DEGRADATION OF H-ACID BY FREE AND IMMOBILIZED CELLS OF *ALCALIGENES LATUS*

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ABSTRACT

*Alcaligenes latus*, isolated from industrial effluent, was able to grow in mineral salts medium with 50 ppm (0.15 mM) of H-acid as a sole source of carbon. Immobilization of *Alcaligenes latus* in Ca-alginate and polyurethane foam resulted in cells embedded in the matrices. When free cells and immobilized cells were used for biodegradation studies at concentration ranging from 100 ppm (0.3 mM) to 500 ppm (1.15 mM) degradation rate was enhanced with immobilized cells. Cells immobilized in polyurethane foam showed 100% degradation up to 350 ppm (1.05 mM) and 57% degradation at 500 ppm (1.5 mM). Degradation rate of Ca-alginate immobilized cells was less as compared to that of polyurethane foam immobilized cells. With Ca-alginate immobilized cells 100% degradation was recorded up to 200 ppm (0.6 mM) of H-acid and only 33% degradation was recorded at 500 ppm (1.5 mM) of H-acid. Spectral analysis of the products after H-acid utilization showed that the spent medium did not contain any aromatic compounds indicating H-acid degradation by *A. latus*.

Key words: H-acid, biodegradation, immobilized cells, *Alcaligenes latus*

INTRODUCTION

Synthetic dyes have been extensively used in textile, power loom and dyeing industries. Azo dyes constitute more than 50% of the dyes produced globally (22). Almost 10 – 15% of the dyes produced are discharged in the effluents (49). The amino and hydroxy naphthalene sulphonic acids are building blocks of azo dyes and the sulphonic acid group confers a xenobiotic character to this class of chemicals (41). The amino groups as additional substituents add polarity to these xenobiotics, which may further resist biodegradation. Even these xenobiotic compounds are partially degraded by certain bacteria and algae; they utilize naphthalene sulphonic acids as a source of sulfur.

Certain bacteria and algae have been reported to utilize naphthalene sulphonic acids as a source of sulfur (21, 43, 48, 50). Kniefel *et al.* (21) showed that under sulfate limitation, axenic batch cultures of the green alga *Scenedesmus obliquus* could metabolize 1-naphthalene sulfonic acid and partially use the sulfonate as a source of sulfur. A small amount of 1-naphthalene sulfonic acid was desulfonated. The resulting 1-naphthol was mostly transformed into 1-naphthyl β-D-glucopyranoside. According to Soeder *et al.* (43) 1-naphthalene sulfonate was utilized by axenic cultures of *Scenedesmus obliquus* and by five other green microalgae as the sole source of sulfur. 1-naphthol appeared as the major metabolite of 1-naphthalene sulfonate. Hence, they concluded that 1-naphthalene sulfonate underwent a desulfonation. Zurrer *et al.* (50) showed that a *Pseudomonas* sp., an *Arthrobacter* sp. and an unidentified bacterium isolated from sewage could desulfonate at least 16 aromatic compounds none of which served as carbon source. *Pseudomonas* sp. strain S-313

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converted 1-naphthalene sulfonic acid, 2-naphthalene sulfonic acid, 5-amino 1-naphthalene sulfonic acid, benzene sulfonic acid and 3-aminobenzene sulfonic acid to 1-naphthol, 2-naphthol, 5-amino 1-naphthol, phenol and 3-aminophenol respectively.

Feigel and Knackmuss (15) have reported degradation of 4-aminobenzene sulphonylic acid in a co-metabolism by two species of bacteria. Hydrogenophaga deaminated 4-aminobenzene sulphonic acid by regioselective 3,4-dioxygenation. The major part of the metabolite was catechol 4-sulfonate which was further metabolized by Agrobacterium radiobacter. Nortemann et al. (33) showed that *Pseudomonas* sp. BN6 could oxidize 1- and 2-naphthalene sulfonate, 1-hydroxynaphthalene 2-sulfonate, 2,6-naphthalene disulfonate and all monosulfonated naphthalene 2-sulfonates which carry one or two substituents in the positions 4-, 5-, 6-, 7- or 8- of the naphthalene ring system with the exception of 4- or 5-amino and 4-hydroxynaphthalene 2-sulfonates. These compounds were converted to the corresponding salicylates. However the strain BN6 did not oxidize substituted naphthalene 1-sulfonates, naphthalene 3-sulfonates and naphthalene disulfonates. 5-Hydroxyquinoline 2-carboxylic acid was obtained as an end product from the degradation of 5-Amino naphthalene 2-sulfonic acid by *Pseudomonas* sp. (32). The formation of 5-hydroxyquinoline 2-carboxylate prevented NADH regeneration and further oxidation of 5-amino naphthalene 2-sulfonic acid was limited by the internal NADH pool. *Moraxella* sp. isolated from industrial sewage plant could degrade Naphthalene 2, 6 and Naphthalene 1, 6 disulfonic acids (48). Regioselective 1,2-dioxygenation caused desulfonation of the compound resulting in accumulation of 5-sulfosalicylic acid which also could be used as the sole carbon source. 5-Sulfosalicylic acid grown cells exhibited high gentisate 1,2-dioxygenase activity. Bacteria degrading amino hydroxy naphthalene sulphonylic acids have been isolated from river Elbe by Nortemann et al. (31). The complete degradation of 6-aminonaphthalene 2-sulfonic acid was carried out by a mutualistic interaction of two *Pseudomonas* strains. Strain BN6 effected the initial conversion of the compound into 5-aminosalicylate through regioselective attack of the naphthalene skeleton in 1,2- position . 5-aminosalicylate was totally degraded by strain BN9. After prolonged adaptation of strain BN6 to growth on 6-aminonaphthalene 2-sulfonic acid, this organism readily converted all naphthalene 2-sulfonates with OH- or NH2- substituents in 5-, 6-, 7- or 8- position. The corresponding hydroxy or aminosalicylates were excreted in stoichiometric amounts.

H-acid (1-amino 8-hydroxy naphthalene 3, 6-disulfonic acid) is a xenobiotic compound (PAH) used as a precursor in the preparation of several azo dyes and is resistant to degradation by all most all microorganisms. Degradation of H-acid by physico chemical methods has been worked out by few investigators (30, 38, 45). However, there are limited reports on biodegradation of H-acid (40, 27).

Immobilized cells have been defined as cells that are entrapped within or associated with an insoluble matrix. Mattiasson (26) discussed various methods of immobilization: covalent coupling, adsorption, entrapment in a three-dimensional polymer network, confinement in a liquid-liquid emulsion and entrapment within a semi permeable membrane. Entrapment in three-dimensional polymer network is widely used for immobilization studies. Various matrices, such as K-carrageenan, alginate, agar, polyacrylamide-hydrazide and polyurethane foam, have been successfully used for immobilization of microorganisms (7, 8, 16, 26).

Under many conditions, immobilized cells have advantages over either free cells or immobilized enzymes. Immobilization imparts more operational flexibility due to the fact that it prevents biomass washout in continuous flow reactors, allows the use of higher cell densities than those obtainable with free cell systems, facilitates the separation of biomass from the treated effluent and offers the potential for improving wastewater treatment and solves the problems associated with solid-liquid separation in settling tanks (5, 11). Use of immobilized cells permits the operation of bioreactors at flow rates that are independent of the growth rate of the microorganisms employed (34). Catalytic stability can be greater for immobilized cells than for free cells and some
immobilized microorganisms tolerate higher concentrations of toxic compounds than do their non-immobilized counterparts (20, 47).

Bioremediation using immobilized bacterial cells for the degradation of benzene was first shown by Somerville et al. (44) using polyacrylamide as immobilizing matrix. Bioremediation with immobilized cells in various matrices has been widely investigated for numerous toxic chemicals such as phenol (2, 19, 29), pentachlorophenol (6, 35), 4-chlorophenol (1, 47), pyridine (23) and naphthalene (24, 25).

Manohar et al. (25) reported the higher rate of naphthalene degradation by *Pseudomonas* sp. strain NGK1 immobilized in polyurethane foam than other matrices (alginate, agar and polyacrylamide) tested and free cells. Polyurethane foam was used as an excellent support for the immobilization of microbial cells for their use in the production of fuels chemicals (16). Joshi and D’souza (18) reported the immobilization of activated sludge for the degradation of phenol. Mordocco et al. (28) studied the effect of parameters such as pH, temperature and dilution rate and bead diameter on the alginate-immobilized cells of *Pseudomonas putida* for the continuous degradation of phenol at low concentration. There was an enhanced mineralization of pentachlorophenol when *Pseudomonas* sp. UG30 cells were immobilized in K-carrageenan compared to free cells (6).

*Pseudomonas paucimobilis* cells immobilized in Ca-alginate-phytagel was able to degrade sulfanilic acid (4-amino benzene sulphonic acid). The compound was used as sole carbon and nitrogen source. Immobilization in Ca-alginate-phytagel did not alter biodegradation activity of the organism (36). Production rate of CO₂ was checked during degradation of 6-amino-2-naphthalene sulphonlic acid by bacteria immobilized on sand particles. A linear correlation between CO₂ formation and growth rate was found for submerged growing cultures as well as for bacteria immobilized on sand particles (13). The present work reports the enhancement of the degradation of H-acid by *Alcaligenes latus*.

### MATERIALS AND METHODS

#### Chemicals

H-acid and other chemicals used in the present investigation were obtained from Hi-media (Bombay) (Figure 1).

![Figure 1. H-acid (1-Amino 8-Hydroxy Naphthalene 3,6-disulfonic acid)](image)

#### Media for H-acid degradation studies

Mineral salts medium of Brilon et al. (3) (12 g Na₂HPO₄.2H₂O, 2 g KH₂PO₄, 0.5 g NH₄NO₃, 0.1 g MgCl₂.6H₂O, 50 mg Ca(NO₃)₂.4H₂O, 7.5 mg FeCl₃.4H₂O and 0.1 ml trace elements solution per liter of medium (37)) containing 50 ppm (0.15 mM) H-acid, 100 mg/l of yeast extract and 100 mg/l of dextrose was used for the degradation of H-acid by *A. latus*.

For alginate-entrapped cells, the degradation medium contained (g/l) K₂HPO₄ 0.15, MgSO₄.7H₂O 0.2, NH₄Cl 1.0, FeCl₃ 0.05 and CaCl₂ 0.2. The pH of the medium was adjusted to 7 (24). Different concentrations of H-acid (100 ppm (0.3 mM) to 500 ppm (1.5 mM)) were added as the sole carbon source.

#### Immobilization of Bacterial cells

**Immobilization in Ca-alginate:** The alginate entrapment of cells was performed according to the method of Bettman and Rehm (2). Alginate (4% w/v) was dissolved in boiling water and autoclaved at 121°C for 15 min. Bacterial suspension (5% w/v) was added to 100 ml sterilized alginate solution and
mixed by stirring on a magnetic stirrer. This alginate/cell mixture, with stirring, was extruded drop by drop into a cold, sterile 0.2 M CaCl$_2$ solution through a burette connected to a tapered pipette tip, by blowing air from the other end of the burette. Gel beads of approximately 2 mm diameter were obtained. The beads were hardened by resuspending into a fresh CaCl$_2$ solution for 2 h with gentle agitation. Finally these beads were washed with distilled water and used for experimentation.

**Immobilization in Polyurethane foam**

A cell suspension was prepared by mixing 10 g of the cell paste (centrifuged pellet with little bit of moisture) with 20 ml of buffer (carbon-free growth medium). One part of polyurethane pre-polymer was cooled on ice. One part (weight/weight) of buffer was added and the mixture was stirred well for one min. One part of the cell suspension was added, and mixing was continued for an additional one min. An additional part of the cell suspension was then added and mixing was continued for another one min. Cell-free foam was made for use as a control by substituting buffer for the cell suspensions. The reaction vessel was kept on ice for 2 h while the polyurethane foam hardened. The foam was removed from the reaction vessel, rinsed with buffer to remove free cells and stored at 4°C. At the beginning of each experiment the foam was rinsed three times with buffer to remove any free cells released during cutting of the foam (35).

**Biodegradation studies**

**Batch degradation studies:**

**1. With free cells system**

To carry out biodegradation studies with free cells 10 ml (8 x 10$^{10}$ CFU/ml) of *Alcaligenes latus* culture was added to each flask with 100 ml of mineral salts medium, 10 mg of yeast extract and different concentrations of H-acid (100 ppm (0.3 mM) to 500 ppm (1.5 mM)). The flasks were incubated at 37°C and at 180 rpm for 24 h.

**2. With immobilized systems**

For biodegradation experiment with Ca-alginate, 5 g (wet weight) of beads, containing the immobilized *Alcaligenes latus* cells, were placed in 250 ml flasks containing 100 ml mineral salts medium and H-acid was added at concentrations ranging from 100 ppm (0.3 mM) to 500 ppm (1.5 mM). The flasks were placed in a rotary shaker at 180 rpm at 37°C for 24 h.

Similarly 2 g of polyurethane foam with immobilized *Alcaligenes latus* cells were added to 100 ml of mineral salts medium containing H-acid at varying concentrations from 100 ppm (0.3 mM) to 500 ppm (1.5 mM). Flasks were placed in a rotary shaker at 180 rpm at 37°C for 24 h.

To determine whether the H-acid was absorbed by the immobilization matrices or not control flasks were kept for incubation with cell free Ca-alginate beads and polyurethane foams separately with different concentrations of H-acid in 100 ml of mineral medium.

**Repeated Batch degradation studies**

To observe the long-term stability of H-acid degradation by immobilized cells in different matrices, the system was used for repeated batch degradation. After each cycle of incubation period (24 h), the spent medium was decanted and beads/foam were washed with sterile water and transferred into a fresh sterile mineral salts medium (100 ml) containing 200 ppm (0.6 mM) of H-acid. The degradation process was carried out as described above and the spent medium was used for the analysis of H-acid degraded.

**Continuous degradation studies**

The continuous treatment of H-acid was carried out in a continuous flow reactor. The reactor was filled with immobilized *A. latus* cells in different matrices. The degradation process was carried out by continuous supply of sterile mineral salts medium/minimal mineral salts medium containing H-acid at different concentrations with constant flow rate (ml/h) with the help of peristaltic pump (Miclins PP10-4C, India).

**Design of bioreactor for continuous treatment:** A cylindrical glass column (4×50 cm, volume 650 ml) with inlet and outlet facilities was used. The bottom of the column was
packed with a glass wool (4 cm diameter) followed by a porous glass-frit. Then the reactor was packed with the respective immobilized cell matrix to a height of 30 cm. The reactor was attached to a reservoir containing mineral salts medium/minimal mineral salts medium with H-acid. The medium was fed into the column continuously with the help of a peristaltic pump (Miclins PP10-4C, India) through a side arm present near the bottom of the column. The medium after H-acid degradation was continuously removed from the side arm situated just above the packed bed. The detention time was calculated by the following formula.

**Detention time:** void volume / flow rate (ml/h).

Degradation Rate (R) = (Ci - Ce) × D

Where:
- Ci = Concentration of H-acid in the influent
- Ce = Concentration of H-acid in the effluent
- D = Dilution rate = Flow rate (ml/h) / Void volume of the reactor (ml)

**Estimation of per cent degradation of H-acid**

At regular intervals a portion of the cultures were withdrawn and collected from the effluent in case of continuous system, centrifuged to remove cells and debris. The absorbance of clear supernatants was read at 390 nm. The concentrations of the H-acid were calculated by referring to the calibrated curve and the percent of the H-acid degraded were then estimated.

**Cell viability enumeration**

To enumerate viable cells in the Ca-alginate beads, beads were washed in saline and one bead was soaked in 1 ml of saline. Soaked bead was shaken with glass beads for 15 min. Sample was plated onto nutrient agar and viable cells were expressed as CFU/ml after overnight incubation (15 h) at 37°C.

For enumeration of viable cells in the PUF (polyurethane foam), PUF were rinsed with saline. One PUF was then torn into fine pieces using sterile forceps, suspended in saline and vortexed to dislodge the immobilized cells. Sample was then plated onto nutrient agar and viable cells were counted after overnight incubation (15 h) at 37°C.

**Confirmation of H-acid degradation**

In order to confirm degradation of H-acid by *A. latus*, the products were extracted from the large amount of spent medium (after centrifugation at 8000 × g for 10 min) using two volumes of diethyl ether. Thus extracted products were separated and characterized using TLC, 1H NMR and IR analysis.

The separated diethyl ether fractions were dried over anhydrous sodium sulphate and the traces of sulphate from the filtrate were removed by anhydrous barium chloride. The metabolites were isolated and purified by TLC.

Thin layer chromatography was carried out by using a glass plate (20 × 10 cm) coated with silica gel G slurry (1:2 w/v in water), dried and kept in oven at 100°C for 2 h. Solvent systems used were chloroform: acetone (80:20) and cyclohexane: ethylacetate: acetone (4:1:1). Spots were observed after exposing the plates to iodine vapors. H-acid and other aromatic compounds were used as a control to compare with degraded products of H-acid.

The bands obtained on the chromatogram were separated and compounds were eluted with diethyl ether. The compounds were then kept for evaporation and sent to I.I.Sc., Bangalore and CDRI, Lucknow for IR and NMR analysis.

**RESULTS**

The results of H-acid degradation in batch system by free cells are presented in Figures 2 and 3. It can be observed that 100% degradation up to 100 ppm (0.3 mM) of H-acid within 18 h. Further increase in H-acid concentration resulted in decrease in the efficiency of H-acid degradation to 65% at 150 ppm (0.45 mM), 50% at 200 ppm (0.6 mM), and 35% at 300 ppm (0.9 mM) and less than 25% at 350 ppm (1.05 mM) and above. At 500 ppm (1.5 mM), the percent degradation was observed to be 15% only even after 48 h of incubation.

The results of degradation of H-acid in batch systems by *A. latus* immobilized in Ca-alginate and polyurethane foam are shown in Figures 2 and 3. *A. latus* immobilized in polyurethane
foam showed 100% H-acid degradation upto a concentration of 350 ppm (1.05 mM) compared to 200 ppm (0.6 mM) with cells embedded in Ca-alginate and only 100 ppm (0.3 mM) with free cells. Degradation efficiency of polyurethane immobilized cells was observed to decrease at higher concentrations of H-acid, at 400 ppm (1.2 mM) 87%, at 450 ppm (1.35 mM) 80% and a lowest of 57% at 500 ppm (1.5 mM) in 24 h. Efficiency of degradation of H-acid by A. latus immobilized in Ca-alginate continuously decreased above 200 ppm (0.6 mM) to record lowest of 33% degradation at 500 ppm (1.5 mM) in 48 h. However, at 350 ppm (1.05 mM) concentration of H-acid, degradation in free cells was only 23% compared to 60% by cells immobilized in Ca-alginate and 100% with cells immobilized in polyurethane foam.

Figure 2. Batch degradation of H-acid by free and immobilized cells of A. latus
Degradation of H-acid by *A. latus*

Free cells were able to show 100% degradation at 100 ppm (0.3 mM) of H-acid for which 18 h of incubation was required. Further increase in concentration resulted in decrease in per cent degradation and increase in incubation period. From 350 ppm (1.05 mM) onwards the per cent degradation of H-acid was reduced to less than 25% which required more than 30 h of incubation. With Ca-alginate immobilized cells upto 200 ppm (0.6 mM) of H-acid 100% degradation was seen within 18 h of incubation after which the per cent degradation started reducing. At 500 ppm (1.5 mM) of H-acid only 33% degradation was recorded after 42 h. However with polyurethane immobilized cells 100% degradation was recorded upto 350 ppm (1.05 mM) concentration within 18 h and reduction of per cent degradation at further concentration was less. At 500 ppm (1.5 mM of H-acid upto 57% degradation was recorded after 24 h of incubation.

The results of repeated batch H-acid degradation are presented in Figure 4. From the figure it can be observed that Ca-alginate immobilized *A. latus* could be reused without changing its degradation efficiency upto 5 cycles and efficiency decreased to 88% upto 10 cycles, but with further increase in cycles resulted in decrease in degradation.
Degradation of H-acid by *A. latus*

efficiency. However, even after 25 cycles 50% degradation efficiency was retained. PUF immobilized *A. latus* showed 100% degradation up to 7 cycles and decrease in the degradation capacity was observed after 7 cycles. And 87% degradation was recorded after 15 cycles and the efficiency remained 80% even after 25 cycles. From the observations it is clear that the PUF immobilized system was more efficient in biodegradation of H-acid for an extended length of time when compared to cells embedded in Ca-alginate.

During the recycling of immobilized microbial systems there was a leakage of cells from the matrix on repeated washings with minimal mineral salts medium in case of Ca-alginate immobilized cells and mineral salts medium in case of PUF immobilized cells. It was observed that the cell leakage was very higher (1x10^3 CFU/ml) in Ca-alginate matrix than in PUF (2x10^1 CFU/ml) after 19 cycles and 25 cycles respectively.

The results of percent degradation of H-acid by continuous system of Ca-alginate immobilized *A. latus* cells are shown in Figure 5. This figure shows the effect of different concentrations of H-acid on percent degradation of H-acid by *A. latus* at a constant flow rate of 20 ml/h. Complete degradation of H-acid was recorded at flow rates of 20 (dt 180 min) to 40 ml/h (dt 90 min) with 200 ppm (0.6 mM) of H-acid and up to 20 ml/hr at 400 ppm (1.2 mM) of H-acid. On increasing the flow rate gradually from 40 to 100 ml/h (dt 36 min) the percent degradation also decreased, and even at 100 ml/h flow rate percent degradation was 50% and 46% with 200 (0.6mM) and 400 ppm (1.2 mM) respectively. At 1000 ppm (3 mM) it was observed to be 74% degradation at 20 ml/h (dt 180 min) flow rate and decreased to 22% at 100 ml/h (dt 36 min).

The efficiency of PUF immobilized *A. latus* under continuous degradation of H-acid is shown in Figure 6. A 100% degradation was observed at flow rates of 20 (dt 240 min), 40 (dt 120 min) and 60 ml/h (dt 79.8 min) at 200 ppm (0.6 mM) of H-acid. Complete degradation was observed at lower flow rates with higher H-acid concentrations, i.e. up to 400 ppm (1.2 mM) at 20, 40 ml/hr flow rates and at 20 ml/h flow rate up to 600 ppm (1.8 mM) of H-acid. Even at highest concentration of 1000 ppm (3 mM) of H-acid, 88% degradation was achieved at 20 ml/h (dt 240 min) flow rate but only 42% degradation was observed with 100 ml/h (dt 48 min) flow rate.

![Figure 4. Repeated batch degradation of H-acid at 200 ppm concentration of H-acid](image.png)
Figure 5. Continuous degradation of different concentrations of H-acid by Ca-alginate immobilized A. latus system

Figure 6. Continuous degradation of different concentrations of H-acid by PUF immobilized A. latus system
Cell viability was greater in PUF than Ca-alginate beads. After 2 h of immobilization cell viability was found to be $5 \times 10^6$ CFU/ml in PUF whereas it was $4 \times 10^4$ CFU/ml in Ca-alginate bead.

The products of H-acid degradation by *A. latus* were separated and identified by TLC and spectral analyses.

The products of H-acid degradation by *A. latus* were extracted with diethyl ether and subjected to TLC by various solvent systems. With Cyclohexane:Ethyl acetate:Acetone (4:1:1) and Chloroform:Acetone (80:20) as solvent system, a single broad band with tailing was observed not corresponding to the H-acid band. $R_f$ values of H-acid in two different solvent systems were found to be 0.23 and 0.19 and those of the product were found to be 0.33 and 0.28. TLC bands were scraped out, extracted with diethyl ether and subjected to spectral analysis.

The H-acid, before subjecting to degradation, exhibited the IR and $^1$H NMR spectra in concurrence with its structure. The IR showed a broad peak at $n_3499$ cm$^{-1}$ indicating merger of absorption peaks due to the vibrations of $\nu$OH and $\nu$NH$_2$. The broad peak may be due to the hydrogen bonding between oxygen of $-\text{OH}$ and hydrogen of $-\text{NH}_2$. The next significant peak in the IR was at $n_{1598}$ cm$^{-1}$, indicating the absence of $-\text{C}=\text{O}$ function in the molecule (Fig. 7).

The $^1$H NMR of H-acid exhibited distinguishable peaks from 6.6 $\delta$ to 8.0 $\delta$, which is the region for the absorption of aromatic protons, indicating the presence of aromatic character in the molecule. There is a distinguishable singlet seen at 3.3 $\delta$, may be due to the resonance of H of phenolic $-\text{OH}$. A sharp singlet is seen at 10.6 $\delta$ accounting for two protons of primary aromatic amine $-\text{NH}_2$. These spectral evidences support the structure assigned to H-acid (Fig. 8).

The IR spectrum obtained for the degradation product showed a very weak peak at $v_{3309}$ cm$^{-1}$ indicating that percentage presence of molecule present in the mixture containing $-\text{NH}_2/-\text{OH}$ is very less. A very strong absorption peak was seen at $v_{1731}$ cm$^{-1}$ accounting for the presence of a molecule in the product containing $-\text{C}=\text{O}$ functionality (Fig. 9). These observations indicate that the H-acid has undergone oxidative degradation to produce various non-aromatic molecules.

The above observation is further supported by the $^1$HNMR spectrum of the product taken in DMSO-$d_6$. None of the protons of the product resonated in the aromatic region of the spectra supporting the idea of disintegration of the molecule to produce number of aliphatic species (Fig. 10). This is further supported by the evidences that a good number of peaks due to the methyl proton are seen in the spectrum.

![Figure 7. IR Spectrum of H-acid](image-url)
Figure 8. $^1$H NMR Spectrum of H-acid

Figure 9. IR Spectrum of degradation product
DISCUSSION

Results of batch degradation studies revealed that polyurethane foam immobilized cells were able to degrade H-acid completely up to 350 ppm (1.05 mM) compared to 200 ppm (0.6 mM) with Ca-alginate immobilized cells and 100 ppm (0.3 mM) with free cells in 48 h. H-acid degradation efficiency of A. latus was observed to be reduced as the concentration of H-acid increases.

The A. latus cells immobilized in the polyurethane foam showed higher degradation rate than the cells immobilized in Ca-alginate. This high rate of degradation may be attributed to the better availability of substrate (H-acid) to the cells as they are immobilized by adsorption process, whereas, with Ca-alginate matrix the cells are embedded in the matrix, where the slow diffusion of substrate (H-acid) and air into the polymer beads may decrease the rate of degradation. Manohar and Karegoudar (30) and Sharanagouda and Karegouder (42) observed similar results with naphthalene degradation by immobilized Pseudomonas sp. It was also suggested that the storage stability and microbial activity of immobilized cells in polyurethane foam are known to be better than those cells immobilized in Ca-alginate (4, 9).

A. latus immobilized in Ca-alginate and polyurethane foam showed complete degradation of H-acid up to 400 ppm and 600 ppm respectively at a flow rate of 20 ml/h. Increased degradation by immobilized cells was due to the accelerated reaction rates caused by the high local cell density in or on the immobilized matrix and also the immobilization of cells could improve the tolerance capacity against toxicity of high concentration of H-acid. Immobilization also provides a kind of membrane stabilization, which was assumed to be responsible for the cell protection and better degradation rates in immobilized cells. Similar observations have been made by Hall and Rao (16) in the production of fuels and chemicals and Lee et al. (23) in the degradation of pyridine by immobilized Pimetobacter sp. It has been reported that immobilization of cells in a polymer matrix can confer protection from high levels of toxic compounds normally lethal to free cells. The protective effect may be due to adsorption of the toxic compound by the matrix, lowering the available concentration of the toxic compound in solution as observed with polyurethane foam (17), activated carbon (29) and beet chips (10). Alternatively it may be due to alteration in the membrane composition of cell in the matrix, rendering the membrane less permeable to toxic substances (12, 19). Immobilized viable
cells have been shown to have altered metabolism, enhanced enzyme induction, altered macromolecules composition, and reduced specific cell growth and cell yield (14).

Repeated batch degradation studies were conducted to observe the effect of physicochemical factors on the degradation efficiency of *A. latus* immobilized in Ca-alginate and polyurethane foam. The results revealed that the polyurethane foam immobilized cells retained complete H-acid degradation capacity for longer period (7 cycles) compared to cells immobilized in Ca-alginate (5 cycles). Cell leakage was almost three times more in Ca-alginate compared to polyurethane foam immobilized cells after 25 cycles. Polyurethane foam showed higher mechanical stability, resistance to chemicals and biological degradation and lower cell leakage, that made it suitable for a long period reuse. Trevors *et al.* (46) observed mechanical instability and gradual cell leakage from Ca-alginate beads, decreasing the degradation rate. Enhancement in the activity of immobilized cells on reuse has been reported for the other systems (18, 39) and this may be due to post immobilization modifications/acclimatization. In contrast, our results on the reuse of immobilized beads/foam cubes did not show enhancement in the degradation of H-acid but degradation rate decreased after several cycles of reuse. Similar observations were made by Sharanagouda and Karegoudar (42) in the degradation of 2-Methyl naphthalene using *Pseudomonas* sp.

The metabolic products produced from H-acid degradation by *A. latus* were analyzed. Spectral scanning of the product showed a single peak at 260 nm and a single band on TLC which were not corresponding to H-acid peak. Further analysis of TLC band by spectral analysis revealed the presence of a large number of small molecules suggesting the formation of aliphatic molecules during microbial utilization and aromatic byproducts were not observed. The IR spectrum of the products indicated the absence of sulphonic acid group suggesting that complete desulphonation occurs during the degradation of H-acid. Similar reports have been given by Wittich *et al.* (48).

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