ND1 is strongly multiresistant. Although its resistance to Cu^{2+}, Cu^{+}, Pb^{2+}, Mn^{2+}, and Ni^{2+} is not as good as ONBA-17, it owns higher MICs of AgNO_{3}, FeSO_{4}, and FeCl_{3}. Together with previously data, the isolate could be considered as a good candidate for biotreatment of industrial wastewaters containing ONBA.

With GC analysis, we found that no objective substance could be detected in the reaction mixture containing ONBA and the intracellular fraction solution taken at 10 h (data not shown). Besides, there was no downtrend of ONBA content in the mixture containing the extracellular and membrane fraction solution (data not shown). These results show that the enzyme(s) involved in the initial degradation of ONBA in ND1 was intracellular enzyme(s).
Cells induced by ONBA and noninduced cells showed ONBA biodegradation activities of 18.13 ± 3.09 and 17.69 ± 4.97 U (mg protein)-1, respectively, indicating that there was no significant discrepancy in ONBA degradation between induced and noninduced cells. This finding also indicated that there may be a constitutive enzyme(s) in the cells responsible for ONBA biodegradation.

In the previously researches, there are two pathways for o-nitrobenzoate (ONB) degradation (at different substrate concentrations levels) in a Gram-positive Arthrobacter protophormiae strain, RKJ100 (5,12). ONB is reductively degraded to o-hydroxylaminobenzate (2-HABA) which is a substrate for two different enzymes, a reductase and a mutase, that convert HABA to anthranilate (AA) and 3-hydroxyanthranilate (HAA), respectively; these further serve as substrates for ring cleavage enzymes (12). In this work we confirmed the existence of o-hydroxylaminobenzaldehyde (OHABA) (data not shown), and proposed that ONBA degradation pathway of strain ND1 is likely to same with that of strain RKJ100. However, it still needs further research to gain solid proofs.

In summary, we describe a novel ONBA-degrading bacterium isolated from activated sludge in a municipal wastewater treatment plant using a culture enrichment technique. The Alcaligenes species strain ND1 enriches our knowledge on Alcaligenes species that can grow on aromatic compounds. Strain ND1 has a high tolerance to ONBA toxicity and is multiresistant to heavy metals. It owns good potential for biotreatment of ONBA-containing industrial wastewaters or in situ bioremediation of ONBA-contaminated soils.

**Table 1.** Minimal inhibitory concentrations (μM) of strain *Alcaligenes* sp. ND1 and *Pseudomonas* sp. ONBA-17 to 13 heavy metals.

| Metal ion | ND1   | ONBA-17 | Metal ion | ND1   | ONBA-17 |
|-----------|-------|---------|-----------|-------|---------|
| Li⁺       | 3,600 | 3,200   | Zn²⁺      | 6,000 | 6,000   |
| Ag⁺       | 3,200 | 2,400   | Mn²⁺      | 4,800 | 6,400   |
| Cu¹⁺      | 6,000 | 8,000   | Ni²⁺      | 1,600 | 3,200   |
| Cu²⁺      | 6,000 | 8,000   | Fe²⁺      | 7,200 | 6,000   |
| Hg²⁺      | 1,800 | 1,200   | Fe³⁺      | 7,200 | 6,000   |
| Bi³⁺      | 1,280 | 1,280   | Co⁴⁺      | 1,600 | 1,600   |
| Pb²⁺      | 4,800 | 6,000   |           |       |         |

**RESUMO**

Isolamento e caracterização preliminar de *Alcaligenes* sp ND1 degradador de o-nitrobenzaldeído

Esse trabalho relata o isolamento e a caracterização de uma nova bactéria degradadora de o-nitrobenzaldeído (ONBA), *Alcaligenes* sp ND1. A bactéria ND1 decompos todo o ONBA (100mg.L⁻¹) do meio M9 em 36 horas. A enzima-chave envolvida na biodegração inicial foi uma enzima constitutiva intracelular. Esta bactéria apresenta um potencial de aplicação para biorremediação.

**Palavras-chave:** *Alcaligenes* sp, biodegradação, o-nitrobenzaldeído.
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