Isolation and characterization of a bacterial strain that degrades cis-dichloroethene in the absence of aromatic inducers

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Introduction

Tetrachloroethene (PCE) and trichloroethene (TCE) have been widely used as organic solvents in many industries, including the metal processing, semiconductor, and dry cleaning industries. However, unexpected leakages of these solvents into the soil and groundwater from factories have caused problematic environmental contamination throughout the world (ATSDR 2011 substance priority list). Microbial biodegradation is a promising solution to environmental chloroethene contamination. PCE and TCE can be degraded to ethene by reductive dehalogenases found in anaerobic bacteria including those in the genus Dehalococcoides (Fig. 1) (Futagami et al., 2008; Maymo-Gatell et al., 1999). However, incomplete degradation, which sometimes occurs, is known to result in the accumulation of hazardous intermediate substances such as cis-

cDCE degradation. YKD221 did not appear to grow on cDCE in a minimal salt liquid medium. However, YKD221 did exhibit an enhanced increase in cell concentration and volume of cells during growth on minimal salt agar plates with cDCE when first grown in LB medium. This behavior appears to have led us to misinterpret our initial results on YKD221 as an indication of improved growth in the presence of cDCE.

Key Words: cis-dichloroethene; Pseudomonas; toluene dioxygenase; trichloroethene
dichloroethene (cDCE) and vinyl chloride (VC) (Futagami et al., 2008; Maymo-Gatell et al., 1999).

The ability to degrade TCE has been reported in many aerobic bacteria, including the toluene-degrading strains Pseudomonas putida F1 (Wackett and Gibson, 1988), Pseudomonas mendocina KR1 (McClay et al., 1995), and Ralstonia pickettii PKO1 (Leahy et al., 1996), the phenol-degrading strain Burkholderia cepacia G4 (Nelson et al., 1987), the methane-degrading strains Methylosinus trichosporium OB3b (Tsien et al., 1989) and Methylocystis sp. M (Uchiyama et al., 1992), and the ammonium-degrading strain Nitrosomonas europaea (Arciero et al., 1989). These bacteria employ dioxygenases or monooxygenases to degrade not only their target substrates (toluene, phenol, methane, or ammonium) but also TCE. These oxygenases are believed to convert TCE to TCE epoxide, which is further transformed to glyoxylate or formate as illustrated in Fig. 1. Because these oxygenases are induced in the presence of their target substrate, the target substrate must be present for the co-metabolic degradation of TCE. An exception is the phenol-degrading strain Wautersia numazuensis TE26, which has been shown to degrade TCE in the absence of an inducing substrate (Kageyama et al., 2005).

Co-metabolic degradation of cDCE has been observed in toluene-degrading strains, such as F1, KR1, and PKO1, and is believed to proceed via a route similar to that illustrated in Fig. 1 (Clingenpeel et al., 2012; Wackett and Gibson, 1988). Similar degradation routes have also been hypothesized for VC and ethene degradation by alkene monooxygenases in the ethene-accumulating bacteria Mycobacterium sp. JS60 and Nocardiooides sp. JS614 (Fig. 1) (Coleman and Spain, 2003; Mattes et al., 2005).

The same oxygenases are believed to be responsible for both TCE and cDCE degradation via epoxides. Polaronomas sp. JS666, a cDCE-degrading strain, has been reported to grow marginally on cDCE (Coleman et al., 2002). Strains JS666 and TE26 degraded cDCE without any inducing substrates (Coleman et al., 2002; Kageyama et al., 2005). These bacteria appear to be useful for remediating cDCE, which accumulates during anaerobic degradation, from the environment.

In order to develop a strain that can be used for bioremediation without inducing substrates, cDCE-degrading aerobic bacteria were selected from cDCE-contaminated soils in this study. An isolate that exhibited strong degradation activity in the absence of inducing substrates was then characterized.

### Materials and Methods

**Bacterial strains, plasmids, culture conditions, and chemicals.** The bacterial strains and plasmids used in this study are listed in Table 1. YKD221 and E. coli strains were grown at 30°C and 37°C, respectively. Strains were grown in the following media: W medium composed of 1.7 g of KH2PO4, 9.8 g of Na2HPO4, 1.0 g of (NH4)2SO4, 0.1 g of MgSO4·7H2O, 9.5 mg of FeSO4·7H2O, 10.75 mg of MgO, 2.0 mg of CaCO3, 1.44 mg of ZnSO4·7H2O, 1.12 mg of MnSO4·4H2O, 0.25 mg of CuSO4·5H2O, 0.28 mg of CoSO4·7H2O, 0.06 mg of H3BO3, and 51.3 µl of 12 N HCl per liter; LB medium composed of 10 g of Bacto Tryptone, 5 g of yeast extract, and 5 g of NaCl per liter; 1/5 LB medium composed of 2 g of Bacto Tryptone, 1 g of yeast extract, and 5 g of NaCl per liter; or R2A agar composed of 0.5 g of yeast extract, 0.5 g of Proteose Peptone No. 3, 0.5 g of Casamino Acids, 0.5 g of Dextrose, 0.5 g of Soluble Starch, 0.3 g of sodium pyruvate, 0.03 g of dipotassium phosphate, and 0.05 g of magnesium sulfate per liter. If necessary, 15 g of agar, 25 mg of kanamycin, 25 mg of nalidixic acid, or 100 g of sucrose per liter were added to the medium. cDCE, trans-dichloroethene (tDCE), TCE, PCE, and toluene were supplied in vapor as sole carbon sources.

**Isolation of cDCE-degrading bacteria.** Soil samples were collected from Akita, Chiba, Niigata, and Osaka, in Japan. Ten grams of soil were placed in a 120-ml glass vial and 100 µl of W medium, 100 µl of cDCE, 100 µl of a solution containing 167 mM urea and 15 mM KH2PO4, and 50 mg of activated carbon powder were added to grow microorganisms individually in the solid-phase-like conditions. Sample vials were sealed with Teflon-coated butyl rubber stoppers and aluminum caps, then incubated at 30°C with rotary shaking at 15 rpm. Each vial was opened every 15 days to add the amounts of W medium, cDCE, urea, and KH2PO4 described above and to remove 0.5 g of soil. Vials were then resealed, and incubation was continued for a total of six months. A 0.2-µl aliquot of the 0.5-g of soil removed from each vial was suspended in 5 ml of W containing 0.5% (w/v) agar and overlaid on a W medium agar plate. Plates were incubated for approximately 7 days at 30°C with cDCE in a sealed container. Any colonies that appeared on the plates were transferred in duplicate.
to fresh W agar plates and incubated at 30°C with, or without, cDCE in a sealed container. Colonies that exhibited better growth in the presence of cDCE than in the absence of cDCE were subjected to single colony isolation using LB, 1/5 LB, or R2A agar plates. Among the 12 candidate strains that we obtained, we chose YKD221 to investigate further, because this strain showed a superior cDCE degradation activity compared with the other strains.

**Degradation activity assay.** Cells were grown to mid-log phase in LB liquid medium, or overnight on LB agar plates. Toluene-induced cells were grown for 24 hours on W agar plates with toluene supplied as a vapor, then collected by suspending in 5 ml of W liquid medium followed by centrifugation. These cells were washed three times with W liquid medium then suspended in W liquid medium to an OD₆₀₀ of 10. One ml of cell suspension was added to a 20-ml glass vial containing 500 μM of chlorinated ethenes, sealed with a Teflon-coated butyl rubber stopper and an aluminum cap, and incubated at 30°C with shaking at 180 rpm. Autoclaved dead cells were used as a control to avoid interference from chlorinated ethene leakage out of sealed vials as well as adsorption of chlorinated ethenes to cell surfaces and the butyl rubber stopper. Depletion of chlorinated ethenes was measured by quantifying the amount of chlorinated ethenes remaining in the headspace of a sealed vial. Headspace gas of 250 ml was subjected to gas chromatography with flame ionization detection (GC-FID) using an Agilent 7890A GC (Agilent, Palo Alto, CA, USA) equipped with a DB-624 capillary column (30 m by 0.32 mm, J&W Scientific, Folsom, CA, USA). The oven temperature was held for 5 min at 35°C, then increased to 165°C at 30°C/min, and held constant at 165°C for 1 min (total runtime: 10.3 min). The injection and detection temperatures were 200°C and 250°C, respectively. Chlorinated ethene peaks were detected at the following retention times: cDCE, 5.01 min; tDCE, 3.34 min; TCE, 6.82 min; PCE, 8.10 min. Peak areas were auto-integrated by the Agilent ChemStation software (Agilent) to determine the amount of chlorinated ethene remaining in each vial.

To determine the amount of chloride ion released, precultured cells were washed three times with 10 mM phosphate buffer (pH 7.0) and resuspended to an OD₆₀₀ of 10 in 10 mM phosphate buffer (pH 7.0) instead of W liquid medium, which contains a large amount of chloride ions. A sealed vial containing 1 ml of cell suspension and 500 μM of chlorinated ethenes was incubated at 30°C with shaking at 180 rpm. Heat-killed cells were used as a control to avoid interference from chloride ions released by the cells. Depletion of chlorinated ethenes was determined as described above; release of chloride ion was quantified by ion chromatography (IC) using an Ion Analyzer IA-300 instrument (DKK-TOA, Tokyo, Japan). After GC-FID analysis, the cell suspension was centrifuged, and the supernatant was filtered through a 0.22 μm syringe filter. A 100 μl aliquot was diluted with 900 μl of filtered MilliQ water and subjected to IC analysis according to the manufacturer’s instructions. IC calibration was performed using a NaCl solution of known concentration.

**Determination of the number and size of cells during growth on cDCE.** YKD221 cells grown in LB liquid medium were washed three times with W liquid medium and resuspended in W liquid medium to a concentration of approximately 5.0 × 10⁷ cells/ml. One ml of cell suspension was placed in a 100-ml glass vial containing 500 μM of cDCE to keep enough oxygen for an oxygenase reaction, sealed with a Teflon-coated butyl rubber stopper and an aluminum cap, and incubated at 30°C without shaking. Every three days, a 100-μl aliquot of the culture was transferred to a 100-ml glass vial containing 900 μl of fresh W liquid medium with, or without, 500 μM of cDCE and incubated at 30°C without shaking. An aliquot of cell suspension was diluted and spread onto LB agar plates, and the concentration of living cells was determined by colony counting after incubation. The sizes of YKD221 cells after incubation with, or without, 500 μM of cDCE were determined by microscopic observation after Gram staining. Gram staining of cells was performed according to the standard procedure (Murray et al., 1994).

**Sequence analysis.** The 16S rRNA gene of YKD221 was amplified by PCR using the universal primer pair (10F and 1500R, Table S1) and sequenced using the ABI 3730 DNA Analyzer (Applied Biosystems, Foster City, CA, USA). The draft genome of YKD221 was sequenced using a combined strategy of shotgun sequencing with 454 GS FLX Titanium (Roche Diagnostics, Branford, CT, USA) and paired-end sequencing with MiSeq 1000 (llumina, Inc., San Diego, CA, USA). The obtained data were assembled using Newbler version 2.6. The assembled draft sequence of the YKD221 genome was analyzed using the Microbial Genome Annotation Pipeline (MiGAP, http://www.migap.org/), which predicts protein-coding.

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**Table 1.** Strains and plasmids used in this study.

| Strain or Plasmid | Relevant characteristic(s) | Reference or source |
|-------------------|-----------------------------|---------------------|
| *Pseudomonas* sp. YKD221 | A wild-type cDCE degrader, Nal⁺, Tol⁺ | This study |
| *Pseudomonas* sp. YKD221 ΔtodC1 | A derivative of YKD221 with the 1,323-bp todC1 deletion, Nal⁺, Tol⁺ | This study |
| Escherichia coli S17-1 | recA, pro, hsdR, RP4-2-TC::Mu in the chromosome | Simon et al. (1983) |
| Escherichia coli JM109 | recA1, supE44, endA1, hsdR17, gyrA96, relA1, thi, Δ(lac-proAB) F’ [traD36 proAR’ lacF’ lacZAM15] | Yanisch-Perron et al. (1985) |

| Plasmids | Relevant characteristic(s) | Reference or source |
|----------|-----------------------------|---------------------|
| pK18mobsacB | pMB1 ori, oriT (RP4), sacB, Kan’ | Schilfer et al. (1994) |
| pK18ΔtodC1 | pK18mobsacB carrying the ΔtodC1 construct | |

Tol⁺, growth on toluene, Tol⁻, no growth on toluene, Kan’, resistance to kanamycin, Nal’, resistance to nalidixic acid.
tRNA, and rRNA genes. Functional annotations of the predicted protein-coding genes were assigned based on UniProt, InterPro, HAMAP, and an in-house database composed of manually curated microbial genome sequences, as reported previously (Shintani et al., 2013).

Nucleotide sequences of the housekeeping genes gyrB, rpoD, and recA, as well as genes involved in toluene degradation, were extracted from the draft genome sequence of YKD221. The draft genome sequence was then aligned with reference sequences obtained from the National Center for Biotechnology Information (NCBI) and the Biological Resource Center of the National Institute of Technology and Evaluation (NBRC) databases using the ClustalW algorithm within MEGA (version 5.0, http://www.megasoftware.net/index.php, Tamura et al., 2011). Phylogenetic trees based on the 16S rRNA and gyrB gene sequences of YKD221 and related type strains were drawn using the neighbor-joining and maximum-likelihood methods (Saitou and Nei, 1987). The 16S rRNA, gyrB, rpoD, and recA, and the toluene-degradation genes were analyzed against the NCBI BLASTN database (http://blast.ncbi.nlm.nih.gov/Blast.cgi). The average nucleotide identity (ANI) was calculated using JSpecies with the default settings for ANI based on BLAST (http://www.imedea.uib.es/jspecies). Single nucleotide polymorphisms (SNPs) between YKD221 and P. putida F1 were identified by mapping YKD221 raw read sequences to F1 complete genome sequence (Accession No. CP000712) using GS Reference Mapper version 2.8 (Roche Diagnostics). The draft genome sequence of YKD221 was deposited in the GenBank nucleotide sequence database under the accession number of BBNC01000001–BBNC01000091.

**Construction of a gene deletion mutant.** DNA manipulations were performed essentially as described by Ausubel et al. (1990) and Sambrook et al. (1989). The region containing the upstream terminal 15-bp sequences of and the connecting upstream 1,045-bp sequences from todC1 were amplified from YKD221 genomic DNA by PCR using the primer pair, FtodC1_FP and FtodC1_RP. A vector plasmid, pK18mobsacB, was linearized by EcoRI digestion and ligated with the upstream and downstream PCR products using an In-Fusion HD Cloning kit (Takara Bio, Otsu, Japan) according to the manufacturer’s protocols. The structure of the resulting recombinant plasmid, pK18AtodC1, was confirmed by restriction enzyme analysis. pK18AtodC1 was introduced into YKD221 by conjugal transfer using E. coli S17-1, and kanamycin-resistant transconjugants were selected in the presence of kanamycin and nalidixic acid. The todC1-deletion mutant with sucrose resistance, which results from allelic exchange, was selected in the presence of sucrose as described previously (van der Geize et al., 2001). The deletion of todC1 in the resulting mutant was confirmed by colony PCR using the primers todC1C2_F and todC1C2_R (Table S1) as well as the inability to grow on toluene. This mutant was designated YKD221AtodC1.

**Results and Discussion**

**Isolation and identification of the selected strain**

Screening of cDCE-degrading bacteria was performed by selecting the candidates able to grow on minimal salt medium plates in the presence of cDCE, to obtain bacteria able to degrade cDCE in the absence of inducing substances. The isolate YKD221 formed a larger colony in the presence of cDCE than in its absence, suggesting that YKD221 grows better in the presence of cDCE than in the absence of cDCE. YKD221 exhibited good cDCE degradation activity as described in the next section. To determine the taxonomic position of YKD221, the 16S rRNA gene sequence of YKD221 was amplified by PCR, sequenced, and compared with 16S rRNA sequences from *Pseudomonas* type strains to generate the phylogenetic tree depicted in Fig. 2A using the neighbor-joining method. The phylogenetic tree created using the maximum-likelihood method was essentially the same as Fig. 2A. These results indicated that the 16S rRNA gene sequence of YKD221 is most similar to *Pseudomonas plecoglossicida* type strain NBRC 103162 with a 99.8% identity (Table S2). However, YKD221 also showed a high identity with the type strains P. monteilii NBRC 103158 and P. putida NBRC 14164 (99.6% and 99.1%, respectively). To verify the taxonomic position of YKD221, draft genome sequencing of YKD221 was performed, and the sequence of gyrB, a DNA gyrase, was used to generate another phylogenetic tree of YKD221 and *Pseudomonas* type strains (Fig. 2B). The gyrB phylogenetic tree suggests that YKD221 is more related to *P. monteilii* and *P. putida* than *P. plecoglossicida*. Comparison of the rpoD and recA sequences encoding RNA polymerase sigma factor and recombinase A suggests the same notion (Table S2). Then, ANI analysis of YKD221 and *Pseudomonas* type strains was performed to determine definitively the taxonomic position of YKD221 (Fig. 2C). We included the *P. putida* strains F1 and KT2440 in the ANI analysis because F1 is a well-known degrader of toluene and TCE and KT2440 is a versatile host strain derived from the famous toluene-degrading *P. putida* strain mt-2. The phylogenetic tree generated by ANI analysis indicated that the strains F1, KT2440, and YKD221 are closely related and that they are slightly more related to *P. putida* NBRC 14164 than to *P. monteilii* NBRC 103158. However, the ANI percentage values between YKD221 and *P. putida* NBRC 14164 and between YKD221 and *P. monteilii* NBRC 103158 are less than 90%. These results suggest that YKD221 belongs to neither *P. putida* nor *P. monteilii*, as an ANI percentage value of 95% has been proposed to be the border between species (Auch et al., 2010; Goris et al., 2007). ANI percentages between YKD221 and F1 and between YKD221 and KT2440 are approximately 100% and 97%, respectively, suggesting that YKD221, F1, and KT2440 form a new species. These results also reveal that genome sequences are very similar between F1 and YKD221. Between them 15 SNPs were found, four of which are located in coding regions. The YKD221 genome has four insertions of 81-, 100-, 113-, and 191-bp compared with the F1 genome. Because these SNPs and insertions are not located around aromatic or alkane degradation genes.

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it is difficult to deduce the sequence difference that is responsible for the difference in chloroethene degradation activity between them. It is noteworthy that strains isolated from Japan and the United States share very similar genome sequences.

Degradation activity of YKD221 on chloroethenes

Resting YKD221 cells, which were grown in LB medium and suspended in 1 ml minimal salt medium, depleted nearly 500 nmol of cDCE within 48 hours (data not shown). These results indicated good YKD221 degradation activity on cDCE. Autoclaved dead cells of YKD221 appeared to deplete a small amount of cDCE, suggesting that a small fraction of cDCE adsorbs to dead YKD221 cells. The release of chloride ions during cDCE depletion was examined in 1 ml phosphate buffer to estimate the proportion of released chloride ion to the amount of cDCE depleted (Fig. 3). The degradation activity of YKD221 decreased approximately 50% in phosphate buffer, resulting in the depletion of 267 nmol of cDCE in 48 hours. Because 82 nmol of cDCE adsorbed to autoclaved cells, cDCE degradation by living YKD221 cells was estimated to be 185 nmol. Over 48 hours, 722 nmol of chloride ions was released by living YKD221 cells, while 211 nmol of chloride ions was released from the autoclaved cells (data not shown). These data suggest that 511 nmol of chloride ions released was associated with cDCE degradation (Fig. 3). A cDCE molecule has two chlorine atoms, so the amount of chlorine atoms released from cDCE should be twice the amount of cDCE that is depleted. This estimation does not agree with our results, however. Probably cDCE molecules adsorbed to dead cells were degraded in the living cells and it is deduced that a total of 267 nmol...

Fig. 2. Phylogenetic trees based on the 16S-rRNA gene sequences (A), the gyrB gene sequences (B), and the ANI analysis (C) of the strain YKD221 and the Pseudomonas type strains.

The scale bars in the panels A and B stand for substitutions per nucleotide position. Numbers at the nodes indicate levels of bootstrap support (%) based on a neighbor-joining analysis of 1,000 re-sampled datasets; only values above 50% are given. The scale bar in the panel C represents the ANI value. The accession numbers of the 16S rRNA and gyrB gene sequences are: P. moorei RW10, NR_042542 and AM293560; P. mosselli BCRC17518, AP072688 and FJ418640; P. taiwanensis BCRC17751, FJ418634 and EU103629, respectively. The nucleotide sequences of the other strains used were obtained from their genome sequences. The accession numbers of the genome sequences are: P. plecoglossicida NBRC103162, BBIV01000001–BBIV01000097; P. monteilii NBRC103158, BBIS01000001–BBIS01000132; P. putida NBRC14164, AP013070; P. putida KT2440, AE015451; P. putida F1, CP000712; P. fulva NBRC16637, BBQ01000001–BBQ01000046; P. parafulva NBRC16636, BBU01000001–BBU01000055; P. japonica NBRC103040, BBIR01000001–BBIR01000187; P. cremoricolorata NBRC16634, BBIP01000001–BBIP01000041; P. oryzihabitans NBRC102199, BBIT01000001–BBIT01000034; P. entomophila LA48, CT573326; P. aeruginosa PA01, AE004091; Halomonas elongata NBRC15536, FN869568.

Fig. 3. Depletion of cDCE and release of chloride ions from cDCE by YKD221.

Cells suspended in 1 ml of 10 mM phosphate buffer (pH 7.0) at OD₆₀₀ of 10.0 were encapsulated with 500 nmol cDCE in a sealed vial and the amount of cDCE depletion was estimated by headspace GC-FID. The release of chloride ions in the cell suspension was estimated by IC. The net depletion of cDCE by living cells, which was estimated by subtracting the background cDCE depletion with autoclaved cells, is represented by open circles. The overall depletion of cDCE by living cells is represented by filled circles. The net release of chloride ions from cDCE, which was estimated by subtracting indigenous chloride ions observed with autoclaved cells, is indicated by open diamonds. Each value is the average ± standard deviation (error bars) based on three independent experiments.
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of cDCE was degraded to form 511 nmol of chloride ions by living YKD221 in 48 hours. In this case, the amount of released chloride ions (511 nmol) is 1.91-fold greater than the amount of cDCE depleted (267 nmol). Regardless, the amount of chloride ions released specifically from living cells was about twice as much as the amount of cDCE depleted by the living cells, indicating that YKD221 efficiently degrades cDCE and removes most of the chlorine atoms from cDCE.

To determine the range of substrates that YKD221 can degrade, the degradation activity of YKD221 on cDCE, TCE, and PCE was examined. YKD221 resting cells in 1 ml minimal salt medium degraded approximately 150 nmol out of 500 nmol TCE within four days, but the cells were unable to degrade tDCE and PCE (data not shown). Additionally, the degradation activity of YKD221 on cDCE was far greater than that on TCE.

**Toluene-dependent cDCE degradation activity of YKD221**

It was reported that the toluene-inducible toluene dioxygenase encoded by *todC1C2BA* is responsible for TCE degradation in *P. putida* F1 (Wackett and Gibson, 1988). Because the genome sequence of YKD221 is very similar to the sequence of F1, toluene-inducible toluene dioxygenase may be involved in TCE degradation in YKD221. To examine this hypothesis, the growth of YKD221 on toluene and the degradation of TCE by YKD221 after growth on toluene were investigated. YKD221 grew well on toluene (data not shown) and ex-
Involvement of YKD221. This study reports the first evidence of the poor that it was difficult to detect apparent colonies. We minimal salt agar plates in the presence of cDCE was so growth on minimal salt agar plates in the presence of cDCE, we expected that YKD221 assimilated cDCE as a degradation activity on both TCE and cDCE is determined by spreading the cell suspension on LB agar plates after appropriate dilutions. In the first round of growth, the cell concentrations in the presence and absence of cDCE unexpectedly increased by 7.3 and 3.8 fold, respectively, compared with that at the start, 4.8 × 10^7 CFU/ml (Fig. 6). In the next five rounds of growth, however, cultures grown in the absence and presence of cDCE reached similar concentrations, which are significantly lower than those of the first round of growth. Cell concentrations in the following two rounds of growth decreased to around 1 × 10^7 CFU/ml and those in the last three rounds of growth converged to around 2 × 10^6 CFU/ml independently with the presence of cDCE. The cDCE independent cell concentrations in the next five rounds of growth indicate that YKD221 does not assimilate cDCE. Although each culture was diluted to one tenth with fresh W-minimal salt medium on each transfer, the cell concentrations after three days of cultivation were largely similar in the last three rounds of growth. YKD221 might have assimilated carbon dioxide using the acetyl-CoA pathway enzymes for CO2 fixation; the presence of enzymes of this pathway was suggested by the YKD221 genome sequence. Unexpected growth in the first round appears to be supported by the residual nutrient from the preculture. The 1.9-fold higher cell concentration in the presence of cDCE indicates better growth and more frequent cell division in the presence of cDCE.

When we measured the sizes of cells in the first round of growth following the transfer from LB medium, the average cell size in the presence of cDCE was 2.0 ± 0.5 μm in length by 1.5 ± 0.4 μm in width, while the average cell size in the absence of cDCE was 1.9 ± 0.4 μm in length by 1.2 ± 0.2 μm in width. The average cell volume in the presence of cDCE was estimated to be 1.7-fold larger than the volume in the absence of cDCE. The higher cell concentration and larger cell volume in the presence of cDCE appeared to result in larger colonies. Although a reason remains to be determined, it is suggested that the presence of cDCE and the residual nutrient from the soil positively affects the first round of growth of YKD221 in minimal salt medium during screening experiments.

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**Supplementary Materials**

Supplementary tables are available in our J-STAGE site (http://www.jstage.jst.go.jp/browse/jgam).

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