Aerobic Degradation of N-Methyl-4-Nitroaniline (MNA) by Pseudomonas sp. Strain FK357 Isolated from Soil

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Abstract

N-Methyl-4-nitroaniline (MNA) is used as an additive to lower the melting temperature of energetic materials in the synthesis of insensitive explosives. Although the biotransformation of MNA under anaerobic condition has been reported, its aerobic microbial degradation has not been documented yet. A soil microcosms study showed the efficient aerobic degradation of MNA by the inhabitant soil microorganisms. An aerobic bacterium, Pseudomonas sp. strain FK357, able to utilize MNA as the sole carbon, nitrogen, and energy source, was isolated from soil microcosms. HPLC and GC-MS analysis of the samples obtained from growth and resting cell studies showed the formation of 4-nitroaniline (4-NA), 4-aminophenol (4-AP), and 1, 2, 4-benzenetriol (BT) as major metabolic intermediates in the MNA degradation pathway. Enzymatic assay carried out on cell-free lysates of MNA grown cells confirmed N-demethylation reaction is the first step of MNA degradation with the formation of 4-NA and formaldehyde products. Flavin-dependent transformation of 4-NA to 4-AP in cell extracts demonstrated that the second step of MNA degradation is a monoxygenation. Furthermore, conversion of 4-AP to BT by MNA grown cells indicates the involvement of oxidative deamination (release of NH₂ substituent) reaction in third step of MNA degradation. Subsequent degradation of BT occurs by the action of benzenetriol 1, 2-dioxygenase as reported for the degradation of 4-nitrophenol. This is the first report on aerobic degradation of MNA by a single bacterium along with elucidation of metabolic pathway.

Introduction

N-Methyl-4-nitroaniline (C₇H₈N₂O₂, MNA) is a nitroaromatic compound, used as an additive for lowering the melting temperature of highly energetic compounds such as 2, 4-dinitroanisole (DNAN), that helps in easier synthesis of insensitive explosives [1,2,3]. During the process of manufacturing, packaging, and handling of insensitive explosives, it may get released into the environment through various waste streams [2]. Presence of electron withdrawing nitro group (–NO₂) in the benzene ring results in acute toxicity to human and other life forms [2,4]. As a result of toxicity and recalcitrant nature of nitroaromatic compounds, public concern has prompted the need to develop clean-up technologies for restoration of environments. Microbial degradation/or transformation could be used as an effective mechanism for the elimination of nitroaromatic compounds from the environment. Microbial degradation of nitroaromatic compounds can take place in both aerobic and anaerobic conditions, however, due to electron withdrawing effect of nitro group, only nitro group is reduced sequentially into aryl nitroso (Ar-NO), aryl hydroxylamine (Ar-NHOH), and aryl amine (Ar-NH₂) without any aromatic ring cleavage [5,6]. These reductive transformations have been shown to be slow and the reductively transformed products are known to be carcinogenic [7,8,9]. The oxidative degradation of recalcitrant compounds under aerobic conditions is expected to be one of the most effective mechanisms for rapid and complete mineralization, which occurs by non-specific enzymes [10,11,12]. There are few reports on biological transformation of MNA in anaerobic fluidized-bed bioreactors, where MNA was reductively transformed into 4-methoxy-phenylenediamine that gets accumulated as dead end product followed by dimer formation [1,2]. Till date there is no information available on aerobic transformation/or degradation of MNA by means of bacterial isolates. Elucidation of catabolic pathways involved in degradation of xenobiotic compounds is critical for both ‘basic understanding’ and ‘development of remediation technologies’ [13,14,15]. Therefore, studies on isolation of MNA degrading aerobic bacteria and elucidation of corresponding metabolic pathways are of great ecological significance. In this communication we report isolation of a bacterial strain Pseudomonas sp. strain FK357, isolated from soil that utilizes MNA as the sole source of carbon, nitrogen, and energy. The degradation of MNA by strain FK357 occurs via the formation of 4-NA, 4-AP, and BT as the major metabolic intermediates. We propose that strain FK357 could be used as a model system for studying the molecular and biochemical mechanism of aerobic degradation of MNA as well as for the development of MNA bioremediation technology.
Materials and Methods

Chemicals and Growth Media

N-Methyl-4-nitroaniline (MNA), 4-nitroaniline (4-NA), 4-aminophenol (4-AP), and benzenetriol (BT) were purchased from Sigma-Aldrich (St, Louis, MO, USA). Minimal salt medium (MSM) used in the present study was prepared as described earlier [16] with slight modification i.e. absence of nitrogen source [(NH4)2SO4]. Stock solution (10 mM) of MNA prepared in HPLC grade methanol was added to an empty Erlenmeyer flask to obtain the working concentrations. Further, the residual methanol in the flask was evaporated under stream of air to leave the dry crystal of MNA in the bottom of flask. Nutrient agar (NA) and nutrient broth (NB) both at one-quarter strength (1/4th) were used as rich media for bacterial growth and culture maintenance.

Soil Microcosms Study

The non-polluted soil sample was collected from the lawn of Institute of Microbial Technology, Chandigarh, India. The pH of the soil was 5.9, and it consisted of 8.2% moisture, 3.5% total organic carbon and 1.7% total nitrogen content, respectively. The soil samples were sieved through 2 mm mesh to remove stones and debris. Soil microcosms experiment was carried out as described previously with slight modification [17]. Subsequently, 5 gm soil sample was suspended in 50 ml MSM in 250 ml Erlenmeyer flask supplemented with 100 μM MNA. Further, the above microcosms were each supplemented with or without (i) 10 mM glucose and 5 mM succinate as the sole carbon source, or (ii) 0.8 g L−1 NH4Cl as the sole nitrogen source, or (iii) 10 mM glucose, 5 mM succinate and 0.8 g L−1 NH4Cl as the sole carbon and nitrogen source, respectively. A control was also set up containing 5 gm soil inoculated with (1%, v/v) seed culture grown in 1/4-NB. The flasks were incubated in dark at 30°C under shaking condition (150 rpm). At regular time intervals, samples were withdrawn from the test and control flasks and analyzed for the disappearance of MNA. All the experiments were carried out in triplicates.

Isolation and Characterization of MNA Degrading Bacteria

Upon complete depletion of MNA in the soil microcosms, the serially diluted soil slurries was spread plated on selective medium (MSM-agar containing 700 μM MNA) and incubated at 30°C for one week. The bacterial colonies appeared on the plate were selected for characterization and were tested for MNA degradation. The selected bacterial isolates were used for primary screening by inoculating in 10 ml vials containing 3 ml carbon-free MSM supplemented with different concentrations (50–700 μM) of MNA. The positive bacterial isolates were selected on the basis of increase in growth accompanied by substrate depletion and release of ammonia (NH4+) and nitrite (NO2−) ions as measured colorimetrically at wavelength of 340 and 540 nm, respectively. Purity of the positive isolates was checked by streaking on 1/4-NA plates. Further, the efficient MNA degrading isolate was identified using polyphasic taxonomy and 16S rRNA gene sequencing [17]. The 16S rRNA gene sequence (1431 base pairs) of strain FK357 was compared to those of the type strains of Pseudomonas using the BLAST search.

Growth Studies and Degradation of MNA by Strain FK357

The growth studies of strain FK357 and degradation of MNA were performed in 50 ml carbon-free MSM supplemented with varying concentrations of MNA ranging from 50 to 700 μM by inoculating (1%, v/v) seed culture grown in 1/4-NB. The flasks were incubated at 30°C under shaking condition of 200 rpm. At every 8 hours, culture was withdrawn and optical cell density at 600 nm (OD600 nm) was measured using Lambda EZ 201 UV-visible spectrophotometer (Perkin-Elmer Inc, USA). The bacterial growth was also monitored by measuring the total protein of the culture with the help of Pierce BCA protein assay kit (Thermo Scientific, USA). Culture fluid samples (2.0 ml) were centrifuged at 8,000 × g for 10 min to obtain cell-free supernatants which were used for analysis of the amount of NH4+ and NO2− released. Subsequently, the above supernatants were also used for the quantitative determination of MNA disappearance and identification of metabolic intermediates by HPLC. Non-inoculated and inoculated flasks with heat killed cells of strain FK357 were used as abiotic and negative controls, respectively. Another control experiment was also conducted by inoculating strain FK357 in carbon-free MSM (not supplemented with MNA).

Resting Cell Studies

Resting cell studies on MNA degradation were carried out according to the method described elsewhere [18]. The overnight 1/4-NB grown seed culture (6%, v/v) of strain FK357 was inoculated into 1.0 L of 1/4-NB supplemented with MNA (150 μM) and incubated at 30°C under shaking at 200 rpm up to 24 hours. Similarly, to obtain un-induced cells, the strain FK357 was grown in 1/4-NB only. When the optical cell density of the culture reached to OD600 ranging between 1.2–1.5, the cells (induced and un-induced) were harvested by centrifugation at 8,000 g at 4°C for 10 min, washed twice with phosphate buffer (20 mM, pH 7.2) and suspended in 100 ml carbon-free MSM. In 25 ml cell suspension 150 μM each of MNA and 4-NA were supplemented separately. Similarly, the degradation of MNA and 4-NA by un-induced cells were also carried out by suspending the un-induced cells in 25 ml MSM supplemented with 150 μM each of MNA and 4-NA, separately. Non-inoculated and inoculated flasks with heat killed cells were used as abiotic and negative controls, respectively. Each flask was incubated at 30°C with shaking at 150 rpm. Samples (2.0 ml) were withdrawn from both control and experimental flasks at regular time intervals of 2 hours and were analyzed for the amount of NO2−, and NH4+ released, followed by High Performance Liquid Chromatography (HPLC) and Gas-Chromatography Mass-Spectroscopy (GC-MS) analysis (methods described later).

Enzyme Assays with Cell-free Lysates

MNA-induced cells of strain FK357 were harvested by centrifugation and washed twice with phosphate buffer (20 mM, pH 7.2) and re-suspended in phosphate buffer. The cell suspensions lysed by passages through a French pressure cell (20,000 lb/in2) were centrifuged at 12,000 rpm for 30 min at 4°C and supernatant was separated to obtain cell-free enzyme extract, which was subsequently used for the enzyme assays. Protein content within cell-free extracts was determined with Pierce BCA protein assay kit (Thermo Scientific, USA). The cell-free extract was used for determining activities of N-demethylase, monoxygenase, and 1, 2, 4-benzenetriol 1, 2-dioxygenase, respectively as described below.

Enzyme assay for N-demethylase. Demethylation reaction i.e. removal of N-methyl group from MNA in cell-free lysates was carried out according to the method of Summers et al. [19]. The reaction was carried out in a total volume of 10 ml phosphate buffer (50 mM, pH 7.5) containing cell-free protein (2.0 mg ml−1) 150 μM NADH, 25 mM Fe (NH4)2SO4, 6 H2O and 150 μM MNA. The enzymatic reaction was initiated by addition of 150 μM MNA and incubated at 30°C under shaking (200 rpm). The control reactions lacking either substrate or cell-free protein
were also used during the above reaction. The N-demethylase activity was determined by measuring time dependent depletion of substrate and formation of N-demethylated product by HPLC. The formation of formaldehyde (HCHO) in the demethylation reaction was quantitatively determined by using an assay based on the Hantzsch reaction [20]. Briefly, in 0.5 ml of culture supernatant equal volume of reagent B (2 M ammonium acetate, 0.05 M acetic acid and 0.02 M acetylacetone) was added and incubated at 38°C for 5 min. Presence of HCHO in the sample was determined colorimetrically by calculating the absorbance at 412 nm as the HCHO adduct dicaetethylhydrolutidine.

**Enzyme assay for monoxygenase.** The monoxygenase mediated conversion of 4-NA into 4-AP was carried out in cell-free lysates prepared from MNA grown cells of strain FK357 as described previously [21]. The reaction was carried out in a total volume of 10 ml phosphate buffer (20 mM, pH 7.0) containing cell-free protein (2.0 mg ml⁻¹) 200 μM NADPH, 150 μM FMN. The reaction was initiated by addition of 100 μM 4-NA and incubated at 28°C. The control reactions lacking either substrate or cell-free protein or cofactors (NADPH or FMN) were also used during the above reaction. The monoxygenase activity was determined by measuring time dependent depletion of substrate and formation of product by HPLC. Similarly, qualitative determination of NO₂⁻ ions released in the above reaction was also estimated colorimetrically.

**Aniline dioxygenase assay.** The dioxygenation reaction involved in the conversion of 4-AP to BT by strain FK357 was measured with an oxygen electrode (YSI, Ohio, USA), according to the method as described previously [22,23]. MNA-induced and un-induced cells (as described in the resting cell study) were harvested by centrifugation at 7,500xg at 4°C for 15 min. Pellets were washed twice with phosphate buffer (20 mM, pH 7.2) and re-suspended in the same buffer. This suspension was used for the assay of dioxygenase by measuring oxygen uptake at 30°C. The reaction was carried out in 1.85 ml volume air-saturated phosphate buffer (20 mM, pH 7.2) containing substrates (70 μM), and cells (0.25 mg of protein).

1, 2, 4-Benz(en)etriol 1, 2-dioxygenase enzyme assay. It is known that the degradation of terminal intermediate 1, 2, 4-benz(en)etriol (BT) starts by the action of 1, 2, 4-benz(en)etriol 1, 2-dioxygenase with the formation of lower pathway intermediate i.e. maleylacetate (MA) [24]. 1, 2, 4-Benz(en)etriol 1, 2-dioxygenase activity was determined spectrophotometrically using Lambda EZ 201 UV-visible spectrophotometer (Perkin-Elmer Inc, Massachusetts, USA) [21]. The reaction mixture consisted of 50 mM sodium phosphate buffer (pH 7.0), 10 mM Fe (NH₄)₂(SO₄)₂.6 H₂O, and 0.5 mg soluble protein of cell-free lysates, in the total volume reaction of 1.0 ml. Reaction was initiated by the addition of 70 μM of BT to the reaction mixture. Enzyme activity was monitored with wavelength scan over a range of 220–360 nm at an interval of 1 min.

**Analytical Methods**
The release of nitrite ions (NO₂⁻) was monitored with a colorimetric method using N-(1-naphthyl) ethylene-diamine-dihydrochloride and sulfanilic acid reagent as described earlier [25]. The NO₂⁻ assay was carried out by mixing equal volume (0.1 ml) of reagent A [0.1% (w/v) sulfanilic acid in 30% (v/v) acetic acid] and reagent B [0.1% (w/v) naphthylethendiamine in 30% acetic acid] to 0.1 ml culture supernatant. Presence of NO₂⁻ in the sample was indicated by the appearance of purple colour and quantified by calculating the absorbance at 540 nm. Ammonia (NH₄⁺) concentrations were also monitored with a colorimetric method using ‘Ammonia Estimation Kit’ (Sigma Aldrich, USA) according to the manufacturers’ recommendation. Standard plots generated with known concentrations of (NH₄)₂SO₄ and NaNO₂ were used to determine the concentrations of NH₄⁺ and NO₂⁻ ions. The quantitative determination of MNA and its metabolic intermediates were analyzed by HPLC [21]. The samples collected from growth, resting cell studies and enzyme assays were analyzed by HPLC using Waters-HPLC 2489 model (Waters, USA) equipped with a UV detector and a C-18 reverse phase column at 30°C [21]. The mobile phase used for the separation of substrate and intermediates consisted of acetonitrile: water (30:70, v/v) under isocratic condition with the flow rate of 1.0 ml min⁻¹. The samples were injected with a constant injection volume of 20 μl. The peaks of eluents were monitored with UV detector over a wavelength scan of 220–290 nm. For the GC-MS analysis, samples were prepared by mixing equal volume of ethyl acetate to the cell-free aqueous culture and liquid-liquid extraction performed by layer separation sequentially at neutral and acidic pH. Extracted organic phase was pooled, dried under nitrogen flow using Rotavapor II (BUCHI, Switzerland). The samples were analyzed by GC-MS using QP2010S (Shimadzu Scientific Instruments, USA) as reported earlier [6]. Briefly, the chromatographic separation of substrate and intermediates was carried out with constant temperature of injector, oven, and detector at 280, 200, and 250°C, respectively. Positive molecular ion mass spectra were scanned in mass/charge (m/z) range of 0–160. Identity of different metabolic intermediates was ascertained by comparison of mass fragmentation pattern and subsequent mass spectral database match from the National Institute of Standards and Technology library (NIST).

**Nucleotide Sequence**
A total of 1431 base-pairs were sequenced for 16S rRNA gene of strain FK357; which has been deposited to GenBank under the accession no. KF011496.

**Results**

**Soil Microcosms Study**
The rate of MNA degradation was slightly higher in nitrogen (NH₄Cl) amended soil microcosms as compared to unamended soil microcosms (Figure 1). The higher rate of MNA degradation in nitrogen-amended soil microcosms, suggests the availability of most favorable nitrogen source from NH₄Cl, which makes it easier for soil microorganisms to utilize N-methyl group as the carbon source. However, in case of soil microcosms amended with a combination of carbon and nitrogen, the rate of MNA degradation was completely inhibited. The inhibition of MNA degradation in the above soil microcosms was likely because the soil inhabitant microorganisms utilized more favorable carbon (succinate and glucose) and nitrogen (NH₄Cl) source (Figure 1). There was no transformation of MNA in the sterile soil (Figure 1).

**Isolation and Characterization of N-methyl-4-nitroaniline Degrading Bacteria**
After 15 days of incubation, when complete degradation of MNA occurred, the serially diluted soil slurries from microcosms was spread plated on MNA selective plates. Total 35 morphologically different isolates were selected and further screened for the degradation of MNA. Among these isolates, strain FK357 was found to be an efficient MNA degrader. The strain FK357 was selected for the identification and study of MNA degradation along with elucidation of metabolic pathways. Strain FK357 was identified as a facultative anaerobe, Gram-negative, non-spore forming, non-motile, catalase and oxidase positive, rod-shaped
Figure 1. Biotransformation of MNA in soil microcosms under aerobic conditions: (●), nitrogen-amended; (○), unamended; (▲), carbon-amended; (□), carbon and nitrogen-amended; (■), sterile. Values are presented as arithmetic mean of data obtained from experiments carried out in triplicate; error bars represent standard deviation. doi:10.1371/journal.pone.0075046.g001

bacterium. It showed optimal growth on nutrient rich medium (NA, NB) at 30°C; however it was also able to grow at 37°C. Strain FK357 was found negative for H2S production. This strain showed acid production from D-fructose, D-glucose, maltose, D-mannose, D-ribose, trehalose, and L-alanine, whereas, negative on glycerol, D-lactose, D-galactose, D-cellobiose, D-raffinose, and L-serine. Strain FK357 also utilizes acetate and succinate as sole carbon sources. The partial 16S rRNA gene sequence (1431 base pair) of strain FK357 showed 99% sequence similarity to that of several Pseudomonas putida strains. Thus, based on the biochemical tests, morphological and physiological characteristics, and phylogenetic analysis, strain FK357 was identified as Pseudomonas sp. strain FK357. The strain Pseudomonas sp. strain FK357 will be made available to other researchers upon reasonable request.

Growth Study and Degradation of N-methyl-4-nitroaniline by Strain FK357

A growth study of strain FK357 was carried out using different concentrations of MNA ranging from 50 to 700 μM. Growth of strain FK357 was completely abolished at an MNA concentration of 500 μM, whereas 250 μM of MNA was found to be optimal for its growth with subsequent degradation of the parent compound (Figure 2A and B). Strain FK357 grew best on 250 μM concentration of MNA at 30°C and pH 7.2. Degradation of MNA during the growth study occurred with the stoichiometric production of nitrite ions and lesser amount of NH4+ (Figure 3). The complete depletion of MNA was observed at 96 hours of incubation (Figure 5). Depletion of MNA was accompanied with an increase in growth of strain FK357 as determined by measuring the total cell proteins up to the value of 10.64 μg ml⁻¹. A transient accumulation of 4-NA occurred in the media during the degradation of MNA (Figure 3). We were unable to detect formaldehyde (HCHO) during the growth study, while transient accumulation of 4-NA, suggested that the first step of MNA degradation starts via the release of –CH3 group in the form of HCHO, followed by the removal of –NO2 and –NH2 groups, respectively. In growth media the accumulation of released NO2 was stoichiometrically up to the value of 249.56±0.17 μM, whereas, a slight accumulation of NH4+ [51.06 μM] was observed. Thus, non-stoichiometric release of NH4+ suggested that it might possibly be utilized as a preferential nitrogen source for the growth of strain FK357. The above results are in close agreement with the results reported earlier on the degradation of amino or nitro containing aromatic compounds by different genera of bacteria including Pseudomonas [21,26,27,28,29,30,31]. The growth yield of strain FK357 on MNA was found to be 0.55 g of cells/g of MNA. The rate of MNA degradation by strain FK357 was calculated to be 3.26±0.08 nmol MNA min⁻¹ mg of protein⁻¹. Strain FK357 was unable to grow in MSM in the absence of MNA. The above results indicated that the strain FK357 utilizes MNA as the sole source of carbon, nitrogen, and energy. Growth of strain FK357 on 4-NA was also checked. Strain FK357 also utilizes 4-NA and the degradation of 4-NA occurred via stoichiometric accumulation of NO2⁻ and slight accumulation of NH4⁺ in the media (data not shown). The growth yield of strain on 4-NA and the rate of 4-NA degradation came out to be 0.43 g of cells/g of 4-NA and 2.73±0.13 nmol 4-NA min⁻¹ mg of protein⁻¹, respectively. In growth study, we identified only 4-NA as an intermediate; further metabolic intermediates were not identified. Based on the accumulation of NO2⁻ and NH4⁺ in the growth media, it could be proposed that the subsequent degradation of 4-NA intermediate may initiate either direct 'oxidative deamination' of aromatic nucleus or oxidative denitration with the formation of 4-nitrophenol or 4-aminophenol as the putative metabolic product, respectively.

Resting Cell Studies

HPLC analysis of the samples from resting cell study, showed the formation of 4-NA, 4-AP, and BT intermediates as confirmed by comparison with authentic standards. The identified intermediates by HPLC were also confirmed by GC-MS analysis. The

Figure 2. Growth characteristics and degradation kinetics of Pseudomonas sp. strain FK357 in carbon-free MSM supplemented with different concentrations of MNA. (●), 50 μM; (▲), 150 μM; (Δ), 250 μM; (○), 350 μM; (■), 500 μM; (□), 700 μM. (A) Growth of strain FK357 on different concentrations of MNA. (B) Degradation of MNA at different concentrations by strain FK357. doi:10.1371/journal.pone.0075046.g002
mass-fragmentation patterns of the above intermediates viz; 4-NA, 4-AP, and BT were matched with the known authentic standards (Figure 4). Based on the above identified intermediates, we hypothesized that the degradation of MNA occurs via the formation of 4-NA metabolite as a result of demethylation of N–CH$_3$ group, followed by oxidative removal of –NO$_2$ group with the formation of 4-AP. Subsequent degradation of 4-AP may occur by oxidative deamination of –NH$_2$ group with the formation of BT as a terminal intermediate.

The resting cell study showed that the MNA-induced cells eliminate the lag phase for the depletion of MNA as observed in case of un-induced cells (Figure 5A). Complete degradation of MNA by the induced cells occurred within 6 hours of incubation with the rate of 0.35±0.07 nmol MNA min$^{-1}$ mg of protein$^{-1}$, whereas, un-induced cells degraded MNA after 15 hours of incubation with the rate of 0.17±0.12 nmol MNA min$^{-1}$ mg of protein$^{-1}$. The above results showed that the rate of MNA degradation by induced cells was two times higher compared to the rate of degradation by un-induced cells. The degradation of MNA occurs with the appearance followed by disappearance of 4-NA, 4-AP, and BT intermediates (Figure 5B). The cells pre-exposed with MNA also degraded 4-NA with the formation 4-AP and BT as intermediates and the complete degradation of 4-NA occurred after 10 hours of incubation (data not shown).

N-Demethylase Activity (Removal of -CH$_3$)

NADH-dependent enzyme assay in the crude cell extract prepared from MNA grown cells catalyzed stoichiometric transformation of MNA to 4-NA (Figure 6A). The conversion of MNA to 4-NA occurred by N-demethylation reaction with the removal of –CH$_3$ group as determined by stoichiometric formation of HCHO (Figure 6A). The specific activity of N-demethylation reaction was found to be 12.33±0.15 nmol min$^{-1}$ mg of protein$^{-1}$. The identification of 4-NA product from MNA in the enzyme assay confirmed that the first step of MNA degradation is an N-demethylation reaction.

Monooxygenase Activity (Removal of -NO$_2$)

The subsequent enzymatic transformation of 4-NA, which was identified as the first metabolite, was also carried out in the above crude cell extract. In the presence of NADPH and FMN cofactor, the crude cell extract showed stoichiometric transformation of 4-NA to 4-AP along with stoichiometric production of NO$_2^-$ (Figure 6B). The specific activity for the conversion of 4-NA to 4-AP was determined to be 2.13±0.12 nmol min$^{-1}$ mg of protein$^{-1}$.
AP was found to be 3.55 ± 0.08 nmol min⁻¹ mg⁻¹ of protein. No monooxygenase activity was observed in the control reactions that lacked either NADPH or FMN or cell-free protein. The above results showed that the second step of MNA degradation is a flavin-dependent monooxygenation reaction.

Oxidative Deaminase Activity (Removal of –NH₂)

The rate of oxygen consumption by 4-AP, 4-chlorophenol (4-CP), and aniline in MNA grown cells of strain FK357 was higher as compared to the MNA and 4-NA (Table 1). However, negligible oxygen consumption was observed by 4-AP, 4-CP and aniline when 1/4-NB grown cells were incubated with these substrates suggesting the inducible nature of enzyme. In contrast, the lesser amount of oxygen consumption by MNA and 4-NA in 1/4-NB grown cells demonstrated that the pathway enzymes involved in the initiation of MNA degradation have moderate constitutive activities. The inducible nature of enzyme involved in the oxidative deamination of –NH₂ from aminophenols and anilines are in close agreement with the earlier reports [22,23,32,33]. The stoichiometric release of NH₄⁺ and formation of BT, chlorocatechol and catechol products from 4-AP, 4-CP and aniline suggested that the deamination of –NH₂ group is a dioxygenation reaction and also it is inducible in nature.

Discussion

Structurally MNA is an analogue to 4-nitroanisole, which contains O-methyl group (O-CH₃) and a nitro group (–NO₂), whereas MNA contains an N-methyl group (N-CH₃) and a nitro group (–NO₂). The mineralization of 4-nitroanisole has been earlier reported by *Rhodococcus* strain AS2 and AS3 [34]. The degradation of 4-nitroanisole starts with O-demethylation of O-CH₃ with the formation of 4-nitrophenol, which subsequently get degraded as a result of ring cleavage via the formation of 4-nitrocatechol intermediate [34]. Soil microcosms study demonstrated the possibility of MNA degradation in aerobic conditions. Nitrogen (NH₄Cl) amended soil microcosms showed the higher rate of MNA degradation as compared to that of unamended soil microcosms. The above result indicates that soil microbes utilizes...
Table 1. Oxygen uptake by MNA-grown and 1/4-NB grown cells.

| Substrates                  | Oxygen uptake (nmol O2/min/mg of protein) by cells grown in: |
|-----------------------------|-------------------------------------------------------------|
|                            | MNA                                         | 1/4-NB                                      |
| N-Methyl-4-nitroaniline     | 65.22±0.7                                     | 33.19±0.1                                  |
| 4-Nitroaniline              | 79.50±1.2                                     | 28.70±1.1                                  |
| 4-aminophenol               | 184.74±0.5                                    | 0.27±0.7                                   |
| 4-chloroanilinopropenol     | 173.34±1.3                                    | 0.78±0.4                                   |
| Aniline                     | 190.10±2.1                                    | 0.50±0.1                                   |

*The reaction was carried in 1.85 ml volume air-saturated phosphate buffer (20 mM, pH 7.2) containing substrates (70 μM), and cells (0.25 mg of protein). Data represents means of at least three separate experiments.

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$\text{N-CH}_3$ group from MNA as the preferential single carbon source i.e. the easiest and first to be used, in the presence of most favourable nitrogen source ($\text{NH}_3\cdot\text{Cl}$). Soil microcosms study suggested that microorganisms utilized MNA as sole source of carbon, nitrogen, and energy. Based on the identified intermediates during the growth study, resting cell study and enzyme assays, we proposed a novel MNA degradation pathway in *Pseudomonas* sp. strain FK357 (Figure 8). In this pathway, conversion of MNA into 4-NA is expected to be catalysed by a putative demethylase enzyme capable of removing $\text{N-CH}_3$ via oxidative pathway. O-demethylation reactions are widely distributed among microorganisms capable of degrading aromatic compounds containing $\text{O-CH}_3$ group, the removal of $\text{N-CH}_3$ group from aromatic compounds has been shown only in few reports. *Pseudomonas putida* CBB5 has been isolated and characterized for enzyme capable of removing $\text{N-CH}_3$ group from aromatic compounds oxidized to HCHO [19,35,36,37,38,39,40,41]. There are various bacterial strains which have been characterized to utilize HCHO as the sole carbon source [20,33,42,43,44]. In the presence of nitrogen source, the strain FK357 also utilized HCHO as the sole carbon source (data not shown). Summers et al. [19] characterized $\text{N}$-demethylase ($\text{Ndm}$) gene from *Pseudomonas putida* CBB5, which showed the broad substrate activity on several purine alkaloids. Thus, based on the above study, we proposed the involvement of a similar bacterial $\text{N}$-demethylation reaction as the first step of MNA degradation. Subsequent degradation of MNA presumably initiated as a result of removal of substituted –NO$_2$ or –NH$_2$ group from the benzene ring. The mechanism for the removal of –NO$_2$ and –NH$_2$ groups from benzene ring poses an interesting question and challenge to the microbial systems for the preferential removal of these groups. Usually there are two reaction mechanisms for the removal of –NO$_2$ group, one is oxidative and another is reductive [5]. In the oxidative reaction, the removal of –NO$_2$ group as nitrite ions ($\text{NO}_2^-$) takes place by hydroxylation, while –NO$_2$ group get sequentially reduced to nitroso (–NO), hydroxylamine (–NHOH), and amino (–NH$_2$) group in the reductive reaction [6,45,46]. Another reaction mechanism involves dioxygenation which removes –NO$_2$ group by adding two hydroxyl groups simultaneously, one at the nitro-substituted position and the other at the adjacent position [5,47]. Removal of –NO$_2$ group by dioxygenation reaction has been reported earlier in the degradation pathway of nitrobenzene by *Comamonas* sp. strain JS763 [5,48]. The dioxygenation of –NO$_2$ group from nitrobenzene resulted in the formation of 1, 2-cis-dihydrodiol product [47,49]. Similarly, the removal of –NH$_2$ group from the benzene ring by dioxygenation reaction has also been reported for the degradation of anilines, diphenylamine, and chloroanilines [22,23,32,50]. Aniline dioxygenase is one of the most commonly characterized enzyme reported for the removal of –NH$_2$ group during the aniline and chloroaniline degradation pathways [22,23,27,33,50,51].

The crude cell extracts showed conversion of 4-NA into 4-AP with stoichiometric production of NO$_2^-$ and the removal of –NO$_2$ from 4-NA occurs via a hydroxylation reaction catalyzed by flavin-dependent monooxygenase (Figure 6A). This reaction is very common for aerobic microbial degradation of nitro or chloro containing aromatic compounds such as 2-chloro-4-nitrophenol, 2-chloro-4-nitroaniline, 4-nitroaniline, 4-nitrophenol, 2,4-dichlorophenol, and 4-chlorophenol, respectively [6,21,33,52,53]. Oxygen uptake studies showed the conversion of 4-AP into BT product. 

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with stoichiometric release of NH$_4^+$, suggesting that a dioxygenase enzyme catalyses the reaction by introducing two oxygen atoms on the benzene ring. Similar reports have also been shown in the case of aniline degradation by strain *Pseudomonas putida* mt-2, *Pseudomonas acidovorans* CA28, *Delftia* sp. AN3, and *Delftia tsuruhatensis* AD9 which degrades aniline via the formation of catechol as the first metabolic product [22,23,29,54]. *Moraxella* sp. strain G is known to mineralize 4-chloroaniline by dioxygenation reaction with the formation of chlorocatechol as a metabolic product along with stoichiometric release of NH$_4^+$ [55]. Furthermore, the crude cell extracts prepared from MNA grown cells also transforms BT to MA, suggesting the involvement of a ring cleavage enzyme i.e. 1, 2, 4-benzenetriol 1, 2-dioxygenase. Formation of MA from BT is commonly reported as a ring cleavage step in 4-nitrophenol degradation pathway [56,57,58]. Based on the above identified metabolites, the degradation pathway has been elucidated, showing N-demethylation of N-CH$_3$ group as the first reaction followed by monooxygenase mediated removal of –NO$_2$ (Figure 8). Further degradation of 4-AP occurs by the dioxygenation reaction resulting in formation of BT which subsequently gets ring cleaved by 1, 2, 4-benzenetriol 1, 2-dioxygenase enzyme.

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