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Novel Regenerated Solvent Extraction Processes for the Recovery of Carboxylic Acids or Ammonia from Aqueous Solutions Part I. Regeneration of Amine-Carboxylic Acid Extracts

L.J. Poole* and C.J. King

(*Ph.D. Thesis)

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NOVEL REGENERATED SOLVENT EXTRACTION PROCESSES FOR THE RECOVERY OF CARBOXYLIC ACIDS OR AMMONIA FROM AQUEOUS SOLUTIONS

PART I. REGENERATION OF AMINE-CARBOXYLIC ACID EXTRACTS

Loree Joanne Poole^{*} and C. Judson King

 $(*Ph.D.$ Thesis)

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March 1990

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NOVEL REGENERATED SOLVENT EXTRACTION PROCESSES FOR THE RECOVERY OF CARBOXYLIC ACIDS OR AMMONIA FROM AQUEOUS SOLUTIONS

by

Loree J. Poole

ABSTRACT

Two novel regenerated solvent extraction processes are examined. The first process has the potential to reduce the energy costs inherent in the recovery of low-volatility carboxylic acids from dilute aqueous solutions. The second process has the potential for reducing the energy costs required for separate recovery of ammonia and acid gases (e.g. CO_2 and H_2S) from industrial sour waters.

The recovery of carboxylic acids from dilute aqueous solution can be achieved by extraction with tertiary amines. An approach for regeneration and product recovery from such extracts is to back-extract the carboxylic acid. with a water-soluble, volatile tertiary amine, such as trimethylamine. The resulting trimethylammonium carboxylate solution can be concentrated and thermally decomposed, yielding the product acid and the volatile amine for recycle.

Experimental work was performed with lactic acid, succinic acid, and fumaric acid. Equilibrium data show near-stoichiometric recovery of the carboxylic acids from an organic solution of Alamine 336 into aqueous solutions of trimethylamine. For fumaric and succinic acids, partial

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evaporation of the aqueous back extract decomposes the carboxylate and yields the acid product in crystalline form. The decomposition of aqueous solutions of trimethylammonium lactates was not carried out to completion, due to the high water solubility of lactic acid and the tendency of the acid to self-associate.

The separate recovery of ammonia and acid gases from sour waters can be achieved by combining steam-stripping of the acid gases with simultaneous removal of ammonia by extraction with a liquid cation exchanger. The use of di-2,4,4-trimethylpentyl phosphinic acid as the liquid cation exchanger is explored in this work.

Batch extraction experiments were carried out to measure the equilibrium distribution ratio of ammonia between an aqueous buffer solution and an organic solution of the phosphinic acid (O.2N) in Norpar 12. The concentration-based distribution ratios increase from 0.11 to 0.46 as the aqueous phase pH increases from 7.18 to 8.15. Regeneration of the organic extractant solution was carried out by stripping at elevated temperatures to remove the ammonia, with 99% recovery of the ammonia being obtained at 125°C.

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CHAPTER 1. INTRODUCTION

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PREFACE

The research presented in this dissertation is a study of two novel regenerated solvent extraction processes. The feasibility of each process is explored through experimental work designed to simulate process steps.

In the first process, carboxylic acids of low volatility are recovered from dilute aqueous solution by extraction with tertiary amines. The organic extract is then regenerated by back-extraction of the acid into aqueous amine solution, followed by thermal decomposition of the alkylammonium carboxylates. The principal applications are for the recovery of organic acids from fermentation broths and waste streams. This process is examined in Chapters 1 through 6.

In the second process, ammonia is recovered from aqueous buffer solutions in the pH range 7.5-9.5 by extraction with an organic liquid cation exchanger. The organic extract is then regenerated by stripping at elevated temperature to remove the ammonia. The principal application is for the treatment of industrial sour waters. This process is examined in Chapters 7 through 12.

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PART I. REGENERATION OF AMINE-CARBOXYLIC ACID EXTRACTS CHAPTER 1. INTRODUCTION

The regeneration of amine-carboxylic acid extracts is a necessary step in the recovery of carboxylic acids from dilute aqueous solutions by extraction with tertiary (e.g., trioctyl) amines. The need to recover carboxylic acids from dilute aqueous solutions arises in many industrial processes. One important example is the production of organic acids by fermentation, in which a major cost is the recovery and purification of the product. Another application is in the processing of aqueous industrial waste or recycle streams.

1.1 Acids Examined

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The acids examined in this work are lactic acid, succinic acid, and fumaric acid. Their formulas and properties are given in Table 1-1. These acids were chosen because they have strong potential for being produced commercially by fermentation $(1,2,3)$. In order for large-scale industrial production of these acids by fermentation to be economically feasible, energy efficient process schemes for recovery of the acids from the broth must be developed (4,5,6).

The production of chemicals by fermentation generally results in dilute aqueous solutions of the product; the product is present at less than 10 wt.% concentration, and usually substantially lower (6). The optimum pH for a particular fermentation process varies according to the particular

Table 1-1. Lactic Acid, Succinic Acid, and Fumaric Acid

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microorganism and substrate used; however, for most fermentations, the pH of the broth must be at or above the first pK_a of the acid.

The recovery of fermentation product organic acids has traditionally been accomplished by the precipitation of the calcium salt of the acid. The salt is formed by the addition of calcium hydroxide to the fermentation broth. The salt crystals precipitate out of solution and are filtered and washed. The salt is then converted to the free acid form by the addition of aqueous sulfuric acid. Calcium sulfate precipitates out of solution and is removed by filtration. The aqueous solution containing the free organic acid is then purified by techniques such as carbon treatment and ion exchange. Acid crystals are recovered from the purified solution upon evaporation of the water. (1)

At the present time, citric acid is the only commodity chemical that is primarily manufactured using a fermentation process (10,11). An alternative acid recovery process in which citric acid is extracted from the fermentation broth into an organic amine solution has recently come into large-scale use (11). The organic extract is regenerated by back-extraction of the acid into water at an elevated temperature. Citric acid crystals are then obtained through evaporative crystallization.

There is a large market for citric acid; it is widely used in the food and pharmaceutical industries (3) and is increasingly being used as a detergent builder (12,13,14). In 1981, the U.S. production of citric acid was 235 million pounds (15).

1.1.1 Lactic Acid

Currently, about half of the world production of lactic acid is by fermentation (16). Before 1950, lactic acid was produced in the United States primarily by fermentation (1); after 1950, production shifted to a chemical synthesis route based on the hydrolysis of lactonitrile (16). Today, the United States imports most of the domestic market demand. In the industrial production of lactic acid by fermentation, concentrations of 12-15 wt.% can be obtained at a pH of between 5 and 7, depending on the microorganism used (16). Lactic acid can be produced at higher concentrations than other fermentation product acids.

More than 50% of all lactic acid produced is used in the food industry, as a food acidulent and as a preservative (16). Lactic acid is also used in the pharmaceutical industry (16) and has potential to be widely used in the manufacture of biodegradable plastics (2,16). The market demand for lactic acid in the U.S. in 1985 was approximately 50 million pounds (2).

1.1.2 Succinic Acid

Succinic acid is not currently being produced commercially by fermentation. Commercial production of succinic acid is generally by the hydrogenation of maleic anhydride or fumaric acid, which are produced by vapor phase oxidation of benzene, butene, or butane with oxygen over V_2O_3 catalyst (3). Currently, all U.S. production is butane-based (17). Studies of

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the fermentative production of succinic acid are being carried out in Japan (3,18,19).

Succinic acid is used in the manufacture of foods and pharmaceuticals, as an intermediate in the synthesis of complex organic compounds, and in the manufacture of resins, pesticides, and other products (3). U.S. production of succinic acid is of the order of one million pounds per year $(4).$

1.1.3 Fumaric Acid

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In the past, fumaric acid has been produced by fermentation in the United States (1). At the present time it is produced by the catalytic oxidation of butane in the vapor phase; either directly or by the initial production of maleic acid followed by isomerization to form fumaric acid (3,17). The fermentative production of fumaric acid resulted in solutions of pH 5 to 6.5 with approximately 3.2 wt.% fumaric acid (20).

In the United States, the major uses of fumaric acid are in the manufacture of paper size resins (40% of use), as a food acidulent (20%), and in the manufacture of unsaturated polyester resins (15%). Due to its lower price, fumaric acid (77.5 cents per pound) competes with citric acid (83.5-89.5 cents per pound [21]) as a food acidulent; however, the lower solubility of fumaric acid limits its use. The current U.S. production capacity for fumaric acid is 39 million pounds per year. The market demand in the U.S. in 1988 was approximately 31 million pounds. (22,23)

1.2 Separation from Dilute Aqueous Solution

Separation by conventional distillation is not economically favorable because the carboxylic acids have low volatilities relative to water. Since the acids are present at low concentration, there would be a high energy cost due to the need to vaporize large quantities of water. Furthermore, there would be no separation of the product acid from other high-boiling impurities.

Separation of the acid from aqueous solution by extraction with conventional solvents is difficult because the aqueous-phase activity coefficient of the acid is low due to its hydrophilicity and its ionic character. In previous investigations, extraction by reversible chemical complexation has been found to be a favorable method for the removal of carboxylic acids from dilute aqueous solution (24,25).

1.3 Extraction by Reversible Chemical Complexation

In extraction by reversible chemical complexation, the extractant pulls the solute into the solvent phase by forming a complex with a functional group of the solute. As long as the stoichiometric ratio is favorable, this complex formation often enables a higher distribution ratio of the acid into the solvent phase than can be achieved using extraction with conventional solvents. The extractant is dissolved in an organic solvent or diluent. Diluents which have functional groups can interact with the complex and affect the distribution ratio of the acid between the aqueous and organic phases.

Long-chain aliphatic amines have been found to be good extractants for carboxylic acids (25,26,27). Alamine 336 (a long-chain tertiary amine, Henkel Corp.) has been used as the extractant in this work because of low water solubility, thermal stability, and commercial availablity. Methyl isobutyl ketone (MiBK) is used as the diluent.

Extraction of an acid by reversible chemical complexation can be described by the following reaction, in which B is the extractant in the organic phase and A is the un-ionized acid (originally present in the aqueous phase):

pA (aqueous) + qB (organic) =
$$
A_p B_q
$$
 (organic) (1-1)

Reversible complexation is driven by the amount of un-ionized acid in the aqueous solution and is therefore favored at low pH, the optimum pH being related to the pK_a of the acid.

The heterogeneous equilibrium constant for the reaction, $K_{p,q}$, can be defined as follows:

$$
K_{p,q} = \frac{[A_p B_q]_{org}}{[A]_{aq}^p [B]_{org}^q}
$$
 (1-2)

In MiBK diluent, and for the particular acids being examined, carboxylic acid/ Alamine 336 complexes with more than one acid molecule per molecule of amine are formed (24) . Table 1-2 gives the values of the equilibrium constants determined by Tamada (24) for the systems examined in this work.

1.4 Regeneration Methods

Once the acid has been extracted into the organic solvent phase, a second step or series of steps is needed to recover the acid in product form and to regenerate the solvent phase so that it can be recycled back to the extractor. Distillation can be used for sufficiently volatile carboxylic acids, such as acetic (25), but is not suitable for the much less volatile acids considered here. Two possible regeneration methods (27) studied in more detail by Tamada (24) involve swings of temperature and/or diluent composition. In these methods, the acid is back-extracted into an aqueous phase by taking advantage of the change in the equilibrium distribution ratio that occurs with changes in temperature and/or composition of the diluent.

1.4.1 Temperature Swing

It has been found that the values of the equilibrium constants decrease with an increase in temperature (24,27). Therefore a process can be run, as shown in Figure I-I, in which the extraction takes place at ambient temperature and back-extraction into water takes place at a higher temperature. The end-product is a somewhat more concentrated aqueous solution of the acid which can be sent to an evaporative crystallizer. In the production of citric acid by fermentation, recovery of the product can

Table 1-2. Equilibrium Constants for Carboxylic Acid/ Alamine 336 (O.3M) in MiBK Systems

 $(K_{p,q}$ values have units of $[mol/l]^{1-(p+q)}$

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(p,q) = (1,1) (1,2) (2,1) (2,2) (3,1) Lactic Acid 1.31 1.52 1.17 ^t Succinic Acid 1.39 1.91 , ; [~] Fumaric Acid --- 4.33 4.81 7.49 .,

(from Tamada, 1989)

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Figure 1-1. Temperature-Swing Regeneration

Figure 1-2. Diluent-Swing Regeneration

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be obtained by extraction into a solvent mixture of trilaurylamine, noctanol and a C_{10} or C_{11} isoparaffin, followed by back-extraction of the acid into water at an elevated temperature (27).

1.4.2 Diluent Composition Swing

The type of diluent that is used affects the structures of the complexes that are formed between the acid and the extractant, as well as the values of the equilibrium constants. A change in the composition of the diluent will change the overall equilibrium distribution of the acid between the aqueous and organic phases. In one form of diluent-swing regeneration, as shown in Figure 1-2, the diluent used in the solvent stream entering the extractor is a combination of two components; one that promotes the forward extraction and one that is an inert diluent. When the solvent stream enters the regenerator, the diluent consists of only the inert component. The composition of the diluent is altered between the extractor and the regenerator by use of a distillation column.

A desirable goal for both of these regeneration processes, temperature-swing and diluent-swing, is to increase the concentration of the carboxylic acid in the aqueous phase. The diluent-swing process can incorporate the temperature-swing effects (by running the two columns at different temperatures) to achieve a greater concentration of the acid.

1.4.3 Displacing Acid

Another regeneration alternative is one proposed by Rückl et. al. (28) for the recovery of citric acid. A tertiary amine extractant in an organic diluent is used to extract citric acid from an aqueous feed stream. The organic solvent phase containing the citric acid is then contacted in a reextraction column with an aqueous phase containing a volatile acid, e.g., acetic acid. There is a transfer of citric acid and acetic acid between the two phases. The acetic acid is then removed from the solvent phase by distillation and recycled back to the reextraction column. The regenerated solvent phase can be reused to extract citric acid from the aqueous feed stream.

1.4.4 Precipitation

Starr (29) has explored a regeneration method in which extracted acids with low solubility in water, e.g. fumaric acid, are recovered from the organic phase by precipitatioq from the extract. solution is heated and precipitation of the acid occurs as the volatile diluent is driven off. The organic extract

1.4.5 pH-Swing

Regeneration of amine-carboxylic acid extracts can be achieved by back-extraction of the acid into an aqueous solution of a strong base such as NaOH; however, this approach results in a salt of the acid rather than 12

what is usually the desired product of the pure acid. Another acid could be added to the aqueous solution of the salt to convert the salt back to the free acid form. However, this regeneration method has the disadvantages of consuming both an acid and a base and producing a salty waste stream.

1.4.6 pH-Swing with a Volatile Base

In this work, a regeneration method involving recycle of a volatile base is examined. As shown in Figure 1-3, the acid is back-extracted from the loaded organic phase into an aqueous trimethylamine solution. The aqueous solution of the trimethylammonium salt is then heated to drive off the water and the trimethylamine, leaving the acid in crystalline form. Trimethylamine can then be recycled, thereby avoiding costs for make-up chemicals and disposal of by-products.

In order to determine the feasibility of this regeneration method, experiments were carried out in three areas. The results of experiments in which the acids were back-extracted from organic amine solutions into aqueous solutions of trimethylamine are discussed in Chapter 4. Chapter 5 discusses the results of concentration and heating experiments in which the water and trimethylamine are driven off from aqueous solutions of the trimethylammonium carboxylates, as well as the work done in washing and further purification of the crystalline acid product. The experimental procedures and analytical methods used are described in Chapter 3.

Figure 1-3. Regeneration using Recycle of a Volatile Base

In Chapter 2, the criteria used in selecting trimethylamine as the volatile base to be used in the process are reviewed. In Chapter 6, the economic viability of this regeneration method is examined and compared to that of other regeneration alternatives.

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CHAPTER 2. SELECTION OF THE VOLATILE BASE

This chapter discusses the criteria used in selecting an appropriate volatile base for back-extraction of carboxylic acids. Various classes of low molecular weight amine compounds are examined with respect to the desired characteristics of the volatile base. Low molecular weight tertiary amines possess the desired characteristics, with trimethylamine being suitable for exploratory use in the regeneration process.

2.1 Desired Characteristics

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The regeneration method examined in this work involves backextraction of the acid into an aqueous solution of a volatile base, followed by concentration and thermal decomposition of the resulting salt. The base used must be water-soluble, have a substantially higher volatility than that of the acid, and not form a thermodynamically irreversible bond with the acid. The base must have sufficient basic strength to compete effectively with the organic extractant (Alamine 336) for the acid. The base must be thermally stable at whatever temperature is required to crack the salt.

2.2 Ammonia

Ammonia is a relatively strong $(pK_b = 4.75$ at 25°C [1]), watersoluble, volatile base. However, the heating of a solution of ammonia and a carboxylic acid results in amide formation (2). Upon further heating, nitrile formation occurs (3). Shukla et al. (4) prepared lactamide by passing a stream of dry ammonia through molten lactic acid at 150°C:

$$
\text{CH}_{3}\text{-}\text{CHOH-COOH} + \text{NH}_{3} \rightarrow \text{CH}_{3}\text{-}\text{CHOH-CONH}_{2} + \text{H}_{2}\text{O} \tag{2-1}
$$

In the present work, lactamide was produced by heating a concentrated aqueous solution of ammonium lactate. An aqueous solution containing 6.28M ammonium lactate was heated at 120°C for 30 minutes. Samples of the solution were analyzed before and after heating by high performance liquid chromatography (HPLC) using an organic acid analysis column (Biorad AMINEX Ion Exclusion HPX-87H) and a differential refractometer detector (Waters, R40l). Fifteen percent of the lactic acid was converted to lactamide during the heating process.

Ammonia is therefore not a desirable choice for the volatile base because it forms a covalent bond with the acid. The covalent amide bond can be broken by reaction with nitrous acid (2):

$$
R\text{-CONH}_2 + HNO_2 \overset{H_2SO_4}{\rightarrow} R\text{-COOH} + N_2 + H_2O \tag{2-2}
$$

or hydrolysis of the amide can be catalyzed by acid or base under "vigorous conditions" (2,5):

$$
R\text{-CONH}_{2} \xrightarrow{H_{2}SO_{4}} R\text{-COOH} + NH_{3}
$$
\n
$$
H_{2}O
$$
\n(2-3)

The first approach consumes nitrous acid and does not regenerate ammonia. The second would be expensive in practice.

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2.3 Primary and Secondary Amines

Primary and secondary amines would also react with carboxylic acids to form amides under the anticipated conditions of the regeneration process. Mitchell and Reid (3) formed amides by passing dimethylamine gas through heated aliphatic acids. Again, the production of a covalent bond makes these compounds undesirable for use as the volatile base.

2.4 Tertiary Amines

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Tertiary amines cannot form amides with carboxylic acids. Urbas (6,7) discusses the cracking of various trialkylammonium salts, with the tertiary amine being either tributylamine or dicyclohexylmethylamine and the acid being either acetic, propionic, butyric, lactic, or citric acid. Depending on the relative volatilities, either the amine or the acid was driven off from concentrated aqueous solutions of the trialkylammonium. salt or from a chloroform extract of the salt.

In the regeneration method examined in this work, the first step is the back-extraction of the acid from the organic solvent phase into the aqueous basic solution. It is desirable that the acid be concentrated during this step. From this standpoint, the best low molecular weight tertiary amine for use in this process is the one which is the most water soluble; i.e., trimethylamine. Trimethylamine is commercially available as a 25 wt.% aqueous solution.
Trimethylamine (TMA) also has the desired characteristics of high volatility (b.p. = 2.9°C [1]), and strong basicity (pK_b = 4.1 at 25°C [8]). The thermal stability of TMA was explored by Kaufman (9) who found that thermal decomposition (loss of a methyl group) occurs above 350° C (9,10). As will be seen, that high a temperature is not required to crack the trialkylammonium salts of TMA and succinic acid or fumaric acid.

In the carboxylic acid recovery process shown in Figure 1-3, TMA is to be recycled from the evaporative crystallizer to the back-extraction column. The evaporative crystallizer should be run under a nitrogen atmosphere and/or at temperatures low enough so that oxidation of TMA will not occur. The conditions under which oxidation of TMA can occur were investigated by Jones and Gesser (10) and Cullis and Waddington (11).

Cullis and Waddington (II) carried out gaseous oxidation of TMA at temperatures of 165°C and 170°C using initial partial pressures of: 100 mm TMA with 100 mm O_2 ; 50 mm TMA with 250 mm O_2 ; and other combinations. The reaction products were formaldehyde and dimethylamine. Jones and Gesser (10) carried out the oxidation of TMA at 200°C with initial partial pressures of 50 mm O_2 and 200 mm of an equilmolar mixture of TMA and deuterated TMA.

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CHAPTER 3. EXPERIMENTAL PROCEDURES AND EQUIPMENT

Three types of experiments were performed:

1. experiments in which the acids were back-extracted from organic amine solutions into aqueous solutions of trimethylamine;

2. concentration and heating experiments in which the water and trimethylamine were driven off from aqueous solutions of the trimethylammonium carboxylates; and

3. experiments aimed at washing and further purifying the crystalline acid product.

This chapter describes the experimental procedures and analytical methods used in these experiments.

3.1 Reagents Used

The reagents used and their sources are given in Table 3-1. Distilled water that had been passed through a Milli-Q water purification system (Millipore Corp.) was used to dilute the lactic acid and trimethylamine (TMA) solutions when necessary and to make aqueous solutions of succinic and fumaric acid. The 25 wt.% aqueous TMA solution was stored in the original 3.5 liter glass container at 4°C to prevent volatilization of the TMA. Before each use, the aqueous TMA solution was brought to room temperature, and the TMA concentration was measured by first adding an excess of $6N$ H₂SO₄ to a 5 ml sample of the TMA solution and then backtitrating with 0.1N NaOH. Methyl red was used as an indicator.

Table 3-1. Source and Description of Chemicals Used

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 $\tilde{\mathbf{g}}_0^{(1,1)}$

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 \mathcal{A}

 \bar{z}

3.2 **Equipment**

High performance liquid chromatography (HPLC) analysis was carried out using a Spectra-Physics SP 8000B chromatograph and a differential refractometer detector (Waters, R40I). Either an organic acid analysis column (Biorad AMINEX Ion Exclusion HPX-87H) or a C₁₈ Radial-Pak μ -Bondapak column (Waters) contained in a radial compression module (Waters, RCM-100) was used. The mobile phase was $0.01N H_2SO_4$.

Gas chromatography (GC) analysis was carried out using a Varian model 3700 chromatograph with an FID detector and a Hewlett-Packard 3390A integrator. Samples $(1 \mu l)$ were injected into a GC column packed with 5% silicone 0V-17 on acid-washed, DMCS-treated, 80/100 mesh Chromosorb W (Alltech Associates). The column was 5 ft. stainless steel tubing with an outer diameter· of 1/8 in. The injector temperature was 290°C. The column temperature was held at 35°C for 2 minutes and then increased to 240°C at a rate of 10°C/min. Nitrogen was used as the carrier gas.

Aqueous phase pH values were determined using an Orion 701A pH meter equipped with an Orion semimicro Ross pH electrode. Flasks containing two-phase extraction systems were placed in a shaker bath- either a Fisher Scientific, Versa-Bath S or a Lab-Line, Orbit Water Bath Shaker.

3.3 Dilution and Heating of Aqueous Lactic Acid Solutions

Lactic acid forms intermolecular esters in concentrated aqueous solution. The hydroxyl group on the 2nd carbon reacts with the acidic COOH group of another acid to form lactoyllactic acid. Reaction with additional acid molecules results in polymer formation. Figure 3-1 shows the equilibrium composition of aqueous lactic acid solutions of varying concentrations (1).

Below 20 wt.%, primarily the lactic acid monomer is present. (At 30 wt.% total lactic acid, 0.86 wt.% lactoyllactic acid is present; while at 15 wt.% total lactic acid, only 0.55 wt.% lactoyllactic acid is present.) If an aqueous solution of the monomer is desired, it can be produced by dilution of the 85 wt.% aqueous solution. However, the kinetics of hydrolysis of the esters are slow. It has been found that heating of the diluted solution increases the rate of hydrolysis and that the equilibrium composition can be reached by boiling the freshly diluted solution for 12 hours. (1)

In the present work, 2.0M lactic acid in water was made by dilution of the 85 wt.% aqueous solution. The resulting solution was boiled with total reflux for 12 hours. The complete hydrolysis of the esters was confirmed by HPLC analysis using both the organic acid analysis column and the C_{18} column.

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Figure 3-1. Equilibrium Composition of Aqueous Lactic Acid (from Holten (1))

Concentration in wt % in terms of equivalent quantities of lactic acid

- A: Free lactic acid
- B: Total polylactic acids
- C: Lactoyllactic acid (dimer)
- D: Lactoyllactoyllactic acid (trimer)
- E: Sum of higher polylactic acids

3.4 Preparation of Amine-Carboxylic Acid Extracts for use in Back-Extraction Experiments

Before the back-extraction experiments could be performed, it was necessary to prepare the organic amine-carboxylic acid extracts. This was done at 25°C by contacting equal volumes of an aqueous solution of the acid with an organic solution of 0.3M Alamine 336 in methyl isobutyl ketone (MiBK), using the experimental results of Tamada (2) or Starr (3) as a guide to predicting the resultant concentration of acid in the organic phase.

For lactic acid, the two phases were contacted in a separatory funnel and then transferred to Erlenmeyer flasks which were placed in a 25°C shaker bath. The flasks were left in the bath (set at approximately 40 oscillations per minute) for 2 days to allow the phases to settle. Vigorous shaking in a separatory funnel of an aqueous solution of succinic acid with the organic amine solution was found to result in emulsion formation; therefore, for both succinic acid and fumaric acid, the two phases were placed directly in Erlenmeyer flasks. The flasks were put in a 25°C shaker bath set at approximately 80 oscillations per minute for 2 to 3 days to allow transfer of the acid to occur across the interface.

In order to obtain sufficiently high final concentrations of fumaric acid, it was necessary for the amount of fumaric acid present initially to exceed the aqueous solubility limit. Thus, when the flasks were placed in the shaker bath, the aqueous phase contained undissolved fumaric acid crystals. As the fumaric acid was transferred from the aqueous to the

organic phase, these crystals dissolved into the aqueous (bottom) phase.

Upon removal of the flasks from the shaker bath, the phases were separated by removal of each phase by pipet. The pH of the aqueous phase was measured. The amount of acid remaining in the aqueous phase was determined by titration with O.OIN or O.IN NaOH, using phenolphthalein as an indicator. The concentration of acid in the organic phase was determined by two-phase titration with O.OIN or O.IN NaOH, using phenolpthalein as an indicator. A blank titration of 0.3M Alamine 336 in MiBK was also performed. Material balances were within 1-2.5%.

3.5 Back-Extraction Experiments

Experiments were carried out in which the acid was back-extracted from the Alamine 336 extract into aqueous solutions of varying TMA concentration. These back-extraction experiments were performed at 25°C. The two phases were contacted in a separatory funnel using an organic to aqueous phase ratio of $8/3$ (v/v). The two phases were then transferred to Erlenmeyer flasks. The flasks were placed in a 25°C shaker bath for 2 days to allow the phases to settle.

Upon removal of the flasks from the shaker bath, the phases were separated by pipet. GC analysis of the organic phase was used to determine if any TMA had been transferred from the aqueous to the organic phase, and if so how much. The pH of the aqueous phase was measured, and the concentration of acid in the aqueous phase was determined by HPLC analysis using the C_{18} column.

The concentration of acid remaining in the organic phase was determined by HPLC analysis of a NaOH extract of the organic phase. The C_{18} column was used. The NaOH extract was obtained by contacting a 5 ml sample of the organic phase with 10 ml of O.3N, 0.34N or 0.53N NaOH in a centrifuge tube. magnetic stir bar. The two phases were contacted by stirring with a The two phase system was then centrifuged for 30 minutes at 2000 rpm to separate the phases. The NaOH solution was concentrated enough to effect essentially complete transfer of the acid from the organic phase to the aqueous phase, as verified by tests in which acid was back-extracted from organic solutions of known acid concentration.

3.6 Concentration and Thermal Decomposition of Alkylammonium

Carboxylates

Aqueous solutions of trimethylammonium carboxylates were heated to vaporize the water and TMA and recover the carboxylic acid in crystalline form. The heating was performed under nitrogen to prevent possible oxidation of the TMA. The apparatus used in these experiments is shown in Figure 3-2.

The aqueous solution (60 ml) was placed in a 3-neck 100 ml roundbottom flask and stirred with a magnetic stir bar. The flask was heated using a heating mantle (Glas-Col Apparatus Co.) attached to a voltage regulator (The Superior Electric Co., Powerstat). The water was driven off

 $\ddot{}$

from the solution, and collected in a graduated cylinder. collected in an absorber flask containing dilute H_2SO_4 . The TMA was

÷,

During the experiment, the rates of evolution of water and TMA were to be determined; therefore, the heating was performed under vacuum to permit separate recovery of water and TMA for data-taking purposes. Under vacuum, after the TMA was removed from solution, it absorbed only slightly (see below) in the water that was present in the condenser and in the graduated cylinder.

At the beginning of an experimental run, the absorber flask was filled with 400 ml of H_2SO_4 concentrated enough to neutralize 10% of the TMA initially present in the aqueous trimethylammonium carboxylate solution. Methyl red was added to the solution to serve as an indicator. When 10% of the TMA had been absorbed, the solution changed in color from red to clear. At this point, the time was recorded and then 5 ml of concentrated $H₂SO₄$ (80 times the concentration of the initial absorber solution) were injected into the absorber flask, causing the solution to return to red in color. This process was repeated throughout the experimental run.

Nitrogen was bubbled through the aqueous trimethylammonium carboxylate solution. The nitrogen flow rate was measured with a rotameter encased in a plastic shield (Omega Engineering, Inc., model FL-Ill). To keep the system under vacuum, a General Electric 1/4 hp vacuum pump and a vacuum regulator (Weiss) were used.

Throughout the experimental run, the volume of water collected was recorded, as a function of time. The pressure in the system was monitored with a mercury manometer at two points -- after the rotameter; and after the cylinder used to collect the water. Any fluctuations in the rotameter reading were recorded.

The temperature of the aqueous trimethylammonium carboxylate solution was monitored throughout the run. The temperature was allowed to increase gradually (reflecting the increase in boiling temperature of the solution) until a point was reached when most of the water had been removed. At this point, a sharp increase in temperature was observed. In order to prevent thermal decomposition of the acid itself, the voltage regulator was adjusted as needed to keep the temperature below 130°C.

At the end of the run, a sample of the water collected was titrated with $0.03N$ H₂SO₄ in order to determine the amount of TMA that had absorbed into the water. When the experiment was conducted under vacuum (absolute pressure of system ranging between 14 and 15 in. Hg), less than 2% of the total TMA was present in the water. A sample of the final absorber solution was titrated with O.OlN or O.IN NaOH in order to determine the amount of TMA that had been absorbed between the last color change and the end of the experimental run.

The moles of TMA collected versus time, volume of water collected versus time, and nitrogen flow rate were used to determine the partial pressures of water vapor and TMA over the aqueous trimethylammonium carboxylate solution as functions of temperature and of the concentrations of salt and free acid in solution. These values may be close to equilibrium, but no special efforts were made to assure that. The data were also used to determine rates of TMA and water removal from the solution as functions of time.

3.7 Washing and Further Purification of Acid Crystals

After an experimental run in which a trimethylammonium carboxylate salt (of either succinic acid or fumaric acid) was concentrated and thermally decomposed as described above, there remained behind a mixture- - light-yellow in color -- of acid crystals, water, and residual TMA. Various techniques were used to wash and further purify the crystals.

One method tried was to hold the mixture at 120°C under vacuum for 6 hours, in an attempt to drive off the remaining water and TMA from the mixture. However, this treatment resulted in solidification of the viscous liquid containing the acid crystals. An elemental analysis of this material was carried out to determine its composition and in particular the amount of residual TMA. The amount of residual TMA determined by this method was confirmed by a material balance (i.e., the amount of TMA originally present in the aqueous solution, minus that amount collected in the absorber flask and the water collection flask).

The elemental analysis of the solidified material was conducted by V. Tashinian of the Microanalytical Laboratory, College of Chemistry, at the University of California, Berkeley. The carbon, hydrogen, and nitrogen contents were measured. The analysis method involved use of a Perkin-Elmer Model 240 Elemental Analyzer in which samples were burned with an excess of oxygen to form CO_2 from carbon, H_2O from hydrogen, and N_2 from nitrogen. The concentrations of these compounds were determined using a thermal conductivity detector.

Another method used to clean the crystals was to wash the mixture with various organic solvents. The mixture was transferred to a piece of filter paper (Whatman, #50) contained in a buchner funnel. The buchner funnel was placed on top of a filter flask as shown in Figure 3-3. The mixture was washed with solvent until the crystals were white in color. The crystals were then dried and weighed and sent to the Microanalytical Laboratory for elemental analysis.

The disadvantage of this washing method was that in order to clean the crystals effectively, it was necessary to use solvents in which the acid had a significant solubility. During the washing process, a portion of the acid crystals dissolved into the solvent and was washed through the filter paper. The solvent was evaporated under vacuum from the solvent/acid mixture present in the filter flask, and then the acid crystals were scraped off the bottom of the filter flask. These crystals were then placed on the filter paper and washed to obtain a subsequent batch of purified crystals. This process had to be repeated up to 6 times (generating 6 separate batches of crystals, each batch weighing between 5-30% of the total acid) in order to obtain a good (greater than 70 wt.%) recovery of the acid in

Figure 3-3. Equipment used to wash the Acid Crystals

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purified crystalline form.

The third, and most successful, method used to treat the crystals was to dissolve the mixture in MiBK, then drive off the solvent, water, and residual TMA by heating under vacuum. The mixture was first dissolved in acetone and then transferred to the filter flask used previously. The buchner funnel was placed on the flask and then the acetone was removed by suction through the vacuum line.

The material (acid, water, and TMA) left behind after the acetone was removed was· dissolved in approximately 200 ml of MiBK. The filter flask was placed on a combination heater/stirrer (Thermolyne, Nuova II), and the solution was stirred with a magnetic stir bar. The solution was heated to 80°C, and, as the solvent evaporated, it was pulled through the vacuum line. The crystals that remained after the solvent was removed were dry; there was no visual evidence of water being present. The crystals were collected and sent to the Microanalytical Laboratory for elemental analysis.

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CHAPTER 4. RESULTS OF THE BACK-EXTRACTION EXPERIMENTS

This chapter discusses the experimental results for the backextraction of lactic, succinic, and fumaric acids from organic extract solutions into aqueous trimethylamine (TMA) solutions. The extract solutions consisted of the acid, the extractant, and the diluent. The extractant was Alamine 336 (Henkel), a tertiary amine with aliphatic chains of 8 to 10 carbons in length, and the diluent was methyl isobutyl ketone (MiBK). During the back-extraction experiments, the concentration of acid in the initial organic phase was held constant while the TMA concentration was varied. For all three acids, essentially 100% of the acid was backextracted into the aqueous phase at conditions in which there was one molecule of TMA for every equivalent weight of acid. This indicates that the basicity of the aqueous TMA is much stronger than that of the organic Alamine 336.

4.1 Modeling of the Back-Extraction

The back-extraction of the acid from the organic phase into an aqueous solution of trimethylamine can be modeled by solving simultaneously the chemical equilibrium and mass balance equations that describe the system. The chemical equilibrium equations describe the complex formation that occurs in the organic phase as discussed in Section 3 of Chapter I, and the dissociation of the acid and of TMA in the aqueous phase.

The solubility of the acid in the amine-free diluent can be accounted for by use of the partition coefficient, P, which is defined as follows:

$$
P = \frac{\text{concentration of uncomplexed acid in organic phase}}{(4-1)}
$$
\n
$$
P = \frac{(4-1)(4-1)}{(4-1)}
$$

where both concentrations have units of moles per liter. The partition coefficients for the three acids between water and MiBK are given in Table 4-1.

The organic solvent phase used in the back-extraction experiments consisted of 0.3M Alamine 336 in MiBK. Acid is present in the organic extract solution as a result of two interactions, acid-diluent and acid-amine. The acid concentration resulting from the acid-diluent interaction is taken to be equal to the product of the partition coefficient, the concentration of un-ionized acid in the aqueous phase, and a correction factor, Φ . The correction factor assumed and used in this work is the volume fraction diluent in the solvent mixture.

4.1.1 Lactic Acid

The back-extraction at 25°C of lactic acid from the organic extract solution into aqueous TMA solutions is described by the following equations:

(i) Equations describing transfer of lactic acid between aqueous and organic phases (1):

$$
HA + \overline{A336} = (\overline{HA})\overline{A336} \qquad \log K_{1,1} = 1.31 \tag{4-2}
$$

$$
2HA + \overline{A336} = (\overline{HA})_2 A336 \qquad \log K_{2,1} = 1.52 \tag{4-3}
$$

$$
3HA + \overline{A336} = (\overline{HA})_3 A336 \qquad \log K_{3,1} = 1.17 \tag{4-4}
$$

$$
P_1 = \text{partition of } \text{lactic acid} = \frac{(HA)}{(HA)} = 0.11 \qquad (4-5)
$$
\n
$$
\frac{(HA)}{(HA)}
$$

where HA denotes the un-ionized acid, A336 denotes the extractant (Alamine 336), the overbar denotes the organic phase, and the $K_{p,q}$ values are the equilibrium constants of the complex formation as determined by Tamada (1) and listed in Table 1-2.

(ii) Equations describing equilibria in the aqueous phase:

$$
HA = H^{+} + A^{-} \qquad \qquad -\log K_{a} = 3.85 \text{ (3)} \tag{4-6}
$$

$$
R_3NH^+ = R_3N + H^+ \qquad \text{-log } K_a = 9.8 \quad (3)
$$
 (4-7)

$$
(H+) + (R3NH+) = (A-) + (OH-)
$$
 (4-8)

where R_3N denotes TMA and Equation 4-8 describes the charge balance in the aqueous phase. The parentheses in Equations 4-5 and 4-8 denote concentrations.

The concentration of un-ionized acid in the aqueous phase can be expressed as the product of the total acid in the aqueous phase and the quantity α_1 , where:

$$
\alpha_1 = \frac{1}{1 + \frac{K_{a, \text{acid}}}{(H^+)}} \tag{4-9}
$$

Then the mass balance for the lactic acid can be written as follows:

$$
0 = [HA]_{init} - K_{1,1}[B](HA) - 2K_{2,1}[B](HA)^{2} - 3K_{3,1}[B](HA)^{3}
$$

$$
-\Phi P_1(HA) - WS(HA)/\alpha_1 \qquad (4-10)
$$

where the first term is the initial concentration of acid in the organic phase, the parentheses denote concentrations in the aqueous phase, the brackets denote concentrations in the organic phase, B denotes the uncomplexed Alamine 336 molecule, Φ is equal to the volume fraction of MiBK (0.85) in the organic solvent, and WS is equal to the water to solvent ratio (3/8) used in the back-extraction experiments.

The mass balance for the Alamine 336 (essentially all present in the organic phase) can be written as follows:

Uncomplexed A336 = Total A336 - Complexed A336
\n[B] =
$$
[B_{total}] - K_{1,1}[B](HA) - K_{2,1}[B](HA)^2 - K_{3,1}[B](HA)^3
$$
 (4-11)

or:

$$
[B] = \frac{[B_{\text{total}}]}{1 + K_{1,1}(HA) + K_{2,1}(HA)^{2} + K_{3,1}(HA)^{3}}
$$
(4-12)

Equation 4-12 can be substituted into Equation 4-10 to give a mass balance for the acid in terms of only two unknown values, the pH of the aqueous raffinate (incorporated into α_1) and the concentration of un-ionized acid in the aqueous phase. A computer program (given in Appendix C) was used to solve the resulting equation for (HA) at set pH values using a Newton Raphson minimization routine.

For each pH value, the recovery of the acid into the aqueous phase was calculated as:

$$
Recovery = \frac{\text{WS}(HA)/\alpha_1}{[HA]_{init}}
$$
 (4-13)

and the ratio of moles of TMA to moles of acid originally present in the organic phase was calculated where:

$$
(R_3N)_{\text{total}} = (R_3N) + (R_3NH^+) = (R_3NH^+)/\beta \tag{4-14}
$$

where:

$$
\beta = \frac{1}{1 + \frac{K_{a,R3NH}^{+}}{(H^{+})}}
$$
(4-15)

and the value of (R_3NH^+) is determined using Equations 4-6 and 4-8. In this manner, a theoretical back-extraction curve for lactic acid was generated as shown in Figure 4-1.

4.1.2 Succinic Acid

The back-extraction at 25°C of succinic acid from the organic extract solution into aqueous TMA solutions is described by the following equations:

(i) Equations describing transfer of succinic acid between aqueous and organic phases (I):

$$
H_2A + A_336 = (H_2A)A_336 \qquad \log K_{1,1} = 1.39 \tag{4-16}
$$

$$
2H_2A + A_336 = (H_2A)_2A_336 \qquad \log K_{2,1} = 1.91 \tag{4-17}
$$

$$
P_2 = \text{partition of succinct acid} = \frac{(H_2 A)}{(H_2 A)} = 0.19 \qquad (4-18)
$$

between water and MiBK
$$
\frac{(H_2 A)}{(H_2 A)}
$$

where H_2A denotes the un-ionized succinic acid, the overbar denotes the organic phase, and the $K_{p,q}$ values are the equilibrium constants of the complex formation as determined by Tamada (I) and listed in Table 1-2.

(ii) Equations describing equilibria in the aqueous phase:

$$
H_2A = H^+ + HA^-
$$
 -log K_{a1} = 4.21 (3) (4-19)

$$
HA^{-} = H^{+} + A^{2-} \qquad \qquad -\log K_{a2} = 5.60 (3) \qquad (4-20)
$$

$$
R_3NH^+ = R_3N + H^+ \t -log K_a = 9.8 \t (3) \t (4-21)
$$

$$
(H+) + (R3NH+) = 2(A2-) + (HA-) + (OH-)
$$
 (4-22)

Equation 4-22 describes the charge balance in the aqueous phase. The parentheses in Equations 4-18 and 4-22 denote concentrations.

The concentration of un-ionized succinic acid in the aqueous phase can be expressed as the product of the total succinic acid in the aqueous phase and the quantity α_2 , where:

$$
\alpha_2 = \frac{1}{1 + \frac{K_{a1}}{(H^+)} + \frac{K_{a1}K_{a2}}{(H^+)^2}}
$$
(4-23)

Then the mass balance for the succinic acid can be written as follows:

$$
0 = [H_2A]_{init} - K_{1,1}[B](H_2A) - 2K_{2,1}[B](H_2A)^2
$$

$$
-\Phi P_2(H_2A) - WS(H_2A)/\alpha_2 \quad (4-24)
$$

where the first term is the initial concentration of acid in the organic phase, the parentheses denote concentrations in the aqueous phase, the

brackets denote concentrations in the organic phase, B denotes the uncomplexed Alamine 336 molecule, Φ is equal to the volume fraction of MiBK (0.85) in the organic solvent, and WS is equal to the water to solvent ratio (3/8) used in the back-extraction experiments.

The mass balance for the Alamine 336 (essentially all present in the organic phase) can be written as follows:

Uncomplexed A336 = Total A336 - Complexed A336

$$
[B] = [Btotal] - K1.1[B](H2A) - K2.1[B](H2A)2
$$
 (4-25)

or:

$$
[B] = \frac{[B_{\text{total}}]}{1 + K_{1,1}(H_2A) + K_{2,1}(H_2A)^2}
$$
 (4-26)

Equation 4-26 can be substituted into Equation 4-24 to give a mass balance for the acid in terms of only two unknown values, the pH of the aqueous raffinate (incorporated into α_2) and the concentration of un-ionized acid in the aqueous phase. A computer program (given in Appendix C) was used to solve the resulting equation for (H_2A) at set pH values using a Newton Raphson minimization routine.

For each pH value, the recovery of the acid into the aqueous phase was calculated as:

$$
Recovery = \frac{WS(H_2A)/\alpha_2}{[H_2A]_{init}} \tag{4-27}
$$

and the ratio of moles of TMA to moles of acid originally present in the organic phase was calculated where, as before (Equation 4-14):

$$
(R_3N)_{\text{total}} = (R_3N) + (R_3NH^+) = (R_3NH^+)/\beta \tag{4-28}
$$

and the value of (R_3NH^+) is determined using Equations 4-19, 4-20, and 4-22 (β was defined in Equation 4-15). In this manner, a theoretical backextraction curve for succinic acid was generated as shown in Figure 4-2.

4.1.3 Fumaric Acid

The back-extraction at 25°C of fumaric acid from the organic extract solution into aqueous TMA solutions is described by the following equations:

(i) Equations describing transfer of fumaric acid between aqueous

and organic phases (I):

$$
H_2A + 2A336 = (H_2A)(A336)_2 \qquad \log K_{1,2} = 4.33 \tag{4-29}
$$

$$
2H_2A + A_336 = (H_2A)_2A_336 \qquad \log K_{2,1} = 4.81 \tag{4-30}
$$

$$
2H_2A + 2A336 = (H_2A)_2(A336)_2 \text{ log K}_{2,2} = 7.49 \tag{4-31}
$$

$$
P_3 = \text{partition of } f \text{umaric acid} = \frac{(H_2 A)}{(H_2 A)} = 2.2 \tag{4-32}
$$

between water and MiBK

where H_2A denotes the un-ionized fumaric acid, the overbar denotes the organic phase, and the $K_{p,q}$ values are the equilibrium constants of the complex formation as determined by Tamada (l) and listed in Table 1-2.

(ii) Equations describing equilibria in the aqueous phase:

$$
H_2A = H^+ + HA^-
$$
 -log K_{a1} = 3.10 (3) (4-33)

$$
HA^{-} = H^{+} + A^{2}
$$
 -log K_{a2} = 4.60 (3) (4-34)

$$
R_3NH^+ = R_3N + H^+ \t -log K_a = 9.8 \t(3)
$$
 (4-35)

$$
(H^{+}) + (R_{3}NH^{+}) = 2(A^{2-}) + (HA^{-}) + (OH^{-})
$$
\n(4-36)

Equation 4-36 describes the charge balance in the aqueous phase. The parentheses in Equation 4-32 and 4-36 denote concentrations.

The concentration of un-ionized fumaric acid in the aqueous phase can be expressed as the product of the total fumaric acid in the aqueous phase and the quantity α_3 , where:

$$
\alpha_3 = \frac{1}{1 + \frac{K_{a1}}{H^+} + \frac{K_{a1}K_{a2}}{(H^+)} \tag{4-37}
$$

where $K_{a,1}$ and $K_{a,2}$ are the acid dissociation constants of fumaric acid, as given in Equations 4-33 and 4-34. Then the mass balance for the fumaric acid can be written as follows:

o = Total Acid in System Acid in Organic Phase Acid in Aqueous Phase

$$
0 = [H_2A]_{init} - K_{1,2}[B]^2(H_2A) - 2K_{2,1}[B](H_2A)^2 - 2K_{2,2}[B]^2(H_2A)^2
$$

 $-\Phi P_3(H_2A) - WS(H_2A)/\alpha_3$ (4-38)

where the first term is the initial concentration of acid in the organic phase, the parentheses denote concentrations in the aqueous phase, the brackets denote concentrations in the organic phase, B denotes the uncomplexed Alamine 336 molecule, Φ is equal to the volume fraction of MiBK (0.85) in the organic solvent, and WS is equal to the water to solvent ratio (3/8) used in the back-extraction experiments.

The mass balance for the Alamine 336 (essentially all present in the organic phase) can be written as follows:

Uncomplexed *A336* = Total *A336* - Complexed *A336*

$$
[B] = [Btotal] - 2K1,2[B]2(H2A) - K2,1[B](H2A)2 - 2K2,2[B]2(H2A)2
$$
 (4-39)

Equation 4-39 is second order in terms of [B] and can be solved for [B] using the quadratic equation. The positive root of Equation 4-39 is:

$$
[B] = \frac{-K_{2,1}(H_2A)^2 - 1}{2(2K_{1,2}(H_2A) + 2K_{2,2}(H_2A)^2)}
$$

$$
\sqrt{\frac{(K_{2,1}(H_2A)^2 + 1)^2 + 4[B_{\text{total}}](2K_{1,2}(H_2A) + 2K_{2,2}(H_2A)^2)}{2(2K_{1,2}(H_2A) + 2K_{2,2}(H_2A)^2)}}
$$
(4-40)

Equation 4-40 can be substituted into Equation 4-38 to give a mass balance for the acid in terms of only two unknown values, the pH of the aqueous raffinate (incorporated into α_3) and the concentration of un-ionized acid in the aqueous phase. This equation is of the form:

$$
f\{pH,(H_2A)\}=0
$$
 (4-41)

Two computer programs were used sequentially to solve this equation for

 $(H₂A)$ at set pH values. In the first program, at a given pH value, the value of the function, f, was calculated for a range of (H_2A) values in order to determine an interval within which the function is equal to zero. The interval is bounded by two (H_2A) values, one that results in a negative value for f and one that results in a positive value for f. In the second program, for the selected pH value, a bisection routine was used to determine the value of (H_2A) within this interval for which f is equal to zero. These two computer programs, "second" and "fumaric", are given in Appendix C.

For each pair of pH and $(H₂A)$ values, the recovery of the acid into the aqueous phase was calculated as:

$$
Recovery = \frac{WS(H_2A)/\alpha_3}{[H_2A]_{init}}
$$
 (4-42)

and the ratio of moles of TMA to moles of acid originally present in the organic phase was calculated where, as before (Equation 4-14):

$$
(R_3N)_{total} = (R_3N) + (R_3NH^+) = (R_3NH^+)/\beta \tag{4-43}
$$

and the value of (R_3NH^+) is determined using Equations 4-33, 4-34, and 4- 36 (β was defined in Equation 4-15). In this manner, a theoretical backextraction curve for fumaric acid was generated as shown in Figure 4-3.

4.2 Results of the Back-Extraction Experiments

Figure 4-1 shows the experimental data for the back-extraction of lactic acid from a solution of O.3M Alamine 336 in MiBK into aqueous solutions of TMA. The theoretical curve generated using the computer program is also shown. Essentially complete removal of the acid from the organic phase is achieved when one mole of TMA is present for every mole of acid. There is reasonable agreement between theory and experiment.

Figure 4-2 shows similar results for the back-extraction of succinic acid. Succinic acid is a di-carboxylic acid; complete removal is achieved when two moles of TMA are present for every mole of succinic acid. The data compare favorably with the predicted curve.

The data for the back-extraction of fumaric acid are shown *in* Figure 4-3. Fumaric acid is a di-carboxylic acid, and complete removal is achieved when two moles of TMA are present for every mole of fumaric acid. Again, the data compare favorably with the predicted curve.

The experimental results for all three acids show that the ionized, aqueous TMA is a much stronger base than the complexing, organic Alamine 336, i.e., the shapes of the curves correspond to the available aqueous TMA pairing stoichiometrically with the back-extracted acid, even when this results in a very low acid concentration in the organic phase.

Figure 4-l. Back-extraction of Lactic Acid from O.3M Alamine 336 in MiBK into Aqueous Solutions of Varying TMA Concentration

Figure 4-2. Back-extraction of Succinic Acid from O.3M Alamine 336 in MiBK into Aqueous Solutions of Varying TMA Concentration

Figure 4-3. Back-extraction of Fumaric Acid from O.3M Alamine 336 in MiBK into Aqueous Solutions of Varying TMA Concentration

In the back-extraction experiments, TMA did not partition significantly (greater than 0.0005 wt.%) into the organic phase at molar ratios of TMA to acid less than or equal to the ratio required to achieve 100% removal of the acid from the organic phase. Under these conditions, essentially all of the TMA is ionized and paired with the acid. For example, in an aqueous solution containing 2 moles of TMA for every mole of succinic acid, less than 2 mol.% of the TMA is in the un-ionized form (see results of computer programs, Appendix C). Essentially only the un-ionized TMA will partition into the organic phase.

When TMA was present in excess of the ratio required to achieve 100% removal of the acid from the organic phase, transfer of TMA into the organic phase occurred as shown in Table 4-2; the larger the excess of TMA, the greater the amount transferred into the organic phase.

Table 4-2. TMA Transferred to Organic Phase

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CHAPTER 5. RESULTS OF THE THERMAL REGENERATION EXPERIMENTS

This chapter discusses the results of concentration and heating experiments in which water and trimethylamine are driven off from aqueous solutions of trimethylammonium carboxylates. For both succinic acid and fumaric acid, removal of water and trimethylamine was essentially complete, leaving behind the acid in crystalline form. The results of work done to wash and further purify these crystals are also discussed.

5.1 Thermal Regeneration Experiments

As shown in the carboxylic acid recovery process diagram (Figure 1- 3), water and trimethylamine (TMA) are to be removed from aqueous solutions of TMA and a low-volatility carboxylic acid. In this final step of the recovery process, complete removal of TMA is desired in order to generate the acid in product form. The regenerated back-extractant (the TMA) is to be recycled within the process. The apparatus used in the thermal regeneration experiments was discussed in Chapter 3, Section 6.

5.1.1 Lactic Acid

As discussed in Chapter 3, Section 3, lactic acid forms intermolecular esters in concentrated aqueous solution. Lactic acid also has a high solubility in water. As an aqueous solution of lactic acid is concentrated by heating, crystalline lactic acid does not precipitate out of solution, but rather the viscosity of the solution increases steadily as polymerization of

the lactic acid molecules occurs. This effect was illustrated in Figure 3-1.

Crystalline lactic acid is usually obtained by fractional vacuum distillation of the commercial syrup (85-90 wt.% lactic acid) which contains both free lactic acid and polymeric forms (1,2). The free lactic acid is removed overhead in the distillation. The middle fraction is collected and cooled in an ice bath whereupon lactic acid crystals form. If separation of the optical isomers is desired, this crystalline material (or the distillate itself, without prior cooling) is then dissolved in an equal volume of a mixture of equal parts of ethyl and isopropyl ethers (1,2,3). Fractional crystallization separates the less soluble L-Iactic acid from the mixture of L- and DL-Iactic acid.

Crystalline lactic acid is not thermodynamically stable at room temperature. If held at room temperature, the crystalline acid "will spontaneously commence intermolecular reactions with the formation of water and esters" (I). Lactic acid is offered commercially as aqueous solutions of varying assay strengths. The general assay levels are 44 , 50 , 80, and 88 wt.% lactic acid (4,5).

In the thermal regeneration experiments carried out with aqueous solutions of trimethylammonium lactate, the goal was to remove essentially all the TMA, leaving behind a concentrated aqueous solution of lactic acid and lactic acid polyesters, similar to the commercial syrup. Figure 5-1 summarizes the results of the experiments. Only 62% of the TMA present in the initial aqueous solution was removed, leaving behind a viscous

Figure 5-1. Heating of Lactic Acid/TMA/H₂O Solutions

aqueous solution of lactic acid and TMA, and also, no doubt, TMA/lactate and lactic acid polyesters. It is possible that after a large portion of the water and TMA are removed, the formation of intermolecular esters could prevent further TMA removal due to transport limitations. Also, the increasing ratio of lactic acid to TMA in the residue probably serves to suppress the volatility of TMA, because of the chemical interaction between lactic acid and TMA.

5.1.2 Succinic Acid and Fumaric Acid

Figure 5-2a shows the rates of water and TMA removal from an aqueous TMA/succinate salt solution initially containing 2 moles of TMA for every mole of succinic acid. Figure 5-2b shows the temperature rise during the experimental run. The temperature rise initially corresponds to the rise in boiling point of the solution. The temperature is held constant at 112°C at the end of the run, as was discussed in Chapter 3.

Although TMA is more volatile than water, it is held in solution due to complexation or ion-pair formation with the acid. The salt is decomposed when a high enough temperature is reached and/or as crystallization of the acid occurs, driving the decomposition reaction. TMA is released after most of the water has been removed. The extra release of TMA at the start of the experimental run (as shown in Figure 5-2a) occurred because there was a slight excess of TMA present in the initial aqueous solution.

Figure 5-2. Concentration and Decomposition of a TMA/Succinate Salt (starting with a O.6M aqueous solution of the di-salt of the acid)

Figure 5-2a. Rate of Removal of H_2O and TMA with Time

Figure 5-2h. Variation of Temperature with Time

The data shown in Figure 5 are from the fifth experimental run conducted with a TMA/succinate salt solution. The results of all experimental runs are reported in Appendix A. As explained in Chapter 3, experimental run #5 was conducted under vacuum for data-taking purposes. Under these conditions, the solution started boiling at 61° C and at the end of the run, the temperature was held at 112° C. Experimental run #3 was conducted at slightly above atmospheric pressure (1.07 atm), and boiling commenced at 101°C. The temperature in the heating flask at the end of run $\#3$ was equal to 172°C. It is necessary to keep the temperature in the heating flask considerably below the decomposition temperature of the acid $(235^{\circ}$ C for succinic acid, 230 $^{\circ}$ C for fumaric acid [6]).

The partial pressures of TMA, water, and nitrogen above the solution as a function of TMA left in solution are shown in Figure 5-3. These data are from experimental run $#6$. The partial pressure values for TMA and water are functions of the nitrogen flow rate used during the experimental run. The nitrogen flow rate used (0.0137-0.0158 moles/min) in experimental runs $#5$ and $#6$ was high enough to purge the heating flask continually of vaporized water and TMA.

The rates of water and TMA removal shown in Figure 5-2a are functions of the rate at which heat is supplied to the heating flask. Although the values of the data points shown in Figures 5-2a and 5-3 are functions of the experimental conditions, the order in which water and TMA are vaporized, illustrated in these figures, is a characteristic of the TMA/water/carboxylic acid system.

Figure 5-3. Partial Pressures above Succinic Acid/TMA/H₂O System during Thermal Regeneration Experimental Run

Figures 5-4a and 5-4b show the results of the thermal regeneration \mathcal{L}^{\pm} experiments carried out with aqueous solutions of the di-trimethylammonium salt of fumaric acid. As with succinic acid, the TMA is released after most of the water is removed, and again an end-product containing the acid in crystalline form is obtained.

5.2 Washing and Purification of Crystalline Product

At the end of an experimental run in which a salt of TMA and succinic acid or fumaric acid was. concentrated and thermally decomposed, there remained in the heating flask a mixture (yellowish in color) of water, acid crystals, dissolved acid, and residual TMA. Various methods described in Chapter 3 were used to wash and further purify the acid crystals. Depending on the washing technique used, recoveries of between 67% and 77% purified succinic acid crystals were achieved.

The first two treatment methods (washing with acetone and chloroform, and washing with acetone and carbon tetrachloride) served to separate the acid crystals from the viscous aqueous solution consisting of TMA and dissolved acid present in roughly equilmolar amounts. The third treatment method -- dissolution of the mixture of solid crystals and aqueous solution into methyl isobutyl ketone (MiBK), followed by evaporation of the MiBK --yielded the highest recovery of succinic acid and was used to treat the fumaric acid crystals. A recovery of 81% of the purified fumaric acid crystals was obtained. With improved washing techniques, it is possible that an even greater percent recovery can be achieved.

Figure 5-4. Concentration and Decomposition of a TMA/Fumarate Salt (starting with a O.IM aqueous solution of the di-salt of the acid)

Figure 5-4a. Rate of Removal of H_2O and TMA with Time

Figure 5-4b. Variation of Temperature with Time

The preliminary work done to determine the best method of washing the crystals was performed on succinic acid crystals from a number of different thermal regeneration experiments. The composition of the mixture remaining in the heating flask at the end of a typical succinic acid experimental run was 40.5 wt % C, 6.6 wt % H, and 3.6 wt % N. The ratio of moles of TMA per mole of acid in the overall mixture (solid crystals and aqueous solution) was therefore equal to 0.4. Table 5-1 shows the degree to which the various washing techniques removed this residual TMA.

Three successive recrystallizations of the succinic acid from MiBK resulted in relatively pure (containing between 0.05 and 0.11 wt % N) crystals. The fumaric acid crystals were recrystallized from MiBK only one time and were of lower purity (containing 0.61 wt % N). Apparently successive recrystallizations are required to remove the residual TMA.

Dissolution of the concentrated TMA/water/carboxylic acid mixture into MiBK serves to lower the concentrations of the I-I TMA/acid salt $(R_3NH^+HA^-)$ and of the un-ionized TMA (R_3N) and the un-ionized carboxylic acid $(H₂A)$. In addition, in an MiBK diluent, the un-ionized form of the TMA molecule would be favored. The combination of these two effects would drive the reaction for the formation of the TMA/acid ion pair:

$$
R_3N + H_2A = R_3NH^+HA
$$
 (5-1)

with:

$$
K = \frac{(R_3NH^+HA^-)}{(R_3N)(H_2A)}
$$
 (5-2)

 $\ddot{}$

Table 5-1. Washing of Succinic and Fumaric Acid Crystals

to the left, thereby increasing the proportion of TMA that is present in volatile form.

After dissolution of the mixture into MiBK, the solution was heated under vacuum to remove the solvent. As the solvent was removed from the mixture, the water and a portion of the TMA were also removed, leaving behind crystalline acid as well as a small amount of solid TMA/acid salt. Successive recrystallizations served to break apart this salt and liberate TMA.

It is possible that the TMA remaining in solution after the heating and concentration of an aqueous solution of TMA and lactic acid could be removed by this dissolution/heating technique, leaving behind a mixture of the free acid and polymeric forms of the acid. Urbas (7) states that the trialkylammonium salt of lactic acid and either tributylamine or dicyclohexylmethylamine can be thermally decomposed to obtain the acid and the tertiary amine; however, no experimental details are given. Further studies on the separation of TMA from aqueous solutions of TMA and lactic acid are left for future work.

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CHAPTER 6. PROCESS DESIGN CONSIDERATIONS

This chapter examines the implications of the experimental results reported in previous chapters. for the design of a carboxylic acid recovery process in which back-extraction into and recycle of a volatile base is the method of regeneration of the amine-carboxylic acid extract. Each step of the process is first examined separately. and then the economic viability of this regeneration method is compared to that of the temperature-swing and diluent composition-swing regeneration methods.

6.1 The Carboxylic Acid Recovery Process

A process diagram for the recovery of low-solubility carboxylic acids from dilute aqueous solutions using the regeneration method explored in this work is shown in Figure 6-1. The acid is extracted by reversible chemical complexation from the aqueous feed into an organic solution of a tertiary amine extractant. The acid is then back-extracted from the loaded organic phase into an aqueous trimethylamine (TMA) solution. The aqueous solution of the trimethylammonium salt is heated in an evaporative crystallizer to drive off the water and the TMA. which are recycled to the process. The acid product is then sent to a final purification step.

6.1.1 Extraction of the Carboxylic Acid by a Tertiary Amine Extractant

The first step of the carboxylic acid recovery process is the extraction of the acid into the solvent phase to produce an amine-carboxylic acid extract and a relatively acid-free aqueous raffinate. Figure 6-2 shows a McCabe-Thiele operating diagram for the extraction at 25°C of succinic acid from dilute aqueous solution into O.3M Alamine 336 in methyl isobutyl ketone (MiBK). The vertical axis is the concentration of acid in the organic phase while the horizontal axis is the concentration of acid in the aqueous phase. The equilibrium curve was determined experimentally by Tamada (1).

In a McCabe-Thiele operating diagram, the equilibrium curve relates the compositions of the two streams leaving a stage in a multi-stage extraction process. The operating curve relates the compositions of streams flowing past each other between stages. Three stages are shown on the diagram in Figure $6-2$. For dilute solutions, the operating curve is a straight line with a slope equal to the water to solvent flow ratio (W/S). (2)

During the extraction process, the equilibrium curve must lie above the operating line. Limiting conditions occur at points where the equilibrium curve and operating line intersect. These 'pinch points' usually occur at the ends of a multi-stage extractor as is shown in Figure 6-2; however, depending on the shape of the equilibrium curve, internal pinch points may occur. (2)

Figure 6-2. Extraction of Succinic Acid from Aqueous Solution into O.3M Alamine 336 in MiBK

Figure 6-2a. Multi-Stage Extractor

The endpoints of the operating line are the pairs of compositions in the streams passing each other at each end of the column. The locations of these points are dictated by the feed composition, the desired removal of the acid from the feed, and the flow ratio used in the extractor. Flow ratios for industrial extractors are typically in the range $1 < W/S < 10$ (2).

Figures 6-3 and 6-4 show equilibrium curves for the extraction at 25°C of lactic acid (I) and fumaric acid (3) from dilute aqueous solution into O.3M Alamine-336 in MiBK. A thorough discussion of the extraction of carboxylic acids by amine extractants, including the effects of changes in temperature or diluent composition on the shape of the equilibrium curve, is given by Tamada (I).

For succinic acid or fumaric acid, vigorous mixing of an aqueous acid solution with an organic solution of Alamine 336 in MiBK results in emulsion formation. Therefore, the multi-stage extractor used in the process should be one that suppresses emulsion formation by either allowing only gentle mixing and/or promoting phase separation.

6.1.2 Back-Extraction of the Acid into Aqueous TMA Solution

Due to the strong basicity of TMA, nearly complete removal of the acid from the organic extract solution can be achieved when one mole of TMA is present for every equivalent weight of acid. Figures 6-5 and 6-6 show equilibrium curves for fumaric acid partitioning between an organic phase of O.3M Alamine 336 in MiBK and an aqueous phase of O.4M TMA or

(from Starr [3])

..

Figure 6-6. Equilibrium Curve for Fumaric Acid at 2S·C Organic Phase: O.3M Alamine 336 in MiBK Aqueous Phase: O.76M TMA (Calculated using computer programs: "second" and "equilr')

O.76M TMA (the vertical axis being the concentration of acid in the aqueous phase). When fumaric acid is present in an amount equal to or less than 1/2 the amount of TMA, essentially all of the acid will be present in the aqueous phase. When fumaric acid is present in an amount greater than 1/2 the amount of TMA, it will partition between the organic and aqueous phases.

A counter-current multi-stage .extractor should not be used to carry out the back-extraction of the acid into aqueous TMA solution. In a counter-current multi-stage extractor, the exiting acid-depleted organic phase would contact an entering aqueous TMA phase containing little or no acid, and as discussed in Chapter 4, TMA transfer into the organic phase would occur. The organic phase would leave the extractor saturated with TMA and another process step would be required to remove the TMA.

In general, there is little incentive to use a multi-stage extractor for two-phase systems with equilibrium curves of the shape shown in Figures 6- 5 and 6-6, because due to the shape of the equilibrium curve each additional stage used in an extractor would serve to increase the concentration of the solute in the extractant phase to only a small degree. In this particular case, the equilibrium curves shown would not be valid for the entire length of a counter-current column, due to the change in TMA concentration in the aqueous phase.

Therefore, a one-stage extractor should be used for the back-extraction of acid from the organic phase into aqueous TMA solution. Emulsion formation was not encountered during the experimental work done with these two-phase systems, and thus a standard mixer/settler extractor could probably be used.

Typical phase ratios for industrial one-stage extractors range from $0.1²$ to 10. These ratios can be achieved by recycle of one of the phases, if necessary. Since it is desirable to concentrate the acid during this backextraction step, the organic to aqueous (S/W) phase ratio should be greater than 1. To avoid TMA transfer to the organic phase, the product of the concentration of TMA and the flowrate for the aqueous phase should be somewhat less than or equal to the product of the concentration $-\frac{1}{2}$ in terms of equivalent weight per volume -- of the acid and the flowrate for the organic solvent phase leaving the forward extractor.

Although 100% recovery of the acid from the organic extract solution can be achieved without TMA transfer to the organic phase, for control purposes, a recovery of about 90% could be used, thereby allowing for slight fluctuations in the acid concentration of the organic solvent stream leaving the forward extractor. The acid not removed from the organic phase would be sent back through the forward extractor. A significant decrease in the concentration of acid in the feed to the carboxylic acid recovery process would require a corresponding decrease in the TMA concentration and/or aqueous-phase flow rate in order to avoid TMA transfer to the organic phase.

The maximum concentration of acid attainable in the final aqueous phase depends on the solubility of TMA in water at the temperature and pressure of operation. Although the solubility of TMA in water at 20°C and atmospheric pressure is approximately equal to 41 wt.% $(4,5)$, TMA is commercially available as a 25 wt.% aqueous solution. This commercial product has a density of 0.933 *glml* at 20°C (4) and therefore has a TMA concentration of 3.95 mol/I.

6.1.3 Degree of Concentration of the Acid

The economic feasibility of the carboxylic acid recovery process depends in part on the degree of concentration of acid that can be achieved during the forward and back-extraction steps; i.e., on the amount of water that must be removed in the evaporative crystallizer. For a given acid, if a recycle stream is not to be used in the back-extractor, then the degree of concentration that can be achieved depends on the maximum possible concentration of extractant in the organic solvent phase (which in turn is dependent on the diluent used), the maximum solubility of TMA In water, and the flow ratios permissible in industrial equipment. If one allows for the possibility of recycle to the back-extractor of a portion of the aqueous stream leaving the back-extractor, then the degree of concentration that can be achieved depends only on the solubility of TMA in water at the temperature and pressure of operation. The conditions that would require recycle of the aqueous phase in the back-extractor are discussed in Section 2.2 of this chapter.

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A sample calculation follows for the degree of concentration attainable during the recovery of succinic acid from dilute solution when the organic solvent phase consists of Alamine 336 in a MiBK diluent and the aqueous feed solution is 3 wt. % succinic acid. The calculation is summarized in Table 6-1. According to Tamada (1,6), 1.0M Alamine 336 in MiBK is the maximum concentration that should be used in the extraction due to problems of third-phase formation and high solvent viscosity at higher concentrations.

For the forward extractor, the assumption of infinite stages was used; the loaded organic solvent stream leaving the extractor is then in equilibrium with the aqueous feed solution and likewise the organic solvent stream entering the extractor is in equilibrium with the aqueous raffinate stream leaving the extractor. For the back-extractor, in order to obtain the maximum concentration of acid in the aqueous TMA solution, the acid recovery from the organic phase was set at 100%. The regenerated organic solvent, containing essentially no acid, is sent back to the forward extractor. Since this regenerated organic solvent stream is in equilibrium with the acid-depleted aqueous raffinate, essentially 100% removal of the acid from the aqueous feed can occur in the forward extractor.

The concentration of succinic acid in the organic phase leaving the forward extractor is calculated as follows:

$$
[H2A] = K1,1[B](H2A) + 2K2,1[B](H2A)2 + \Phi P(H2A)
$$
 (6-1)

where H_2A denotes the succinic acid, B denotes the uncomplexed Alamine

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Table 6-1. Sample Calculation for Degree of Concentration of Succinic Acid

FORWARD EXTRACTOR

Aqueous Feed: 3 wt % succinic acid (0.250M) $pH = 2.49$

Inlet Organic Solvent Stream: 1.0M Alamine 336 in MiBK

Aqueous Raffinate: water containing little acid

Outlet Organic Solvent Stream: 1.35M succinic acid and 1.0M Alamine 336 in MiBK

Water/Solvent Flow Ratio: 5.40

BACK-EXTRACTOR

Initial Organic Phase: 1.35M succinic acid and 1.0M Alamine 336 in MiBK

Initial Aqueous Phase: 3.95M TMA

Final Organic Phase: 1.0M Alamine 336 in MiBK

Final Aqueous Phase: 1.97M succinic acid and 3.95M TMA

Solvent/Water Phase Ratio: 1.46

Concentration Factor: $1.97/0.250 = 7.88$

336 molecule, the parentheses denote concentrations in the aqueous phase, the brackets denote concentrations in the organic phase, the $K_{p,q}$ values are the equilibrium constants of the complex formation as listed in Table 1- 2, Φ is equal to the volume percent of MiBK (0.50) in the organic solvent, and P (0.19) is the partition coefficient for succinic acid between water and MiBK as given in Table 4-1.

In order to solve Equation 6-1, the values of $[B]$ and (H_2A) must first be calculated as follows:

$$
[B] = \frac{[B_{\text{total}}]}{1 + K_{1,1}(H_2A) + K_{2,1}(H_2A)^2}
$$
(6-2)

$$
(H2A) = \frac{(H2Atotal)}{1 + Ka,1 + Ka,1Ka,2 \frac{(6-3)}{(H+)}
$$

where the numerator in Equation 6-3 is the concentration (0.250M) of succinic acid in the aqueous feed solution, $K_{a,1}$ and $K_{a,2}$ are the first and second dissociation constants $(10^{-4.21}$ and $10^{-5.60}$) of succinic acid, and $(H⁺)$ is the hydrogen ion concentration $(10^{-2.49})$ in the aqueous feed solution as determined experimentally.

As shown in Table 6-1, the concentration of acid in the loaded organic phase was calculated to be 1.35M. A material balance for the succinic acid can be used to calculate the water/solvent flow ratio in the forward extractor. This value, 5.40, is within the desired range.

In the back-extractor, the commercially available 25 wt. % (3.95M) aqueous TMA solution is used as the aqueous phase. The phase ratio is set so that the number of moles of TMA in the back-extractor is equal to twice the number of moles of succinic acid, so as to obtain 100% recovery of the acid from the organic phase while at the same time avoiding transfer of TMA to the organic phase. This solvent/water phase ratio is equal to 1.46, which is within the desired range. The acid concentration in the final aqueous phase is then equal to half the TMA concentration or 1.97M.

The degree of concentration achieved during the forward and backextraction steps is equal to the final aqueous acid concentration (1.97M) divided by the acid concentration in the aqueous feed solution (0.250M). In this example, the degree of concentration -- or concentration factor -- is equal to 7.88, or approximately 8. In order to obtain a greater degree of concentration, a more concentrated aqueous TMA solution must be used.

6.1.4 Thermal Decomposition of the Trimethylammonium Carboxylates

After the carboxylic acid has been back-extracted into an aqueous solution of TMA, the aqueous solution of the trimethylammonium carboxylate is heated in an evaporative crystallizer to drive off the water and TMA. As discussed in Chapter 2, oxidation of TMA during this step is undesirable. The evaporative crystallizer should be run under a nitrogen atmosphere and/or at a temperature low enough so that oxidation of TMA will not occur. The temperature should also be kept considerably below the thermal decomposition temperature of the carboxylic acid (235°C for 84

succinic acid, 230°C for fumaric acid [7]).

The energy consumption for this heating and concentration step is given by the sum of the sensible heat required to raise the temperature of the aqueous trimethylammonium carboxylate solution to the initial boiling temperature, the sensible heat required to sustain boiling as the boiling temperature increases during concentration, and the latent heat required to vaporize the water. For a given weight of acid product, the higher the concentration of the initial aqueous trimethylammonium carboxylate solution, the lower the energy consumption during this step.

6.1.5 Final Purification Step

As discussed in Chapter 5, a final purification step is needed to remove residual TMA from the crystalline acid product. This step could consist of dissolution in MiBK, or another solvent, of the material left at the end of the evaporative crystallization step followed by heating (possibly under vacuum) to drive off the MiBK, and the residual TMA and water. If this method is used, a vapor stream of MiBK, TMA, and water would be produced.

In order to avoid chemical consumption, separation of the MiBK (or other solvent) from the vapor stream would be required so that the MiBK could be reused in this purification step, and the TMA and water could be recycled to the back extractor. This separation could be achieved by passing the MiBK/TMA/water vapor stream through a condenser. If the

system is under sufficient vacuum (see Chapter 3, Section 6) the TMA will remain in the vapor phase. The MIBK and water will condense and can be collected in a trap. Once condensed, the MiBK and the water will form two different phases and can be separated by decantation. Stripping can be used to remove residual MiBK from the water phase, if this is necessary. The TMA vapor stream can then be passed through water under atmospheric pressure to allow absorption to occur, producing an aqueous TMA solution.

6.2 Comparison of pH-Swing Regeneration to Temperature-Swing

Regeneration and Diluent Composition-Swing Regeneration

This work examines a carboxylic acid recovery process in which regeneration of an amine-carboxylic acid extract is achieved by backextraction of the acid into an aqueous solution of a volatile base followed by a heating and concentration step to obtain the acid product. In Chapter I, Section 4, alternative regeneration methods were discussed. Two of the regeneration methods, temperature-swing and diluent composition-swing also consist of a back-extraction step which would need to be followed by a heating and concentration step in order to obtain the acid in product form.

Important factors in determining the economic viability of a process are the energy consumption, chemical consumption, and capital and other operating costs of the process. Table 6-2 compares these three regeneration methods with regard to these factors. All three regeneration methods involve full recycle of chemicals; the only chemical consumption would be through entrainment, solubility, or degradation losses.

Table 6-2. Comparison of pH-Swing, Temperature-Swing and Diluent Composition-Swing Regeneration Methods

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6.2.1 Energy Consumption

During the back-extraction step, the energy consumption would be higher for the temperature-swing and diluent composition-swing regeneration methods than for the pH-swing regeneration. In temperature-swing regeneration (see Figure 1-1), energy would be required to raise the temperature of the inlet streams to the back-extractor; however, some of this energy could be provided by heat exchange between the loaded organic solvent stream and the regenerated organic solvent stream. In diluent composition-swing regeneration (see Figure 1-2) energy is required to alter the diluent composition by distillation.

After the back-extraction step, all three regeneration methods have a concentration and heating step during which water (water and TMA for the pH-swing process) is removed to generate the acid in product form- either in crystalline form or as a concentrated aqueous solution. The energy consumption during this step would depend on the degree of concentration of the acid that had already been achieved during the forward and back-extraction steps; i.e. on the amount of further concentration required. The degree of concentration of acid that can be attained during the forward and back-extraction steps is compared below for the three regeneration methods.

The pH-swing regeneration method requires a final purification step that is not needed for the temperature-swing or diluent composition-swing methods. The energy consumption for this step is given by the sum of the sensible heat required to raise the temperature of the MiBK/TMA/water/acid mixture to the boiling point of MiBK (117-118°C at atmospheric pressure [7]), and the heat required to vaporize the MiBK and the water.

6.2.2 **Comparision of Degree of Concentration**

The degree of concentration of the acid that can be achieved during the forward and back-extraction steps was compared by Tamada (I) for the temperature-swing and diluent composition-swing processes. The degree of concentration (concentration factor) is shown in Figure 6-7 as a function of the acid concentration of the aqueous feed to the carboxylic acid recovery process. These curves were generated for aqueous feed solutions of succinic acid.

The organic solvent phase used in the temperature-swing process was O.3M Alamine 336 in MiBK. For the diluent composition-swing process, the organic solvent phase used in the forward extractor consisted of 15% (v/v) Alamine 336 (0.3M Alamine 336), 55% (v/v) chloroform, and 30% (v/v) heptane, while the organic solvent phase used in the back-extractor consisted of 30% (v/v) Alamine 336 $(0.6M)$ Alamine 336), 10% (v/v) chloroform, and 60% (v/v) heptane.

Also shown in Figure 6-7 is the degree of concentration attainable when pH-swing regeneration is used. This curve was calculated as detailed in Section 1.3 of this chapter. The organic solvent phase used in the calculations was 0.3M Alamine 336 in MiBK. The following assumptions of

Figure 6-7. Comparison of the Maximum Possible Concentration Factor for Temperature- (1), Diluent Composition- (1) and pH-Swing Regeneration

Tamada (I) were used in the calculations: infinite stages in the forward extractor; and 95% recovery of the acid from the aqueous feed. The pH values for aqueous solutions of succinic acid -- needed to solve equation 6- 3 -- of varying concentration were taken from Tamada (1).

During the back-extraction step, the commercially available (3.95M) aqueous TMA solution was used for the calculations involving the higher values of aqueous feed concentration. However, below about 0.05M succinic acid in the aqueous feed, the concentration of acid in the loaded organic phase was low enough that if $3.95M$ TMA was used, the phase ratio (W/S) in the back-extractor would need to be less than 0.1 in order to prevent TMA transfer into the organic phase. (Higher values for the phase ratio would result in 2-phase systems in which there was an excess of TMA- greater than the amount required to recover 100% of the acid from the organic phase.) Therefore, for low concentrations of acid in the aqueous feed solution, the phase ratio in the back-extractor was set at 0.1, and TMA solutions less concentrated than 3.95M were used.

For feed solutions below about 0.05M succinic acid, a recycle stream of the aqueous TMA phase could be used in the back-extractor to achieve W/S equal to or greater than 0.1. This use of a recycle stream would prevent the necessity of using TMA solutions less concentrated than 3.95M, and would thereby result in concentration factors for feed solutions below 0.05M succinic acid greater than those shown in Figure 6-7.

In order to determine the TMA concentration needed (or for higher values of aqueous feed concentration, the phase ratio needed) in the backextraction step, the following equation was used:

$$
[H2A] \times TMA \text{ to acid ratio required } = (TMA) \times W/S
$$
 (6-4)
to achieve desired recovery
in the back-extractor

where $[H_2A]$ is the concentration of acid in the loaded organic phase leaving the forward extractor: (TMA) is the aqueous TMA concentration, and the TMA/acid ratio needed was determined from Figure 4-2. The recovery of acid obtained in the forward extractor was set at 95% as given by Tamada (I); therefore, the recovery of acid in the back-extractor is that needed to produce a regenerated organic phase with a succinic acid concentration in equilibrium with the concentration of succinic acid in the aqueous phase leaving the forward extractor.

For the range of feed concentrations shown in Figure 6-7, the concentration factor is. highest for the pH-swing regeneration method. The energy required during the concentration and heating step of the carboxylic acid recovery process would therefore be lowest for a process using pHswing regeneration.

6.2.3 Capital **and** Operating Costs

The capital costs for the back-extraction step of the carboxylic acid recovery process using pH-swing regeneration would probably be the lowest of the three regeneration methods due to the lower capital costs for a one-
stage extractor than for the multi-stage extractors that would be needed for the other two regeneration methods. In addition, the temperature-swing regeneration would include capital costs for a heat-exchanger, while diluent composition-swing regeneration would include the capital and operating costs of the distillation column. However, the pH-swing regeneration method must bear the capital and operating costs of the equipment needed for the final purification step, which is not required when the other two regeneration methods are used.

6.3 Summary and Conclusions

A carboxylic acid recovery process is proposed in which regeneration of an amine-carboxylic acid extract is achieved by back-extraction of the acid into an aqueous solution of trimethylamine. The resultant aqueous solution is concentrated by evaporation and the trimethylammonium carboxylate is decomposed thermally, yielding the volatile trimethylamine for recycle and the acid in product form.

The feasibility of this process has been explored through experimental work with lactic acid, succinic acid, and fumaric acid. Process calculations using data from forward and back-extraction experiments show that a high degree of concentration of the acid can be achieved by extraction with a high molecular-weight tertiary amine in an organic diluent, followed by back-extraction into aqueous trimethylamine solution. Experiments involving the thermal decomposition of the trimethylammonium carboxylates yielded approximately 80% recovery of succinic acid and fumaric acid in crystalline

form. These recovery fractions can probably be increased. experimental results and an examination of the overall process design demonstrate that the carboxylic acid recovery process examined in this work should be viable for acids of low volatility. The

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APPENDIX A. EXPERIMENTAL DATA

A.1 Back-Extraction Experiments

This section gives the experimental results for the back-extraction of lactic, succinic, and fumaric acids from organic extract solutions into aqueous trimethylamine (TMA) solutions.

Table A-I. Back-Extraction of Lactic Acid

Initial Organic Phase: 0.71M lactic acid and 0.3M Alamine 336 in methyl isobutyl ketone (MiBK)

Lactic Acid

in Final

in Final TMA in Initial in Final in Final in Final Aqueous Phase Final Aqueous Aqueous Phase Organic Phase (M) pH (M) (M) 0 2.04 0.61 0.465 0.38 3.77 0.78 0.413 1.14 4.87 1.52 0.272 1.33 5.27 1.28 0.234 1.51 5.42 1.32 0.213 1.70 5.96 1.40 0.0916 1.89 6.28 1.60 0.0511 3.92 9.62 2.11 $\overline{}$.

Organic to Aqueous (S/W) Phase Ratio: $8/3$ (v/v)

Table *A-2.* Back-Extraction of Succinic Acid

Initial Organic Phase: 0.45M succinic acid and 0.3M Alamine 336 in methyl isobutyl ketone (MiBK)

Organic to Aqueous (S/W) Phase Ratio: 8/3 (v/v)

Initial Organic Phase: 0.44M succinic acid and 0.3M Alamine 336 in methyl isobutyl ketone (MiBK)

Organic to Aqueous (S/W) Phase Ratio: 8/3 (v/v)

Table *A-3.* Back-Extraction of Fumaric Acid

Initial Organic Phase: 0.286M fumaric acid and 0.3M Alamine 336 in methyl isobutyl ketone (MiBK)

Organic to Aqueous (S/W) Phase Ratio: 8/3 (v/v)

A.2 Thermal Regeneration Experiments

This section gives the experimental results of concentration and heating experiments in which water and trimethylamine are driven off from aqueous solutions of trimethylammonium carboxylates.

A.2.1 Preliminary Regeneration Experiments conducted with

Aqueous Trimethylammonium Succinate Solutions

Table A-4. Regeneration Run #1

Initial Aqueous Solution: 60 ml of 0.6M succinic acid and 1.2M TMA

Treatment: The solution was heated under vacuum. No nitrogen stream was used. Temperature of solution: SO°C - 126°C Pressure of System: 7 - 12 in. Hg

Final Conditions: 1. heating flask contained viscous yellowish liquid mixed with white crystals. The crystals consisted of 40.55 wt.% C, 5.17 wt.% H, and 0.61 wt.% N. The liquid contained approximately 20% of the initial TMA.

- 2. water collection flask contained 3S.2 ml of TMA/water (43.7% of the initial TMA).
- 3. absorber flask contained 30.6% of the initial TMA.

Table A-5. Regeneration Run #2

Initial Aqueous Solution: 60 ml of 0.6M succinic acid and 1.2M TMA

Treatment: The solution was heated at 1.05 atm. A nitrogen stream was used. Temperature of solution: 99°C - 220°C

Final Conditions: I. heating flask contained dark brown solid material; decomposition of the succinic acid had probably occurred.

Table A-6. Regeneration Run #3

Initial Aqueous Solution: 60 ml of 0.6M succinic acid and 1.2M TMA

Treatment: The solution was heated at 1.07 atm. A nitrogen stream was used. Temperature of solution: 101°C - 172°C

Final Conditions: 1. heating flask contained viscous yellowish liquid mixed with white crystals.

> 2. water collection flask contained 43 ml of TMA/water (3.6% of the initial TMA).

3. absorber flask contained 32% of the initial TMA.

Conclusion: TMA was lost through leaks in the experimental apparatus.

Table *A-7.* Regeneration Run #4

A.2.2 Thermal Regeneration Experiments conducted with Aqueous Trimethylammonium Succinate and Trimethylammonium Fumarate

Table A-S. Regeneration Run #5

Initial Aqueous Solution: 60 ml of 0.6M succinic acid and 1.2M TMA

Treatment: The solution was heated under vacuum. Nitrogen flowrate: 0.0137 - 0.0158 mOles/min Temperature of solution: 61°C - 112°C Pressure of system: 14.8 in. Hg

- 2. water collection flask contained 43.5 ml of TMA/water (1% of the initial TMA).
- 3. absorber flask contained 80% of the initial TMA.

Time (hrs)	cumulative volume (ml) of H ₂ O collected	cumulative moles of TMA collected
$\bf{0}$	$\mathbf 0$	$\bf{0}$
0.17	$\mathbf{1}$	
0.43	3	
0.79	5.5	0.0072
2.08	17	
2.52	21	
2.72	23	
2.80	24	
2.90	25	
2.97	26	0.0144
3.07	27	
3.17	28	
3.27	29	
3.35	30	
3.46	31	
3.57	32	
3.63	33	
3.75	34	
3.83	35	0.0216
3.93	36	
3.98	37	
4.10	38	
4.20	39	
4.24	39.5	0.0288
4.28	40	
4.42	41	
4.48	41.5	
4.50	41.75	0.0360
4.55	42	
4.68	43	0.0432
4.76	43.5	
5.08	43.5	0.0504
6.26	43.5	0.0574

Table A-S. (continued)

Table A-9. Regeneration Run #6

Initial Aqueous Solution: 60 ml of 0.6M succinic acid and 1.2M TMA

Treatment: The solution was heated under vacuum. Nitrogen flowrate: 0.0137-0.0158 moles/min Temperature of solution: 61°C - 121.5°C Pressure of system: 14.9 in. Hg

Final Conditions: I. heating flask contained viscous yellowish liquid mixed with white crystals.

- 2. water collection flask contained 44.75 ml of TMA/water (1.25% of the initial TMA).
- 3. absorber flask contained 80% of the initial TMA.

Table A-tO. Regeneration Run #7

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APPENDIX B. NOMENCLATURE

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APPENDIX C. COMPUTER PROGRAMS

This appendix gives the computer programs used to model the partitioning of either lactic, succinic, or fumaric acid between an organic solution of Alamine 336 in methyl isobutyl ketone (MiBK) and aqueous trimethylamine solutions. The equations describing these systems and the numerical methods used to solve the equations were discussed in Chapter 4, Section 1. The computer programs used for lactic and succinic acid were written using the software package, Lotus 123 (Lotus Development Corp., 1983). The computer programs used for fumaric acid were written in the FORTRAN computing language.

The theoretical curve shown in Figure 4-1 was plotted using the results given in columns K and L below. The equations used to generate the numbers in columns B through N are given after the table of results. The variable z is the loading -- the number of moles of acid per mole of Alamine 336 -- in the initial organic phase.

(REGEN31.UK1)

simulation of regeneration by back extraction with tms


```
B21: (S1) 1+10^(A21-$B$10)+10^(2*A21-$B$10-$R$11)
    C21: (S1) 1+10^\circ(9B$10-A21)+10^\circ(AB1-8B$11)D21: (51) 1+10^($B$10+$B$11-2*A21)+10^($B$11-A21)
    E211 (51) +H21 
    F21: (S1) 4**B*8*B21**B*18*E21^3+3*(B21**B#18**B*7-B*9*B*17*B*8+3**B*B*B*B*B
*E21^2+2*(*B*6*B21**B$18-B$17*B$9*B$7+2**B$7*$B$9)*E21+(B21**B$18-B$17*B$9*B$6+$
B$9*$B$6+B$19)
    G21: (F5) ($B$8*B21*$B$18)*E21^4+(B21*$B$18*$B$7-B$17*B$9*B$8+3*$B$9+$B$8)*E
21^3+($B$6*B21*$B$18-B$9*B$17*B$7+2*$B$7*$B$9)*E21^2+(B21*$B$18-B$17*B$9*B$6+$B$
3**B*6+B*19) *E21-*B*17**B*9
    H21. (51) +E21-G21/F21 
    121. +H21*B21 
    J21: (1+10"(A21-B$5))*(-10"-A21+10"(A21-14)+I21/C21+2*I21/D21)
    K21: (F2) +J21/#B#17/#B#9##B#18
    L21: (F2) +I21/($B$9*$B$17)*$B$18
    ~1. +8.17*(L21-1) 
    N21: @LOG(H21)
```
C2 Succinic Acid

The theoretical curve shown in Figure 4-2 was plotted using the results given in columns K and L below. The equations used to generate the numbers in columns B through N are the same as those given above for lactic acid. The variable *z* is the loading -- the number of moles of acid per mole of Alamine 336 -- in the initial organic phase.

(REGEN31.WK1)

simulation of regeneration by back extraction with tms

pKb = 9.80
K11 = 24.55 $K1! = 24.55 24.55$
 $K2! = 81.28 81.28$ K21 = 81.28 81.28
K31 = 0.00 0.00 0.00
 0.30 BTOT = 0.30
pKa1 = 4.21 pKa1 = 4.21
pKa2 = 5.60 pIC.2 • 5.60 **.. --_....** gu ... pI4 1/.1ph. 1/.1ph.' 1/.1ph.2 G df/dx N ... _. Aq Actd TM TMper " Acid Load log(A)


```
FILE: SECOND
               DATA
                         \blacksquareVH/78 SP2 - CMS Release 5.5 1/11/90 BB06A
C THIS PROGRAM WAS USED TO DETERMINE (H2A) VALUES THAT BOUND AN INTERVAL
C WITHIN WHICH FIPP, (H2A)), I.F., FOUATION 4-41, IS FOUAL TO ZERO.
r
      INTEGER K
      DITURLE PRECISION PH.ALPHA.X.AFREE.PFREE.ATOT.Y.KF.KS.
         KT, SOL, KAF, KAS, SW, HPLIIS, RTOT, RR, AA, ANEW.FA.AFO
     \bulletRATIO.THATCT.ANSWER
C THE PROGRAM WAS PIN AT SET VALUES OF AQUEOUS-PHASE PH.
      READ(5,*) PH
      HPLUS=10.**(C.-PH)
C
C VALUES OF THE EQUILIBRIUM CONSTANTS. ACID DISSOCIATION CONSTANTS
C AND PARTITION COFFFICIENT ARE SET
      KF=10.004.33
      KS=10.***. #1
      K7=10.**7.49
      K#F=10.**(0.-3.10)
      K#S=10.**(0.-4.60)
      57L = 2.200SML=SOL#0.85PM
c
C THE INITIAL CONCENTRATION OF EXTRACTANT AND OF ACID IN THE ORGANIC
C PHASE AND THE ORGANIC TO AQUEDUS PHASE PATIO ARE SET
      8707=0.300
      ATOT=0.075D0
      WRITE(6.0) 'ATNT FOUALS'.ATNT
      $W=0.00/3.00C
C SLPHA (EQUATION 4-37) IS CALCULATED
      REPHA=1./(1.+WAF/HPLUS+(WAF#WAS/fHPLUS##2.)))
C X IS THE INITIAL GUESS FOR THE CONCENTRATION OF THE TOTAL AMOUNT OF
C ACTO THE ROUFFUS PHASE AT FOUTLINGTIM
C.
      X = 0.15C ANFULTS THE INITTAL GUESS FOR THE CONCENTRATION OF UNIONIZED ACID
C TN THE AQUENUS PHASE, I.E., FH241, AS CALCULATED RELOW
         ANEW-ALPHA®X
      HRITE(6,*) 'PH IS', PH, 'HPLUS IS', HPLUS
      WRITE(6,0) .ALPHA IS', ALPHA, 'GUESS IS', X
C
C IN THIS LOOP, VALUES OF F (EQUATION 4-41) ARE CALCULATED FOR
C VARYING VALUES OF IH2A)
      DN 10 K=1,100
         AFREE=ANEW
         WRITE(6,*) *AFREE EQUALS*,AFREE
         AEO=AFREE/ALPHA
         WRITE(6.0) '
                           AFO FOUALS', AFO
c
C THE CONCENTRATION OF INCOMPLEXED EXTRACTANT IS CALCULATED
         SFREE=(0.-(KS#(AFREE##2.)+11)+
     ≠ T(KS#(AFRFE##2,}+1,}##2,+4##TNT#{2,#KF#AFRFE+2#KT#(AFPFF##2,}})
     * *<sub>00</sub>.5
         BFREE=BFRFF/(4.0(KF*AFPFF+KT*(AFPFF**2.)))
```

```
VH/YA <P? - CMS Release 5.5 1/11/90 BROEA
FILE: SECOND
               DATA
                         \blacksquareWRITE(6.0) 'PEPEE EQUATS', REPEE
C
C THE VALUE OF F (FOUATION 4-43) IS CALCULATED
         BB=KF*(BFPFF##2.)#SH+SNL#SW+1./ALPHA
         AA=2.0*KS*PFPEF*SW+2.*KT*(PFPFF**2.0)*SW
         FA={{0,-PP}+{{PR##2,}+4,#ATNT#SH#AA}##0,5}/{2#AA}-AFRFE
                             FA FOUALS', FA
         WRITE(6, 0) \rightarrowFA IS POSITIVE!
         IF (FA .GF. O.) WRITER6, *) *
\mathbf cC THE TOTAL CONCENTRATION OF TRIMETHYLAMINE (ALL IN THE AQUECUS PHASE)
C IS CALCULATED
         THATOT={1.+10.==(PH-9.pt))={10.==(PH-14.)={10.=={0,-PH)}+
        {10**{0.-3.10}}*ANEW*{10***H}*2*{10.**{0.-4.60-3.10*2*PH}*ANEW}}
     \bulletWRITE(6.*) 'THATOT EQUALS'.THATOT
c
C THE RINTIN OF MOLFS OF THA TO MINES OF ACID IN THE SYSTEM IS
C. CALCULATED
         RATIO={TWATOT/ATOT}/SW
         WRITE(6,*) 'PATIO EQUALS', PATIO
         IF (FA .LT. 0.) GO TO 20
\mathbf{c}C A NEW GUESS FOR THE CONCENTRATION OF THE TOTAL AMOUNT OF ACID
C IN THE AQUENUS PHASE IS SET
         AEO-AEO+0.01
\mathbf{c}C A NEW GUESS FOR THE CONCENTRATION OF INTONIZED ACID IN THE
C AQUEDUS PHASE IS SET, AS CALCULATED RELIN
         ANEW=AFO*ALPHA
10
      CENTINUE
20
      CONTINUE
         NRITE(6,*) "LAST ANEW FOURLS", ANFW
         WRITE(6.0) 'K FOUALS'.W
         STOP
         END
```

```
FILE: FUMARIC DATA
                             VH/YA SP2 - CHS Release 5.5 1/11/90 8806A
                         \mathbf{A}C THIS FROGRAM WAS USED TO GENERATE THE THEORETICAL BACK-EXTRACTION
C CURVE SHOWN IN FIGURE 4-3
£.
      INTEGER K.N
      DOUBLE PRECISION PH, ALPHA, AFPEF, REPEE, ATOT, Y, TOL, KE, KS,
         KT.SOL.KIF.KAS.SH.HPLUS.ATNT.AR.AA.Z.FA.FY.
     \bulletRATIO, THATPT, ANSWER, UPPFR, LINER, BY, BRY, AAY, TEST,
     \bullet\bulletTLOWER.TUPPFP.RFCOVER
c
C VALUES OF THE EQUILIBRIUM CONSTANTS, ACID DISSOCIATION CONSTANTS,
      KF=10.*** .33
      K5=10.***. 81
      KT = 10.497.49KAF=10.**(0.-3.10)
      KhS=10.**(0.-4.60)
      SOL=2.200
      SML=SOL*0.85
      TOL = 0.0000001e
C THE INITIAL CONCENTRATION OF FITRACTANT AND OF ACID IN THE OPGANIC
C PHASE AND THE DREAMIC TO AQUEDUS PHASE RATIO APE SET
      RTN=0.300
      AF07=0.28575P0
      SP = 8.00/3.00c
C TN THTS LOOP, PH VALUES ARE SET. THE UPPER- AND LOWER-ROUND VALUES
C FOR THE TOTAL CONCENTRATION OF ACTO IN THE AQUENUS PHASE AS
C DETERMINED IN THE PROGRAM, "SECOND", ARE READ IN
      0030 N=1,16READ(5.*) PH.TLOWER.TUPPER
        HPLUS=10.*****-PH)
        WRITE(6, *) *\bulletWRITE(6,*) 'PH FOUALS', PH, 'HPLUS FOUALS', HPLUS
        WRITE(6, *) 'PARTITION FOUALS', SML
t
C ALPHA (FOUATION 4-37) IS CALCULATED
        ALPHA=1./(1.+KAF/HPLUS+fKAF*KAS/fHPLUS**2.)))
        WRITE(6,*) 'ALPHA IS', ALPHA
        WRITE(6,0) 'CUESSES ARE', TINKER, TUPPER
C
C IN THIS LOOP, FOR THE SET VALUE OF PH, FOUATION 4-41 IS SOLVED, I.E.,
 THE VALUE OF (H2A) THAT RESULTS IN F = 0 IS DETERMINED.
\mathbf cDR 10 K=1,100
\mathbf cC THE UPPER- AND LOWER-POUND VALUES OF (H2A) ARE CALCULATED
        LOWER=ALPHA*TLOWER
        UPPER=ALPHA*TUPPER
         AFPFE=LOWFP
C
   CALCULATING F(A), I.F., FEPH, (H2AILTWER)), FOUATION 4-41
\mathbf cc
         BFPEE=(0.-(FS*(AFPEE**7.)+1))+
     * ${KS*{AFRFE##2.}+1.}##2.+4##TNT#{2.*KF#AFRFE+2#KT*{AFPEF##2.}}}
     * * * 0.5
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FFLF: FUMARIC DATA
                              V^{\mu}/7A SP2 - CMS Release 5.5 1/11/90 BRO6A
         BFREE=9FRFF/(4.*(KF*AFPFF+KT*(AFRFF**2.)))
          RR=KF*(RFPFF##2.)*SW+SPL*SW+1./AIPHA
          AA=2.0#KS#PFPFF#SH+2.#WT#{RFPFF##2.0)#SH
          FA=((0.-RP)+((PR$$2.)+4.$ATOT$SH$4A)$$0.5)/(2$AA)-AFDFF
\mathbf cCALCULATING FY. I.E.. F FPH. (M2A:IIDDER/2+LOWER/2)]. FOUATION 4-41
\mathbf c\mathbf{r}Y=LOWER+(UPPFR-LOWER)/2.
          PY=(0.-(KS#fY##2.)+1))+
       | ${KS#{Y##2, }+1, }##2,+4#RTNT#{?,#KF#Y+2#KT#{Y##?_}}}
     * 10.5RY=BY/(4.*(KF*Y+KT*(Y**?.)))
          887=KF*{RY002.}#SW+SOL#SW+1./ALPHA
          AAY=2.0*K <* PY*SH+2.*KT*(*****2.0)*SH
         FY={{0,-PPY}+{{BRY##2,}+4,#ATOT#SH#AAY}##0,5}/{2#AAY}-Y
         IF (FY .FR. 0.) GO TO 21
\mathbf cCOMPERING OID AND NEW GUESSES
r
c
          7 = 1TUPPER-TLNWER)/2.
          IF (Z .LE. TOL) GO TO 24
          TEST-FASEY
c
C A NEW VALUE OF ETTHER THE LINER- OR UPPER-ROUND IS SET
          IF (TEST .CT. 0.) TLOWED=V/ALDHA<br>IF (TEST .LT. 0.) TUPPFR=V/ALDHA
      CONTINUE
16
         WRITE(6,*) 'DOESN''T WORK, K FOUALS',K
21CRNTTNUF
         WRITE(6.0) 'FY FOUALS ZERO'
24
       ETNT I NUE
         WRITE(6,*) 'ITERATIONS FOUALS'.K
         WRITE(6.0) *AFREE FOUATS'.Y
C THE TOTAL AQUEDUS-PHASE CONCENTRATION OF ACID AT EQUILIBRIUM IS
C CALCUEATED AS (H2A)/ALPHA, RELOW
         ANSWER-Y/ALPHA
         WRITE(6,*) 'ATOT,EO,AD FONALS',ANSHER
\mathbf cC THE TOTAL CONCENTRATION OF TRIMETHYLANINE (ALL IN THE AOUETUS
C PHASET IS CALCULATED
          THATOT={1.+10.00(PH-9.Pl))#{{JO##(PH-14.))-{10##{0.-PH}}+
         (10**(0*-3_*10)) aya(10***0*10**(10**60-4_*60-3_*10*2*0H)\bulletWRITE(6, *) *THATOT EQUALS*, THATOT
c
C THE RILTIN OF MOLES OF THA TO MOLES OF ACTO IN THE SYSTEM IS
C EALCULATED
         RATIO=(TWATOT/ATOT)/SW
         WRITE(6,*) 'PATIO EQUALS', PATIO
          RECOVER=(ANSWER/ATOT)/SW
         HRITE(6,0) PRECOVERY FOULLS' JRECOVER
          CONTINUE
30
          STRP
         FND
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Output from the program, "fumaric" :

PH EQUALS 9.56000000000000005 HPLUS EQUALS 0.275422870333816626E-09 PARTITION EQUALS 1.87000005245208722 ALPHA IS 0.380185805558739118E-11 6UESSES ARE 0.7599999999999999995 0.76299999999999999 ITERATIONS EQUALS 15 AFREE EQUALS 0.290081734834267825E-11 ATOI,£O,AQ EQUALS 0.7b2999908~~1265~19 IMATOT EQUALS 2.38417483681268627 RATIO ~QUALS *3.12883836852058561* RECOVERY EQUALS 1.00131221581005958

PH E4UALS 9.38999999999999990 HPLUS EQUALS 0.407380277804112837E-09 PARTITION EQUALS 1.87000005245208722 ALPHA IS 0.831751545956632751E-11 CUESSES ARE 0.7599999999999999995 0.7629999999999999998
ITERATIONS EQUALS 15 **ITERATIONS EQUALS** AFREE EQUALS 0.634626353415782408E-11 ATOT, EQ, AQ EQUALS 0.762999908447265590 TMATOT ŁQUALS 2.10618450724493989 RATIU EQUALS 2.1b~02166305110208 RECOVERY EQUALS 1.00131221581C05980

PH EGUALS 9.19999999999999996 HPLUS EQUALS 0.630957344480193315E-09 PARTITION EQUALS 1.810000052~5208722 ALPHA IS O.199521523019981211E-l~ GUESSES ARE U.159999999999999995 0.162999999999999996 ITERATIONS EQUALS 15 AFREE EQUALS O.151951468535213842E-IC ATDT.EQ,AQ EQUALS 0.161609405517516110 TMATOT EQUALS 1.89711979264111497 RATIO EQUALS 2.48965852052639747 **RECOVERY EQUALS U.999467408815716669**

PH EQUALS 9.00000000000000000 HPLUS EQUALS 0.10000000000000000IE-08 PARTITION EQUALS 1.87000005245208722 ALPHA IS 0.501166043545598904E-10 ~UfSSES AH~ 0.7~999~999999999~95 0.762999999999999998 ITERATIONS EQUALS AFREE EQUALS 0.3P1733847117881006E-10 ATOT.EQ.AQ EQUALS 0.761688323974608825 TMATUT EQUALS 1.75929563952453250 RATIO EQUALS 2.30878692850988521 RECUVERY EQUALS 0.999590976344631024

PH EQUALS 8.0uOOOOOOOOOOOOOOO HPlUS EQUALS 0.10000000000000000ZE-07 PARTITION EQUALS 1.87000005245208722 ALPHA IS C.500988545660584509E-08 GUESSES ARE 0.7599999999999999995 0.7629999999999999998
TTERATINNS FAILLES 15 **ITERATIONS EQUALS** AFREE EQUALS 0.381743440993960973E-08 ATOf,EQ.AU EQUALS 0.76198037119726~136 TMATUT EQUALS 1.54725679461296739 RATIO EQUALS 2.03052072783853976 RECUVERY EQUALS 0.999974248290372861

PH EQUALS 1.0000000000000000u HPlUS EQUALS O.999999999999999955E-07 PARTITION EQUALS 1.87000005245208722 ALPHA IS C.499200391302131503E-Cb GUESSES ARE 0.75600000000000000000005 0.76199999999999999997
ITERATIONS EQUALS 16 **ITERATIONS EQUALS** AFREE EQUALS 0.379426132963017C77E-06 ATOT.EQ.AQ EQUALS C.760067779541015329 TMATUT EQUALS 1.51947029464512817 RATIO EQUALS 1.99~055504783632&v RECUVENY EQUALS 0.997464277612880992

PH EGUALS 6.21999999999999997 HPLUS EQUALS 0.602559586074357777E-06 PARTITION EQUALS 1.87000005245208722 ALPHA IS 0.177704296609843232E-04 LUlSSfS ARE v.lv1000000000000003 0.71199999999999999' ITERATIONS EQUALS AFREE EQUALS 0.126280321293523223E-04 ATOT.ŁQ.AQ EQUALS 0.710620529174804547 THATUT EQUALS: 1.40492894214835329 RATlu EQUALS 1.~'313B76922356073 RECUVERY EQUALS 0.932572872932814365

PH EGUALS 5.27000000000000002 HPLUS EQUALS 0.537031796370252712E-05 PARTITIUN EQUALS 1.87000005245208722 ALPHA IS C.118942745931729504E-02 ~UESSES ARE 0.699999999999999991 0.711999999999999994 ITERATIONS EQUALS 17 AFREE EQUALS 0.846872242138577722E-03 ATOT, EQ, AQ EGUALS 0.711999908447265531 TMATOT EQUALS 1.29101635230516912 RATIO EQUALS 1.70219993740914588 RECOVERY EQUALS 0.934383081951792024

PH EQUALS 5.00000000000000000 HPLUS EQUALS 0.999999999999999999997E-05 PARTITIUN EQUALS 1.87000005245208722 ALPHA IS 0.357195560435231768E-02 GUE SSES ARE 0.23700000000000000002 0.2410000000000000006 ITERATIUNS EQUALS 16 AFREE EQUALS 0.658596401548687412E-03 ATUT, EU, AC EUUALS 0.240371520996v93616 TMATOT EQUALS U.410821451237148999 RATIL EQUALS 0.539135762778410771 RECUVERY EQUALS 0.315448190283587421

PH EQUALS 4.9499999999999996 HPLUS EQUALS 0.112201845430196356E-04 PARTITION EQUALS 1.87000005245208722 ALPHA 15 C.434247165188559377E-02 'U~SSES AW~ 0.£17999999999999999 0.222000000000000003 **ITERATIONS EQUALS** AFREE EQUALS 0.959611757861260372E-03 ATUT, EQ.AQ EQUALS v.220982849121093616 TMATUT EQUALS 0.372105065538926041 RATIU EQUALS 0.488326857662632577 RECUVERY EQUALS G.290003739004059860

PH EQUALS 4.6999999999999996 HPLUS EQUALS 0.199526231496887998E-04 PARTITIUM EQUALS 1.07000005245208722 ALPHA IS 0.109975514125581365E-01 GUESSES ARE v.iJ7999999999999998 0.139999999999999999 **ITERATIONS EQUALS** AFREE EQUALS u.153965652652087007E-02 ATOT, EU, AC EQUALS 0.139999938964843723 TMATUT EQUALS 0.215607469649707520 RATIO EGUALS 0.282949435235836641 RECUVERY EUUALS 0.183726954022104622

PH EQUALS 4.600000000000000009 HPLUS EQUALS 0.251188643150957948E-04 PARTITIUN EQUALS 1.87000005245208722 ALPHA IS 0.1556529753291689C8E-01 GUESSES ARE 0.113000000000000003 0.115000000000000005 ITERATIONS EQUALS AFREE EQUALS 0.179C00826625507528E-02 ATUT,EY,AY EQUALS 0.114999938964843729 TMATUT EQUALS 0.169790777826511460 RATIO EQUALS 0.222822543079411364 RECOVERY EQUALS 0.150918555071973395

PH EQUALS 4.54000000000000004 HPLUS EQUALS 0.288403150312660563E-04 PARTITION EQUALS 1.87000005245208722 ALPHA IS 0.190305389721107621E-01 GUESSES ARE 0.9900000000000000047E-01 0.102999999999999999 ITEMATJUNS EQUALS Ib AFREE EQUALS 0.195011007016773400E-02 ATOT, EQ, AQ EQUALS C.102440368652343669 TMATOT EQUALS 0.147241998412571518 RATIO EQUALS 0.193230969C45369447 RECUVERY EWUAL> v.134436179333764336

PH EQUALS 4.330000000000000007 HPLUS EQUALS 0.467735141287198128E-04 PARTITIUN EQUALS 1.87000005245208722 ALPHA IS C.368969418562646086t-01 GUESSES ARE 0.64000000000000000013t-01 0.68000000000000000000000000019E-01 **ITERATIONS EQUALS** AFREE EQUALS 0.245567119098119227E-02 ATOT, EW, AW EQUALS C.665548706054686634E-OI TMAIUT EQUALS v.664486551669359088E-01 RATIU EQUALS 0.113449678696766282 RECUVERY EQUALS 0.873423496759431233E-01

PH EGUALS 3.95999999999999996 HPLUS EQUALS 0.109647819614318512E-03 PARTITIUN EQUALS 1.87000005245208722 ALPHA 15 0.100969936798364399 GUESSES ANE O.279999999999999997E-Ol 0.319999999999999998E-Cl ITERATIONS EGUALS 16 AFREE EQUALS 0.311219000201320815E-02 ATOT,EQ,AQ EQUALS O.308229370117187~13E-01 TMATOI EQUALS 0.327660850492828620E-01 RATIO EQUALS 0.430001116132321011E-01 RECUVERY EQUALS 0.404500485718093678E-01

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PH EGUALS 2.82000000000000006 HPL
PARTITION ENJALS 1.87000005245208722 HPLUS EQUALS 0.151356124843620793E-02 ALPHA IS C.652095950581069950 GUESSES ARE U.4U0C00U00000000008E-02 0.7000000000000000015E-02 ITERATIONS EQUALS $\overline{15}$ AFREE EQUALS 0.368748240218568125E-02 ATOT, EQ, AQ EQUALS 0.565481567382811884E-02 \sim TMATUI EQUALS 0.485888644620219691E-03 RATIU EQUALS 0.637649139921548150E-03 RECOVERY EQUALS 0.742101794465632389E-02

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FTLF: EQUILE
                DATA
                               VM/YA SD2 - CMS Release 5.5 1/11/90 BROEA
                          \blacksquareC THIS PROGRAM WAS USED TO GENEPATE THE EQUILIPRIUM CUPVES SHOWN IN
C FIGURES 6-5 AND 6-6
r
       INTEGER K.N
      DRUBLE PRECISION PH.ALPHA.AFPFF.AFPFF.ATOT.Y.TOL.KF.KS.
          KT, SOL, KAF, KAS, SH, HPLUS, RTOT, RR, AA, Z, FA, FY,
     \bulletRATIO, THATPT, ANSWER, UPPFR, LOWER, PY, BBY, AAY, TEST,
     \bullet\bulletTLOWER, TUPPER, PECNYER, TOLL, TESTR, O, OTE, TTUPPER, TTLOWER
c
C VALUES OF THE EQUITIBRIUM CONSTANTS, ACID DISSOCIATION CONSTANTS,
C AND PARTITION COFFFICIENT APF SET
      KP=10.***.33
      K5=10.***.81
       KF=10.**7.49
      KAF=10.**(0.-3.10)
       K#5=10.**(0.-4.60)
       501*2.200SOL=SOL#0.85PP
       TUL=0.0000001
       TOLL=0.00001
\mathbf cC THE INITIAL CONCENTRATION OF EXTRACTANT IN THE ORGANIC PHASE AND C THE ORGANIC TO APIEOUS PHASE RATIO ARE SET
      870T=0.300
       59=8.00/3.00
\mathbf cC THE PN, AMOUNT OF ACID IN THE SYSTEM, AND THE UPPER- AND LOWER-
C BOUND VALUES AS PETERMINED IN THE PROGRAM, "SECOND", APF READ IN
       READ(5.*) ATPT.PH.TTLOWER.TTIMPPF9.NTF
\mathbf cC IN THIS LOOP, EQUATION 4-41 IS SPLVED FOR THE GIVEN VALUES OF TOTAL
  ACTO AND AQUEOUS-PHASE PH
r
       DU 30 N=1.100
         HPLUS=10.##fr.-PH)
         WRITE(6.0) '
         WRITE(6.*) 'ATNT EQUALS',ATNT
         WRITE(6,*) *PH FOUALS*, PH,*HPLUS FOUALS*, HPLUS
         URITE(6,0) 'PARTITION FOUALS', SML
 ALPHA (FOUATION 4-37) IS CALCULATED
\mathbf cALPHA=1./(1.*KAF/HPLUS+(WAF#KAS/fHPLUS**2.)))
         WRITE(6,*) 'ALPHA IS', ALPHA
         WRITE(6,*) . . CUESSES ARE., TTLOWER, TTUPPER
         TLOWER=TTL CHER
         TUPPER=TTUPPFP
      DR 10 K=1,100
C
C THE VALUES OF (H2A), I.F., AGUEDUS-PHASE UNIONIZED ACID CONCENTRATION,
C ARE CALCULATED
          LOWER=ALPMA#TLOWER
          UPPER=ALPHA#TUPPER
           AFREE-LOWFP
c
   CALCULATING F(A), I.F., FOUATION 4-41, WITH (H2A) = LOWER
\mathbf cC
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FILE: FOUILE
                              VM/YA SD2 - CMS Release 5.5 1/11/90 BRO6A
                DATA
                          \blacksquareRFREE=(0.-(KS#(AFREE##2.1+1))+
     * T{KS*(AFREE**2.)+1.}**2.+4********/2.*KF*AFREF+2*KT*(AFPFF**2.))}
     \bullet*0.5BFREE=BFRFF/(4, *{KF*#FPFF+YT*(AFPEF**2.)))
          BR=KF*(BFPFF##2.)*SW+SPL*SW+1./ALPHA
          AA=2.0*KS*PFREF*SW+2.*YT*(REPFF**2.0)*SW
          FA=({0,-RP}+({PR##2,}+4,#ATOT#SW#AA}##0,5}/(2#AA}-AFRFF
\mathbf c\mathbf{c}CALCULATING FY. T.F.. EQUATTON 4-41. WITH (H2A) = LOWER/2 + UPPER/2
r
          Y=LOWER+ftPPFR-LOWER)/2.
          RY=(0.-(KS*(Y**Z.)+1))+
     \bullet${KS*(Y**2.)+1.)##2.+4*RTNT#{2.*KF*Y+2*KT*{Y**2.)))
     \bullet**0.5
          BY=BY/(4, *{FF*Y+KT*(Y**?.)))
          RRY=KF*fPY##2.J*SW+SDL*SW+1./ALPHA
          AAY=2.0*KS#AY*SH+2.*KT*(RY**2.0)*SH
          FY={{0,-RPY}+{{RBY**2,}+4,*ATTT#SI*AAY}**0,5}/{2*AAY}-Y
          IF (FY .FO. 0.) GO TO 21
\mathbf cCOMPREING OLD AND NEW GUESSES
r.
Ċ
          Z = (TU^{\text{op}} E R - T1U^{\text{WFR}})/2.
          IF (Z .LF. TOL) GO TO 24
          TEST=FA*FY
r.
C A NEW VALUE FOR FITHER THE UPPER OR LOWER BOUND IS SET
         IF (TEST .CT. 0.) TLOWER-Y/ALPHA
      CRNTINUE
10
         WRITE(6.0) 'DOESN''T WOOK, K FOUALS'.K
      CBNTINUE
21
         WRITE(6,*) 'FY EQUALS 7FPN'
24EFWT TNUF
         WRITE(6,*) 'ITERATIONS FOUALS', K
          WRITE(6.*) 'AFPEF FOUALS'.Y
\mathbf cC THE TITTAL CONCENTRATION OF ACTO THE AQUEOUS PHASE AT EQUILIBRIUM
C IS CAECULATED
          ANSHER=Y/ALPHA
         WRITE(6,*) VATNT, EO,AO FOUALS', ANSWER
\mathbf{c}C THE TITAL CONCENTRATION OF TRIMETHYLAMINE CALL IN THE AQUEOUS
C PHASE? IS CALCULATED
         THATOT={1,+10,00(PH-9,P1))0{{1000{PH-14,}}-{1000{0,-PH}}+
         (1000(0,-3,10)) eya(1000PH) +20(10.00 (0.-4.60-3.10+20PH)ay))
     \bulletWRITE(6, *) ***ATOT EQUALS', TWATOT
c
C THIS VALUE OF THE CONCENTRATION IS COMPARED TO THE DESIRED VALUE.
C NHICH, IN THIS EXAMPLE, IS 0.76 MML/L
         0 = DABS(THATNT-0.76)
         IF (Q .LE. TOLL) GO TO 34
         TESTR=THATFT-0.76
C IF THE CALCULATED THE CONCENTRATION IS NOT THE DESTRED VALUE. THE
```
FILE: EQUILE DATA VP/YA SP2 - CMS Release 5.5 1/11/90 8806A \blacktriangle C VALUE OF THE LOUFFUS-PHASE PH IS ANJUSTED, AND THE CALCULATIONS ARE **C REPEATED** IF (TESTR .CT. 0.) PH=PH=NTF
IF (TESTR .LT. 0.) GO TN 36 **CONTINUE** 30 WRITE(6.*) 'DDFSN''T WORK. N ='.W **CONTINUE** 36 WRITE(6,*) 'DOESN''T WORK, TESTR =',TESTR $3⁴$ **CONTINUE** URITE(6,*) 'THATOT FOUALS'.THATOT c C THE RILTIO OF HOLFS OF THA TO ACTO IN THE SYSTEM IS CALCULATED RATIO=(THE TOT/ATOT)/SW WRITE(6,*) 'RATIO EQUALS', RATIO RECOVER=CANSHER/ATOT)/SW **WRITE(6,*) 'RECOVERY EQUALS'.RECOVER STOP**

END

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