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#### CONTINUOUS CHLOROSULFONATION OF BENZENE

#### I. PHASE RELATIONS

BY

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A THESIS
SUBMITTED TO THE FACULTY OF
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#### ABSTRACT

This work presents liquid-liquid equilibria data on a system suitable for continuous chlorosulfonation of benzene. Complete information is presented for the ternary, chlorosulfonic acid - sulfuric acid - carbon tetrachloride. A partial investigation of the ternary systems, chlorosulfonic acid - sulfuric acid - benzene-sulfonyl chloride; chlorosulfonic acid - carbon tetrachloride - benzenesulfonyl chloride; and the quaternary system, carbon tetrachloride - benzenesulfonyl chloride - sulfuric acid - chlorosulfonic acid is also presented.

#### APPROVAL OF THESIS

FOR

# DEPARTMENT OF CHEMICAL ENGINEERING NEWARK COLLEGE OF ENGINEERING

BY

#### FACULTY COMMITTEE

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#### INTRODUCTION

#### A. Purpose and Scope

Chlorosulfonation is a common organic reaction.

Industrially it finds application in the preparation of dyes and drugs through organic syntheses (e.g. chlorosulfonation of acetanilide for sulfa drugs, o-toluenesulfonyl chloride for saccharin). Commercial processes are batchwise, the economics of which require efficient recovery and utilization of hydrochloric acid from spent, excess chlorosulfonic acid.

This work was initiated to develop a process for continuous chlorosulfonation. It was envisioned that the technique would involve continuous reaction and extraction of the organic product with recycle of the excess chlorosulfonic acid. Such a process may have economic advantages over the batch method.

Benzene to benzenesulfonyl chloride was chosen for study because of the availability of process information and the relative simplicity of the reaction. The problem herein is concerned primarily with the phase relationships between chlorosulfonic acid, benzenesulfonyl chloride, sulfuric acid and carbon tetrachloride at 25°C. It is planned that further work will utilize the information obtained here to develop the process kinetics and reaction scheme.

#### B. Historical and Theoretical

Chlorosulfonation of benzene (7), (11), (15) toluene (12), naphthalene (10) and related compounds (3), (6), (8), (10), (13), (19), have been reported in the liter-The classical laboratory procedure for the preparation of benzenesulfonyl chloride is described by Gilman and Blatt (7). Industrial methods do not differ much from the laboratory procedure. The German patent (12) describes a process calling for the slow continuous introduction of the hydrocarbons into chlorosulfonic acid at -10°C. The common practice is to carry out the reaction of benzene and toluene at 20-25°C. Use of an inert solvent in chlorosulfonation has also been reported (10), (18). The latter reference is a report of a German technique, discussed below. Data on continuous chlorosulfonation is noticeably lacking. Discussions on continuous sulfonations are available (9), (23) which may serve as a guide to equipment design.

<u>Chlorosulfonation</u>. Chlorosulfonic acid reacts with both aliphatic and aromatic compounds. The reaction may be represented:

$$R \cdot H + C1S0_3H ---- RS0_2OH + HC1 (1)$$
  
 $R \cdot S0_2OH + C1S0_3H ==== RS0_2C1 + H_2S0_{J_1} (2)$ 

To drive the reaction of the second step as far as possible to the right, an excess (50+150%) of chlorosulfonic acid over the two theoretical moles is always required (10). For chlorosulfonation of benzene, the reaction may be carried out by adding the benzene to the acid, a three to five molar excess of the acid being employed at temperatures of 20-25°C. Hydrochloric acid is liberated throughout the reaction. The addition usually requires two to three hours. At completion, the mixture is quenched on ice and the organic material is extracted from the aqueous acid by carbon tetrachloride or another inert solvent. Recovery of the sulfonyl chloride can be accomplished by distillation. Under 10 mm pressure the sulfonyl chloride boils at 112°C. Properties of this compound are given in Table I. Yields vary from 75-77% based on the benzene charged (7). The excess of chlorosulfonic acid is important, since with small excesses, diphenyl sulfone formation becomes more favorable. German report (18) shows that increased yields of benzenesulfonyl chloride may be obtained by the addition of sodium chloride to the reaction. The salt removes sulfuric acid as the sodium acid sulfate, driving the reaction to completion. With carbon tetrachloride present, this salt concentration may be reduced. The data are summarized below. (Table II)

TABLE I

#### Physical Properties of Benzenesulfonyl Chloride

Molecular Weight	176.6
Melting Point	14.5°C.
Boiling Point	251.5°C.
d15	1.3842

Colorless, oily liquid, with penetrating odor.

Insoluble and stable in cold water.

Soluble in ether and alcohol.

#### Vapor Pressure Data (14)

toc.	p (mm Hg)
65.9	1
96.5	5
112.0	10
174.5	100
198.0	200
224.0	400
251.5	760

The rate of hydrolysis has been determined (16) at 25°C. with:

$$0.053 \text{ N H}_2\text{SO}_{\frac{1}{4}} - 0.01$$
  
 $0.136 \text{ N H}_2\text{SO}_{\frac{1}{4}} - 0.01$   
 $0.26$  N H $_2$  SO $_{\frac{1}{4}} - 0.01$ 

TABLE II

Effect of Additives in Chlorosulfonation of Benzene

Method	Ben	zene		cosul-	Na Cl	øs0;	>Cl
	Mols	<u>Grams</u>	Mols	Grams	Grams	Grams	<u>Jo</u>
Gilman and Blatt (7)	2	156	6	700	none	272	76-77
NaCl added	2	156	10.7	1,250	120	<b>31</b> 8	90
NaCl - CCl4	2	156	6	700	120	318	90

Solvents. To develop a continuous chlorosulfonation process, it is desirable to perform the reaction in the presence of an inert solvent. The role of the solvent is to extract the product from the reaction medium as it is formed. The solvent should have a high affinity for the organic product and should separate from spent acids without difficulty. Butual solubilities of acids and solvent must be low. Recovery of the product should be possible by simple means (e.g. solvent evaporation) with recycle of the inert solvent to the reaction. The literature discusses carbon tetrachloride, carbon disulfide and chloroform as solvents for chlorosulfonation (10).

Liquid-liquid equilibria. Some knowledge of the liquid-liquid equilibria in this system is necessary. The kinetics and subsequent process development depend, to a large extent, on the extraction operation. Hence, attention must be given to liquid-liquid relationships. For this work, interest centers on the ternary and quaternary systems. The ternary formed from two pairs of partially miscible liquids is of special concern. Theoretical consideration of both the ternary and quaternary liquid systems are discussed by Treybal (21). Various methods for obtaining and presenting data are also given. Data on four component systems are relatively limited.

A recent review of liquid-liquid extraction (22) cited nine references on four component systems. Brancker, et al. (1). (2) presented data on chloroform-water-acetic acid-acetone at 25°C. A tetrahedral representation was employed to represent the four component equilibrium condition. Chang and Moulton presented data on the system, benzene-ethyl isovalerate-ethyl alcohol-water (4). Other methods of representation have been presented. Cruikshank, et al. (5) described a graphical method using cartesian coordinates where a ternary of three completely miscible liquids is treated with a solvent which is immiscible with one of them. A method that may have application to the four components concerned with herein was published by Prince, et al. (17). The paper is concerned with the correlation of phase compositions in systems where one component is partially miscible with two others and completely miscible with the third. The correlations may have application to the four component system under consideration here namely, carbon tetrachloride, chlorosulfonic acid, sulfuric acid, benzenesulfonyl chloride.

#### **EXPERIMENTAL**

#### A. Apparatus

Extraction and separation of each mixture was carried out in a cylindrical, graduated all-glass separatory funnel. The funnel was graduated to 25 milliliters in half milliliter divisions. Both the top and bottom of the cylinder contained 24/25 tapered joints. The outlet terminated in a fine capillary tip, giving excellent control when separations were made. An air escape line was provided from the top of the funnel (below the point of the stopper) to the tapered joint below the outlet cock. The apparatus was similar to the equipment described in Scientific Glass Company catalogue (item number J 1867).

Additions of the materials to the extractor were made from micropipettes. These were graduated in tenths of a milliliter to a total of 10 milliliters.

Titrations required the use of standard 25 or 50 milliliter burettes.

Chloride titrations were carried out using a Beckman Instruments Inc. automatic titrator (described in Bulletin 239A of the Beckman Instruments Inc.). A titrating accuracy in the range of ±0.5% was obtainable with the instrument.

#### B. Materials

The materials employed were:

Carbon Tetrachloride Merck - Merck & Co., Inc., Rahway,

N. J.

C. P. grade

Sulfuric Acid Merck - Merck & Co., Inc., Rahway,
N. J.

Reagent grade 95-98%

Fuming Sulfuric Acid Merck - Merck & Co., Inc., Rahway,

N. J.

Free SO<sub>2</sub> - 20.0% - 23.0%

Benzenesulfonyl Chloride - Matheson Co. Inc., East

B.P. 102-103º/5 mm

Rutherford, N. J.

Chlorosulfonic Acid - DuPont and Co. Inc.,
Wilmington, Del.

100% Sulfuric Acid was prepared from reagent sulfuric acid and fuming sulfuric acid.

#### C. Procedure\*

All materials employed in the experiments were kept in a constant temperature bath at 23-25°C. Prior to mixing, carbon tetrachloride and sulfuric acid were transferred to microburettes and it was assumed that no change in temperature took place. Great care was used to keep the sulfuric acid from contacting the atmosphere. Chlorosulfonic acid and benzenesulfonyl chloride were taken directly from the constant temperature storage by pipettes. The first component added to the extractor in each case was the least dense one. As an example, when working with carbon tetrachloride, chlorosulfonic acid, sulfuric acid, the solvent carbon tetrachloride was added first. The weight of the first component was accurately recorded on an analytical balance. The second component was added in a similar manner and the mixture well shaken. example above, chlorosulfonic acid would have been added next). When proceeding from the one phase system to the two phase system, a technique similar to the cloud point method was used (21). The third, or fourth component was added dropwise with continuous shaking until a

<sup>\*</sup> The chemicals employed here are very corrosive and the utmost care was required in handling them.

cloudiness appeared. At this cloud point, one or two drops more of the component was added to bring about the phase separation. The point of phase change was selected as the equilibrium condition rather than the cloud point because it was more reproducible. The weight of this component was recorded accurately. At the equilibrium point, the mixture was shaken for five to ten minutes then allowed to come to rest. Equilibrium appeared to be reached within a matter of minutes. When obtaining tie line data, the mixture was allowed to settle for fifteen to thirty minutes. After settling, the bottom layer was drained into a weighed twenty-five milliliter, stoppered flask. The top layer was drained into a simi-The weight of both layers was then accurately lar flask\*. determined on the analytical balance. Analysis of either layer could be made.

<sup>\*</sup> The interface was readily observed because the chlorosulfonic acid imparted a slight straw color to the acid layer.

#### D. Analytical Methods

Analytical techniques were essentially the same regardless of the system under investigation. To analyze the bottom, or predominately acid layer, a one or two gram aliquot was taken. This aliquot was carefully added dropwise to a known amount of standard 1 normal caustic. Back titration of the excess alkali with standard 1 normal sulfuric acid provided data on total acidity due to both acids. A Volhard analysis using the titrator gave the accurate estimate of chlorosulfonic acid present. With the three component system, chlorosulfonic acid. sulfuric acid and carbon tetrachloride, the weight percentage of each component in the entire bottom layer could be calculated. The upper solvent layer was analyzed in the same manner but the entire layer was used (0.5-15 grams). Carbon tetrachloride was always determined by difference.

The analytical methods for the four component system differed from the above. Here only the top solvent layer was analyzed. The entire layer (0.5-2.0 grams), after weighing, was cooled in an ice bath, then drowned with fifteen to twenty milliliters of deionized water. The total acid content was rapidly determined by titration

with standard 1 normal caustic, using a phenolphthalein indicator. Following the titration, fifteen to thirty milliliters of standard 1 normal alkali was added and the mixture heated to reflux for half an hour. This caused the carbon tetrachloride to boil off and the benzenesulfonyl chloride to react with the alkali. Back titration of the excess caustic at room temperature with sulfuric acid provided the data on increased acidity due to the sulfonyl chloride. A Volhard titration with the titrator gave total chloride due to the sulfonyl chloride and to chlorosulfonic acid. Quantities of all materials could then be determined with carbon tetrachloride found by difference. Knowledge of one layer provides the estimate for the other layer.

In the vast majority of runs, material balances were better than 97%. Losses of material occurred when the extractor was not rinsed down after the layers had been cut.

In the titrations as employed above, the acid equivalent of chlorosulfonic was one-third of the molecular weight. The acid equivalent of the sulfonyl chloride was half the molecular weight. The equation involved here may be given as follows:

 $H_2SO_{\downarrow\downarrow} + \phi SO_2C1 + C1SO_3H \xrightarrow{H_2O}$ 

 $H_2SO_4 + H_2SO_4 + HC1 + ØSO_2C1 \xrightarrow{NaOH}$ 

 $Na_2SO_4 + Na_2SO_4 + NaCl + ØSO_2Cl \xrightarrow{NaOH}$ 

 $Na_2SO_{14} + Na_2SO_{14} + ØSO_3Na + NaCl + NaOH$ 

#### E. Results

The experimental data are given in the following tables. Information is presented on the ternary systems and the quaternary system. Graphical illustrations are presented in Figures  $1 - \frac{1}{4}$ .

TABLE III

## Solvent Selection

Vol. % ClS03H	Solvent Added	Vol. %	Phases	Comments
50.0	Chloroform	50.0	1	
61.5	Chloroform	38.5	l	
28.6	Chloroform	71.4	1.	
50.0	Skelly B	50.0	1	Reaction and darkening of mixture
50.0	Tetrachlorethane	50.0	1	Slow reaction
67.0	Ethylene Dichloride	33.0	1	
50.0	Ethylene Dichloride	50.0	1	
30.0	Ethylene Dichloride	70.0	1	
75.0	Ethylene Dichloride	25.0	1	
30.0	Cyclohexane	70.0	2	Slow reaction
30.0	Cyclohexane (pure)	70.0	2	No reaction
10.0	Carbon Tetrachloride	90.0	2	
25.0	Carbon Tetrachloride	75.0	2	

It appears as if carbon tetrachloride is the most suitable solvent.

TABLE IV

# Chlorosulfonic Acid - Sulfuric Acid - Carbon Tetrachloride

Nun I		
Weight	Weight	70

 Total Weight
 Actual Weight
 Weight 5

 62.0829 gms
 0.7665 gms
 10.00
 H2S04

 61.3164 gms
 3.1701 gms
 41.25
 CIso3H

 58.1463 gms
 3.7690 gms
 48.80
 CC14

54.3773 tare 7.7056 total

Produced a 2 phase system.

#### Run 2

Total Weig	ht Actual	Weight	Weight %	
65.5406 gm 63.9158 gm 56.9481 gm	1.6248 7.4677 2.0630	gms	14.50 67.00 18.50	CISO3H CISO3H
54.3851 ta	re 11.1555	total		

Produced a 2 phase system.

## Run 3

Total Weight	Actual Weight	Weight %
67.2300 gms	0.7933 gms	6.15 H <sub>2</sub> SO <sub>1</sub> ,
66.4367 gms	4.4678 gms	34.62 CC1 <sub>1</sub> ,
61.9689 gms	7.6366 gms	59.22 C1SO <sub>3</sub> H

54.3323 tare 12.8977 total

Produced a 2 phase system.

Run 4

Added more H2SO4 to run 3.

Total Weight	Actual Weight	Weight %	
74.7764 gms	8.3397 gms 4.4678 gms 7.6366 gms	40.65 21.80 37.35	C120 <sup>3</sup> H CC1 <sup>1</sup> †
	20.4441 gms		

Produced a 2 phase system.

#### Run 5

Total Weight	Actual Weight	Weight %	
66.3903 gms 66.2364 gms 61.8280 gms	0.1539 gms 4.4084 gms 7.14487 gms	1.30 36.70 62.00	н <sub>2</sub> 2014 сст <sup>1</sup>
54.3793 tare	12.0110 total		

Produced a 2 phase system.

# Run 6

Employing an acid mixture of 87.03 weight per cent C1SO3H and 12.97 weight per cent H2SO4.

Total Weight	Actual Weight	Weight %	
104.9056 gms 103.3860 acid mixture	12.916 gms 1.925 gms 1.519 gms	78•94 11•76 9•29	<b>с</b> 1803н Н2804 СС14
88.5446 tare	16.360 total		

Produced a 1 phase system.

Run 7

Added more CCI4 to run 6.

Total Weight	Actual Weight	Weight 5	
106.4347 gms	12.916 gms 1.925 gms 3.048 gms	72.19 10.76 17.04	C1803H H2804 CC114
	17.889 total		

Produced a 1 phase system.

## Run 8

Added more CCl1 to run 7.

Total Weight	Actual Weight	Weight 5	
107.9514 gms	12.916 gms 1.925 gms 4.565 gms	66.55 9.92 23.52	CC17 HS20/t CC17 CC17
	19.406 total		

Produced a 2 phase system. Top layer just visible.

Run 9

Using acid mixture as in run 6.

Total Wei	<u>lght</u>	Actual V	Velsht	Weight 5	
110.8396 96.9545	gma gms	13.8851 6.9953 1.0425	gms gms	63.34 31.91 4.75	CCI), C1SO3H H2SO4
88.9166	tare	21.9229	total		

Produced a 2 phase system.

#### Run 9 (continued)

#### Equilibrium Points

Acid Layer	Actual Weight	Weight %	
	1.7856 gms 6.6344 gms 0.9690 gms	19.02 70.66 10.32	cci <sub>li</sub> ciso <sub>3</sub> h h <sub>2</sub> so <sub>1</sub>
	9.3890 total		
Solvent Layer	12.0995 gms 0.2609 gms 0.0735 gms	97.31 1.77 0.82	ссі <sub>і</sub> сі sо <sub>3</sub> н н <sub>2</sub> sо <sub>4</sub>
	12.4339 total		

#### Run 10

Using acid mixture as in run 6.

Total Weight	Actual We	eight Weight S	
106.4346 gms 99.2579 gms 88.9103 tare	7.1767 g 9.0055 g 1.3421 g	ms 40.95 ms 51.39 ms 7.65	ссі). с1303н н <sup>5</sup> 201
	17.5243 t	otal	

Produced a 2 phase system.

## Tie Line Data

Acid Layer - 13.3547 gms
Solvent Layer - 3.9470 gms

Assay of Solvent Layer - 2.6 meq total acid 29.68 mg Cl-

# Run 10 (continued)

# Equilibrium Points

Solvent Layer	Actual Weight	Weight %	
	0.097 gms 0.042 gms 3.808 gms	2.46 1.06 96.48	C1SO3H H2SO4 CC14
	3.947 total		
Acid Layer	3.3687 gms 8.9580 gms 1.3000 gms	24.72 65.74 9.50	стго <sup>3</sup> н н <sup>5</sup> го <sup>1</sup>
	13.6267 total		

#### Run 11

Using an acid mixture as in run 6.

Total Weight	Actual Weight	Weight %	
107.1845 gms 102.5017 gms 88.9113 tare	4.6828 gms 11.8277 gms 1.7627 gms	25.63 64.73 9.64	ссі <sub>ј</sub> сізб <sub>3</sub> н н <sub>2</sub> зој
	18.2732 total		

Produced a 2 phase system.

# Equilibrium Points

Acid Layer	Actual Weight	Weight %	
	11.7947 gms 1.7027 gms 3.6361 gms	68.60 10.25 21.15	cc1/† H520/† C1203H
	17.1335 total		

Run 12
Using an acid mixture 67.49% C1S03H - 32.50% H2S04.

Total Weight	Actual Weight	Weight %	
114.2622 gms 106.7000 gms 88.0677 tare	7.5622 gms 12.5758 gms 6.0562 gms	28.87 48.00 23.12	C1SO3H C1SO3H
	26.1945 total		

Produced a 2 phase system.

#### Equilibrium Points

Solvent Layer	Actual Weight	Weight %	
	6.0537 gms 0.2110 gms 0.0583 gms	95•74 3•34 0•92	ссі <sub>ј</sub> стаозн пааој
	6.3230 total		
Acid Layer	1.5085 gms 12.3648 gms 5.9979 gms	7.59 62.22 30.18	ссіј сізозн н <sub>2</sub> зој
	19.8712 total		

#### Run 13

Using acid mixture as in run 12.

Total Weight	Actual Weight	Weight %	
103.6673 gms 91.1390 gms 88.0540 tare	8.4559 gms 4.0722 gms 3.0850 gms	54.16 26.08 19.76	C1S03H H2SO4 CC14
	15.6131 total		

Produced a 2 phase system.

Run 13 (continued)

Equi	lil	ori	um l	Po:	ints

Solvent Layer	Actual Weight	Weight %	
	1.6193 gms 0.0300 gms 0.0050 gms	97.88 1.81 0.31	CC1/ <sub>1</sub> C1SO <sub>3</sub> II H <sub>2</sub> SO <sub>1</sub>
	1.6543 total		
Acid Layer	1.4657 gms 8.4250 gms 4.0672 gms	10.50 60.36 29.1/ <sub>1</sub>	cc1 <sup>6</sup> c120 <sup>3</sup> H H <sup>2</sup> 20 <sup>4</sup>
	13.9579 total		

# Run 14

Total Wei	<u>lght</u>	Actual V	<u>Velght</u>	Weight %	
111.7956 109.1784 92.5400	gms gms	2.6172 16.6384 4.4800	gms	11.03 70.10 18.87	cci <sup>†</sup> ci so <sup>‡</sup> H
88.0600	tere	23.7356	total		

Produced a 2 phase system, just barely separating.

# <u>Run 15</u>

Total Weight	Actual Weight	Weight %	
105.4060 gms 105.2148 gms 94.1078 gms	0.1912 gms 11.1070 gms 6.0516 gms	1.10 64.38 34.88	сст <sub>4</sub> стзо <sup>3</sup> н
88.0562 tar	e 17.3498 total		

Equilibrium point - appearance of 2 phases.

Run	16

Total Weight	Actual Weight	Weight %	
116.9440 gms 110.0148 gms 91.1830 gms	6.9292 gms 18.8318 gms 3.1407 gms	23.97 65.16 10.87	H <sub>2</sub> SO <sub>1</sub> С1SO <sub>3</sub> н СС1 <sub>14</sub>
88.0423 tare	28.9017 total		

Equilibrium point - appearances of 2 phases.

# Run 17

Total Weight	Actual Weight	Weight %	
120.0300 gms 119.44.14 gms 97.9607 gms	0.5886 gms 21.4807 gms 9.9207 gms	1.73 67.22 31.05	сст <sub>4</sub> стгозн стгозн
88.0400 tare	31.9555 total		

Equilibrium point - appearance of 2 phases.

# <u>Run 18</u>

Total Weight	Actual Weight	Weight %	
143.1388 gms 105.9388 gms 89.6246 gms	37.2000 gms 16.3142 gms 1.5712 gms	67.60 29.60 2. <b>7</b> 0	п <sub>2</sub> sо <sub>4</sub> стзозн сст <sub>4</sub>
88.0534 tare	55.0854 total		

Produced a 2 phase system.

## Run 19

Total Weight	Actual Weight	Weight %	
101.2800 gms 92.6715 gms	8.6085 gm <b>s</b> 4.6304 gms	65.02 34.98	clso <sub>3</sub> H cc1 <sub>4</sub>
88.0411 tare	13.2389 total		

Borderline on 2 phases.

Run 20

Added to run 19.

Total Weight	Actual Weight	Weight %	
103.9218 gms 103.44683 gms 92.6715 gms	0.4535 gms 10.7968 gms 4.6304 gms	2.86 67.98 29.16	ciso3H cci <sup>7</sup>
88.0411 tare	15.8807 total		

Equilibrium point - appearance of 2 phases.

#### Run 21

Total Wei	ght	Actual V	Weight	Weight %	
100.3228 97.7910 89.6400		2.5318 8.1510 1.5926	gms	20.63 66.40 12.97	CI SO3H CI SO3H
88.0474	tare	12.2754	total		

Equilibrium point - appearance of 2 phases.

# <u>Run 22</u>

Total Weight	Actual Weight	Weight %	
106.2820 gms 100.8810 gms 89.6660 gms	5.4010 gms 11.2150 gms 1.6250 gms	29.61 61.48 8.91	H <sub>2</sub> SO <sub>1</sub> CISO3H CC1 <sub>4</sub>
88.0110 tare	12.8/100 total		

Equilibrium point - appearance of 2 phases.

## Run 23

Total Weight	Actual Weight	Weight %	
114.2430 gms 105.8284 gms 90.1490 gms	8.4146 gms 15.6794 gms 2.0982 gms	32.13 59.86 8.01	H <sub>2</sub> SO <sub>]</sub> С1SO <sub>3</sub> H ССІ <sub>]</sub>
88.0508 tare	26,1922 total		

Equilibrium point - appearance of 2 phases.

#### Run 24

Total Wei	ght	Actual V	Weight	Weight %	
103.5822 101.9200 90.7630		1.6622 11.1570 2.7146		10.70 71.82 17.48	cci <sup>1†</sup> cieo3h H <sup>5</sup> eo <sup>1</sup>
88.0484	tare	15.5338	total		

Equilibrium point - appearance of 2 phases.

## Run 25\*

Total Weight	Actual Weight	Weight %	
94.3320 gms 93.8640 gms 89.3063 gms	0.4600 gms 4.5577 gms 6.3181 gms	4.12 40.18 55.70	ccij H2SOU ciso3H
82.9882 tare	11.3438 total		

Equilibrium point - appearance of 2 phases.

# Run 26

Total Weight	Actual Weight	Weight %	
90.0860 gms 89.5260 gms	0.5600 gms 6.5460 gms	7.88 92.12	CISO3H
82.9800 tare	7.1060 total		

Produced a 2 phase system.

## Run 27

Added ClSO3H to run 26.

Total Weight	Actual Weight	Weight %	
92.4360 gms	2.9100 gms 6.5lt60 gms	30.77 69.23	clso3H
	9.4560 total		

Produced a 2 phase system.

<sup>#</sup> Runs 25 - 38 employed 99+% H<sub>2</sub>SO<sub>4</sub>.

Run 28

Total Weight	Actual We	eight	Weight %	
90.8245 gms 90.6196 gms 87.2876 gms	0.2049 gr 3.3320 gr 4.2981 gr		2.61 42.53 54.86	CCI) H2SO1 CISO3H
82.9895 tare	7.8350 to	otal		

Equilibrium point - appearance of 2 phases.

## Run 29

Total Weight	Actual Weight	Weight %	
92.3880 gms 91.6760 gms 88.5431 gms	0.7120 gms 5.5274 gms 3.1329 gms	7.60 59.00 33.40	ccil ciso3H H <sup>S</sup> so <sup>†</sup>
83.0157 tare	9.3723 total		

Equilibrium point - appearance of 2 phases.

# <u>Run 30</u>

Total Weight	Actual Weight	Weight %	
91.2810 gms 91.0950 gms	0.1860 gms 7.2263 gms	2.51 97.49	clso3H
83.8687 tare	7.4123 total		

Equilibrium point - appearance of 2 phases.

# Run 31

Total Weight	Actual Weight	Weight %	
94.4934 gms 94.3715 gms	0.1219 gms 10.5263 gms	1.1 98.9	H220/4
83.8452 tare	10.6482 total		

Equilibrium point - appearance of 2 phases.

Run	3	2

Total Weight	Actual Weight	Weight %	
107.4300 gms 103.9150 gms 90.0020 gms	3.5150 gms 13.9130 gms 6.1153 gms	14.93 59.09 25.97	C1S03H H2SO4 CC14
83.8867 tare	23.5433 total		

Produced a 2 phase system.

#### Run 33

Total Weight	Actual Weight	Weight %	
100.0920 gms 95.0261 gms 85.9330 gms	5.0659 gms 9.0931 gms 2.0655 gms	31.22 56.04 17.73	cci <sup>l4</sup> ciso <sup>3</sup> H cci <sup>14</sup>
83.8675 tare	16.22/15 total		

Produced a 2 phase system. Total volume 9.6 c.c. Bottom layer - 9.25 c.c.

## Run 34

Added more H2SO4 to run 33.

Total Weight	Actual Weight	Weight %	
105.7200 gms	10.6939 gms 9.0931 gms 2.0655 gms	49.00 41.60 9.40	CCI) <sup>†</sup> CISO3H COI) <sup>†</sup>
	21.8525 total		

Produced a 2 phase system. Total volume 13 c.c.
Bottom layer - 12 c.c.

Run 35

Added more H2SO14 to run 34.

Total Weight	Actual Weight	Weight %	
115.9712 gms	20.9451 gms 9.0931 gms 2.0655 gms	65.24 28.32 6.43	н <sub>2</sub> so <sub>ј</sub> c1so <sub>3</sub> н cc1 <sub>4</sub>
	32.1037 total		

Produced a 2 phase system. Total volume 18.5 c.c. Bottom 17.5 c.c. Bottom layer appears turbid.

Run 36

Added more H2SO4 to run 35.

Total Weight	Actual Weight	Weight %	
125.7152 gms	30.6893 gms 9.0931 gms 2.0655 gms	73.34 21.73 4.93	H <sub>2</sub> SO <sub>]</sub> CI SO <sub>3</sub> H CCI <sub>4</sub>
	41.8479 total		

Produced a 2 phase system.

## Run 37

Total Weight	Actual Weight	Weight %	
103.2603 gms 103.0390 gms 101.2892 gms	0.2213 gms 1.7498 gms 17.3462 gms	1.10 9.15 89.75	сот Стао <sup>3</sup> н Н <sup>5</sup> 20 <sup>1</sup>
83.9430 tare	19.3173 total		

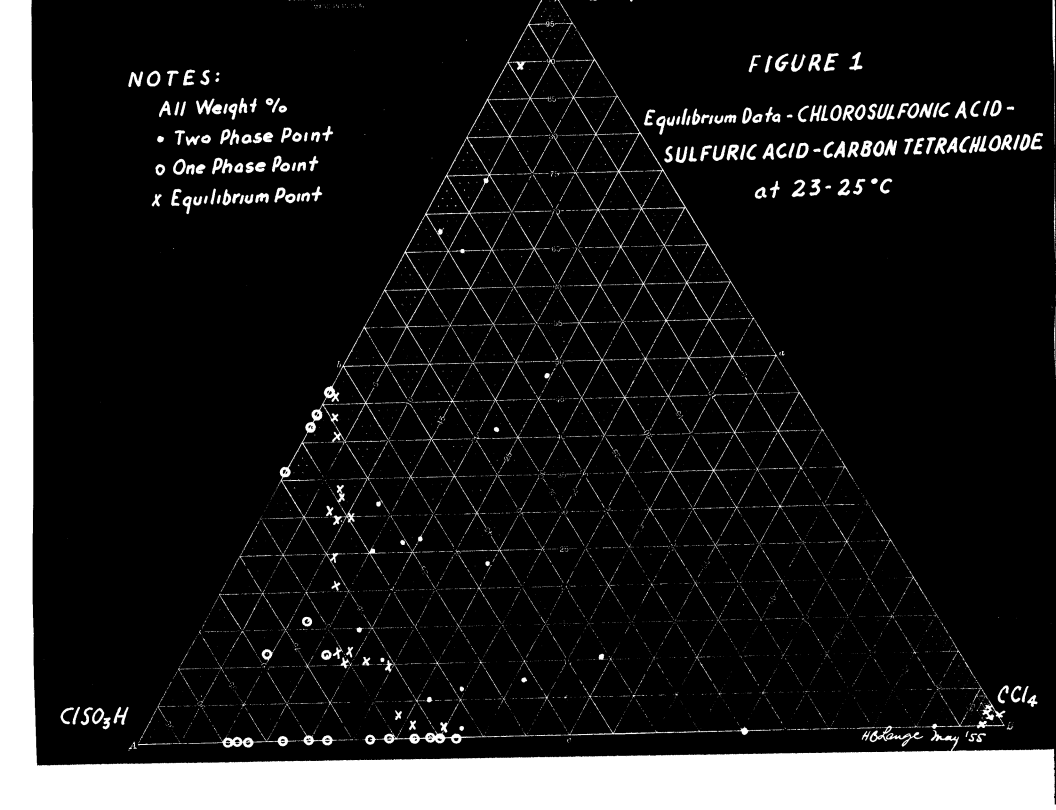
Equilibrium point - appearance of 2 phases.

Run 38

Total Weight	Actual Weight	Weight %	
102.5986 gms 102.2595 gms	0.3391 gms 18.3152 gms	1.82 C 98.18 H	CI[ 2SO[4
83.9443 tare	18.6543 total		

Added CCll dropwise.

Equilibrium point - appearance of 2 phases.



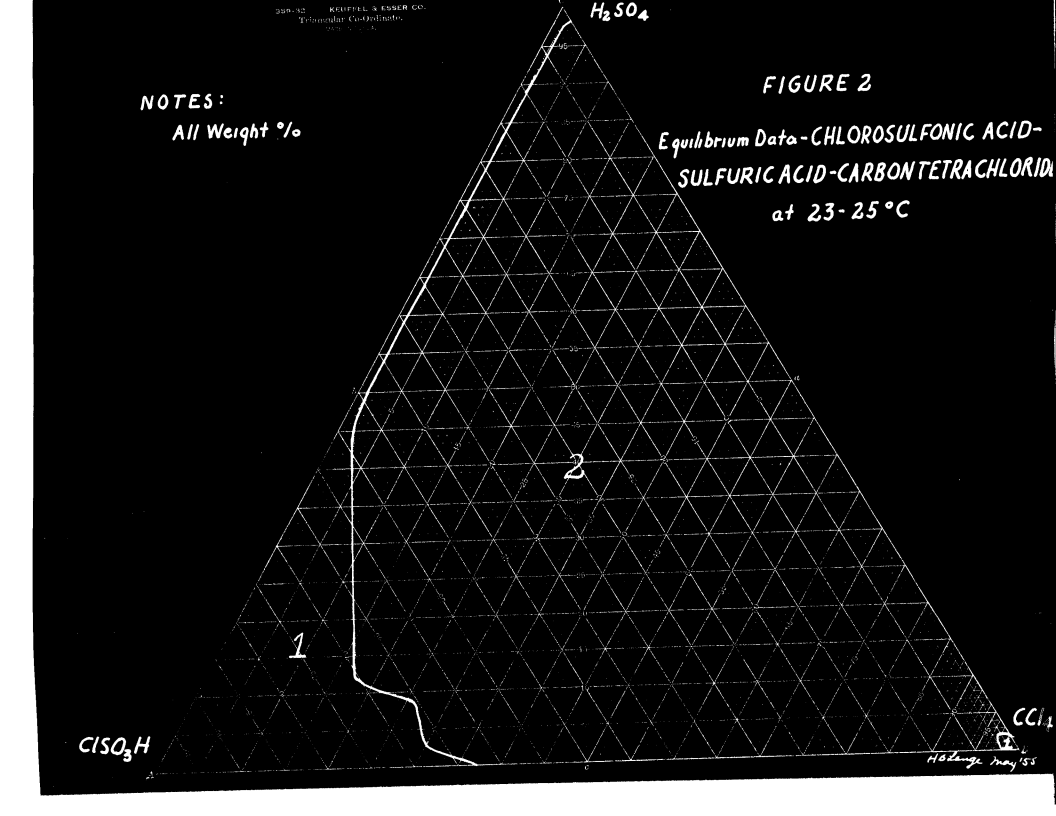


TABLE V

# Chlorosulfonic Acid - Sulfuric Acid - Benzenesulfonyl Chloride

#### <u>Run 39</u>

Total Weight	Actual Weight	Weight %	
106.8585 gms 97.7085 gms 94.4000 gms	9.1500 gms 6.3464 gms 3.3085 gms	48.70 33.72 17.58	H <sub>2</sub> SO <sub>]</sub> C1SO <sub>3</sub> H ØSO <sub>2</sub> C1
88,0536 tare	18.8049 total		•

Produced a 1 phase system.

# Run 40

Added more H2SO4 to run 39.

Total Weight	Actual Weight	Weight S	
143.9585 gms	46.2500 gms 6.3464 gms 3.3085 gms	79.13 13.72 7.15	п <sub>2</sub> 30) <b>с1</b> 30311 ø302 <b>с1</b>
	55.9049 total		

Produced a 1 phase system.

## Run 41

Total Wes	<u>lght</u>	Actual	Weight	Weight %	
109.4343 100.2830 95.9140	gms gms	9.1513 7.8574 4.3690	gms gms gms	42.80 36.85 20.35	H2SO) ØSO2CI CISO3H
88.0566	tare	21.3777	total		

Run 42

Added more H2SO4 to run 41.

Total Weight	Actual Weight	Weight %
127.7343 gms	27.4513 gms 7.8574 gms 4.3690 gms	69.18 H <sub>2</sub> SO <sub>1</sub> 19.80 ØSO <sub>2</sub> C1 11.01 C1SO <sub>3</sub> H

Produced a 1 phase system.

## Run 43

Total Weight	Actual Weight	Weight %	
97.6358 gms 96.9215 gms 94.1700 gms	0.7143 gms 2.7515 gms 6.1153 gms	7.45 28.72 63.83	C1S03H H2SOL ØS02C1
88.0547 tare	9.5811 total		

 $\emptyset$ SO<sub>2</sub>Cl and H<sub>2</sub>SO<sub>1</sub> produced a two phase system titrated with ClSO<sub>3</sub>H to 1 phase.

# Run 114

Total Weight	Actual Weight	Weight %	
97.0528 gms 96.4910 gms 95.6132 gms	0.5618 gms 0.8778 gms 7.5286 gms	6.50 9.80 83.70	C1S03H H2SOJ ØS02C1
88.0846 tare	8.9682 total		

 $\emptyset$ SO<sub>2</sub>Cl and H<sub>2</sub>SO<sub>4</sub> produced a 2 phase system titrated with ClSO<sub>3</sub>H to 1 phase.

# Run 45

Total Weigh	Actual	Weight Weigh	1t %
106.3208 gm 105.6854 gm 92.4311 gm	1.1344 18 13.2543 14.3721	gms 6.40 gms 70.60 gms 23.00	
88.0590 ta	re 18.7608	total	

 $\emptyset$ SO<sub>2</sub>Cl and H<sub>2</sub>SO<sub>4</sub> produced a 2 phase system titrated with ClSO<sub>3</sub>H to 1 phase.

# Run 46

Total Weight	Actual Weight	Weight %	
102.7072 gms 102.5662 gms 90.2786 gms	0.1410 gms 12.2876 gms 2.1978 gms	0.96 83.90 15.04	C1S03H H2SOL ØS02C1
88.0808 tare	14.6264 total		

 $\emptyset SO_2C1$  and  $H_2SO_{\parallel}$  produced a 2 phase system titrated with  $C1SO_3H$  to 1 phase.

# Run 47

Total Weight	Actual Weight	Weight %	
99.6200 gms 98.6910 gms 94.1388 gms	0.9290 gms 4.5522 gms 6.0611 gms	8.05 39.42 52.43	C1S03H H2SO[ ØS02C1
88.0777 tare	11.5h23 total		

 $\emptyset$ SO<sub>2</sub>Cl and H<sub>2</sub>SO<sub>4</sub> produced a 2 phase system titrated with ClSO<sub>3</sub>H to 1 phase.

#### Run 48

Total Weight	Actual Wei	ght Weight S	2
105.1400 gms 103.6035 gms 94.8702 gms	1.5365 gr 8.7333 cm 6.8312 gr	ıs 51.02	C1SO <sub>3</sub> H H2SO <u>[</u> ØSO <sub>2</sub> C1
63.0390 tar	e 17.1010 to	tal	

 $\emptyset$ SO<sub>2</sub>Cl and H<sub>2</sub>SO<sub>1</sub> produced a 2 phase system titrated with ClSO<sub>3</sub>H to 1 phase.

## Run 49

Total Weight	Actual Weight	Weight %	
99.7845 gms 99.2453 gms 98.3421 gms	0.5392 gms 0.9032 gms 10.2937 gms	4.60 7.60 87.70	C1SO3H H2SOL ØSO2C1
88.0484 tare	11.7361 total		

 $\emptyset$ SO<sub>2</sub>Cl and H<sub>2</sub>SO<sub>4</sub> produced a 2 phase system titrated with ClSO<sub>3</sub>H to 1 phase.

## Run 50\*

Total Weight	Actual Weight	Weight 5	
89.0076 gms	0.1006 gms	1.67	C1S03H
88.9070 gms	0.8805 gms	14.61	H2S01
88.0265 gms	5.0453 gms	83.72	ØS02C1

 $\emptyset$ SO<sub>2</sub>Cl and H<sub>2</sub>SO<sub>1</sub> produced a 2 phase system titrated with ClSO<sub>3</sub>H to 1 phase.

<sup>%</sup> Runs 50 - 62 used 99+%  $H_2SO_{4}$ , all others used 98-98.5%  $H_2SO_{4}$ .

Run	51

Total Weight	Actual Weight	Weight %	
89.4109 gms 89.3773 gms 87.5795 gms	0.0336 gms 1.7978 gms 4.5665 gms	0.52 28.10 71.37	CISO3H H2SOL ØSO2CI
83.0130 tare	6.3979 total		

 $\# SO_2Cl$  and  $\mathrm{H}_2SO_{L\!\!\!/}$  produced a 2 phase system titrated with ClSO\_3H to 1 phase.

## Run 52

Total Weight	Actual Weight	Weight %	
91.2330 gms 91.1916 gms 88.0585 gms	0.0414 gms 3.1331 gms 5.0507 gms	0.50 38.09 61.41	C1S03H H2S0[ ØS02C1
83.1078 tare	8.2252 total		

 $\emptyset SO_2Cl$  and  $H_2SO_{l\downarrow}$  produced a 2 phase system titrated with  $ClSO_3H$  to 1 phase.

## Run 53

Total Weight	Actual Weight	Weight %	
93.7810 gms 87.2800 gms	6.5330 gms 4.2380 gms	60.65 39.35	M22 <b>0</b> ] Ø302 <b>0</b> 1
83.0100 tare	10.7710 total		

Produced a 1 phase system.

## Run 54

Total Weight	Actual Weight	Weight %	
86.3171 gms 85.3611 gms	0.9560 gms 2.0999 gms	34.25 65.75	#28 <b>0</b> ] \$80 <b>201</b>
83.2612 tare	3.0559 total		

Run 55

Added more H2SO4 to run 54.

Total Weight	Actual Weight	Weight %	
87.4861 gms	2.1250 gms 2.0999 gms	50.35 49.65	H2SO4 ØSO2 <b>C1</b>
	4.2249 total		

Produced a 2 phase system.

# Run 56

Added more H2SO4 to run 55.

Total Weight	Actual Weight	Weight %	
90.3297 gms	14.9686 gms 2.0999 gms	70.40 29.60	\$202 <b>C1</b>
	7.0685 total		

Produced a 1 phase system.

# <u>Run 57</u>

Total Weight	Actual Weight	Weight %	
104.8210 gms 103.3490 gms	20.0878 gms 1.4720 gms	93.50 6.50	H2SOL ØSO2 <b>C1</b>
83.2612 tare	21.5598 total		

Produced a 1 phase system.

# <u>Run 58</u>

Added more ØSO2Cl to run 57.

Total Weight	Actual Weight	Weight %	
106.8550 gms	20.0878 gms 3.5060 gms	85.20 14.80	H <sub>2</sub> SO <sub>1</sub> ØSO <u>2</u> C1
	23.5938 total		

			<i>ا</i> د
	Run 59		
Total Weight	Actual Weight	Weight %	
94.6140 gms 86.2813 gms	8.3327 gms 2.4238 gms	77.l47 22.53	H <sub>2</sub> SO <sub>2</sub> C1
83.8575 tare	10.7565 total		
Produ <b>c</b> ed a	1 phase system.		
	<u>Run 60</u>		
Added more	ØS02Cl to run 59.		
Total Weight	Actual Weight	Weight %	
99.4900 gms	7.2998 gms 8.3327 gms	46.70 53.30	ØS02 <b>C1</b> H2S <b>0</b> [4
	15.6325 total		
Produced a	l phase system.		
	<u>Run 61</u>		
Total Weight	Actual Weight	Weight %	
105.8775 gms	13.7893 gms 8.3327 gms	62.33 37.67	øs02 <b>c1</b> H2S0 <sub>4</sub>
	22.1220 total		
Produced a	2 phase system.		
	<u>Run 62</u>		
Added ClSO	3H to run 61.		
Total Weight	Actual Weight	Weight %	
105.9205 gms	0.0430 gms 13.7895 gms 8.3327 gms	0.19 62.21 37.60	c1303H &805c1 H <sup>2</sup> 20 <sup>l†</sup>

22.1652 total
Produced a 1 phase system.

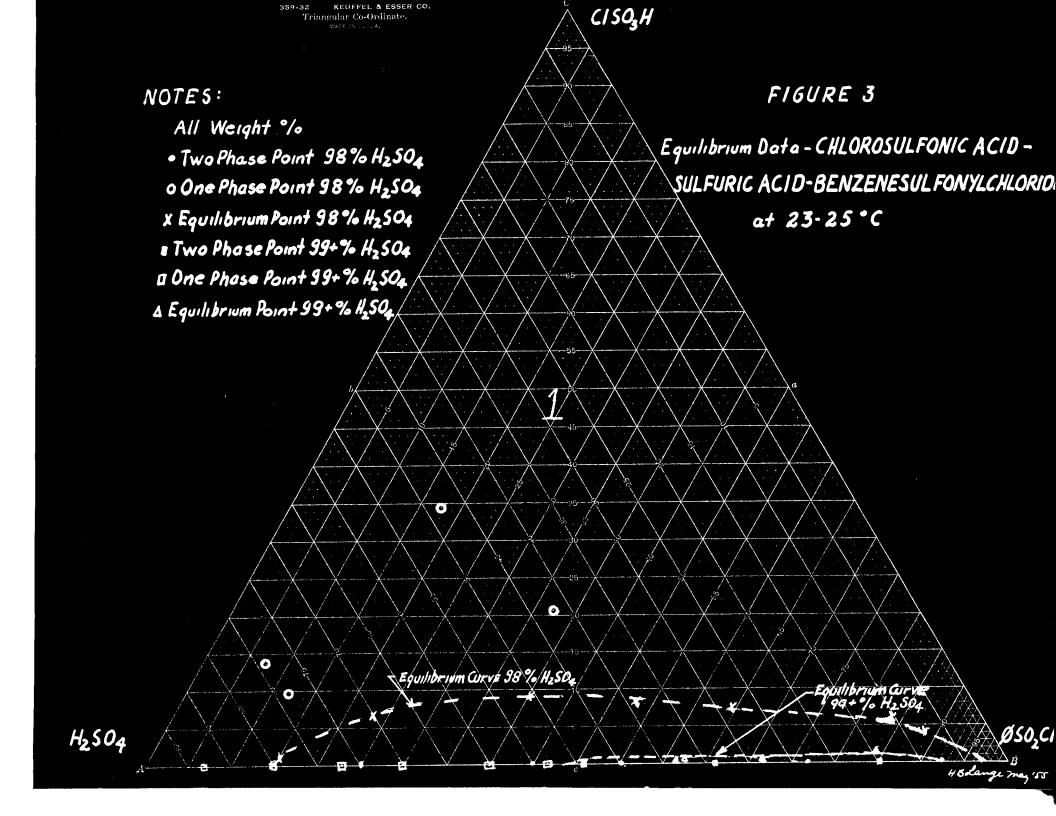


TABLE VI

# Chlorosulfonic Acid - Carbon Tetrachloride - Benzenesulfonyl Chloride

Run	63
	-

Total Weight	Actual Weight	Weight %	
92.9590 gms 90.3600 gms 87.9073 gms	2.5990 gms 2.4527 gms 4.0313 gms	28.60 27.00 14.140	ciso <sub>3</sub> H øso <sub>2</sub> ci cci <sub>l</sub>
83.8760 tare	9.0830 total		

Produced a 1 phase system.

# <u>Run 64</u>

Added more CCl4 to run 63.

Total Weight	Actual Weight	Weight %	
97.7391 gms	2.5990 gms 2.4527 gms 8.8115 gms	18.78 17.71 63.51	C1S03H ØS02C1 CC1/4
	13.8632 total		

Produced a 1 phase system.

# Run 65

Added more CCll4 to run 64.

Total Weight	Actual Weight	Weight %	
100.1202 gms	2.5990 gms 2.4527 gms 11.1926 gms	16.01 15.04 68.85	C1S03H ØS02C1 CC14
	16.2443 total		

Run 66
Added more ClS03II to run 65.

Total Weight	Actual Weight	Weight %	
110.6624 gms	13.1412 gms 2.4527 gms 11.1926 gms	49.15 9.15 41.70	ccilt ccilt
	26.7865 total		

Produced a 1 phase system.

# Run 67

Total Weight	Actual Weight	Weight %	
103.5012 gms 96.9613 gms 87.9574 gms	6.5399 gms 9.0039 gms 31.8614 gms	13.80 19.00 67.20	c180311 &205c1 cc1 <sup>1</sup>
56.0960 tare	47.4052 total		

Produced a 1 phase system.

## Run 68

Total Weight	Actual Weight	Weight 5	
110.2308 gms 103.2522 gms 90.2245 gms	6.9786 gms 13.0277 gms 36.8734 gms	12.27 22.89 64.83	øso <sub>2</sub> ci ciso <sub>3</sub> h cci <sub>li</sub>
53.3461 tare	56.8847 total		

Run 69
Added more Cls03H to run 68.

Total Weight	Actual Weight	Weight %	
122.4862 gms	6.9786 gms 25.2831 gms 36.8784 gms	10.10 36.60 53.30	øso <sub>2</sub> ci ciso <sub>3</sub> ii cci <sub>l;</sub>
	69.1401 total		

Produced a 1 phase system.

#### Run 70

Total Weight	Actual Weight	Weight %	
96.0167 gms 95.5409 gms 87.9127 gms	0.4758 gms 7.6282 gms 29.38 <b>7</b> 5 gms	1.27 20.35 78.47	CC1/1 CC1/1 CC1/1
58.5252 tare	37.4915 total		

Produced a 2 phase system.

# Run 71

Added ØS02Cl to run 70.

Total Weight	Actual Weight	Weight %	
96.6667 gms	1.1258 gms 7.6282 gms 29.3875 gms	2.95 20.00 77.05	ocit clso3H ccit
	38.11:15 total		

Run 72

From 4 component data - run 82.

Actual Weight		Weight 5	
1.9253 gms 2.3150 gms 2.3130 gms	4	29.38 35.22 35.20	c1303H cc1 <sup>††</sup> &205c1
6.5533 total			

Produced a 1 phase system.

## Run 73

From 4 component data - run 83.

Actual Weight	Weight %	
1.4082 gms 3.7028 gms 1.6804 gms	20.70 54.40 24.70	olso311 colf øso5ci
6.7914 total		

Produced a 1 phase system.

# Run 74

From 4 component data - run 84.

Actual Weight	Weight %	
1.0035 gms 5.1245 gms 0.8454 gms	14.40 73.50 12.10	01803H 001/1 080201
6.9734 total		

Run 75

From 4 component data - run 85.

Actual Weight	Weight %	
2.5175 gms 2.6262 gms 3.2439 gms	30.00 31.35 38.65	ст so зн сст <sub>и</sub> øso 2 ст
8.3876 total		

Produced a 1 phase system.

## Run 76

Total Weight	Actual Weight	Weight %	
99.2360 gms 98.9375 gms 96.7075 gms	0.2985 gms 2.2300 gms 12.7635 gms	2.05 14.58 83.47	ØS02C1 C1S03H CC1/4
83.9440 tare	15.2920 total		

Produced a 2 phase system.

# <u>Run 77</u>

Added more \$50201 to run 76.

Total Weight	Actual Weight	Weight %	
99.4133 gms	0.4758 gms 2.2300 gms 12.7635 gms	3.08 14.42 82.51	gsozci ciso3h cci <sup>†</sup>
	15.4693 total		

Equilibrium point - appearance of 1 phase.

Run 78

Added more CCl1 to run 77.

Total Weight	Actual Weight	Weight %	
102.3615 gms	0.4758 gms 2.2300 gms 15.6123 gms	2.60 12.17 85.23	øso <sub>2</sub> ci cci <sub>4</sub>
	18.3181 total		

Produced 2 phases near equilibrium point.

# Run 79

Added ØS02Cl to run 78.

Total Weight	Actual Weight	Weight 5	
102.3900 gms	0.4863 gms 2.2300 gms 15.6123 gms	2.65 12.17 85.28	øso <sub>2</sub> ci ciso <sub>3</sub> h cci <sub>l</sub>
	18,3286 total		

Equilibrium point - appearance of 1 phase.

# Run 80

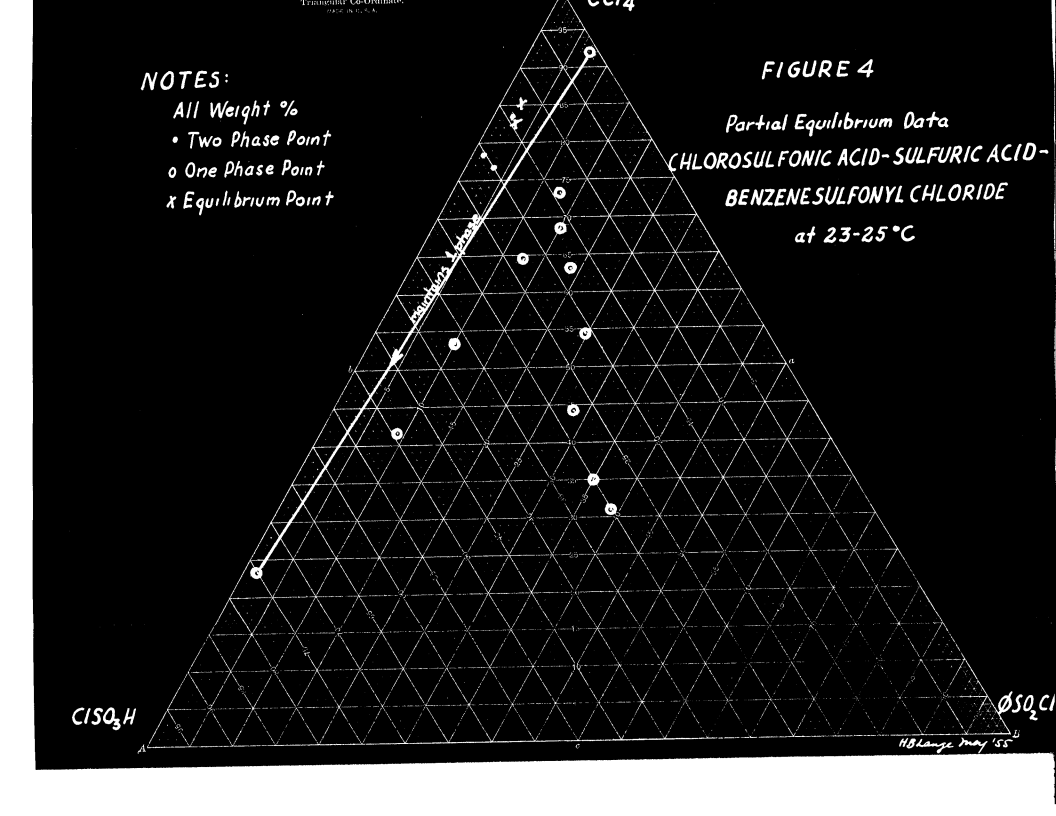
Total Weight	Actual Weight	Weight %	
94.7819 gms 94.6907 gms 93.9402 gms	0.0912 gms 0.7505 gms 10.0202 gms	0.84 6.91 92.25	C1S03H ØS02C1 CC1/4
83.9200 tare	10.8619 total		

<u>Run 81</u>

Added ClS03H to run 80.

Total Weight	Actual Weight	Weight %	
126.6202 gms	31.9295 gms 0.7505 gms 10.0202 gms	74.78 1.75 23.47	C1303H CC17 CC17
	42.7002 total		

Remained a 1 phase system throughout addition.



#### TABLE VII

# Carbon Tetrachloride - Benzenesulfonyl Chloride Sulfuric Acid - Chlorosulfonic Acid

Run 82

Total Weight	Actual Weight	Weight %	
97.5660 gms 94.6133 gms 92.6880 gms 90.3730 gms	2.9533 gms 1.9253 gms 2.3150 gms 2.3130 gms	31.00 20.22 24.38 24.30	820501 621 <sup>17</sup> 820 <sup>2</sup> H
88.0600 tare	9.5066 total	•	

Bottom three components gave a 1 phase system. All four gave a 2 phase system. Total volume 6 c.c., bottom layer 4 c.c.

Run 83

Total Weight	Actual Weight	Weight 5	
98.5120 gms 94.8494 gms 93.4412 gms 89.7384 gms	3.6626 gms 1.4082 gms 3.7028 gms 1.6804 gms	35.04 13.47 35.20 16.07	и <sub>2</sub> so <sub>l</sub> ciso3н cci <sub>l</sub> øso2ci
88.0580 tare	10.4540 total		

Bottom three components gave a 1 phase system. Four components gave a 2 phase system. Total volume 6.7 c.c., bottom 3.5 c.c.

Run 84

Total Weight	Actual Weight	Weight %	
96.9456 gms 95.0280 gms 94.0245 gms 88.9000 gms	1.9176 gms 1.0035 gms 5.1245 gms 0.8454 gms	21.57 11.28 57.64 9.50	H2SO4 CISO3H CC14 ØSO2C1
08.0546 tare	8.8910 total		

Bottom three components produced a 1 phase system.

All four gave a 2 phase system.

#### Extracted Layers

## Assay of Solvent Layer

Cold titration required 6.8 c.c. of 1 N NaOH.

Added 10 c.c. of 1 N NaOH and heated to boiling for half hour. Back titration - 3.4 c.c. of 0.987 N H<sub>2</sub>SO<sub>4</sub>. Chloride assay required 49.0 c.c. of standard 3.59 mg Cl/ml solution.

### Equilibrium Phases

Solvent Layer	Actual Weight	Weight %	
	0.095 gms 0.187 gms 0.593 gms 4.622 gms	1.72 3.40 10.78 84.00	H <sub>2</sub> SO <sub>]</sub> c1SO <sub>3</sub> H øSO <sub>2</sub> C1 cC1 <sub>4</sub>
	5.497 total		

## Run 84 (continued)

Acid Layer	Actual Weight	Weight %	
	1.8226 gms 0.8165 gms 0.2524 gms 0.5025 gms	53.75 24.00 7.14 14.81	H <sub>2</sub> SO <sub>4</sub> CISO <sub>3</sub> H ØSO <sub>2</sub> CI CCI <sub>[4</sub>
	3.3940 total		

## Run 85

Total Weight	Actual Weight	Weight %	
92.4096 gms 91.3932 gms 88.8757 gms 86.2495 gms	1.0164 gms 2.5175 gms 2.6262 gms 3.2439 gms	10.80 26.77 27.93 34.49	H <sub>2</sub> SO] <sub>1</sub> СС1 <sub>]1</sub> ФSO <sub>2</sub> С1
83.0056 tare	9.3040 total		

Produced a 1 phase system throughout.

# Run 86

Added more H2SO14 to run 85.

Total Weight	Actual Weight	Weight %	
93.3138 gms	1.9206 gms 2.5175 gms 2.6262 gms 3.2439 gms	18.63 24.42 25.48 31.47	H <sub>2</sub> SO] <sub>1</sub> C1SO3H CC1] <sub>1</sub> ØSO2C1
	10.3082 total		

Run 87

Added more H2SOlt to run 86.

Total Weight	Actual Weight	Weight %	
94.4015 gms	3.0083 gms 2.5175 gms 2.6262 gms 3.2439 gms	26.39 22.10 23.04 28.46	H <sub>2</sub> SO <sub>1</sub> C1SO <sub>3</sub> H CC1] <sub>1</sub> ØSO2C1
	11.3959 total		

Produced 2 phases. Total volume 7.4 c.c., bottom 4.75 c.c.

#### Extracted Layers

Acid Layer	G - 61.2240 gms T - 53.7705 gms N - 7.4535 gms
Solvent Layer	G - 57.7115 gms T - 53.9564 gms N - 3.7551 gms

## Assay of Solvent Layer

Cold titration required 25.8 c.c. of 1 N NaOH. Added 30 c.c. of 1 N NaOH. Heated to boiling for twenty minutes. Back titration - 16.4 c.c. of 0.987 N H<sub>2</sub>SO<sub>4</sub>. Chloride assay 116.95 c.c. of standard 3.59 mg Cl/ml solution.

## Equilibrium Phases

Solvent Layer	Actual Weight	Weight %	
	0.543 gms 0.574 gms 1.428 gms 1.210 gms	14.47 15.28 38.02 32.22	H <sub>2</sub> SO CISO3H CC1 <sub>1</sub> ØSO2C1
	3.755 total		

## Run 87 (continued)

Acid Layer	Actual Weight	Weight %	
	2.4653 gms 1.9435 gms 1.1981 gms 2.0339 gms	32,27 25,42 15,70 26,60	H2SO4 C1SO3H CC14 ØSO2C1
	7.6/108 total		

## Run 88

Total Weight	Actual Weight	Weight %	
89.8855 gms 89.44995 gms 86.6223 gms 84.5450 gms	0.3860 gms 2.0773 gms 2.8772 gms 1.5520 gms	5.60 41.74 30.14 22.52	H <sub>2</sub> SO <sub> </sub> C1SO <sub>3</sub> H CC1, ØSO <sub>2</sub> C1
82.9930 tare	6.8925 total		

Produced a 1 phase system.

# Run 89

Added more H2SO14 to run 88.

Total Weight	Actual Weight	Weight %	
90.215l <sub>+</sub> gms	0.7159 gms 2.0773 gms 2.8772 gms 1.5520 gms	9.91 39.84 28.76 21.49	&205c1 C1203H C2201 M2201
	7.2224 total		

Produced a 1 phase system.

## Run 90

Added more H2SOL to run 89.

Total Weight	Actual Weight	Weight %	
90.6762 gms	1.1767 gms 2.0773 gms 2.8772 gms 1.5520 gms	15.31 37.45 27.04 20.20	H2SOL C1SO3H CC1L ØSO2C1
	7.6832 total		

Run 91

Added more H2SO14 to run 89.

Total Weight	Actual Weight	Weight %	
91.4300 gms	1.9305 gms 2.0773 gms 2.8772 gms 1.5520 gms	22.88 34.10 24.62 18.40	H2SO  C1SO3H CC1  ØSO2C1
	8.4370 total		

Produced a 2 phase system.

#### Extracted Layers

Acid Layer	G - 62.8047 gms T - 55.5536 gms N - 7.2511 gms
Solvent Layer	G - 55.1210 gms T - 54.1530 gms N - 0.9680 gms

### Assay of Solvent Layer

Cold titration required 4.4 c.c. of 1 N NaOH. Added 15 c.c. 1 N NaOH. Heated to boiling for half hour. Back titration - 12.6 c.c. of 0.987 N H<sub>2</sub>SO<sub>4</sub>. Chloride assay - 21.4 ml of 3.59 mg Cl/ml solution.

## Equilibrium Phases

Solvent Layer	Actual Weight	Weight %	
	0.573 gms 0.066 gms 0.226 gms 0.103 gms	59.19 6.82 23.35 10.64	CC1) <sub>1</sub> ØSO2C1 C1SO3H

# Run 91 (continued)

Acid Layer	Actual Weight	Weight %	
	1.5043 gms	20.14	ccij
	1.8645 gms	24.96	H280]
	1.3260 gms	17.75	Ø802C1
	2.7742 gms	37.14	C1803H

# Run 92

Total Weig	ht	Actual V	<u>Weight</u>	Weight %	
104.1853 g 95.0570 g 91.9142 g 81.5078 g	m <b>s</b>	9.1283 3.1428 10.14064 24.8480	gms gms	19.20 6.60 21.85 52.25	CC1/4 CC1/4 CC1/4 CC1/4
56.6598 t	are	47.5255	total		

#### DISCUSSION

In the initial phases of this work, two batch chlorosulfonations of benzene employing carbon tetrachloride as an inert solvent were made. The method of Gilman and Blatt (7) was employed. In the first run, the weight ratio of carbon tetrachloride to chlorosulfonic acid was 63%-37% respectively. The phase separation was rapid at the conclusion of the reaction period but the carbon tetrachloride layer was the same as the volume of solvent charged. It was suspected that, (a) benzenesulfonyl chloride was not in the organic layer, (b) the carbon tetrachloride had a high solubility in the acids or (c) a combination of both (a) and (b) ex-The second run was performed using a ratio of 54% chlorosulfonic acid to 46% carbon tetrachloride. this case, phase separation was extremely slow and the carbon tetrachloride layer was again the same volume as that charged. These disturbing conditions prompted investigation into solvent systems and into an understanding of phase relationships. Some knowledge of liquid-liquid equilibria was necessary prior to a study of continuous operation.

#### A. Solvent Systems

A preliminary screening of possible solvents was undertaken. They were selected on the basis of density, relative inertness, availability and cost. However, the relative immiscibility with chlorosulfonic acid was the deciding factor in adoption of the solvent. Table III lists the solvent pairs in the ratios mixed and observed. Only carbon tetrachloride and cyclohexane showed some immiscibility. The commercial grade of cyclohexane which contains from 1% - 5% benzene was not acceptable because of the slow reaction with chlorosulfonic acid. Pure cyclohexane, spectrographic grade, behaved as the commercial grade but without the reaction. Due to the cost of this material, it was not considered further at this time. Carbon tetrachloride was adopted for further study.

It is of interest to note that the list of possible solvents is much larger than shown here. Selection of a more favorable solvent, from the liquid equilibria standpoint, may simplify any process. For a study in liquid-liquid equilibria other solvents are worth investigation. Trichloroethylene, perchloroethylene and fluorohydrocarbons are worthy of investigation.

# B. <u>Chlorosulfonic Acid - Sulfuric Acid - Carbon Tetra-</u> chloride System

The equilibrium relationships between chlorosulfonic acid, sulfuric acid and carbon tetrachloride are illustrated in Figures 1 and 2. The experimental runs are given in Table IV.

The data show that the ternary is binodal. One component is partially miscible in the other two. That is, carbon tetrachloride is partially miscible in sulfuric acid and partially in chlorosulfonic acid.

The small one phase region at high carbon tetrachloride concentrations was difficult to obtain with
high accuracy. It is known to exist because of repeated
experimental data showing some sulfuric acid and chlorosulfonic present in carbon tetrachloride. Practically,
this region would be of little concern. Recovery of the
acids from carbon tetrachloride would probably be uneconomical.

The large one phase region at high chlorosulfonic acid concentrations is very interesting. The shape of the equilibrium line between the one phase and two phase region is different from that usually observed for binodal systems. Of particular interest is the region between

65 - 70 weight per cent chlorosulfonic acid and 10 - 25 weight per cent sulfuric acid. Small changes in sulfuric acid concentration have a large influence on the concentration of carbon tetrachloride necessary to maintain the one phase region. Such a condition might be suspected since sulfuric acid and carbon tetrachloride are practically mutually insoluble. The experimental data in this region were carefully determined and checked. Three approaches were employed. Mixtures of carbon tetrachloridechlorosulfonic acid were prepared and titrated slowly with sulfuric acid till the first appearance of the 2 phases. In Figure 1. it can be seen how well these points line up with the starting point and 100% sulfuric acid. second approach was to start with the one phase mixtures of the acids and titrate with carbon tetrachloride. These points lie on the line contacting the starting point and 100% carbon tetrachloride. The final method was to use the tie line technique and analyze each phase of a two phase mixture.

These points fitted the expected curve. The fact that the experimental points plotted so well also indicates the reliability of the data under these experimental conditions. The small part of the one phase region lying along the 1% carbon tetrachloride line is to be expected from the rest of the curve. It is known that 2 phase

points in the high sulfuric acid area must have an equilibrium point to match with that in the high carbon tetrachloride region. The equilibrium point at 53% chlorosulfonic acid, 45% sulfuric acid and 1% carbon tetrachloride indicates that the region above it may be very small. The data show this very small region. It is interesting to observe that runs 34 and 35 in Table IV show that no increase in the volume of the carbon tetrachloride was observed (in the high sulfuric acid area) as the sulfuric acid concentration was increased. This is what would happen with the equilibrium line as shown in Figure 2.

During the course of the experimental work, it was found that some of the sulfuric acid used was less than 99% acid. Fortunately no difference was observed in equilibrium points as a result of 98 or 99% sulfuric acid, (runs 25 - 38, Table IV). This may be due to the fact that carbon tetrachloride is equally insoluble in water or sulfuric acid-water mixtures.

# C. Chlorosulfonic Acid - Sulfuric Acid - Benzenesulfonyl Chloride System

Table V and Figure 3 summarize the data obtained with this system. The first experiments (runs 39 - 49) were made with 98-98.5% sulfuric acid. This was not intentional and it was not discovered until the data shown in Figure 3 was obtained. It can be seen that with this small amount of water present a 2 phase region occurs. Changing to acid of 99+% assay, the 2 phase region almost disappeared (Figure 3). Actually it was found that the 2 phase region occurred at high concentrations of benzenesulfonyl chloride and that small amounts of chlorosulfonic acid (0.5-2.0% by weight) were sufficient to produce a 1 phase system.

The only apparent reason for the difference in the equilibrium curves is in the acid strength. This may be explained by the fact that benzenesulfonyl chloride is insoluble in the sulfuric acid-water-hydrate and enough was present to upset the normal equilibria or to act as a fourth component. When sulfuric acid and the benzenesulfonyl chloride are mixed prior to the addition of chlorosulfonic acid the presence of water tended toward the immiscibility condition. However, if chlorosulfonic

acid was added before the benzenesulfonyl chloride, it would react with the water and form sulfuric and hydrochloric acids. The data with 99+% acid tends toward a completely miscible three component system.

# D. <u>Chlorosulfonic Acid - Carbon Tetrachloride - Benzene-</u> sulfonyl Chloride System

Table VI and Figure 4 contain the data for this ternary. Although the study is not extensive, some conclusions may be drawn.

It is known that benzenesulfonyl chloride is miscible in carbon tetrachloride and in chlorosulfonic acid. However, it has been shown that carbon tetrachloride and chlorosulfonic acid form a partially miscible pair.

Hence, it is anticipated that this ternary would contain a 2 phase region. This is demonstrated (in part) by the data. It is also conceivable that this two phase region would lie near the high carbon tetrachloride concentrations. This too is shown (in part) by the data. The 2 phase region cannot cover much of the area, high carbon tetrachloride - low chlorosulfonic acid. Runs 80 and 81, Table VI, illustrate that a one phase region is maintained, going from 1 weight per cent acid to 75 weight per cent along the line to 100% chlorosulfonic acid (Figure 4).

# E. <u>Carbon Tetrachloride - Benzenesulfonyl Chloride -</u> Sulfuric Acid - Chlorosulfonic Acid System

Analysis of a four component system requires extensive data. The data assembled here can serve only as a first approximation of the entire system.

The first factor that appears important is the sulfuric acid concentration. In all experiments where a 2
phase region was obtained, the sulfuric acid concentration
was over 20 weight per cent. The concentration was also
greater than that of chlorosulfonic acid (except run 91,
Table VII). Run 91, which had a higher chlorosulfonic
acid concentration than sulfuric acid, gave a phase separation but an unfavorable distribution of sulfonyl chloride.
The solvent layer retained only 15 weight per cent of the
sulfonyl chloride charged.

The second factor is the solvent concentration. No direction is apparent from the limited number of runs reported here. However, a high recovery of benzenesulfonyl chloride in the organic layer was obtained in run 84, Table VII. Here the organic solvent retained 70% weight of the initial sulfonyl chloride. The initial mixture contained 58 weight per cent solvent. Conditions of this experiment gave a more favorable distribution of

sulfonyl chloride than run 87, Table VII. Under conditions of the latter only a 37 weight per cent recovery of the chloride compound was in the organic stream. The evidence from these experiments is, that an interaction exists between sulfuric acid and carbon tetrachloride concentration for maximum chloride recovery. It is conceivable that conditions such as 50-60% carbon tetrachloride, 15-20% sulfuric acid, 10-15% chlorosulfonic acid and 5-10% sulfonyl chloride would give a favorable distribution of all materials.

#### CONCLUSIONS

In an effort to develop a process for the continuous chlorosulfonation of benzene, liquid-liquid equilibria data for a proposed system were studied. Ternary data for chlorosulfonic acid - sulfuric acid - carbon tetrachloride were developed at 23-25°C. The system is composed of two single phase regions and a two phase region. ternary chlorosulfonic acid - sulfuric acid - benzenesulfonyl chloride is mostly a single phase system. very small two phase region was found, the size of which is determined by the purity of sulfuric acid. The ternary. chlorosulfonic acid - carbon tetrachloride - benzenesulfonyl chloride has a small two phase region. However, this system was not completely developed. The quaternary. carbon tetrachloride - benzenesulfonyl chloride - sulfuric acid - chlorosulfonic acid shows that high concentrations of sulfuric acid are necessary for a favorable distribution of benzenesulfonyl chloride in the solvent layer.

The work herein can serve as a basis for the study of continuous chlorosulfonation of benzene.

### RECOMMENDATIONS

It is recommended that further work be carried out to complete the quaternary system study begun here and to develop the continuous chlorosulfonation process.

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