INTRODUCTION

Nitrilases are enzymes widely expressed in prokaryotes and eukaryotes that hydrolyse non-peptide carbon–nitrogen bonds [1]. They are important industrial enzymes used to convert nitriles directly into corresponding acids and ammonia [2]. Nitrilases have attracted interest as a valuable alternative to chemical hydrolysis in organic chemical processes because of their mild reaction conditions and reduced output of environmental pollution. Previous studies have reported the occurrence, applicability and action of nitrilases, and the cloning of the nitrilase genes from Alcaligenes faecalis JM3, Arabidopsis thaliana, Brassica napus, Gordonia terrae MA-1, Klebsiella pneumoniae subsp. ozaenae, Rhodococcus rhodochrous K22 and R. rhodochrous J1 [1,3–5]. Nitrilases have different substrate specificities, making them useful for the hydrolysis of a large number of nitriles [5]. Nitrilases have been classified into three major categories on the basis of substrate specificity, although some nitrilases exhibit broad substrate specificity. Aromatic nitrilases act on aromatic or heterocyclic nitriles, aliphatic nitrilases act on aliphatic nitriles, and arylacetonitrilases act on arylacetonitriles [5,6].

The catalytic triad Cys-Glu-Lys is conserved in all members of the nitrilase superfamily [1,4,7–9], and there are currently no crystal structures available for nitrilases, but there are for other members of the nitrilase superfamily: N-carbamyl-D-amino acid amidohydrolase from Agrobacterium [10], nitrilase–fragile histidine triad fusion protein (NifH/Ant1) from Caenorhabditis elegans [11] and a homologous protein of unknown function from Saccharomyces cerevisiae [12]. A recent article on molecular modelling of nitrilase [13] described interactions between the structural subunits of nitrilase, but did not provide details about its active site. In spite of many studies on substrate specificity of nitrilases, the mechanism of substrate specificity has not been elucidated using structural and mutational analysis, except for study of the substrate specificity of arylacetonitrilase [14].

In the present paper, we performed sequence alignment, mutational analysis and molecular modelling of nitrilase in order to find a determinant residue for the substrate specificity of nitrilase from Rhodococcus rhodochrous ATCC 33 278. Our results demonstrate that aromaticity of the amino acid at position 142 is required for reaction with aliphatic substrates.

MATERIALS AND METHODS

Materials

The kits for PCR product purification, gel extraction and plasmid preparation, as well as the DNA-modifying enzymes, were purchased from Promega. The aromatic nitriles (benzonitrile, m-tolunitrile and 2-cyanopyridine) and aliphatic nitriles (adiponitrile, glutaronitrile, sebacanitrile and succinonitrile) were purchased from Sigma Chemical Company.

Bacterial strains, plasmids and growth conditions

Rhodococcus rhodochrous ATCC 33 278, Escherichia coli ER2566, and the plasmid pET-28a (+) (Novagen) were used as a source of genomic DNA of nitrilase, host cells and expression vector respectively. R. rhodochrous ATCC 33 278 was grown aerobically at 26 °C in yeast malt extract broth (1.0% glucose, 0.5% peptone, 0.3% yeast extract and 0.3% malt extract). The recombinant E. coli for expression of the enzyme was cultivated in 500 ml of LB (Luria–Bertani) medium (1.0% tryptone, 0.5% yeast extract and 1.0% NaCl) in a 2 litre flask containing 20 μg/ml kanamycin at 37 °C with shaking at 200 rev./min.

Abbreviation used: LB, Luria–Bertani.

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When the $D_{600}$ of the culture reached 0.6, IPTG (isopropyl β-D-thiogalactoside) was added to a final concentration of 0.1 mM to induce nitrilase expression, and the culture was incubated at 16°C for 16 h.

Gene cloning and site-directed mutagenesis of nitrilase

The nitrilase gene was amplified by PCR using genomic DNA isolated from R. rhodochrous ATCC 33278 as a template. Oligonucleotide primer sequences used for gene cloning were designed using the published DNA sequence for nitrilase from R. rhodochrous J1 (GenBank® accession number D11425). Forward (5′-TTCATATGGTGCTGAAATACACAAAC-3′) and reverse (5′-TACAAGCTTTCAAGTGGGGCGTG-3′) primers were designed for introduction of the underlined NdeI and HindIII restriction sites respectively. The amplified DNA fragment was purified using a Promega PCR purification kit and was ligated into the NdeI and HindIII sites of pET-28a(+). The resulting plasmid was used to transform the E. coli ER2566 strain. E. coli cells containing the plasmid were grown on LB medium. The expression of the gene encoding nitrilase was determined by both SDS/PAGE (12% gels) and the assay of enzyme activity. Site-directed mutagenesis was performed using the overlap extension PCR method [15].

Purification of nitrilase

The recombinant cells were harvested from the culture broth by centrifugation at 6000 g for 30 min at 4°C, washed twice with 0.85% NaCl and then resuspended in buffer A composed of 50 mM sodium monophosphate, 300 mM NaCl, 10 mM imidazole and 0.1 mM PMSF as protease inhibitors. The resuspended cells were disrupted by ultrasonication (Fisher Scientific) on ice. The cell debris was removed by centrifugation at 13000 g for 20 min at 4°C, and the supernatant was filtered through a 0.45 μm pore-size filter. The filtrate was applied to HisTrap HP chromatography column (GE Healthcare) equilibrated with buffer A. The bound protein was eluted with a linear gradient between 10 and 250 mM imidazole in buffer A. The active fraction was dialysed at 4°C for 24 h against 50 mM potassium phosphate buffer (pH 7.5), and the resulting solution was used as purified enzyme. All purification steps using columns were carried out with an FPLC system (Bio-Rad Laboratories) in a cold room. The protein concentration was quantified by the Bradford method [16]. The purified proteins were confirmed by SDS/PAGE (12% gels).

Determination of molecular mass

The subunit molecular mass of nitrilase were examined by SDS/PAGE (12% gels) under denaturing conditions, using the proteins of a pre-stained ladder (MBI Fermentas) as reference proteins. All protein bands were stained with Coomassie Blue for visualization. The molecular mass of the native enzyme was determined by gel-filtration chromatography on a Sephacryl S-300 preparative-grade column HR 16/60 (GE Healthcare). The purified enzyme was activated by adding benzonitrile as substrate for 24 h. The enzyme solution was analyzed by SDS/PAGE with Coomassie Blue staining. The molecular mass of the native enzyme was calculated by comparing with the migration length of reference proteins.

Measurement of activity and determination of kinetic parameters of wild-type and mutant enzymes

The activity of nitrilase was measured by assaying the release of ammonia as described previously [17,18]. The specific activity was determined after the enzymatic reaction was performed in 50 mM potassium phosphate buffer (pH 7.5) containing 1 mM substrate and 10% (v/v) methanol at 25°C for 20 min. In the kinetic study, various amounts of nitriles (0.1–0.5 mM) were incubated in 50 mM potassium phosphate buffer (pH 7.5) containing nitrilase enzyme and 10% (v/v) methanol at 25°C for 20 min. The reaction was stopped by adding HCl to the reaction mixture at a final concentration of 200 mM. The kinetic parameters were determined by fitting the data to the Michaelis–Menten equation.

Modelling

The three-dimensional structure of R. rhodochrous ATCC 33278 nitrilase was generated using the Accelrys Discovery Studio Modeller [19]. The aim of our comparative modelling was to generate the most probable structure of the query protein through alignment with template sequences that simultaneously satisfied spatial restraints and local molecular geometry. A BLAST search of the Protein Databank demonstrated strongest similarity between R. rhodochrous ATCC 33278 nitrilase and two N-carbamyl-D-amino acid amidohydrolase PDB entries, 1ERZ and 1FO6. These similar proteins were used as templates for modelling, after alignment with the ClustalW program [20]. The structure generated was improved further by refinement of the loop conformations by assessing the compatibility of their amino acid sequences with known PDB structures using the Protein Health module in Discovery Studio. All simulation experiments were carried out using a HP XW6200 Workstation with dual Intel Xeon 3.2 GHz processors.

Nitriles were docked in the models of wild-type and mutant nitrilases using the Suresurf docking program (Tripos) [21]. The active site was defined as a collection of amino acid residues enclosed within a sphere of 4.5 Å (1 Å = 0.1 nm) radius centred on the catalytic triad [11]. Each docking run consisted of 100 independent docks with 1000 iteration cycles. A random start was used to generate the substrate position within the docking box. The substrate orientation giving the lowest interaction energy was chosen for additional rounds of docking.

RESULTS

Cloning and molecular mass of nitrilase

The nitrilase gene was obtained from chromosomal DNA of R. rhodochrous ATCC 33278 using primers designed from the published genomic DNA sequence of R. rhodochrous J1 (GenBank® accession number D11425). The cloned nitrilase gene contained a sequence of 1101 bp encoding a 366-amino-acid protein and was the same as that of the published R. rhodochrous J1-A6 sequence (GenBank® accession number EF467367).

The subunit molecular masses of the purified wild-type and mutant enzymes in SDS/PAGE were approx. 40 kDa. They were similar to the calculated value of 41338 Da based on the 366-residue protein and six histidine residues, as determined with the Compute pI/Mw tool [22]. After the purified enzyme was activated by adding benzonitrile as substrate for 24 h, the active form of the enzyme with substrate eluted as a peak between ferritin (440 kDa) and thyroglobulin (669 kDa), corresponding to a molecular mass...
A determinant residue in nitrilase of substrate specificity

Figure 1 Determination of molecular mass of nitrilase from *R. rhodochrous* ATCC 33278 by gel-filtration chromatography

The reference proteins were thyroglobulin (669 kDa), ferritin (440 kDa), aldolase (158 kDa) and conalbumin (75 kDa). Nitrilase eluted at a position corresponding to 497 kDa. Results are means ± S.D. for three experiments.

of 496 kDa (Figure 1). These results indicate that the enzyme migrates as a dodecamer in gel filtration and thus may also be active as a dodecamer in solution.

Selection of the individually conserved residues for aliphatic and aromatic nitriles by sequence alignment

Sequence data for the nitrilases were obtained using the ExPASy proteomics server. Sequence alignment was performed with aliphatic nitrilases (EC 3.5.5.7) of *R. rhodochrous* ATCC 33278, J1 and K22, with aromatic nitrilases (EC 3.5.5.1) from *A. thaliana* (NIT1, NIT2, NIT3 and NIT4), *Nicotiana tabacum* (NIT4), and *Bacillus* sp., and with N-carbamyl-D-amino acid amidohydrolase from *Agrobacterium* as a template for modelling using the ClustalW program (Figure 2). All nitrilases displayed a conserved catalytic triad consisting of Glu-48, Cys-165 and Lys-131 of *R. rhodochrous* J1, as well as the individually conserved amino acid residues Lys-9, Leu-79, Arg-129, Tyr-142 and Tyr-175 for aliphatic and aromatic nitrilases. These conserved residues were selected as candidate determinant residues for substrate specificity for aliphatic and aromatic nitriles.

Alanine substitutions of the individually conserved residues for aliphatic and aromatic nitriles and the catalytic triad

The aromatic nitrile substrates benzonitrile, *m*-tolunitrile and 2-cyanopyridine were selected on the basis of their functional group, and the aliphatic nitriles adiponitrile, glutaronitrile, sebaconitrile and succinonitrile were chosen on the basis of their side-chain carbon number. The chemical structures of the substrates are shown in Figure 3. Each individually conserved residue and the catalytic triad were sequentially replaced with alanine to produce nitrilase mutants. Enzyme activities towards the aromatic...
and aliphatic nitrile substrates were determined for the alanine-substituted mutants and were compared with those of wild-type nitrilase. The mutants with an alanine-replaced catalytic triad exhibited no activity towards any nitrile. Activities towards aromatic and/or aliphatic nitriles were displayed by the wild-type enzyme and all of the alanine-substituted mutant enzymes, with the exception of R129A. The kinetic parameters of the wild-type enzyme and the K9A, L20A and Y142A mutant enzymes are shown in Table 1, but the kinetic parameters of the Y175A mutant enzyme are not shown owing to low activity (less than ∼6 μmol/min per mg). The alanine substitution at position 129 completely abolished enzymatic activity towards all nitriles. Among various amino acids, only the positively charged amino acids (arginine, lysine or histidine) at position 129 had activity. We are currently investigating the role of positively charged amino acids at position 129. The Y142A mutant hydrolysed aromatic, but not aliphatic nitriles, indicating that the amino acid at position 142 might be crucial for determining substrate specificity.

**Site-directed mutagenesis at position 142**

The tyrosine residue at position 142 of the nitrilase enzyme was replaced with respective non-polar aliphatic amino acids, including glycine, alanine, valine, leucine and isoleucine. Although they exhibited no activity towards any of the tested aliphatic nitriles, all of the mutant enzymes, except the glycine-substituted mutant, displayed activities towards aromatic substrates, including benzoniitrile, m-tolunitrile and 2-cyanopyridine (Figure 4). The less hydrophobic 2-cyanopyridine (having a nitrogen atom on its ring) was a poorer substrate for the nitrilase. Thus the enzymatic reaction of nitrilase probably involves hydrophobic interactions between the enzyme and a substrate. Among the mutants, the Y142A mutant showed the highest

### Table 1 Kinetic parameters of wild-type and alanine-substituted mutant enzymes of the individually conserved residues for aliphatic and aromatic nitrilases

| Substrate | Enzyme | $K_m$ (μM) | $k_{cat}$ (min$^{-1} \times 10^{-4}$) | $k_{cat}/K_m$ (min$^{-1} \cdot M^{-1} \times 10^{-2}$) |
|-----------|--------|------------|----------------------------------|------------------------------------------|
| **Aromatic nitriles** | | | | |
| Benzonitrile | Wild-type | 1070 ± 10 | 94 ± 1 | 880 ± 20 |
| K9A | 1020 ± 10 | 94 ± 1 | 920 ± 10 |
| L79A | 910 ± 20 | 126 ± 2 | 1380 ± 50 |
| Y142A | 1580 ± 150 | 173 ± 8 | 1090 ± 150 |
| m-Tolunitrile | Wild-type | 1850 ± 140 | 143 ± 12 | 770 ± 120 |
| K9A | 1620 ± 20 | 105 ± 3 | 650 ± 30 |
| L79A | 1790 ± 10 | 166 ± 2 | 930 ± 20 |
| Y142A | 612 ± 10 | 113 ± 6 | 1850 ± 130 |
| 2-Cyanopyridine | Wild-type | 1550 ± 30 | 106 ± 1 | 690 ± 20 |
| K9A | 1250 ± 10 | 113 ± 3 | 900 ± 30 |
| L79A | 1500 ± 20 | 151 ± 6 | 1010 ± 50 |
| Y142A | 819 ± 10 | 185 ± 1 | 2260 ± 40 |
| **Aliphatic nitriles** | | | | |
| Glutaronitrile | Wild-type | 1240 ± 50 | 173 ± 3 | 1400 ± 80 |
| K9A | 1190 ± 10 | 196 ± 2 | 1650 ± 30 |
| L79A | 1150 ± 20 | 189 ± 3 | 1630 ± 50 |
| Y142A | – | – | – |
| Adiponitrile | Wild-type | 933 ± 100 | 276 ± 24 | 2960 ± 570 |
| K9A | 851 ± 20 | 343 ± 3 | 4040 ± 130 |
| L79A | 921 ± 10 | 298 ± 3 | 3240 ± 70 |
| Y142A | – | – | – |

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activity towards aromatic substrates. As the molecular size of the non-polar aliphatic amino acid side chain increased, the activities of the mutants containing non-polar aliphatic residues, except glycine, towards aromatic substrates decreased. The far-UV CD spectra of the Y142G mutant and the wild-type enzyme were similar (results not shown); nevertheless, this mutant, which contained the smallest residue (glycine), showed no activity.

When Tyr-142 was replaced with a charged polar amino acid such as asparagine, arginine, aspartate or glutamate, the mutants exhibited no activity towards any tested substrate. The Y142S mutant, however, displayed activity towards aromatic, but not aliphatic, nitriles, showing substrate specificity similar to that of the mutants containing non-polar aliphatic amino acids. When Tyr-142 was replaced with the aromatic amino acid phenylalanine, the mutant retained activity towards both aromatic and aliphatic nitriles.

### Kinetic analysis of wild-type and mutant enzymes

The kinetic parameters of the purified wild-type and mutant enzymes at position 142 were determined for the hydrolysis of aromatic and aliphatic substrates (Table 2). The kinetic parameters of the wild-type and mutant enzymes for the hydrolysis of succinonitrile and sebaconitrile, and the Y142L and Y142I mutants for aromatic nitriles were not determined because of their low specific activities (less than ∼6 μmol/min per mg). Among the mutants containing non-polar aliphatic residues at position 142, the Y142A mutant showed the highest kcat values for aromatic nitriles such as benzonitrile, m-tolunitrile and 2-cyanopyridine. Compared with the wild-type enzyme, the Y142A mutant showed slightly lower kcat values for aromatic nitriles such as benzonitrile, m-tolunitrile and 2-cyanopyridine. The far-UV CD spectra of the Y142G mutant and the wild-type enzyme were similar (results not shown); nevertheless, this mutant, which contained the smallest residue (glycine), showed no activity.

**DISCUSSION**

A homology model of nitrilase from *R. rhodochrous* ATCC 33 278 was constructed based on the crystal structure of templates of two N-carbamyl-D-amino acid amidohydrolase PDB entries (1ERZ
and 1FO6). Although the level of sequence identity between nitrilase and template was relatively low (\(\sim 15.3 \% \) identity and 38.6\% similarity), the catalytic triads (Glu-48, Cys-165 and Lys-131) of \( \textit{R. rhodochrous} \) nitrilase and the \( \textit{N-carbamyl-D-} \) 
amino acid amidohydrolase of the same nitrilase superfamily were completely conserved as shown in Figure 2. The alanine substitution of the catalytic triad of \( \textit{R. rhodochrous} \) nitrilase exhibited no activity towards any nitrile (Table 1), suggesting that the catalytic triad was involved in the active site. Moreover, the activities of the mutants at position 142 for various substrates indicated that the position was closely related to substrate specificity. Thus the active-site structure of the homology model, including the catalytic triad and position 142, can be used in docking studies.

The molecular docking studies of \( \textit{R. rhodochrous} \) ATCC 33 278 nitrilase show substrate interactions in the active site. In the catalytic triad, Cys-165 attacks the cyano group of nitriles as a nucleophile, Glu-48 assumes the role of a general base, and Lys-131 is involved in stabilization of a tetrahedral intermediate structure \([10]\). Position 142 is located at one side of the catalytic triad (Figure 5). When the aromatic benzonitrile was docked in the active site, the wild-type nitrilase showed a slightly longer distance between the cyano group of benzonitrile and the sulfur of Cys (3.24 Å), compared with that of the Y142A mutant (2.94 Å) (Figures 5A and 5B). The distances seem to explain the slightly higher \( k_{\text{cat}} \) values of the Y142A mutant for aromatic substrates. The order of the specific activities of the mutants containing non-polar aliphatic residues was Y142A > Y142V > Y142L > Y142I, in increasing order of side-chain size (Figure 5). Thus an increase in the size of the side chain on the amino acid at position 142 caused a decrease in the \( k_{\text{cat}} \) values of the mutants for aromatic nitriles. This may result from increased steric hindrance between the aromatic substrate and the aliphatic residue owing to the increased side-chain size. The model in Figure 5(B) shows that the steric hindrance with each respective mutant correlates with the increase in the size of the non-polar aliphatic residue at position 142. In the active site of the wild-type enzyme, the distance between glutaronitrile, which is an aliphatic substrate, and cysteine (3.03 Å) was shorter than that between benzonitrile and cysteine (3.24 Å) (Figures 5A and 5C). This difference in distance may explain the slightly higher \( k_{\text{cat}} \) for glutaronitrile than for benzonitrile of the wild-type enzyme. However, obtaining structures of the enzyme in complex with these substrates would be necessary to provide further evidence for this conclusion.

The wild-type enzyme exhibited activity towards both classes of aromatic and aliphatic substrates, whereas the mutants containing a non-polar aliphatic residue at position 142 and the Y142S mutant had activities towards aromatic, but not aliphatic, substrates. The loss of specificity for aliphatic nitriles in the mutant enzymes suggests that the aromatic side-chain on the residue at position 142 is important for nitrilase hydrolysis of aliphatic substrates. When Tyr-142 was replaced with phenylaniline, the resultant Y142F mutant retained activities towards both aromatic and aliphatic nitrile substrates, with substrate specificity similar to that of the wild-type enzyme. This suggests that the aromatic rings of tyrosine and phenylaniline at position 142 are involved in nitrilase-catalysed binding and electron transfer for aliphatic substrates. This phenomenon can be explained by electron transfer between enzyme and substrate via a conjugated system \([23]\). We propose that the conjugated \( \pi \)-electron system of the aromatic substrate serves as an electron acceptor, whereas the aliphatic substrate has no conjugated system. As a result, the mutants containing aliphatic residues at position 142 had no activity towards aliphatic substrates owing to their lack of an aromatic ring to provide an electron acceptor. This reinforces our hypothesis further that an aromatic ring at position 142 is used as an electron acceptor during the catalytic reaction between nitrilase and aliphatic nitriles.

The mutants containing charged amino acids such as aspartate, glutamate, arginine and asparagine at position 142 showed no increased activity towards any substrate, probably because the charged amino acids disrupt hydrophobic interactions with the substrate. These results indicate the importance of hydrophobic interactions in the recognition of nitrile substrates. The glycine residue at position 142 would not be suitable for hydrophobic interaction with any substrate, and, consequently, the Y142G mutant displayed no activity towards aromatic or aliphatic substrates. To confirm our molecular modelling and structural analyses, the crystal structure of nitrilase should be determined.

In the present study, the amino acid at position 142 of nitrilase derived from \( \textit{R. rhodochrous} \) ATCC 33 278 was identified as a determinant residue of substrate specificity based on the following results: (i) the mutants containing aromatic amino acids...
retained activity towards both aliphatic and aromatic nitriles; (ii) the mutants containing non-polar aliphatic amino acids showed activity towards aromatic, but not aliphatic, nitriles; (iii) for the mutants containing non-polar aliphatic residues, the activity towards aromatic substrates decreased with increasing size of the amino acid side chain; and (iv) the mutants containing charged amino acids had no activity towards any nitrile. These data enable a more detailed understanding of the roles of selected active site residues in the catalytic process of the \textit{R. rhodochrous} nitrilase.

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