ATRAZINE DEGRADATION BY Bacillus megaterium ISOLATED FROM AGRICULTURAL SOIL<br>El-Sawah, M.M.A. ${ }^{1}$; Samia M.M. Bauoymy ${ }^{1}$; Eman H. Ashour ${ }^{1}$; Lobna A. Moussa ${ }^{2}$ and Samah A. H.Shady ${ }^{1}$<br>${ }^{1}$ Microbiol. Dept., Fac. Agric., Mansoura Univ., Mansoura, Egypt.<br>${ }^{2}$ Soil, Water and Environ. Res. Instit., Agric. Res. Center (ARC), Giza.


#### Abstract

In present study the capability of some microorganisms isolated from Egyptian soil to a triazine herbicide, atriazine, decomposition was assessed. Nine atrazine-degrading bacteria were isolated from soil that received repeated exposures of the commonly used herbicides atrazine. These isolates were belonged to genera Basilllus, Pseudomonas and Micrococcus. One isolate showed good growth and clearing zone on mineral salts agar media amended with atrazine (at 500 ppm ) as a carbon and/or nitrogen source. This most active strain involved in atrazine degradation were selected and identified. The strain was classified as Bacillus megaterium. It degraded $45.8 \%$ of initial concentration of atrazine concurrent with increasing the population size from $10^{5}$ to $10^{8} \mathrm{CFU} / \mathrm{ml}$ culture. Atrazine-degrading enzymes by B. megaterium appear to be induced. No released chloride was detected from B. megaterium culture indicating that the triazine ring may be remained intact and the atrazine-metabolites not hydroxyatrazine. Atrazine metabolites may be deethylatrazine or deisopropylatrazine. These results indicate that $B$. megaterium can play an important role in the bioremediation of atrazine-contaminated sites.


Keywords: atrazine, Basilllus, Pseudomonas and Micrococcus, degradation, Bacillus megaterium, bioremediation.

## INTRODUCTION

Atrazine, 2-chloro-4-(ethylamine)-6-(isopropylamine)-s-triazine, an organic compound consisting of an s-triazine-ring, is a widely used herbicide. Atrazine is one of the most environmentally prevalent s-triazine-ring herbicides through its global use to stop pre- and post-emergence broadleaf and grassy weeds in major crops, e.g., maize, sorghum and sugarcane. It is also used for non-selective weed control in non-cropland and industrial areas (Sparling et al., 1998). Atrazine acts by binding to the plastoquinone-binding protein in photosystem II, inhibiting electron transport (Ackerman, 2007).

Atrazine poses a potential health threat. Long-term exposure to atrazine-contaminated drinking water can potentially cause cancer. However, it has been classified a class C/possible human carcinogen (Loprieno et al. 1980).

Atrazine is though to be persistent especially in aquifers or anaerobic sediments (Widmer and Spalding, 1995). The half-life of atrazine in natural soils ranging from 13 to 261 days (U.S. EPA, 2003). The frequency of the occurrence of atrazine in the environment is related to extensive usage, atrazine's moderate persistence, and its mobility through the soil (Burkart and Kolpin, 1993). However, residues of both the parent compound and its
derivatives have been detected in soils years after application (Schiavon, 1988).

The fate of s-triazine compounds in the environment is directly correlated with the ability of microbes to metabolize them. Atrazine biodegradation can occur by two pathways. Atrazine can be dechlorinated followed by removal the other ring substituents via amidohydrolases by the enzymes AtzA, AtzB, and AtzC. The end product is cyanuric acid. The best characterized organism that performs this pathway is Pseudomonas sp. strain ADP. The second pathway involves dealkylation of the amino groups. subsequent dechlorination yields cyanuric acid. The end result is 2 -chloro-4-hydroxy-6-amino-1,3,5-triazine, which currently has no known path to further degradation. This path occurs in Pseudomonas species and a number of bacteria (Wackett et al., 2002 and Zeng et al., 2004).

Atrazine degrades in soil by the action of microorganisms. Microbial degradation determines the environmental impact and efficacy of an herbicide. Rates of biodegradation affected by atrazine's low solubility. Atrazine itself is a poor energy source due to the highly oxidized carbons in the ring. It is catabolized as a carbon and nitrogen source in limiting environments. Inorganic nitrogen accelerates atrazine catabolism whereas organic nitrogen decreases it. Low concentrations of glucose can decrease the bioavailability, whereas higher concentrations promote the catabolism of atrazine (Ralebitso et al., 2002). A variety of bacteria and fungi that dealkylate or dechlorinate atrazine but do not mineralize the s-triazine ring have been isolated (Donnelly et al. 1993, Behki et al. 1993; Nagy et al. 1995, De Souza et al., 1995 and 1996 and Mougin et al. 1997 and Bouquard et al. 1997). Several microorganisms including members of the genera Pseudomonas, Acinetobacter, and Agrobacterium capable of atrazine mineralization have been isolated from sediments or soils that have frequently contact with this herbicides (Vanderheyden et al. 1997; Struthers et al. 1998; Rousseaux et al. 2001 and Topp 2001).

The objective of this research was to enrich for microorganisms capable of mineralization of high concentration of atrazine and use it as a carbon and energy source.

## MATERIALS AND METHODS

## Chemical

Herbicide atrazine (W.P 80\% KZ) 2-chloro-4-[ethylamino]-6-[isopropylamino]-1,3,5-triazine (Fig. 1) was a gift from pesticides center lab., Egypt.


Fig. 1. Atrazine

## Enrichment and Isolation

One kg dry weight soil was placed into glass beaker, fortified with commercial atrazine at a double of rate field dissolved in water which adds to save soil moisture at approximately $70 \%$ field capacity, thoroughly mixed with the soil and beaker covered with porous aluminum foil to ensure gas exchange. The beaker was, incubated at $30^{\circ} \mathrm{C}$ for three weeks. Each amendment of atrazine was applied a total of 3 times at 3 -weeks intervals, and after the final amendment, the soil remained in the incubator for an additional 2 weeks prior to further analysis. Water was added to replace any water lost during the incubation period.

Ten grams of previously atrazine-treated soil was placed in 250 ml of basal salts medium containing $\mathrm{K}_{2} \mathrm{HPO}_{4}, 0.5 \mathrm{~g}$; $\left(\mathrm{NH}_{4}\right)_{2} \mathrm{SO}_{4}, 0.5 \mathrm{~g}$; $\mathrm{MgSO}_{4} .7 \mathrm{H}_{2} \mathrm{O}, 0.5 \mathrm{~g} ; \mathrm{FeCl}_{3} . \mathrm{H}_{2} \mathrm{O}, 10 \mathrm{mg} ; \mathrm{CaCl}_{2} . \mathrm{H}_{2} \mathrm{O}, 10 \mathrm{mg} ; \mathrm{MnCl}_{2}, 0.1 \mathrm{mg} ;$ $\mathrm{ZnSO} 4,0.01 \mathrm{mg}$; glucose $1 \mathrm{~g} / \mathrm{l}(\mathrm{pH} 6.8)$ prepared as described by (Radosevich et al., 1995). This medium was amended with atrazine as an active ingredient at 50 ppm . The pesticide was solved in acetone as a stock solution and added after autoclaving media.

The enrichment culture was incubated with shaking for 1 week at $30^{\circ} \mathrm{C}, 5 \mathrm{ml}$ was placed in 45 ml of saline solution, and the cultures was isolated by serial dilution and plated onto Petri dishes containing nutrient agar medium. Individual colonies were picked up and purified. Pure isolates were streaked on agar slants and stored at $4^{\circ} \mathrm{C}$.

All isolates were tested for growth on atrazine at 100 and 500 ppm as carbon and/or nitrogen source on mineral salts agar medium. The best isolates which show high, rapid growth and clearing zone on agar plates were picked up to complete the experiments.

## Isolates identification:

The isolates were identified according to Sneath et al. (1986). The growth ability of selected isolates on atrazine concentrations

A fresh cell suspension of selected isolate (OD 0.5 at 620 nm ) was incubated aerobically in $100-\mathrm{ml}$ flasks contain 40 ml of basal salts medium with active ingredient of atrazine as a sole source of both carbon and nitrogen at $0,50,100,300$ and 500 ppm and incubated on a rotary shaker ( 170 rpm ) at $30^{\circ} \mathrm{C}$. The growth rate was measured by OD at $620 \mathrm{~nm}\left(\mathrm{~A}_{620}\right)$ at $0,4,8$, $12,16,20,24,48,72$ and 96 h.

## Atrazine-degradation by growing cells

Selected isolate was cultured in $100-\mathrm{ml}$ flasks contain 40 ml of basal mineral salts medium with active ingredient of atrazine as sole carbon and nitrogen source at 50 ppm . The culture was grown for 30 days on a rotary shaker ( 170 rpm ) at $30^{\circ} \mathrm{C}$. The growth was measured by cell counts and the residue of atrazine was measured by GLC at $0,3,7,10,15,21$ and 30 days. The control was free from any isolates to notice the impact of photodegradation or hydrolytic effect on atrazine.

To determined if the atrazine-degrading enzymes are produced constitutively or induced, basal mineral medium amended with 50 ppm of atrazine as a carbon and nitrogen source and $100 \mu \mathrm{~g}$ of chloramphenicol $\mathrm{ml}^{-1}$ was inoculated with cells and incubated for 10 days. Concentrations of
herbicide were determined by GLC. Population size was determined by plating (Struthers et al., 1998).
Residue analysis
Atrazine were extracted twice by ethyl acetate (1:1, vol/vol). Pooled fractions were dried over sodium sulfate, reduced just to dryness in vacuum, and taken up in a small volume of acetone for GLC analysis (Behki et al., 1993).

## Chloride release

Cell culture of selected isolate was inoculated into chloride, carbon, and nitrogen-free liquid medium with atrazine at 100 ppm . Cultures were incubated for 10 days on a rotary shaker (170 rpm) at $30^{\circ} \mathrm{C}$. Chloride release was determined by $\mathrm{AgNO}_{3}$ (Smith et al., 2005).

## RESULTS AND DISCUSSIONS

## Isolation and selection of atrazine-degrading bacteria

Fifteen individual bacterial isolates were obtained after purification. Nine of the isolated bacteria, which show different morphology according to growth on agar plates and to shape of cells under microscope, were chosen for subsequent experiments. These isolates classified according to Gram stain, morphology and some biochemical characteristics to three genera, Bacillus, Pseudomonas, and Micrococcus. Ayansina and Oso (2006) reported that Bacillus sp. and Pseudomonas sp. were the most frequently isolated bacteria from atrazine treated soils.

These isolates showed a varied ability to grow on mineral salts agar with atrazine ( 100 and 500 ppm ) as a carbon and/or nitrogen sources.

The mineral salts agar medium amended with 100 ppm atrazine was used for the isolation of the individual members, while the medium with higher atrazine concentration ( 500 ppm ) was employed for potential atrazine degradation activity.

The isolates encoding. A1, A5, A8 and A9 showed good growth on atrazine as a carbon and/or nitrogen source at (100 ppm). Only two isolates A8 and A9 were proven to have the activity of degrading and growing on atrazine as a sole carbon and nitrogen at high concentration ( 500 ppm ). Isolate A8 showed clearing zones after 2 weeks while isolate A9 showed clearing zone after 3 week. However, these clearing zones are indicating to atrazine being dissolved and transformed. Since, the isolate A8 was the best one, which showed high ability to grow on atrazine as a carbon and/or nitrogen at high concentration (500 ppm).

## Identification of selected isolate

According to morphological and biochemical characteristics isolate A8 was long rode, Gram-positive, spore-forming ellipsoidal and terminal, motile, catalase positive, hydrolysing of casein, gelatin and starch, utilizing of citrate, produce acid from glucose but not gas, V.P test and indole production were negative. On nutrient agar the growth was creamy, moderately dull. This isolate belongs to genus Bacillus and fall within Bacillus megaterium (Sneth et al., 1986).

## The growth rate on atrazine concentrations

The growth rate of $B$. megaterium showed that $B$. megaterium could to grow on atrazine as a source of carbon and nitrogen at 50 ppm with a specific growth rate $0.0623 \mathrm{hr}^{-1}$ and doubling time was 11.13 hr (Fig. 2). The concentration 100 ppm had similar effect; the specific growth rate was 0.0596 $\mathrm{hr}^{-1}$ with doubling time 11.63 hr . By increasing atrazine concentration to 300 and 500 ppm the specific growth rate was greatly reduced to reach 0.0176, $0.0102 \mathrm{hr}^{-1}$ respectively. Doubling time was 39.38 hr at 300 ppm atrazine and 67.96 hr at 500 ppm atrazine. Noteworthy B. megaterium could tolerate high concentration of atrazine and the stationery phase was continued after 96 hr. Atrazine decomposition by soil microorganisms was recently reported by Marecik et al. (2008)


Fig. 2. The growth rate of $B$. megaterium on concentrations of atrazine.

## Degradation of atrazine

B. megaterium could to grow successfully in liquid media with atrazine as a carbon and nitrogen source at 50 ppm . Under these conditions viable counts increased from $10^{5}$ to $10^{8} \mathrm{CFU} \mathrm{ml}{ }^{-1}$ culture. $45.8 \%$ of atrazine was transformed within 30 days concurrent increase in the population size (Fig. 3). Viable cell count was measured, with initial inocula density of $30.2 \times$ $10^{5} \mathrm{CFU} \mathrm{ml}{ }^{-1}$ culture, where it increased until 10 days to reach $45.1 \times 10^{8}$ CFU ml ${ }^{-1}$ culture then reduced gradually to $33 \times 10^{2} \mathrm{CFU} \mathrm{ml}{ }^{-1}$ culture at 30 days of incubation.

The loss of atrazine in uninoculated treatment was negligible, only $5.0 \%$ of atrazine was disappeared within 30 days. Such loss may be due to hydrolytic effects. This result is in agreement with Behki et al. (1993), Mandelbaum et al. (1993) and Radosevich et al. (1995).

Atrazine-degrading enzymes by $B$. megaterium appear to be induced. The persistence rate of atrazine in medium with $100 \mathrm{\mu g}$ chloramphenicol $\mathrm{ml}^{-1}$ after 10 days was $98.9 \%$ and cells population was decreased from $10^{6}$ to $10^{4} \mathrm{CFU} \mathrm{ml}{ }^{-1}$ culture. However, the persistent rate of
atrazine in medium without chloramphenicol after 10 days was $68.8 \%$ with increase the population size from $10^{6}$ to $10^{8} \mathrm{CFU} \mathrm{ml}^{-1}$ culture.

The persistence rate of atrazine and growth activity for $B$. megaterium results in contrast with what mentioned by Marecik et al. (2008) that the mesophilic bacteria B. megaterium has lower counts of cells was observed in nutrient broth with atrazine at $5.00 \mathrm{mg} \mathrm{l}^{-1}$. The long sequential enrichment technique with high concentration of pesticides used to make an adaptation to the microorganisms. In addition, free- carbon and nitrogen media used obligate the adaptation bacteria to use the pesticide. However, according to Bergey's Manual Systymatic Bacteriology B. megaterium has many strains. Hence, as in our knowledge that was the first report about atrazine-degradation by $B$. megaterium isolated from Egyptian soil.


Fig. 3. Persistence of atrazine and growth rate of B. megaterium.
Agrobacterium radiobacter J14a grown in nitrogen-free medium with citrate and sucrose as carbon sources mineralized $94 \%$ of $50 \mu \mathrm{~g}$ of $\left[{ }^{14} \mathrm{C}-\mathrm{U}-\right.$ ring] atrazine $\mathrm{ml}^{-1}$ in 72 h with increasing in the population size from $7.9 \times 10^{5}$ to $5.0 \times 10^{7}$ cells $\mathrm{ml}^{-1}$. While in the medium with $50 \mu \mathrm{~g}$ of [ $\left.{ }^{14} \mathrm{C}-\mathrm{U}-r i n g\right]$ atrazine $\mathrm{ml}^{-1}$ as a sole carbon and nitrogen source, only $11 \%$ of the atrazine was mineralized. Populations declined from $7.6 \times 10^{5}$ to $3.1 \times 10^{5}$ cells $\mathrm{ml}^{-1}$ after 120 h (Strurhers et al. 1998).

## Chloride release

No released chloride was detected from B. megaterium culture indicating that the chlorohydrolase enzymes may be not found in this bacterium. Also, this result indicating that the triazine ring may be remained intact. Also, this result a signal to that the atrazine-metabolites not hydroxyatrazine, it may be deethylatrazine or deisopropylatrazine.

Microbial utilization of the lateral-chain carbon and of both the lateral and ring nitrogen has been observed in few microorganisms. Rhodococcus strain TE1 metabolized atrazine under aerobic conditions to produce deethyland deisopropylatrazine, which were not degraded further and which
accumulated in the incubation medium (Behki et al. 1993). Bacterial growth on atrazine typically involves mineralization of the alkyl-side chains that can be used as a nitrogen, carbon and energy sources, whereas the triazine ring is fully oxidized and cannot be used as an energy source (Radosevich et al. 1995; Struthers et al. 1998 and Bichat et al. 1999).

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تحليل الأترازين بواسطة بكتيريا باسيلس ميجاتريم المعزولة من التربة الزراعية محمود محمد عوض الله اللسواح 1 وسامية مرسىى بيومى1 1 وإيمان حسين عاشور 1 ولبينى أحمد موسى² وسماح عبدالله حافظ شادى 1
 2 قسم بحوث الميكروبيولوجيا ـ معهـ الأراضى والمياه والبيئة - مركز البحوث الزراعية ـالجيزة .

استهذ البحث دراسة مقدرة بعض الميكروبات المعزولة من التربة المصرية على تحليـل مبيد الحشائش الأترازين ، حبث تم عزل تسعة عزلات بكتيرية محللة للأنترازين من التربــة باستـخدام تكنيك المزار ع المقو اه ، وقد أُنبعت هذه العز لات لأجناس الباسبلس والليبدوموناس والميكروكوكس ، وقد أظهـرت العزلـة أ8 نمـو جيـد ومنطقـة رائقـة علـى بيئـة الآجـار المحتويــة علـى أمـلاح معدنيـة و المدعمة بمبيد الأترازين بتركيز 500 جزء فـى المليون كمصدر للكربون والنيتروجين ، ومـن ثم




 أترازين ، وتدل هذه النتائج على أن بكتيريا باسيلس ميجانتريم قد تلعب دور اً هاماً فى العـلاج الحيوى

