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# A novel and complete gene cluster involved in the degradation of aniline by *Delftia* sp. AN3

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#### Abstract

A recombinant strain, Escherichia coli JM109-AN1, was obtained by constructing of a genomic library of the total DNA of Delftia sp. AN3 in E. coli JM109 and screening for catechol 2,3-dioxygenase activity. This recombinant strain could grow on aniline as sole carbon, nitrogen and energy source. Enzymatic assays revealed that the exogenous genes including aniline dioxygenase (AD) and catechol 2,3-dioxygenase (C23O) genes could well express in the recombinant strain with the activities of AD and C23O up to 0.31 U/mg wet cell and 1.92 U/mg crude proteins, respectively. The AD or C23O of strain AN3 could only catalyze aniline or catechol but not any other substituted substrates. This recombinant strain contained a recombinant plasmid, pKC505-AN1, in which a 29.7-kb DNA fragment from *Delftia* sp. AN3 was inserted. Sequencing and open reading frame (orfs) analysis of this 29.7 kb fragment revealed that it contained at least 27 orfs, among them a gene cluster (consisting of at least 16 genes, named danQTA1A2BRDCEFG1HIJKG2) was responsible for the complete metabolism of aniline to TCA-cycle intermediates. This gene cluster could be divided into two main parts, the upper sequences consisted of 7 genes (danQTA1A2BRD) were predicted to encode a multi-component aniline dioxygenase and a LysR-type regulator, and the central genes (danCEFG1HIJKG2) were expected to encode meta-cleavage pathway enzymes for catechol degradation to TCA-cycle intermediates. Unlike clusters tad from Delftia tsuruhatensis AD9 and tdn from Pseudomonas putida UCC22, in this gene cluster, all the genes were in the same transcriptional direction. There was only one set of C23O gene (danC) and ferredoxin-like protein gene (danD). The presence of only one set of these two genes and specificity of AD and C23O might be the reason for strain AN3 could only degrade aniline. The products of danQTA1A2BRDC showed 99%-100% identity to those from Delftia acidovorans 7N, and 50%-85% identity to those of tad cluster from D. tsuruhatensis AD9 in amino acid residues. Besides this dan cluster, the 29.7 kb fragment also contained genes encoding the trans-membrane transporter and transposases which might be needed for transposition of the gene cluster. Pulsed-field gel electrophoresis (PFGE) and plasmid curing experiments suggested that the dan cluster might be encoded on the chromosome of strain AN3. The GenBank accession number for the dan cluster of Delftia sp. AN3 is DQ661649.

Key words: aniline; biodegradation; Delftia sp. AN3; genomic library; aniline degradative gene cluster

# Introduction

Anilines constitute a group of xenobiotics (pollutants which resemble naturally occurring compounds), that are in use for a long time in the production of dyes, pesticides, plastics, and pharmaceuticals (Meyer, 1981; Kearney and Kaufmann, 1977). This group of refractory compounds is considered as an important environmental hazard and is subject to legislative control by the European Economic Community (EEC) directive and in the Priority Pollutant List of U.S. Environmental Protection Agency (Federal Register, 1979). Xenobiotics are released into the environment through industrial wastewaters and their direct application to soils, resulting in the introduction of carcinogenic and mutagenic chemicals to the environment. Their prevalence in the surroundings leads to

bio-accumulation and can disturb the ecological balance of the microbial system in soil. Such accumulation in the long run may be hazardous to the human's health. Since it is known to be toxic and carcinogenic to living organisms (Lyons et al., 1985), highly aniline-tolerant bacterial strains were isolated from various environments (Meyers, 1992; Loidl et al., 1990; Konopka et al., 1989; Konopka, 1993; Boon et al., 2001; Liu et al., 2002), which are desirable for environmental applications as well as for the biotransformation of aniline and its analogues. Several aniline degradative genes or gene clusters were cloned from aniline-degraders (Meyers, 1992; Fujii et al., 1997; Takeo et al., 1998; Fukumori and Saint, 1997; Uratal et al., 2004; Liang et al., 2005). Most of them were plasmidencoded, including pCIT1 (Pseudomonas sp. strain CIT1, Meyers, 1992), pYA1 (Acinetobacter sp. strain YAA, Takeo et al., 1998), pTDN1 (Pseudomonas putita UCC22,

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Fukumori and Saint, 1997), pNB2 (P. putida SM1443, Bathe, 2004) and these plasmids normally are large ones (about 100 kb). Up to date, only three gene clusters that contained complete genes involved in the pathway of conversion of aniline to intermediates of TCA-cycle have been reported, two were the plasmid-encoded tdn cluster (pTDN1) from P. putita UCC22 (Fukumori and Saint, 1997, 2001) and atd cluster from Acinetobacter sp. YAA (Fujii et al., 1997), and the other one was the chromosomeencoded tad cluster from D. tsuruhatensis AD9 (Liang et al., 2005), and clusters tdn and tad showed significant similarity in nucleotide sequence and genetic organization to each other despite of their different geographic origin (Liang et al., 2005). In both tdn and tad clusters, there were two genes for catechol 2,3-dioxygenase and two genes for ferredoxin-like protein, and this might be the molecular mechanism for both strains P. putita UCC22 and D. tsuruhatensis AD9 could also degrade methylsubstituted aniline in addition to aniline (Fukumori and Saint, 1997; Liang et al., 2005). Delftia sp. AN3 was a highly effective aniline-degrading strain and it could only degrade aniline (Liu et al., 2002). The molecular mechanism of aniline degradation and aniline degradative genes of strain AN3 might be different from those of strains P. putita UCC22 and D. tsuruhatensis AD9.

In this study, we described the cloning of a novel and complete gene cluster that converts aniline to TCA-cycle intermediates from *Delftia* sp. AN3 and the mechanism for aniline degradation by strain AN3 was also discussed in molecular level.

# 1 Materials and methods

# 1.1 Bacterial strains, plasmids, primers, and growth conditions

The bacterial strains, plasmids and primers used in this study are listed in Table 1. The culture media used were Luria-Bertani (LB) medium (Sambrook and Russell, 2001) and mineral medium (MMN) (Liu *et al.*, 2002). *Delftia* sp. AN3, *E. coli* JM109 was grown at 30 and 37°C, respectively. Apramycin (Am) was used in selective media at final concentration of 133 mg/L. The growth of strains was measured by the turbidity at the wavelength of 600 nm.

# 1.2 Plasmid curing and pulsed-field gel electrophoresis (PFGE) detection

For curing the plasmid from *Delftia* sp. AN3, a modified sodium dodecyl sulfate treatment method of El-Mansi *et* 

*al.* (2000) was used. Detection of the mega plasmid in *Delftia* sp. AN3 was carried out according to Barton *et al.* (1995). Separation of chromosome and plasmid DNAs was carried out on an agarose gel (1%) under conditions of 6 V/cm and 70 s for 22 h using a Bio-Rad pulsed-field gel electrophoresis apparatus (PFGE, Bio-Rad, USA).

#### 1.3 Construction of the genomic library of strain AN3

The genomic DNA of strain AN3 was obtained by the method of Sambrook and Russell (2001). The genomic DNA obtained was partially digested with *Sau*3AI or *Mbo*I and separated in a 0.6% (*W/V*) agarose gel. DNA fragments about 30 kb were recovered and purified, then ligated to pKC505 digested by *Hin*dIII and *Bam*HI and packaged by Gigapack III XL Packaging Extract Kit (Stratagene, USA), then introduced into component cells of *E. coli* JM109 by transduction.

#### 1.4 Screening for the positive clone

*Escherichia coli* transformants carrying this library were cultured on LB plate supplemented with Am (133 mg/L) and screened by spraying catechol solution (10 mmol/L dissolved in phosphate buffer) onto the colonies on the LB plate for detection of catechol 2,3-dioxygenase (C23O) activity. Then the positive-candidate clones were further confirmed by C23O assay. Protein concentration was determined by the method of Bradford (1976) using Bio-Rad protein assay kit, with bovine serum albumin as a standard.

#### 1.5 Assays of enzymatic activities

## 1.5.1 Aniline dioxygenase (AD)

Strain AN3 and positive clone (*E. coli* JM109-AN1) were grown on MMN (containing 300 mg/L of aniline for *E. coli* JM109-AN1) till to an  $OD_{600}$  about 0.1 and then cells were harvested by centrifugation at 5,000×g, washed twice with 20 mmol/L phosphate buffer (pH 7.0), and resuspended in the same buffer. This suspension was used to assay aniline dioxygenase. The activity of the enzyme was measured with an oxygen electrode (YSI, Ohio, USA), according to the method of Fukumori and Saint (1997). To estimate the endogenous respiratory rate, 0.3 ml of sterile distilled water instead of aniline solution was used in a parallel experiment.

#### 1.5.2 Catechol 2,3-dioxygenase

Cells (the same cell suspension as used in the aniline dioxygenase assay) were disrupted by sonication using an ultrasonic disruptor (TOMY UD-200; power 6, 2 min, three times on ice). The cellular lysate were centrifuged at  $19,000 \times g$  for 20 min, and the supernatant was used

Table 1 Strains, plasmids and primers used in this study

Strain or plasmid	Relevant characteristics	Reference
Delftia sp. AN3	Wild type that could degrade aniline	Liu et al., 2002
E. coli JM109	$recA1supE44endA1hsdR17gyrA96relA1Thi\Delta(lac-proAB)F' traD36proAB^+lacI^{q}lacZ\DeltaM15$	Sambrook and Russell, 2001
E. coli JM109-AN1	<i>E. coli</i> JM109 containing pKC505-AN1, growing on aniline as sole carbon, nitrogen and energy source	This study
pKC505	Shuttle vector, 18.7 kb, Am <sup>r</sup>	Richardson et al., 1987
pKC505-AN1	Am <sup>r</sup> , a 29.7-kb Sau3AI fragment from genomic DNA of Delftia sp. AN3 in pKC505	This study

– E. coli JM109

300

250

for enzymatic assay. The reaction mixture (total 3.0 ml) contained 2.0 ml phosphate buffer, 0.6 ml of 1 mmol/L catechol, 0.2 ml deionized water and 0.2 ml cellular lysates. The reaction proceeded at 22°C. C23O was determined by measuring the production of 2-hydroxymuconic semialdehyde at 375 nm (Sala-Trepat and Evans, 1971) using a DU-7 spectrophotometer (Beckman, USA). The absorption coefficient of 2-hydroxymuconic semialdehyde was 12,000 mol<sup>-1</sup>cm<sup>-1</sup>.

# 1.6 Plasmid isolation

For isolation of recombined plasmids bearing C23O activity, pKC505-AN1 of the positive clone, a modified alkaline lysis method was used (Thomas et al., 1998).

### 1.7 DNA sequencing, sequence assembly and analysis

The recombinant plasmid pKC505-AN1 containing a DNA fragment about 30 kb from strain AN3 was sequenced with shotgun method by the Beijing Genome Institute (Huada Corp., Beijing, China). Contigs were assembled using the GCG Wisconsin package. The orfs were analyzed using Vector NTI and DNAman software (Lynnon Biosoft, USA). Sequence comparisons were made against the sequences in the GenBank using the BLASTx program (Altschul et al., 1990).

#### 1.8 Construction of phylogenetic tree

Peptide sequences of various enzymes or subunits were extracted from NCBI (Altschul et al., 1990). Phylogenetic trees were generated using the neighbor joining method of Saitou and Nei (1987) with the AlignX software (Informax, Maryland), and multiple sequence alignment was done using ClustalX. (Thompson et al., 1997). The length of each branch pair represents the evolutionary distance between the sequences.

# 1.9 Aniline analysis

Aniline concentration was determined at 230 nm using HPLC (Waters, Millipore, USA) equipped with a C-18 reverse phase column (100 mm long) and a UV-detector. The elution was MeOH/H<sub>2</sub>O (V/V)=75/25 and the elution rate was 1.5 ml/min. Under these conditions the retention time of aniline was about 3.25 min.

# 2 Results

#### 2.1 Plasmid detection and curing experiment

Cells of strain AN3 for PFGE were grown on aniline (MMN) to ca.  $OD_{600}{=}0.6$  and then harvested and washed twice with sucrose buffer (0.3 mol/L). The result (data not shown) showed that there was not any band except of the genomic DNA. Furthermore, this aniline-degrading ability of strain AN3 could not be cured.

# 2.2 Cloning of aniline degradation genes from strain AN3, and the growth of recombinant strain on aniline as sole carbon, nitrogen and energy sources

A genomic library of strain AN3 was constructed as described in Materials and methods. The transformants

Aniline concentration (mg/L) 200 0.08 0.06 8 150 100 0.04 50 0.02 0 60 80 120 20 40 100 0 Time (h) Fig. 1 Growth of recombinant strain E. coli JM109-AN1 in mineral

--- E. coli JM109-AN1 --- Aniline remained

medium (MMN, Liu et al., 2002) using aniline as sole carbon, nitrogen and energy source and the degradation of aniline. The initial aniline concentration was 300 mg/L. The cultivation was conducted out on a shaker at 180 r/min at 30°C.

were spread out on LB plates containing Am (133 mg/L), and then catechol solution was sprayed onto the colonies formed on the plates. One colony turned to brilliant yellow firstly and then to brown after spraying of catechol solution, indicating that there was an accumulation of 2-hydroxymuconic semialdehyde, an intermediate in the aniline degradation pathway via meta-cleavage of catechol (Takeo et al., 1998). This recombinant strain could grow on aniline as sole carbon, nitrogen and energy source up to 0.1 of  $OD_{600}$  and degrade the added aniline completely (Fig.1). This recombinant strain and plasmid were designated as E. coli JM 109-AN1 and pKC505-AN1, respectively.

# 2.3 Expression of AD and C23O of the recombinant strain E. coli JM 109-AN1 growing on aniline

The results in Table 2 show that AD and C23O expressed well in E. coli (recombinant strain E. coli JM 109-AN1). The activities of AD and C23O were as high as 0.31 U/mg wet cell and 1.92 U/mg protein, respectively, despite they were lower than those of original strain AN3 (1.09 U/mg wet cell and 5.22 U/mg protein for AD and C23O, respectively). The results (Table 2) also indicated that the AD and C23O of strain AN3 or E. coli JM109-AN1 were highly specific, only aniline and catechol could be the substrate for AD and C23O, respectively; not any other substituted anilines or catechols could be catalyzed.

#### 2.4 Sequence analysis of the cloned fragment

The DNA fragment cloned in pKC505-AN1 was sequenced, and the nucleotide of the total 29.7 kb region was thereby determined. Sequence and orf analyses of this total 29.7 fragment indicated that there were at least 27 intact orfs (Fig.2a and Table 3). Homology searches for these orfs were performed to identify gene function. It was found that among these 27 intact orfs, at least 16 of which (danQTA1A2BRDCEFG1HIJKG2) were expected in corresponding to the complete metabolism of aniline to TCA-cycle intermediates via meta-cleavage pathway as shown in Fig.2b. As summarized in Table 3, the 6th-10th gene products danQTA1A2B and the danR product showed

0.12

0.10

3-Methylaniline

4-Methylaniline

growing on aniline					
	AD	C230			
Substrate	Activity* (U/mg wet cell)	Substrate	Activity* (U/mg protein)		
Aniline	1.09 (0.31)	Catechol	5.22 (1.92)		
2-Chloroaniline	0 (0)	3-Chlorocatechol	0 (0)		
3-Chloroaniline	0 (0)	4-Chlorocatechol	0 (0)		
4-Chloroaniline	0 (0)	3-Methylcatechol	0 (0)		
2,4-Dichloroaniline	0 (0)	4-Methylcatechol	0 (0)		
2-Methylaniline	0(0)	-			

 Table 2
 Activities of aniline dioxygenase (AD) and catechol 2,3-dioxygenase (C23O) of strain AN3 and recombinant strain *E. coli* JM109-AN1 growing on aniline

\* The activities in parentheses were those of enzymes expressed in recombinant strain *E. coli* JM109-AN1. One unit (U) of AD was defined as the amount consuming 1 mol  $O_2$ /min under the conditions described in Materials and methods. One unit (U) of C23O was defined as the amount of protein needed for oxidation of 1 µmol catechol/min at 22°C under the conditions described in Materials and methods.

0(0)

0 (0)



**Fig. 2** Genetic organization of the aniline degradative *dan* gene cluster and the comparison of it with *tad* cluster (a) and putative aniline degradation pathway (b) of *Delftia* sp. AN3. The sequenced region of 29.7 kb is shown. The function of other genes was explained in Table 3. AD: aniline dioxygenase; C23O: catechol 2,3-dioxygenase; HMSD: 2-hydroxymuconic semialdehyde dehydrogenase; HMSH: 2-hydroxymuconic semialdehyde hydrolase; OCD: 4-oxalocrotonate decarboxylase; OEH: 2-oxopent-4-dienoate hydratase; HOA: 4-hydroxy-2-oxovalerate aldolase; ADA: acetaldehyde dehydrogenase.

significant homology (64%–100%) with multi-component aniline dioxygenases and Lys-type regulators, respectively, present in other aniline degrading strains, *D. acidovorans* 7N, *P. putida* UCC22, *Frateuria* sp. ANA-18 and *D. tsuruhatensis* AD9, especially with *D. acidovorans* 7N. The remaining 11 gene products (DanQDCEFGHI112JKLM) exhibited considerable amino acid identity (64%–100%) to enzymes of central catechol degrading enzymes via *meta*-cleavage pathway found in other aromatics degrading and aniline-degrading bacteria. There are also some other genes (*danWVU*) that encode proteins such as aromatic amino-transferases and transport trans-membrane proteins found in *Polaromonas naphthalenivorans* CJ2 (Copeland et al., unpublished, accession number ZP\_01022585) and Polaromonas sp. JS666 (Lucas et al., unpublished, accession number ZP\_00502922) etc., showed 72%–75% amino acid identity. Besides, on the two sides of the dan gene cluster are the transposases encoded by danX1SN, and danO potential TnpR recombinase. These gene products are believed to function in horizontal gene transfer among the strains. In addition, orfX2YZ were also found in the meta-cleavage of other aniline degraders, such as D. tsuruhatensis AD9 (Liang et al., 2005), and Pseudomonas sp. K82 (Kim et al., 2004), and a phenol phenol degrader, Comamonas testosteroni TA441 (Arai et al., 1998) in addition to P. putida UCC22.

### Table 3 Analytical data on the aniline degradation genes and other genes of *Delftia* sp. AN3\*

ORF	Gene	Position (No. of nt)	Mw/No. of aa <sup>b</sup>	Putative function	Homologous protein (sequence identity)	Source	Accession No.
1	danW	1–1392 (1392)	51 kDa/464 aa	Aromatic Amino- transferase	Aminotran 1, 2 (299/398, 75%) Aminotran 1, 2	Polaromonas sp. JS666 P. naphthalenivorans	YP_549701 ZP_01022585
2	danV	1408–1731 (324)	12 kDa/108 aa	4-Hydroxybenzoyl-	(296/397, 74%) PnaP 1037 (78/115, 67%)	CJ2 P. naphthalenivorans CI2	ZP_01022584
		(324)		Correntoesterase	4HBT (80/115, 69%)	Polaromonas sp. JS666	ZP_00502921
3	danU	1734–2522 (789)	29 kDa/263 aa	Transmembrane protein	MotA ExbB (173/232, 74%)	Polaromonas sp. JS666	ZP_00502922
4	1V1	2(72, 2400	201-D-/272	T	MotA ExbB (169/232, 72%)	P. naphthalenivorans CJ2	ZP_01022583
4	danX I	2672–3490 (819)	30 KDa/273 aa	Transposase	Transposase (116/116, 100%) Transposase 25	D. acidovorans P. naphthalenivorans	AA038205
5	danS	3795–4304	18 kDa/170 aa	Transposase	(36/92, 39%) Transposase 34	CJ2 Burkholderia	ZP_00280163
		(510)			(22/43, 51%) Transposase 34	fungorum LB400 Ralstonia metalli-	ZP_00598173
6	danQ	4762-6264	56 kDa/501 aa	Glutamine	(22/43, 51%) ORF7NA (402/402, 100%)	durans CH34 D. acidovorans 7N	BAD61047
		(1505)		synthetase	(493/493, 100%) TdnQ (487/493, 98%)	D. acidovorans	AAO38206
7	danT	6287–7054 (768)	27 kDa/256 aa	Glutamine amidotransferase	ORF7NB (256/256, 100%)	D. acidovorans 7N	BAD61048
0	dan A I	7167 8444	48 kDa/426 aa	Lance subunit	TadT (155/240, 64%) Tdn A 1	D. tsuruhatensis AD9	AAX47240
0	aanAT	(1278)	48 KDa/420 aa	of dioxygenase	(425/426, 99%) ORF7NC	D. acidovorans	BAD61049
9	danA2	8447–9085	24 kDa/213 aa	Small subunit	(424/426, 99%) ORF7ND	D. acidovorans 7N	BAD61050
		(639)		of dioxygenase	(199/199, 100%) TadA2	D. tsuruhatensis AD9	AAX47242
10	danB	9105–10109 (1005)	36 kDa/335 aa	Aniline dioxygenase reductase	(142/200, 71%) ORF7NE (335/335, 100%)	D. acidovorans 7N	BAD61051
		()			TadB (251/335, 74%)	D. tsuruhatensis AD9	AAX47243
11	danR	10160–10917 (758)	28 kDa/252 aa	LysR-type regu- latory gene	ORF7NF (252/252, 100%)	D. acidovorans 7N	BAD61052
12	danD	11062_11469	14 kDa/136 aa	Small ferredovin-	TadR (190/252, 75%) ORE7NG	D. tsuruhatensis AD9	AAX47244
12	uunD	(408)	14 KDa/150 aa	like protein	(129/129, 100%) CbzT	P. putida UCC22	AAX38580
13	danC	11470–12411	35 kDa/314 aa	Catechol 2,3-	(69/111, 62%) ORF7NH	D. acidovorans 7N	BAD61054
		(942)		dioxygenase	(314/314, 100%) TdnC (260/214, 85%)	P. putida UCC22	CAA42452
14	orfX2	12559–13013 (455)	15 kDa/151 aa	Unknown product	(209/314, 85%) ORF4 (83/112, 74%)	P. putida UCC22	BAB62051
		. ,			ORFV (83/112, 74%)	D. tsuruhatensis AD9	AAX47251
15	danE	13016–14467 (1452)	51 kDa/484 aa	2-HMSD	HMSD (426/484, 88%)	Ralstonia eutropha JMP134	AAZ65034
16	danF	14740-15603	31 kDa/288 aa	2-HMSH	(354/485, 72%) DmpD	P. puttaa UCC22 Pseudomonas putida	САА36993
-		(864)			(202/283, 71%) TadF	D. tsuruhatensis AD9	AAX47253
17	danG1	15471–16412	34 kDa/314 aa	2-Keto-4-pentenoate	(190/277, 68%) CnbE	Comamonas sp. CNB-1	ABB13580
		(942)		hydratase	(261/261, 100%) ORF (226/260, 86%)	Rubrivivax gelati- nosus PM1	ZP_00242167

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Continued

18	danH	16435–17220 (786)	28kDa/262 aa	4-Oxalocrotonate decarboxylase	4-OD (216/262, 82%)	Rubrivivax gelati- nosus PM1	ZP_00242168
				,	CnbF (227/262, 79%)	<i>Comamonas</i> sp. CNB-1	ABB13581
19	orfY	17249–17848 (600)	26kDa/200 aa	Unknown product	ORF3 (135/193, 69%)	Pseudomonas putida UCC22	BAB62048
					ORFDR (56/139, 40%)	Deinococcus radiodurans R1	AAF11685
20	danI	18366–19595 (1230)	44kDa/410 aa	Acetaldehyde dehydrogenase	TdnI (270/300, 90%)	P. putida UCC22	BAB62056
					TadI (269/300, 89%)	D. tsuruhatensis AD9	AAX47255
21	danJ	19612–20649 (1038)	37kDa/346 aa	4-Hydroxy-2- oxovalerate aldolase	TadJ (291/327, 88%)	D. tsuruhatensis AD9	AAX47256
					TdnJ (291/327, 88%)	P. putida UCC22	BAB62057
22	orfZ	20665–21639 (975)	34kDa/325 aa	Putative exported protein	ORFJ (249/326, 76%)	C. testosteroni TA441	BAA88506
					CdoL (189/249, 75%)	Comamonas sp. JS765	AAG17139
23	danK	22252–23571 (1320)	46kDa/440 aa	2-Amino-5- chloromuconate deaminase	CnbH (427/427, 100%)	Comamonas sp. CNB-1	ABB13583
24	danG2	23617–24129 (513)	18kDa/171 aa	2-Keto-4-pentenoate hydratase	CnbI (171/171, 100%)	<i>Comamonas</i> sp. CNB-1	ABB13584
					TdnG (164/165, 99%)	Pseudomonas sp. CT14	ABA25976
25	danL	24410–27376 (2967)	10kDa/989 aa	Tn3 family transposase	TnpA (947/989, 95%)	D. acidovorans MC1	AAP88287
					TnpA (945/989, 95%)	Ralstonia metalli- durans CH34	CAI30244
26	danM	27382–27851 (471)	17kDa/157 aa	TnpR recombinase	TnpR (149/172, 86%)	Ralstonia metalli- durans CH34	NP 943467
					TnpR (147/152, 96%)	N. europaea ATCC 19718	CAI30245
27	danN	28352–29827 (1476)	53kDa/492 aa	Sulfate transporter	STAS (482/492, 97%)	<i>R. metallidurans</i> CH34	ZP_00595727
					STAS (405/491, 82%)	γ-Proteo- bacterium KT71	ZP_01101356

\* The accession number for the nucleotide sequence of the *dan* cluster in GenBank is DQ661649. The amino acid sequences of the products of the genes are available kDa in GenBank under the given accession numbers. No. of nt: The number of the nucleotide position in *dan* cluster; Mw: molecular weight; No. of aa: number of amino acid.

# 2.5 Comparison of the *dan* gene cluster with *tad* gene cluster

As shown in Table 3, the putative products of the dan genes showed identity to those of tad genes of D. tsuruhatensis AD9, which was a chromosome-encoded aniline degradative gene cluster. Hence, the gene organization of the dan gene cluster was compared with that of the tad gene cluster (Fig.2a). It was found that the gene organization of both gene clusters is similar, especially the aniline dioxygenase genes share the same organization and showed significantly identify in their products. However, there are some differences between them. Firstly, in dan gene cluster all of the genes corresponding for the conversion of aniline to TCA intermediates were in the same transcriptional direction. Secondly, there was only one set of C23O gene (danC) and ferredoxin gene (danD) in dan gene cluster, while there were two sets of C23O gene (tadC1C2) and ferredoxin genes (tadD1D2) in tad gene cluster; there were two 2-oxopent-4-dienoate hydratase genes (danG1G2) in dan gene cluster but there was only one (tadG) in tad gene cluster. Thirdly, the dan gene cluster lacks the genes corresponding to orfU, orfS, tadD2

and *tadC2* between *tadC1* and *orfV* of the *tad* cluster, and the genes corresponding to *tadL* (4-oxalocootonate tautomerase) and *orfX* (MarR-type regulator) of *tad* gene cluster. Fourthly, in contrast to *tad* gene cluster, in *dan* gene cluster there are two additional genes, *danY* and *danZ*, and their products showed considerable similarity to ORF3, a putative membrane protein, of *tdn* gene cluster of *Pseudomonas putida* UCC22 and to ORFJ, a putative exported protein, of *aph*, a phenol degradation gene cluster of *Comamonas testosteroni* TA441 (Arai *et al.*, 2000), respectively.

# 2.6 Phylogenetic relationships of the AD, C23O, and other proteins of strain AN3 with other homologous proteins

To clarify the phylogenetic relationship of the AD of strain AN3 with those of other aniline-degrading bacteria, phylogenetic trees were constructed using the amino acid sequences of the glutamine synthetase-like proteins, the large subunit of the dioxygenase component and aniline dioxygenase reductase in the AD enzyme systems. As shown in Figs.3a, 3b and 3c, the phylogenetic trees consisted of two major branches, the Tdn-branch and

a Aniline dioxygenase (AD) glutamine-synthetase-like protein					
100	DanQ	Delftia sp. AN3			
100 BAD61047	TdnQ	D. acidovorans 7N			
LAAO38206	TdnQ	D. acidovorans			
0.1 BAC82524	TdnQ	Frateuria sp. ANA-18			
	TadQ	D. tsuruhatensis AD9			
100 BAA12805	TdnQ	P. putida UCC22			
BAA13010	AtdA1	Acinetobacter sp. YAA			
b Large subunit of oxygenase comp	oonent				
94	DanA1	Delftia sp. AN3			
<u>100</u>  AAO38208	TdnA1	D. acidovorans			
BAD61049	TdnA1	D. acidovorans 7N			
0 1 BAC82526	TdnA1	Frateuria sp. ANA-18			
95 BAA12807	TdnA1	P. putida UCC22			
100 – AAX47241	TadA1	D. tsuruhatensis AD9			
BAA13012	AtdA3	Acinetobacter sp. YAA			
c Aniline dioxygenase reductase					
100	DanB	Delftia sp. AN3			
59 BAD61051	TdnB	D. acidovorans 7N			
AAX47243	TadB	D. tsuruhatensis AD9			
0 1 100 <sup>1</sup> BAA12809	TdnB	P. putida UCC22			
BAC82528	TdnB	Frateuria sp. ANA-18			
BAA23552	AtdA5	Acinetobacter sp. YAA			
d Catechol 2,3-dioxygenase (C23C	))				
99	DanC	Delftia sp. AN3			
1 <u>00</u> BAD61054	ORF7NH	D. acidovorans 7N			
88 AAC79918	CdoE	Comamonas sp. JS765			
<u>0.1</u> BAA34176	AphB	C. testosteroni TA441			
CAA42452	TdnC	P. putida UCC22			
56 100-AAX47246	TadC1	D. tsuruhatensis AD9			
100 AAD0525	) CbzE	P. putida GJ31			
AAF02430	PhnE2	Burkholderia sp. RP007			
ABB13075		Burkholderia sp. 383			
AAF02426	PhnE1	Burkholderia sp. RP007			
BAA23555	AtdB	Acinetobacter sp. YAA			
	0 TadC2	D. tsuruhatensis AD9			
100 BAB6205	0 TdnC2	P. putida UCC22			

**Fig. 3** Phylogenetic trees constructed based on the amino acid sequences of AD glutamine-synthetase (a), large subunit of the dioxygenase component (b), aniline dioxygenase reductase (c) and C23O (d). Sequence accession number and strain names are shown. The numbers at node indicate the bootstrap percentages of 1000 resamples.

the Atd-branch. In these trees, DanQ, DanA1 and DanB are located closest to TdnQ, TdnA1 and TdnB of *D*. *acidovorans* 7N, respectively.

To locate the *meta*-cleavage pathway of strain AN3 among those of other aromatic-compound-degrading bacteria, another phylogenetic tree was constructed using C23O amino acid sequence (Fig.3d). The *meta*-cleavage pathway of strain AN3 contains only one C23O, DanC. In the tree, DanC belongs to a branch including ORF7NH, TdnC and TadC1 from other aniline-degrading bacteria, *D. acidovorans* 7N, *P. putida* UCC22 and *D. tsuruhatensis* AD9, respectively. Fig.3d illustrates that DanC is closely related to TadC1 and TdnC, but not to TadC2 and TdnC2.

# **3 Discussion**

Delftia sp. AN3 was isolated as an aniline-degrader from activated sludge from a municipal wastewater treatment plant, and it could only assimilate aniline (Liu *et al.*, 2002). Up to date, the mechanism of several anilinedegraders has been illustrated in molecular level, including *D. tsuruhatensis* AD9 (Liang *et al.*, 2005), *P. putida* UCC22 (Fukumori and Saint, 1997) and Acinetobacter sp. YAA (Fujii *et al.*, 1997), due to the cloning and study of aniline-degradative gene clusters. Both plasmidencoded and chromosome-encoded gene clusters involving the complete metabolism of aniline to TCA-cycle intermediates have been cloned from *D. tsuruhatensis* AD9 (*tad* cluster, Liang *et al.*, 2005), *P. putida* UCC22 (*tdn*  cluster, Fukumori and Saint, 1997) and Acinetobacter sp. YAA (atd cluster, Fujii et al., 1997), and tad cluster was chromosome-encoded and the other two were plasmid-encoded. Although tad cluster and tdn cluster were differently encoded, they showed similar gene organization and striking identity of their products, and were hypothesized to be evolved from one ancestor (Liang et al., 2005). In this study, we cloned a novel aniline degradation gene cluster (dan cluster) from Delftia sp. AN3, which is responsible for the complete metabolism of aniline to TCA-cycle intermediates via a meta-cleavage pathway. Both the negative results of the PFGE and plasmid curing experiments suggested that dan cluster might be chromosome-encoded. The cloning of dan cluster into E. coli enabled it to grow on aniline as sole carbon, nitrogen and energy sources, and to degrade the added aniline completely (Fig.1). This result suggested that the dan cluster contained all the genes needed for conversion of aniline to TCA-cycle intermediates. This was further confirmed by the analysis of the sequence and orfs (Table 3). Furthermore, the enzymatic assays indicated that the AD and C23O did express well in the recombinant strain, E. coli JM109-AN1 (Table 2).

Despite they were from very close relatives and shared significant identity of most gene products, dan cluster differentiated to *tad* cluster in several aspects including gene organization, gene components, and their transcriptional direction (Fig.2a and Table 3). In addition, the phylogenetic analysis revealed that the components of AD systems of *dan* cluster and *tad* cluster belonged to the two different branches (Figs.3a, 3b, and 3c). These differences might infer that they have evolved from different ancestors. In dan cluster of strain AN3, there was only one set of C23O gene (danC), while there were two sets of C23O gene in tad cluster (tadC1 and tadC2) of D. tsuruhatensis AD9 and *tdn* cluster of *P. putida* UCC22 (*tdn*C and *tdn*C2). The phylogenetic analysis of DanC (Fig.3d) illustrates that it is closely related to TadC1 and TdnC, but not to TadC2 and TdnC2. Fukumori and Saint (2001) reported that TdnC and TdnC2 have distinct substrate specificity: TdnC showed relatively high activity on substituted catechols (3-methylcatechol and 4-methylcatechol), while TdnC2 showed less activity on these substituted catechols. Liang et al. (2005) reported that TadC1 and TadC2 have the similar substrate specificity to that of TdnC and TdnC2. Both strains UCC22 and AD9 can assimilate m-toluidine (3-methylaniline) and p-toluidine (4-methylaniline) in addition to aniline, and the authors assumed that it might be necessary for cells to acquire another C23O for these methylcatechols to expand the assimilation range for toluidines. But DanC has no C23O activity on those chlorocatechols and methylcatechols (Table 2) despite it is closely phylogenetic related to TdnC and TadC1. The presence of only one set of C23O gene and the specificity of AD and C23O (Table 2) of strain AN3 might be the reason for it not to assimilate any substituted anilines (Liu et al., 2002).

In conclusion, all of the above differences between *dan* cluster and other aniline-degradative clusters, especially

the gene organization, presence only one set of C23O gene or absence of some genes and specificity of AD and C230, indicated that *dan* cluster was a new aniline-degradative cluster, despite of they were quite similar and shared high identity of the gene products.

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