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BIODEGRADATION OF MUSTARD



Ronald J. Young

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RESEARCH AND TECHNOLOGY DIRECTORATE

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A literature search to identify microorganisms of potential value for the degradation of mustard was carried out. Selection of micro- organisms was based on tolerance to low pH and chloride ions, conditions that retard mustard hydrolysis. Several bacteria able to degrade organic sulfides and/or sulfonium compounds under these conditions were identified. Fungi and yeasts are also of potential use, as are enzymes from halo- and thermophilic organisms. The major difficulty in the use of microorganisms and enzymes for mustard degradation is the low solubility of mustard in water.						
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The goal of biodegradation of mustard by direct microbiological action on the agent rather than on its hydrolysis product has been impeded by the chemical reactivity and insolubility of the agent. Microorganisms were inactivated by mustard which also was hydrolyzed during incubation in the aqueous medium used for growth of the microorganism. Microorganisms possessing an enzyme system functional against mustard or sulfonium compounds, intermediates in mustard hydrolysis, should be resistant to killing by the agent. With this assumption, a search of the literature was made to identify microorganisms able to degrade organic sulfur compounds and/or to grow under conditions, such as the presence of chloride ions, a relatively high concentration of mustard, and acid pH, in which mustard hydrolysis rate is low.

Some members of the genera Halomonas, Haloanaerobium, Halobacteroides and Sporohalobacter are involved in the degradation of organic sulfides. These microorganisms tolerate NaCl up to 1M and are found in ocean estuaries, along the sea shore, and in anoxic hypersaline waters and their sediments. Unidentified microbes present in anoxic salt marshes and one strain of <u>Pseudomonas</u> (MS strain) are able to degrade sulfonium compounds. Some strains of neutrophilic Thiobacillus and Pseudomonas, present in soil and aquatic regions rich in decaying organic matter, are also active in the degradation of organic sulfides. Although all the above microorganisms are neutrophiles, their ability to degrade organic sulfur compounds, particularly sulfonium compounds, and tolerance of chloride ions fulfills two of the criteria for use.

Less information is available on the degradation of organic sulfides by acidophiles. The acidophile <u>Acidiphilium</u> is currently under intensive study for use in the desulfurization of fossil fuels. Exploration of the application of this bacteria for mustard degradation is indicated. Acidophilic <u>Thiobacillus</u> appear to have limited use for mustard breakdown except for the halotolerant <u>T. prosperus</u>, originally isolated from a shallow geothermally heated seafloor. Acidic marine sites are attractive as sources of isolates for study.

Yeasts and fungi have not been used for the degradation of mustard, and there have been few reports on the utilization of organic sulfides by these microorganisms. This notwithstanding, these microorganisms are attractive candidates for use in degradation of mustard. Yeasts and fungi are acid tolerant and some are halotolerant. Fungi are rather universal in their utilization of organic matter. They are also valuable as a source of macro- and simple organic molecules such as exo- and endoenzymes, surfactants, and polymer, some of which may be

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beneficial in mustard degradation.

The use of microbial enzymes rather than the microorganisms for mustard breakdown is a viable alternative. Enzymes of halophilic and thermophilic microorganisms are able to function in the presence organic solvents thereby alleviating the problem of mustard solubility in water. The ability of extracts or lysates of the thermophilic acidophile <u>S. acidocaldarius</u> to degrade mustard may be a good test of the enzymatic rather than the microbiological approach to mustard demilitarization.

The literature search has identified several microorganisms of potential use in the biodegradation of mustard. However, the critical component in biodegradation of mustard is the dispersion or solubilization of the agent in a medium in which the microorganism remain viable. Defining such a system is a necessary first step in any such endeavors.

PREFACE

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BIODEGRADATION OF MUSTARD

1. INTRODUCTION

Biodegradation of mustard stockpile is an attractive alternative to incineration or chemical treatment because of economic, safety and environmental concerns. Two studies have been carried out on the biological degradation of mustard. Bacteria isolated from soil previously exposed to mustard were either resistant or sensitive to mustard. The first isolates did not degrade mustard (1), while the second isolates, although sensitive to mustard itself, were able to utilize the hydrolysis product of mustard, thiodiglycol, as an energy and carbon source (2). This property of the second isolates, identified as Pseudomonas pickettii and Alcaligenes xylosoxidans ssp. xylosoxidans, coupled with prior alkaline hydrolysis of mustard, is the basis of a method with a high potential for the economical and safe degradation of stockpiles of mustard. There is nevertheless a place for a biological procedure that degrades mustard directly, and completely to environmentally benign compounds. For example, the final products from biological degradation could be sulfate, The reconciliation of the chloride and carbon dioxide. intractable chemical, physical, and toxic properties of mustard with conditions suitable for growth of microorganisms presents a formidable, but perhaps not an insurmountable, obstacle to development of biological methods for its destruction. The lack of success thus far in attempts to degrade mustard directly suggests that exploration of alternate approaches would be Thus microorganisms other than neutrophilic appropriate. bacteria, and from habitats other than mustard treated soils are worthy of consideration. In this report, the properties of mustard pertinent to the use of microorganisms are considered, the result of a survey microorganism of potential use for the biological degradation of mustard is presented, and possible methods for the biological degradation of mustard are discussed.

2. PROPERTIES OF MUSTARD IN RELATION TO MICROORGANISMS

a. Chemical

The crux of any procedure for the direct biological degradation of mustard (I) is the stability and solubility of mustard in water, as growth of microorganisms occurs in aqueous medium. Hydrolysis of mustard is complex (3,4). The rate determining step is the formation of a cyclic sulfonium ion (II) which rapidly reacts further with water to produce thiodiglycol (IV)

as the final product (Figure 1). The end product can react with the cyclic sulfonium intermediates (I, II) to produce branched sulfonium ion intermediates (V-VII). Chloride ion inhibits the formation of the cyclic sulfonium ions, and low pH results in reversal of hydrolysis. Further, hydrolysis of the branched sulfonium intermediates is slow, is dependent on the hydroxide ion concentration, and is negatively correlated with the concentration of the reactant, the sulfonium intermediates (4). Mustard is poorly soluble in water, (VI,VII) and concentrations sufficient to retard hydrolysis require use of agents able to increase its miscibility with water. In this respect an apolar organic solvent or detergent would be desirable as mustard hydrolysis rate is significantly reduced in such solvents or detergent. These considerations point to a desirability for salt tolerant, acidophilic microorganisms that are resistant to organic solvents, neutral surfactants, or surfactants from biological sources.

b. Biological

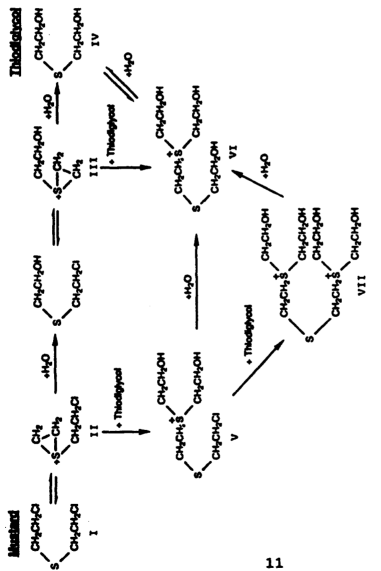
An important consideration in using microorganisms to degrade mustard is the cytotoxic properties of the chemical. The primary target of mustard in the microbial cell is DNA resulting in replication. Other cellular processes are inhibition of unaffected as cells continue to grow, with apparently unbalanced growth as the cause of cell death (5). This seeming insensitivity of proteins to mustard maybe due to a low intracellular concentration of mustard, commensurate with the low concentration of mustard required to induce cell death, and the poor solubility of mustard in water. Microorganisms possessing an enzyme system capable of degrading the agent may continue replication when exposed to low be able to concentrations of mustard. The consequences of exposure of microorganisms to the higher concentrations of mustard that is desirable for the retardation of its hydrolysis is unclear. The intracellular mustard concentration would most likely also be high, but the species of mustard predominately present intracellularly, their stability, and reactivity in a heterogeneous milieu maybe quite different from what has been observed in a homogeneous solution. A careful balance of the conditions to maintain stability of mustard and the viability, even of salt and acid tolerant microorganisms, is necessary.

3. MICROORGANISMS

a. Halophilic Microorganisms

The rate of hydrolysis of mustard in water is retarded six fold in the presence of 0.14M NaCl (6). Higher concentrations of NaCl

Mustard Hydrolysis



sulfonium ion (II). Thiodigyicol (IV) is rapidly formed and this The rate determining step in in turn reacts with the cyclic sulfonium intermediates to give the hydrolysis of mustard (I) is the formation of the cyclic dependent. Chloride ion retards the formation of the cyclic the sulfonium intermediates (V-VII) whose hydrolysis is pH Hydrolysis of mustard. Adapted from reference 2 sulfonium ions. Figure 1.

may retard the rate of hydrolysis further. The beneficial concentration of NaCl is limited, not by the tolerance of microorganisms for salt, but by the extraordinary low solubility of mustard in salt solutions as extremely halophilic bacteria are able to grow in saturated salt solution. The need to maximize the concentration of mustard in order to enhance stability will undoubtedly limit the salt concentration to much less than 1.5 M, conditions suitable for the slightly tolerant category of halophilic bacteria (7). This ceiling on Nacl concentration effectively eliminates consideration of members of the aerobic halophilic archaebacteria family, the family Halobacteriaceae, as the lower range of salt concentration for growth for members of this family is 1-1.5 M (8,9). Halotolerant bacteria of potential interest include those belonging to the genus Halomonas and the genera Haloanaerobium, Halobacteroides and Sporohalobacter as they can be cultured in lower salt concentrations. Members of the genus Halomonas eg H. elongata and <u>H. halodurans</u> show great flexibility for salt and are able to grow in salt concentrations between 0.016 M to 5.5 M (10), those of <u>Haloanaerobium</u>, <u>Halobacteroides</u> whereas and Sporohalobacter require a higher range of salt concentration, 0.45 M to >1.0 M (11). <u>Halomonas</u> species have been isolated from a solar saltern, an estuary and from the Pacific ocean. The Haloanaerobium, Halobacteroides and Sporohalobacter species are obligate anaerobes whose habitats are anoxic hypersaline waters, their sediments, and solar salterns. All microorganism species mentioned above are chemoorganotrophs and have possible applications in the removal of organic waste (10,11). Haloanaerobium praevalens is of particular interest in the present context as the organism degrades methionine producing methylmercaptan as a product (12). <u>Halobacteroides</u> strains have been shown to use methanethiol as the sole sulfur source for growth (13).

56 M

Anaerobes present in anoxic hypersaline environments, anoxic salt marsh and marine-ocean sediments are involved in metabolism of sulfur compounds and sulfur cycling (13,14), and samples from such sources may produce useful isolates. For example ethylated sulfur compounds are converted to ethane by methanogenic bacteria (15a). Dimethylsulfoniopropionate $(CH_3)_2 - S^+ - (CH_2)_2 -$ COOH, found in some algae, is degraded by microorganisms in anoxic salt marsh sediments to dimethylsulfide and acrylic acid, or 3-mercaptopropionic acid and methanethiol (15, 16).Dimethylsulfide and methanthiol are in turn converted to H₂S and methane by sediment microorganisms such as sulfate reducers (13,14,16-18). By analogy, the cyclic sulfonium ion derived from mustard might be degraded to less toxic compounds such as ethylene sulfide, ethylene chloride, ethylene and 2-chloromercaptoethanol, and the sulfonium ion aggregates to a number of compounds including vinyl alcohol, and ethanol among others.

The end products of degradation of an organic sulfur compound by

an anaerobic microorganism are H_2S and methane or a sulfide, and an aerobic microorganism would be necessary to oxidize the sulfide to the more environmentally benign sulfate. Although the goal of degrading mustard to environmentally benign products may not be realized by anaerobic halophilic microorganisms, they nevertheless may, at least, serve in a first detoxifying step, since some species are able to cleave a carbon-sulfur bond, or perhaps more correctly, a methyl-sulfur bond. Mixed cultures may be necessary to achieve the desired degradation to H_2S and methane. As an aside, the enzyme involved in the degradation of dimethylsulfoniopropionate (19) deserves consideration as a device for protection against mustard exposure.

Microorganisms present in hypersaline, marine or aquatic environments are, with few exceptions neutrophilic (20a). The requirement for neutrality is not incompatible with mustard stability as the multimeric sulfonium chloride intermediates in mustard hydrolysis have some degree of stability in neutral conditions particularly in the presence of chloride anions (4). The abnormally high intracellular salt concentration of some eubacteria such as the <u>Haloanaerobium</u>, <u>Halobacteroides</u> and <u>Sporohalobacter</u> species (10) may also contribute to intracellular stability and afford a measure of protection to the toxic action of mustard.

Although many microorganisms tolerant of salt are known, it is unlikely that mustard would be soluble in solvent mixtures at the sodium chloride concentrations discussed above. Mustard hydrolysis is retarded by the chloride anion and the retardation is enhanced under apolar conditions. Thus the chloride concentration need not be high to stabilize mustard under these conditions. Quaternary ammonium chlorides are a source of chloride anions, and they may be an alternative, or even preferred over a very low concentration of sodium chloride because of their higher solubility in organic solvents.

b. Colorless Sulfur Bacteria

Bacteria utilizing reduced sulfur compounds as an energy source for growth are collectively known as colorless sulfur bacteria. The sulfur compounds are oxidized to sulfate either aerobically or anaerobically in the presence of nitrate. Except for the genera <u>Thiobacillus</u>, <u>Sulfolobus</u> and <u>Acidianus</u>, colorless sulfur bacteria are neutrophilic (20). The requirement for neutrality need not be an impediment to the use of colorless sulfur bacteria as hydrolysis of mustard even at neutral pH and in the absence of salt is significantly retarded if the concentration of mustard is sufficiently high. An acid (HCl) environment is preferred as this favors the reversal of both hydrolysis and the formation of the cyclic sulfonium ion. These reactions, however, require strongly acidic conditions.

b.1. Neutrophilic Sulfur bacteria

This group of bacteria encompasses 18 genera whose members display a wide diversity of nutritional requirements and physiological types (20). Their habitats include aquatic regions, sediments, ocean hydrothermal vents, waste water treatment plants, soils, and any environment where reduced sulfur compounds are present. Studies of the use of organic sulfides rather than inorganic sulfides, hydrogen sulfide, or salts such as thiosulfate, as substrates by this group of bacteria appear to be restricted to the genera Hyphomicrobium, Thiobacillus and Pseudomonas. The naturally occurring methyl sulfide is oxidized to sulfate by <u>Hyphomicrobium</u> S sp., Hyphomicrobium EG (21-23), Thiobacillus thioparus TK-m and Thiobacillus MS1 (24,25), but only to dimethyl sulfoxide by Pseudomonas acidovorons DMR-11 (26). Methyl sulfide is also degraded by an obligately chemolithoautotrophic, bacillus T5, isolated from a microbial mat in the North Sea (27), an unidentified bacterium present in surface seawater (28), and facultative chemolithoautotrophs, tentatively identified as Thiobacillus strains E3-E7 isolated from garden compost, cattle manure, marine mud, pond water and moss, respectively (29). Thiobacillus thioparus is an obligate autotroph, and other Thiobacillus autotrophs as well as Thiobacillus facultative heterotrophs (29a) may also be able to use organic sulfur compounds as substrates. Only the Thiobacillus and, perhaps, Pseudomonas strains appear to have potential interest for the degradation of mustard as <u>Hyphomicrobium</u> is an obligate methylotroph. Nevertheless, a <u>Hyphomicrobium</u> species or other unidentified microorganisms present in soil samples were reported to oxidize methyl and higher sulfides to sulfate (30, 31).

Methyl and higher sulfides are the end products of the action of a variety of aquatic and terrestrial microorganisms on sulfur amino acids, and other sulfur containing compounds from decaying organic matter (32,33). The sulfur amino acids are themselves sulfides and these microorganisms may be able to utilize mustard for growth. Thus sediment, soils with decaying organic matter, and aquatic environments appear to hold promise as sources of microorganisms for which mustard maybe a substrate. It should be noted, however, that at neutral pH and in the absence of salt, the predominate chemical species would not be mustard itself, but the sulfonium aggregates. For this reason a strain of Pseudomonas (MS strain) which is able to use trimethylsulfonium chloride as sole carbon source (34-36), deserves consideration for study in the biological degradation of mustard even though substrate C₁-sulfur compound, is and the a dimethylsulfoniopropionate, discussed above, cannot replace trimethylsulfonium chloride as a substrate. Bacterium 5H2 also can grow on trimethylammonfum chloride, and since it is a

facultative methylotroph (37), it is a more attractive candidate.

Mustard sulfoxide is much less toxic than mustard as it does not react with proteins under normal physiological conditions (5). Bacteria oxidizing sulfides to the sulfoxide should receive more than a passing thought with respect to their action on mustard.

b.2. Acidophilic Sulfur Bacteria

Two of the three genera of colorless sulfur bacteria, <u>Sulfolobus</u> and <u>Acidianus</u> are thermophilic and are not considered further. Members of the third, <u>Thiobacillus</u>, are found in soil and water around mineral and coal mines, leaching dumps, acid soils and acid lakes such as volcanic lakes. The aerobic chemolithotropic <u>Acidobacillus</u> converts inorganic sulfides into sulfuric acid. Consequently studies have centered on their corrosive properties and use in mineral leaching rather than on their ability to use organic sulfides as substrates. Some <u>thiobacillus</u> are capable of growing as heterotrophs in the presence of low concentrations of substrate e.g. <u>Thiobacillus rubellus</u>, <u>Thiobacillus delicatus</u> (38), <u>Thiobacillus acidophilus</u> (39), and <u>Thiobacillus cuptinus</u> (40).

c. Other Acidophilic Microorganisms

A new genus of acidophiles, <u>Acidiphilium</u>, has recently been described (41). Members of this genus, originally isolated as contaminants of <u>Thiobacillus</u> species and present in coal and mineral sulfide environments, are mesophilic, acidophilic and heterotrophic. Many species of this genus have been described (39,42-46). The <u>Acidiphilium</u> strains vary a great deal in their ability to utilize organic compounds for growth, but all are heterotrophs, unable to use sulfur or inorganic sulfides for growth. The <u>Thiobacillus</u> and <u>Acidiphilium</u> species appear to be attractive candidates for further study. Possible uses for the latter species are under consideration (46a).

d. Considerations of Acidophily

Acidophiles are capable of growth at a pH of 1 (range 1-4). The cytoplasmic pH is, however, close to neutral, 5.5-6.7 when the external pH is 2-4 (47). In contrast, the pH of the periplasm of acidophilic gram-negative bacteria is thought to be close to the pH of the growth medium (48). The chemiosmotic consequences of this pH gradient is a tendency for exclusion of permeant cations, but not of uncharged chemical species or anions, due to the existence of a positive membrane potential in actively respiring bacteria (47). This means that the sulfonium species are denied entry into the acidophile, but not the uncharged mustard sulfide. The SO_{4}^{2-} anion, resulting from oxidation of sulfide, is non permeant and does not react with mustard. The bacterial membrane is permeable to $Cl^{-}(47,49)$, the anion of importance with respect to mustard stability, and maintenance of the uncharged mustard. Even though membrane polarity is reversed when the external pH is lower than about 2.5 (50), Cl anions are still able to enter because of the chloride porter system (48). Thus, while it is important for Cl anions to be present, the concentration should not be high, <3 (45,51), to avoid inhibition of growth due to a high internal concentration of this anion. Thiobacillus prosperus, isolated from a shallow geothermally heated seafloor, is a halotolerant (6% NaCl) Although this organism acidophile. is an obligate chemolithotroph (20a) other obligate chemolithotrophs have been shown to be able to degrade simple organic sulfides (27). Acidic marine areas appear attractive as potential sites for investigation as sources of isolates. Another possible source of Cl tolerant acidophiles is the stomach as the acidity in this organ is results from HCl. Ruman microorganisms have been shown degrade organic sulfur compounds including sulfonium to compounds (52).

4. COAL AND OIL

Combustion of fossil fuels generates oxides of sulfur resulting in the major environmental problem of acid rain. Microbial removal of sulfur from the fuels before combustion has received a great deal of attention as a means for alleviating this problem (53-59). Aromatic sulfides are the major sources of organic sulfur in the fuels. Coal in addition contains iron sulfide, pyrites, and sulfate. The inorganic sulfide is readily removed by the action of <u>Thiobacillus ferrooxidans</u>, <u>Thiobacillus thiooxidans</u> and <u>Sulfolobus acidocaldarius</u>, but the goal of organic sulfur removal from fossil fuels is still being pursued. Although the structure of the organic sulfides in fossil fuels is quite different from mustard or of the related sulfonium ions, it is instructive to consider the studies with the fossil fuels since the methods used and results obtained may be applicable to the degradation of mustard.

a. Coal

Sulfur in coal is covalently linked to carbon and is an integral component of the structure of coal. Thus it has proved difficult to desulfurize coal by microbial means. Dibenzothiophene (DBT) is universally used as a model compound in studies seeking microorganisms for the removal of sulfur from coal. Many reports of the bacterial degradation of DBT in the presence of a carbon source are available, but there are fewer reports of bacteria using DBT as source of carbon and sulfur. Bacteria cometabolizing DBT include members of the genera <u>Pseudomonas</u> (60-65), <u>Beijerinckia</u> (66), <u>Rhodococcus</u>, and <u>Bacillus</u> (67,68), <u>Corvnevbacterium</u> sp. (69), and unidentified microorganisms (70,71), including yeasts (72). <u>Rhizobium</u> sp., <u>Acinetobacter</u> sp. (59), <u>Brevibacterium</u> sp. (73), <u>Sulfolobus acidocaldarius</u> (74), and an unidentified microorganism isolated from a deep sea thermal vent (75) are able to use DBT or other sulfides as a source of carbon and sulfur.

Two pathways, neither cleaving the carbon-sulfur bond, have been identified for the degradation of DBT: oxidation of the sulfur to the sulfoxide and sulfone, and cleavage of one of the benzene rings (76). All microorganisms studied to date, except for <u>Sulfolobus acidocaldarius</u>, <u>Brevibacterium</u> sp., <u>Corvnevacterium</u> sp., and perhaps several unidentified microorganisms (72,77), degrade DBT by either of the two pathways. The exceptions cleave the carbon-sulfur bond of DBT via the sulfone forming sulfate (69,72,73,77), or incorporate the sulfur into the biomass (68,78). Sources for the isolates were garden soil, environments surrounding petroleum refineries, coal storage areas and strip mines, and oil and coal tar contaminated soil. Enrichment culture techniques, a sulfur bioavailability assay (55), and chemical mutagenesis (55,60) were used for the isolation of the microorganisms.

Although many microorganisms are able to degrade DBT, a suitable process for the microbial desulfurization of coal is not at hand (53-56). One problem is the dissimilarity of the growth conditions of bacteria used for desulfurization: microorganisms for removal of organic sulfur are neutrophilic and those required for inorganic sulfur removal are acidophilic. As a possible solution, attention has recently been focused on the acidophilic heterotroph <u>Acidiphilium</u>, as this bacterium coexists with the acidophilic <u>Thiobacillus</u> used in removal of inorganic sulfur. An interesting approach has been the application of techniques of molecular genetics with the aim of introducing characteristics to enhance desulfurization, such as the ability to degrade organic sulfur compounds, into <u>Acidiphilium</u> (79,80).

b. Oil

DBT and representative organic sulfides present in oil are degraded by species of Pseudomonas, a Flavobacterium sp., a Xanthomonas sp. (81-83), and fungal cultures (84) without releasing sulfur as sulfate. The carbon-sulfur bond of DBT and model compounds are cleaved anaerobically by species of <u>Desulfovibro</u> (85,86). Successful desulfurization of oil has been <u>Desulfovibro</u> (87) unidentified demonstrated with and microorganisms in marine water samples (88). One major obstacle in oil desulfurization is the immiscibility of oil with water. Studies with DBT suggest that degradation would be improved by increasing miscibility with a solvent (81,89). Interesting developments in this area of miscibility is the claim for

desulfurisation of oil by a membrane extract of <u>Bacillus</u> <u>sphaericus</u> and <u>Rhodococcus rhodochrous</u> contained in a organic medium (90), and the use of microemulsions of crude oil and the microorganism or extracts of the microorganism (91).

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c. Application to Mustard Degradation

The studies with oil and coal amply demonstrate the existence of many bacteria with the ability to metabolize or cometabolize organic sulfides. All such bacteria, except for Acidiphilium, are neutrophiles and thus do not possess any advantage over those isolated by Harvey and DeFrank (2) for the degradation of mustard. The acidophile Acidiphilium is deserving of study for mustard degradation. The techniques of selective enrichment in sulfur deficient medium, chemical mutagenesis, and molecular for the introduction of characteristics biology for desulfurization eg. the plasmid-encoded DBT degradation gene (92) into a suitable recipient (acidophiles) are methods with possible application for mustard degradation. Biosurfactants may also be valuable as a solution to problems of mustard-water miscibility.

Bacterial extracts in organic solvents, or microemulsions of mustard and the extracts are other possible approaches. Stabilization of mustard by chloride anions should remain in the forefront of any scheme.

5. YEAST AND FUNGUS

Yeast and fungus have received relatively little attention as microorganisms for the degradation of sulfur compounds. Many studies have made passing mention of the presence of yeast and fungus in isolates of bacteria which degraded sulfur compounds, but it is not clear in these studies if the yeast or fungus possessed the ability to degrade sulfur compounds. Only in isolated instances have the identity of the yeast or fungus been established, presumably because in most studies they constituted a minority of the microorganisms in the isolate. Two yeast and 9 fungal isolates were obtained from oil contaminated soils in the Ogoni area and Ibadan, Nigeria. All were able to grow on petroleum hydrocarbons. The yeasts were not identified and 4 of the 9 fungal isolate were identified as species of Aspergillus, Cochliobolus, Monilia, and Trichoderma (93). Oil was degraded by fungal isolates from soil and the sea from different areas of Kuwait. The isolates were identified as Aspergillus terreus, A. sulphures, Mucor globosus, Fusarium sp. and Penicillium citrinum (94). <u>Aureobasidium pullulans</u>, commonly known as black yeast, has been used to clean oil spills in the ocean and inland waters (95). Although yeast and fungus have been successfully used in the degradation of oil, and solubilization of coal (96-101), it is not clear if the sulfur compounds present in these fuels are broken down and the fate of organic sulfur is unknown.

other fungal species, Mortierella isabellina NRRL 1757, Cunninghamella elegans, and Rhizopus arrhizus are able to oxidize the sulfur of aromatic and heterocyclic sulfides, including the model compound DBT, to the sulfoxide and sulfone (102-104), while dimethyl sulfide was oxidized to the oxide by an unidentified fungus (105). Ligninase, an extracellular enzyme ligninolytic fungus, Phanerochaete produced by the catalyzed the oxidation of thianthrene, a chrysosporium. heterocyclic sulfide present in coal (106). This fungus is able to solubilize coal (107) and it is possible that other oil and coal degrading fungi such as those discussed above may also possesses sulfur oxidation potential. One such fungus, Paecilomyces sp. TLi, solubilized coal under acid conditions, cleaved the carbon-sulfur bonds of DBT degrading it to 2,2'dihydroxybiphenyl, and broke down dibenzylsulfide to a variety of products (108).

The above studies suggest that yeasts and fungi have limited application as microorganisms for the demilitarization of mustard. However, only a few species have been tested for their ability to degrade organic sulfur compounds, and the studies have concentrated on aromatic and heterocyclic sulfides which are recalcitrant to microbial breakdown. The fungi have several attractive attributes that warrant consideration for their use in the degradation of mustard. First, yeasts and fungi are generally acid tolerant and some are halotolerant (109,110). Second, fungi are versatile in substrate utilization, able to metabolize a wide spectrum of foreign substances including chlorinated compounds, polyaromatic hydrocarbons, pesticides and dyes, and have been extensively used in treatment of industrial waste, in waste water treatment, in the degradation of xenobiotics, and in biotechnology (111-114). Third, fungi are the source of a diverse group of substances ranging from macromolecules to simple organic compounds that have industrial use (115). These substances include enzymes and lipids (biosurfactants) some of which are produced extracellularly (116,117), and are of potential use in the breakdown of mustard such as the extracellular enzyme of Phanerochaete chrysosporium (106, 118).

6. **BIOCATALYTIC APPROACHES**

Halophilic and thermophilic microorganisms have not been considered thus far because conditions required for growth present problems such as mustard-water miscibility in the high salt concentrations, and mustard stability in high temperatures, respectively. The enzymes produced by these microorganisms have interesting and unusual properties that encourage a biocatalytic approach to mustard degradation. The intracellular concentrations of salt (KCL and NaCL) in halophilic bacteria are

high and their intra- and extracellular enzymes are able to function under conditions of low water potential. Thus the enzymes are active in aqueous solutions high in salt or organic solvents, and are stable at room temperature (119-123). Enzymes from thermophilic bacteria and fungi are thermostable, show resistance to denaturation by detergents, chaotropes and organic solvents, to proteolytic cleavage, to chemical reactions that result in denaturation, and perhaps, to pH extremes (122-127). These properties suggest that it would be feasible to breakdown mustard enzymatically in non-aqueous media (128,129), or a homogeneous aqueous-organic medium under conditions of pH, salt and mustard concentrations that are consistent with resistance of mustard to hydrolysis. This possibility is open to test with the thermophilic acidophiles <u>S. acidocaldarius</u> and <u>S. solfataricus</u>. These two bacteria degrade sulfides present in coal (130-132), but are unsuitable as organisms for mustard degradation because of the high temperature required for growth. Extracts or lysates of the two bacteria may show activity against mustard when incubated with the agent under suitable conditions in a organic-aqueous medium. Microorganisms whose habitats are rich in reduced sulfur compounds such as thermal sulfur springs, soil in volcanic areas, warm environment surrounding coal mines and coal storage areas, thermal vents in oceans, sludge and waste treatment plants may possess enzyme systems able to degrade organic sulfides (127,133). Screening of extracts or lysates of these microorganisms for mustard degradation activity may be an avenue for exploration.

7. CONCLUSION

The foregoing discussion has identified microorganisms with potential use in the biodegradation of mustard. Success in the biodegradtion of mustard hinges not on tolerance of the microorganism for conditions that enhances mustard stability, but on mustard solubility, and viability of the microorganism in systems that are able to disperse or solubilize mustard. Such systems, whether detergent or organic solvent-based, should be defined and evaluated as a first step. Fungi should receive serious consideration as they often show more resistance to solvent insult than bacteria, and some are producers of detergents and exoenzymes that may be useful in mustard degradation. Enzyme degradation of mustard may eliminate solubility and viability problems.

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