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# Spectrophotometric study of the Boron- Curcumin system

Frank Xavier Pollio

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Spectrophotometric Study Of The Boron-Curcumin System

By

Frank Xavier Pollio

A Thesis

Submitted To The Graduate Faculty

Of

The University of Richmond

In Candidacy

For The Degree of

Master Of Science In Chemistry

August 1958

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Approved:<br>W. Allan Powell

### **ACKNOWLEDGEMENT**

I am sincerely gratetul to Dr. W. Allan Powell tor his personal assistance and guidance in supervising this research program. His many valuable suggestions and observations have been or upmost value.





#### **INTRODUCTION**

Prior to about a half century ago. existing methods for quanitative detection *ot* boron were sensitive enough only for the macrorange or boron concentration. With the introduction of the curcumin colorimetric method at the turn of the century, a potential tool tor miororange determination *ot* boron was in the hands *ot* the analytical chemist.

Throughout the last fifty years or so after the introduction *ot* the method 1 much work has been done in modifying it for various different situations, but very little study has actually been done on better understanding the mechanism of the method itself.

With mechanism and structure in mind, the aim of this research project was to investigate the role played by the various reagents used in the curcumin spectrophotometric method. The reagents were used in varied concentrations and at times some eliminated entirely, in the hope that a greater understanding be gained in the overall mechanism followed by the reaction.

One of the major objectives was to try to determine the structure *ot* the colored substance formed in the reaction. A step towards this goal was gained by obtaining molar ratios of the entities suspected to be an intricate part of the colored substance.

Host of the studies made were studies of absorption using the visible range of the spectrum (550 millimiorons) where the colored substance was found to absorb at its maximum. A photoelectric colorimeter could have been used instead or a spectrophotometer, however the resolution and precision would not have been as great.

A potentiometric titration study was also carried out so as to gain greater insight into the preliminary reaction occurring between boric aoid and oxalic acid prior to the addition *ot* curaumin in the final color forming step of the method.

### HISTORICAL

The practical importance of boron in relation to agriculture is one of the recent findings of science. The reason why this fact remained obscure to science and mankind was in part due to the lack of detection or the minute quantities or boron to be tound in the soils and plants of the universe. Up until the turn of the century, methods existing for quantitative detection *of* boron were sensitive enough only for the macrorange of boron concentration. Since the concentration of boron in plans and soils are confined to a narrow range and in general, the concentration of boron in the soil should not exceed a few parts per million, a mioromethod for the detemination of boron was desirable.

It will suffice to mention at this point that the early methods for quantitative analysis of boron were chiefly gravimetric or volumetric, the latter being the more reliable am accurate or the two. Gravimetric methods were not reliable because a sparingly soluble stoichiometric compound of boron which could serve as a basis for gravimetric detection was not available. (11) All volumetric methods in general follow a similar pattern in which the pH of a boron solution is adjusted with subsequent addition *ot* a polyol and titration with a standard strong base (4).

With a need for a method for microdetermination of boron, the colorimetric methods were investigated. or the colorimetric methods the two most widely employed and investigated were those in which quinalizarin and ourcumin served as the color fonning reagent.

 $-3-$ 

The quinallzarin method was found to be the more rapid and less subject to interference by other trace elements, While the curcumin method the more sensitive of the two. In the quinalizarin method  $(17)$ quinalizarin reacts with boric acid in concentrated sulfuric acid giving a colored substance mich is used as the basis for analysis.

Sinoe the work which will be presented in this research project is mainly concerned with the curcumin method, a brief historical review *ot* the origin of' the use of ourcumin as a boron detection tool and the subsequent important modifications and applications *of* the method shall be presented.

Curoumin, 1, 7-bis(4-hydroxy-3-methoxyphenyl)-1, 6-heptadiene, has as empirical formula  $C_2H_{20}C_6$ . It is an unsaturated hydroxy-ketone and is the coloring principle *of* ourcuma (turmeric) i.e. the dye from the rhizones of curcuma tincturia, which fluorishes in the East. Commercially ourcumin is extracted from this dye by a series *ot* organic extractions. (7) Curcumin still finds a limited use as a dye for silk and is also used as an indicator (gives a brown color with caustic alkali and red coloration with boric acid).

By means of infrared studies, Spicer and Strickland (24) confirmed the enol structure of curcumin given by Ghosh  $(9)$ 



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The accepted structure of curcumin (keto form) given by Yoe and Sarver (28) and most of the recent investigators is given asr



Curcumin in the keto form resembles acetyl acetone of which it most certainly appears to be a derivative.

Ourcumin was first used in a colorimetric method for boron determination at the beginning of the century. In 1902, Hebebrand (11) devised a colorimetric method depending upon the formation of a red color upon the addition or turmeric to a water-alcohol solution containing boron and hydrochloric acid. In the same year Cassal and Gerrans  $(l_1)$  revised the method somewhat introducing oxalic acid which was found to increase the sensitivity of the method considerably.

Another modifier of Hebebrand's method at about the year 1915 was Filippi  $(8)$ . He modified the method so as to eliminate sources of error due mainly to the varying composition of the curcumin used in the tests. Filippi modified the method as follows: to better establish the different intensity or colors he took up the residue containing curcumin and boric acid with 2 ml. of 90% ethyl alcohol pipetting it into very narrow test tubes of the same capacity and then observed the color which he claimed persisted unmodified for 12 hours.

The turmeric or curcumin method actually became popular when the method was revised further by Bertrand and Agulhon (2). Their method was essentially a paper method in which the concentration of boron present in

a sample was proportional to the length which the colored compound formed between boron and curcumin in a strip of turneric paper would travel by capillary action along the length of the paper.

Halphen (10) came out with his method a short while after Bertrand and Agulhon introduced their paper mothod in 1913. His method consisted in collecting boric acid as methyl borate in 0.2 ml. of 1N sodium hydroxide solution. The solution was then evaporated to dryness in a small test tube and the residue dissolved in 1 ml. of water plus 2 ml. of hydrochloric acid. Standards containing known amounts of boric acid were similarly treated. To each sample was then added 1 ml. of an ethyl acetate solution of turneric and the resulting red colors were compared after 50 minutes.

Numerous other investigators studied the Bertrand and Agulhon method extensively in the years to follow adding slight modifications of their own but the method remained essentially the same.

It was not until 1939 that Naftel (16) came out with modifications of the Cassal and Gerrans method  $(l_1)$ . This modification is probably the most popular curcumin procedure used to this date. This is the method with some modification which is followed in this research project. Essentially it involves making the boron sample basic with calcium hydroxide followed by evaporation of the sample to dryness. An alcoholic solution of oxalic acid and curcumin along with hydrochloric acid is then added to the cooled sample and evaporated further to dryness on a water bath. The residue is then extracted with ethyl alcohol and compared with known standards.

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Naftel's method has been used quite extensively and has received some modifications from later investigators. Silverman and Trego (22) dissolved the curcumin dye in oarbitol•aoetone, which solution is stable for months, and used phenolphalein to control the acidity. They used sodium carbonate instead of calcium hydroxide 1n the preliminary' drying step.

Winsor ( 27) extracted boron trom fresh plant tissue and analyzed the boron present in microquantities by another modification of Naftel's method.

Hegedus (12), after a critical review of various methods baaed on the formation of rosocyanin from yellow curoumin under the influence of free boric acid, presented his modified method. It consisted of adding 2 ml. curcumin reagent (prepared by dissolving  $0.1$  g. curcumin in 100 ml. glacial acetic acid saturated with  $\alpha$ xalic acid.) The solution was then evaporated to dryness on a water bath and the residue dissolved in 20 ml. *ot* ethyl alcohol.. The araount of boron present was then determined from the color intensity after illumination for  $3$  minutes. He found that  $\mathbf{F}_1$ , Ti, Zr, Mo, Be, W, in large amounts interfered with the test. Oxidising agents such as nitrate, chlorate, bromate, iodate, or nitrite should be reduced it present, to prevent the oxidation *ot* ourcumin. The method ia most suitable for establishing the boron content in mineral waters of low boron content. The method is sensitive for a range of boron from 0.5 to 8 micrograms.

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Bottini (3) used still another modification of the ourcumin method using a basic instead of acidic media. He acidified a boron solu• tion with acetic acid and added 2 ml. of a 0.1% curcumin solution. After evaporation on a steam bath he separated the excess curcumin by washing with ethyl ether until the washings were colorless. The residue was then treated with methyl acetate (100 ml. methyl acetate  $\div$  1 ml. 0.25N potassium hydroxide). The solution obtained was then compared with standards. The method gave good results in the microrange with a sensitivity of 1 micro• gram.

Rynascewicz, Sleeper and Ryan (20) extracted boron from sodium chloride with ethyl alcohol and estimated the boron present colorimetrioally using curcumin. Although  $0.5$  micrograms of boron can be measured by their method, an increment of only 1 microgram of boron can be detected in concentrated sodium hydroxide by their method. Their procedure is also a moditied version of the Naftel method but is much longer and drawn out.

In 1954 Troug and Berger (26) presented a simplified curcumin procedure of that used by Naftel for plant and soil analysis which r equired only one evaporation and filtration.

Martin and Maes (14) replaced the omoentrated acid used by Naftel with a ltl mixture of glacial acetic acid and acetic anhydride.

With the realization of the powerful deoxidizing ability of elemental boron, its strong Affinity for many gases, potential as a fuel for special appl cations, excellent insulating possibilities, neutron absorbing ability and that it could possibly replace weightly lead and concrete as protection

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around nuclear reactors, boron research intensified and numerous methods of determining boron appear in recent literature, many for boron and curgumin. compounds. Among these some may be mentioned at this point dealing with modifications of the curcumin method applied to particular cases.

Staple, Marshall, Nelson, Simon (25) developed a method for determining boron in  $TF_e-UF_e$ . Their method consists of evaporation, dissolution of the boron containing substance in an alcohol-oxalic acid• hydrochloric acid solution and then addition of curcumin with subsequent spectrophotometric analysis.

De Abinati am Paeques (17) emphasize that it is critically important that the drying time be constant in the colorimetric analysis or boron in uranium compounds and in graphite by means of the red color formed by the action of borio acid on curcumin in the presence *ot* oxalic acid. They state that color intensity is higher with aborter drying periods.

Silverman and Trego (22) determined boron colorimetrioally (cureumin•oxalio aoid method) in uranyl chloride solution without prior re• moval of uranium. They claimed that the interference ascribed to the presen or uranium lies in the tact that uranium preferentially combines with the available oxalic acid. There.fore if a suitable excess *ot* oxalic aoid is added, the uranium forms a soluble uranium-oxalate complex which need not be removed.

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Rynasiewicz and Consalvo (19) describe a procedure for the analysis *ot* boron in 7% U-Zr alloys containing approximately 0 .015% boron. They fused tine metal chips trom the alloy with sodium carbonate. The melt was dissolved in water and then made slightly alkaline, filtered and evaporated to dryness. The boron was then leached f'rom a slurry of the salt using 95% ethyl alcohol and then determined colorimetrically by the curcumin method.

Hot much has been done on a study *ot* the ourcumin method itself, that is in determining the optimum conditions necessary *tor* maximum sensitivity and the effects of varying the conditions of the reaction. Also very little is known about the actual mechanism that the reaction follows. (This is the major aim of the work presented in this thesis.)

Among the first investigators of the curcumin method were Clarke and Jackson (6) back in 1908. They made a study of roaooyanin, the red substance formed by the action *ot* boric acid on turmeric paper (ouraumin). They claimed that rosocyanin and curcumin were merely isomers  $(C_{14}H_{14}O_4)$ , only that one of the two hydroxyls of curcumin was modified, probably to an oxide tonn in rosocyanin. They found that aqueous alkali in the cold converted the curoumin into roaooyanin. They found that white wool was dyed blue by the rosocyanin reaction and alkali. They heated ourcumin with boric aoid and 811lturic acid in alcohol, and when this was poured into water rosocyanin was obtained. This was then purified by washing with ether and reprecipitated from alcohol by dilute sodium chloride solution. No boron was found in rosocyanin and analysis and molecular weight detennination in•

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dicated the formula to be  $C_{14}H_{14}O_4$ . It formed a purple powder with metallic yellowish-green reflex. They found further that ourcumin and boric acid in the cold poured into ether yielded an addition product *ot*  the two, decomposed by water forming rubrocuroumin  $C_{14}H_{14}O_4$ , a vermilion powder, unstable on standing at 240 degrees centigrade or upon heating w11h sulfuric acid, forming rosocyanin.

Silverman and Trego (23) made a study of the effect of oxalic aoid by varying its concentration and holding everything else constant. They found that 2 ml. of a 5% solution to be the best working concentration of oxalic acid for a study *ot* boron in the range *ot* 0 .o to SO micrograms *ot* boron. They also made a study or the hydrochloric acid concentration and found poor results were to be obtained it a great excess of acid were used. They used sodium carbonate to neutralize the acidity and give a solution or residual alkalinity. They claimed that it the solution were not alkaline then significant amounts of boric acid (probably metaborio acid) would be lost by volatilization in the primary evaporation step of the curcumin method. Instead of extracting the residue with  $25$  ml. ether alcohol they used acetone.

Troug and Berger (26) stated that curcumin dissolved in ethyl alcohol gave a solution that is strongly yellow in color while the product of' the curaumin-borate reaction, rosooyanin, ia *o!* a deep red color, giving the test solution a reddish hue. To show that curcumin and rosooyanin are spectrally separable, the absorption curve of each of the two pure substances dissolved in ethyl alcohol was determined using a Beckman DU spectrophotometer. They noticed that the absorption peak for msocyanin falls at approximately  $\mu$ 80 mu. Curcumin did not absorb light over the range of

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maximum absorption of rosocyanin thus eliminating an absorption effect due to excess of reagent in the determination of boron.

Troug and Berger (26) state that Hafford (Ph.D. Thesis, University of Wiseonsin 1942) suggests that the rosocyanin is probably formed by a loose combination of the borate with one of the hydroxyl groupa of the curaumin molecule.

Yoe and Sarver (28) point out that beryllium, aluminum, iron, zirconium and magnesium form colored lakes with curcumin. Kolthoff (13) showed that these lake formations take place in alkali solutions. The excess of oxalic acid present in the regular curoumin procedure tor boron prevents by solution the formation of these interfering lakes, according to Kolthoff.

Yoe and Sarver ( 28) postulate that curcumin probably acts in one of two ways with boron: (a) the hydrogens of the end hydroxyls may be replaced by an equivalent of metal, where upon a ring is closed by coordination with the oxygen of the o-methoxy group (b) it may act like an enolizable acyclic diketone since it also resembles aeetyl aoetone in structure.

Troug and Berger (26) point out that rosocyanin slowly hydrolyzes to ouroumin, hence all colorimetric readings should be made within two hours after solution or the colored residue in alcohol.

Spicer and Strickland  $(24)$  working on a macro scale did some work in elucidating the structure of the compounds formed in the reaction between: (a) boron and curcumin, {b) boron, curcumin and oxalic acid, discussing the probable mechanisms involved in the two reactions.

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Rosooyanin, the compound formed when curcumin and boric acid react in the presence of a mineral acid, has been shown by the two investigators to contain two curcumin molecules coordinated around a boron atom associated with one equivalent of an anion. The probable structural formula and electronic configuration of the molecule have been suggested by them on the basis of spectrophotometric and ion exchange studies. They also claim to  $h^{\text{ave}}_{\lambda}$  isolated the red compound formed when curcumin reacts with boric acid in the presence of oxalic acid and have called this compound rubrocuroumin. The substance is shown to be a compound of ourcumin, boron and oxalate in a lilil ratio. Using analytical results, absorption spectra etc. they suggested a structure similar to that of rosocyanin but containing the curcumin molecules joined by a boro-oxalate bridge. They claimed further that if both oxalic acid and a mineral acid are present then a mixture of rubrocurcumin and rosoeyanin results, although the yield of tha former greatly exceeds that or the latter when the two acids are present in equimolar quantities. Observed also was the effect of water in inhibiting the formation of both rubro• curoumin and rosocyanin but its effect was most marked on the yield of the former.

### EXPERIMENTAL

 $\mathcal{O}(\mathcal{O}_\mathcal{A})$  , where  $\mathcal{O}_\mathcal{A}$  is the set of the  $\mathcal{O}_\mathcal{A}$ 

### I. Apparatus

Beckman Model B Spectrophotometer - Abaorbanoe readings ware obtained at a wavelength of 550 millimicrons (in mostoases) and pyrex cells were used. Ethyl alcohol was used as the standard in absorbance measurements but at times a blank containing all of the experimental reagents minus boron was substituted.

Beckman Model H2 Glass Electrode pH Meter - All pH measurements and potentiometric titrations were carried out using this meter exclusively. A Beckman pH solution (pH  $= 7.0 \pm 0.1$ ) was used to standardize the instru-<br>ment prior to each series of measurements.

Water Bath – For all work requiring dryings at  $55^{\circ} \pm 3^{\circ}$ C. a water bath designed to maintain a constant temperature was used. This bath was designed so as to hold a dozen porcelain evaporation dishes and to keep water vapor avay from samples being evaporated.

Hot Plate - A three temperature range control hot plate was used for all preliminary dryings using the low heat position so as to minimize splattering.

Centrifuge  $\sim$  A motor driven centrifuge was used in separating any residue in the alcoholic extractions before spectrophotometric measure• ments.

Miscellaneous Apearatus - Soft glass was used in all cases in which glassware was required. Porcelain dishes were used for the reaction containers. Polyethylene bottles were used for storage of boron solutions. Volumetric flasks were used for preparation of the curcumin solutions and other reagents required. • A 50 ml. burette was used in conjunction with the pH meter in the potentiomotric titrations.

### II. Reagents

Reagents  $-$  Reagent grade chemicals were used when necessary or possible.

Standard Boron Solution  $-$  A stock solution was prepared containing 100 micrograms of boron per ml. b7 dissolving 0. 5716 gms. of boric acid in water and diluting to one liter. Dilute solutions of boron were prepared from this stock solution.

Calcium Hydroxide Solution - A .1N calcium hydroxide suspension was ' prepared by dissolving 1.853 g. of solute in water and diluting to *500* ml.

Curcumin Solutions - Solutions were prepared by dissolving 0.1 gram of Eastman curoumin in 1000ml. of either acetone or 95% ethyl alcohol depending on which modification was being studied. Solutions were prepared fresh before use.

Oxalic Acid Solution - An acetone or .95% ethyl alcohol solution containing .2 g. oxalic acid per ml. acetone was prepared by dissolving 20 g. *ot* di-hydrated oxalic acid in 100 ml. acetone or *95%* ethyl alcohol depending on the solvent being studied.

Hydrochloric Acid - A 1:1 solution of hydrochloric acid was prepared by mixing equal volumes of concentrated hydrochloric acid and water.

Ethyl Alcohol - The alcohol used throughout was 95%.

Acetone - Reagent grade.

Organic Acids - Eastman organic chemicals were used throughout when a study was made of various organic acids substituted in place of oxalic acid.

Alcoholic Sodium Hydroxide - A .1N solution was prepared by dissolving 2 g. of sodium hydroxide pellets in 95% ethyl alcohol and making it up to 500 ml. in a volumetric flask with the 95% alcohol.

Acid Potassium Phthalate - Primary Standard used to standardize the alcoholic sodium hydroxide. It was previously dried in an oven before use.

### III. Preliminary Investigation

Powell and coworkers  $(18)$ , did a great deal of work to elucidate the mechanism of the reaction involved between boron-curcumin and oxalic aoid. Evidence obtained by them indicates that the red color *of* the ourcumin method is that or a l·l-1 boron•curcumin-oxalio complex. They also found that oxalic acid sensitieea the method, mainly because oxalic aoid increases the rate of reaction and permits equilibrium to be reached before the reaction is frozen by losaot reaction medium. They pestulated a oomplexation between boron and oxalic prior to the main reaction with curcumin. Also studied by them was the ability of other materials, substituted in oxalic acid's place, to sensitize the reaction.

Because of the importance of adherence to Beer's Law and wavelength of maximum abaorbanoe it was decided to check these points as part or the first phase of this project.

The modified Naftel procedure used was as follows: to a boron sample (present as boric acid) in a small porcelain evaporation dish was added 1 ml. of a  $0.1N$  Ca(OH)<sub>2</sub> suspension. This mixture was then evaporated to dryness on a hot plate using the lowest heat possible so as to avoid splattering and consequent loss of some of the boron. To the cooled residue was then added 1 ml. acetone containing 0.2 g. oxalic acid, 0.2 ml. l•l HCl and finally 2 ml. *ot* a 0.1\$ curcumin solution ( .1 g. ourcumin/100 ml. acetone). The solution was then evaporated to dryness on a water bath maintained at *55•* ! 3•c. and heated at this temperature for an additional. 30 mirmtes after complete dryness was observed. The colored residue waa then

extracted with 25 ml. of 95% ethyl alcohol, the resulting solution centrifuged and the absorbance measured at a wavelength of *550* milli• microns using ethyl alcohol as reference solution to set the zero of the instrument.

### A. Determination of Wavelength or Maximum Abaorbance

To determine the wavelength of maximum absorbance a 2 microgram sample of boron was placed in a porcelain dish and run according to the procedure on page 17. Using a sensitivity of  $\mu$  and ethyl alcohol as the standard the absorbance was measured in the wavelength range of  $\frac{1}{7}$  to  $675$ millimicrons, taking readings at increments or 25 millimicrona. The re• sults are shown in the following table.

### TABLE I



.A. maximum absorbanoe was recorded at 550 millimicrons in agreement with previous investigators.

In subsequent work a wavelength of *550* millimicrons was used exclusively for abaorbanoe measurements.

B. Verification of Adherence to Beer's Law

Solutions containing 0.0, 0.5, 1.0, 1.5, 2.0, 2.5 micrograms or boron were treated according to the modified Naftel procedure discussed on page 17. Duplicate samples were used throughout.

### TABLE II

Adherence to Beer's Law

Reference solution - Ethyl Alcohol Sensitivity  $-1$ Wavelength • *550* millimicrons



### Results:

Boron

When micrograms boron vs. average absorbance are plotted, a straight line is obtained as shown in Figure 1. This linear relationship between absorbance and micrograms of boron in the range 0.0-2.5 micrograms of boron is proof that Beer's Law is followed in that low boron concentration range.

It should be pointed out here that the method described here gives very good reproducibility for a series of samples run at a particular time, but the absorbanoe values for solutions containing the same amount of boron have been found to vary when run at different times. Therefore for work where greater than average precision is required in quantitative work,



Figure I

a standard curve should be prepared at the time and under the same conditions as the unknown samples whose boron oontent is to be determined •

The cause of this variation in absorbance cannot be justly explained by one variable alone, but is caused by several variables. To mention a few the atmospheric conditions of the day can be one *ot* the variables influencing the rate ofevaporation. It has been found in much of the preliminary experimentation with the modified Naftel Method that shorter evaporation periods at *55•0.* gives slightly greater absorbanoe readings. Consequently a hot dry day should favor quicker drying periods and correspondingly slightly greater absorbance readings. Curcumin itself is another variable to be considered. If the curcumin reagent is not freshly prepared but used after several days, it will cause the absorbance to be decreased somewhat, probably because there is some decomposition occurring when the ourcumin is in solution.

It also appears that oxalic acid's activity decreases with the age of the solution.

IV. Determination *ot* Function of the Various Reagents

Because of the great advantage to be gained in the interpretation of a possible mechanism of the b oron-curcumin reaction and also of elucidating of the structure *ot* the colored substance formed, a study was made or the function of the various reagents used in the curoumin reaction.

By greater understanding the function and effects or varying or totally eliminating some or the reagents, a keener knowledge of the mode or reaction was hoped to be gained •

Also included in this section of the project is a kinetic in• vestigation of the reaction. Though the kinetic study was not as successful as hoped in elucidating the mechanism of the reaction, it did, however, give important generalizations regarding effects on the overall reaction when reagent quantities were varied or held constant depending on the situation being investigated. Since the kinetic study wasmore successful as a reagent study than on actual mechanism study, it is incorporated more appropriately under this heading.

A. Action of Oxalic Acid and Organic Acids

Oxalic acid is quite effective in increasing *the* sensitivity of the ourcumin method, but little is actually known about the mechanism of its action. A possibility is that oxalic acid forms an intermediate complex with boric acid prior to the color forming reaction with curcumin. The complex formed between boric acid and oxalic acid should have little strain due to the planar structure of boric acid in which the 3 hydroxyl groups are separated by angles of  $120^{\circ}$ . Diol compounds, such as oxalic acid with hydroxyl groups on adjacent carbons have been found to react readily with boric acid even in aqueous solutions whereas beto compounds react to an appreciable extent in non aqueous solvents. A five membered ring should thus result from the interaction of boric acid and oxalic acid. The oxalic-borate complex resulting is probably more susceptible to complexation (or chelation) With curcumin than boric acid would be alone.

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To gain greater insight into the effect of oxalic acid and to investigate other acids as substitutes tor oxalic acid, a study of their effect on the sensitivity of the reaction was first atarted by Powell and coworkers  $(18)$  and continued in this study. The acids used were selected on a basis of similarity to oxalic acid or because or various substituted groups which might show effects 0£ alight changes in structure.

Since the main advantage in using oxalic acid in the Naftel type method used in this project is to increase the sensitivity or the method, a comparison of individual runs using several different organic acids was made. The comparison was made against a run in which no organic acids was introduced at all. Absorbance readings were obtained from the final alcoholic extracts using the procedure on page 17. Absorbance readings were then plotted (y coordinate) vs. micrograms of boron (x coordinate) and the slopes of the resulting curves obtained from the relationship slope  $\cdot \cdot \cdot$ . By subtracting the slope or the standard curve, in which no organic acid was used, from the slope of a particular organic acid run, a "D" value ( $D$  • tendency to sensitize) for the particular organic acid run was obtained. This "D" value could have a positive value or negative Yalue, depending whether the organic acid sensitized or desensitized the method. Essentially the same procedure was used for all the organic acids investi• gated. Solutions containing O.O, *0.\$, 1.0,* 1.5, 2.0, 2.5 micrograms or boron were prepared and treated in porcelain dishes with 1.0 ml. of 0.1N  $Ca(OH)_2$ and evaporated to dryness. To the dry, cooled residues was added l ml. or

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acetone or ethy'l alcohol containing *1.59* millimoles of the particular organic acid being investigated. The value *1.59* millimoles *ot* oxalic aoid was arrived at by Powell (18) as the quantity of oxalic acid which gave the optimum absorbance reading for a particular lquantity of boron. Since it was desirable to compare the organic acids to oxalic acid itself, the same molar quantities of organic acids were used in each particular run.

Next were added  $0.2$  ml. 1:1 HCl and  $2$  ml. of  $0.1\%$  curcumin solution. (The curcumin solution used was made by dissolving  $\lceil .\rceil$  g. curcumin powder in 100 ml. acetone or 95% ethyl alcohol, depending on where the organic acid was dissolved in acetone or ethyl alcohol. The resulting solution was then evaporated to dryness on a water bath at 55<sup>°</sup>  $\pm$  3<sup>°</sup>C. with an additional 30 minute standing period after the solutions reached complete dryness. The residues were then extracted with  $25$  ml. of  $95\%$  ethyl alcohol, the extracts centrifuged and subsequent absorbance readings obtained at a wavelength of  $550$  millimicrons using ethyl alcohol as a standard.

Since most of the acids were not too soluble in acetone, ethyl alcohol was used as the solvent in most cases. When the organic acids were found to be not too soluble in either acetone or ethyl alcohol, a solution for this problem was to dissolve the 1.59 millimoles of organic acid and the required amount of curcumin together in 3 ml. of either acetone or ethyl alcohol. This solution was successful in all the difficult cases encountered. This *3* ml. method was used mostly in the oases of aromatic acids with relatively high molecular weights.

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### TABLE III

Effect of Various Acids on Slope of the Curcumin Standard Curve

Acid	"D" Values	Acid <sup>2</sup>	"D" Values
Oxalic Acid Tartaric Acid o-Phthalic Acid Phthalic Anhydride Salicylic Acid Acetyl-Salicyclic Acid 2,5-Dihydroxy benzoic $2, 4$ -Dihydroxy benzoic p-Hydroxy benzoic acid Gallic Acid <sup>1</sup>	+ 0.237 + 0.004 + 0.004 $-0.033$ $+0.017$ $+0.012$ + 0.052 + 0.014 $+0.012$ + 0.177	Formic Acid Acetic Acid Propionic Butyric Caproic Oxalic Acid Malonic Succinic Adipic Trichloroacetic Dichloroacetic Monochloroacetic -Chloropropionic -Chloropropionic Maleic Fumaric Benzoic	$+0.131$ 0.099 40.046 $-0.050$ $-0.112$ $+0.251$ $+ 0.010$ $-0.101$ $-0.094$ + 0.206 $+0.102$ $+0.099$ $+0.033$ 0.008 $+0.010$ $-0.092$ $+0.180$
Acetone standard curve slope $\approx 0.0450$ Ethyl alcohol " Ħ	0.0178 Ħ	Phenylacetic 1,2,3-Tricarboxylic propane Citric Mandelic Aconitic Aminoacetic Glyoxal Mercaptoacetic	$+0.017$ $+0.122$ $+0.022$ $+0.008$ $-0.086$ $-0.003$ $-0.045$ $-0.122$

 $\ddot{\phantom{a}}$ Gallic acid when extracted with  $25$  ml. EtOH gave a yellow-green solution instead of the usual yellow to orange-red solution.

 $\overline{\mathbf{z}}$ 

 $\bar{A}$ 

Results in this column from W. A. Powell's Tech. Quarterly Report for Callery Co., Callery, Pa. 1956.

From the above results it is readily seen that the best acids from the standpoint of increasing the sensitivity of the modified Naftel method are oxalic, trichloroacetic, benzoic, formic and 1.2,3=tricarboxylic propane. Gallic acid is not here included since when the dried residue of the  $55^{\circ}$   $\pm$  3<sup>°</sup>C. drying was extracted with ethyl alcohol it gave a yellow-green solution, unlike the other acids extracted. Thus a direct comparison with the other acids cannot be made.

It must be remembered that the comparisons were made using conditions optimum tor oxalic acid; possibly some or the other acids could be better under other conditions.

Going through the above list of acids, it can readily be seen that some are mono-acids, di-acids and tri-acids.

In the mono-acid group it can readily be seen that with an increase in acidic strength of the acid, complexation is favored more with boric acid. Thus trichloroacetic has a greater  $"D"$  value than dichloroacetic acid and monoohloroaoetio aoid respectively.

In the di•acid group, chelation with boric acid rather than mere oomplexation seemed to be favored. Thus any group of acids that form relatively strain free rings with boric acid should sensitize to a greater extent. Of all the di-acids exalic acid has the highest "D" value which is easily understood when ring formation is considered. The action between oxalic acid and boric aoid gives riaa to a *5* membered ring. Theoretically any acid which would in a similar way give rise to a  $5$  membered ring as oxalic acid does, should be ideal as a sensitizer. Malonic acid, which has an additional carbon more

than oxalic, upon chelation with boric acid should give rise to a  $6$  membered ring which is less strain free than a 5 membered heterocyclic ring. This explains certainly why its "D" value is less than that for oxalic acid.

Sise seems to be quite important also regarding the organic aoid. Most of the acids that were investigated in this projoot were much bulkier than those investigated by Powell (18). Thus almost all of the  $n_{\text{min}}$  values obtained in my work were much smaller than those obtained by Powell due to steric hinderance which would result. The intermediate chelate fonned between the di-acids and boric acid would be less susceptible to direct attack from curcumin than would be the case for a less bulky organic acid. Tho extra two coordination positions on the boron would probably be covered up or shielded away from the attacking curoumin by the bulky part of the organic acids chelated with boric acid.

The importance of steric hinderance can also be observed in the case of ;maleic and fumaric acid. The carboxylic groups of maleic are ois to each other and therefore chelate more readily with boric acid than does fumario acid, in which case the carboxylic groups are trans to each other.

B. Variation of Reagents in the Color Forming Reaction

After the first drying carried out on the hot plate in the moditied Naftel procedure for boron detennination, a white precipitate, of calcium borate is formed. Boron is thus stabilized and less subject to loss on heating in this state. In the following step of the reaction, namely during the water bath evaporation at 55°C., the color forming step of the reaction occurs.

 $-26 -$ 

Entering into this color formation reaction with the calcium borate are four reagents which play an intricate role in some manner in the formation of the colored substance. Acetone, oxalic acid, HCl, and curcumin together or alone actupon the calcium borate to produce the colored substance. All seem to have a definite role, though of not necessary equal importance in the overall picture. Five different runs, all aimed at solving a certain phase or the boron•ouroumin reaction mechanism were carried out. Two of the color forming step reagents, namely oxalic acid and HCl were eliminated entirely in some of the runs. HCl in one particular run was varied in concentration while the remaining reagents were held constant. Still in another run HCl was added to the final alcoholic extract instead of in its customary spot, namely in the color forming step.

For all the runs, 2 micrograms of boron were previously treated with 1 ml. O.1N Ca(OH)<sub>2</sub> and evaporated to dryness on a hot plate using low heat.

The color developing reagents were then added as indicated below and evaporation continued at  $55^{\circ}$  <u>+</u> 3°C. followed as usual with subsequent extraction with 25 ml. of 95% ethyl alcohol. The pH was also measured for each alcoholic solution, to note any radical change in hydrogen ion concentration, and absorbance readings obtained using ethyl alcohol as the reference solution.

To the hot plate residues the following reagents were addedr

 $- 27 -$ 

Run 1

1 ml. acetone

no oxalic acid

no ltl nc1

2 ml. curcumin solution (.1 g. ourcumin/100 ml. acetone) Run 2

1 ml. acetone

no oxalic acid

0.2 ml. 1:1 HCl

2 ml. curcumin solution (.1 g. curcumin/100 ml. acetone)

 $Run_3$ 

1 ml. acetone (containing .2  $g$ . oxalic acid)

0.2 ml. 1:1 HC1

 $Run \, \mu$ 2 ml. curcumin solution (.1 g. curcumin/100 ml. acetone)

1 ml. acetone (containing .2 g. oxalic acid)

vary quantity of HCl from 0.2 to 2.4 m.e. (milli-equivalents)

2 ml. curcumin solution (.1 g. curcumin/100 ml. acetone)

Run 5

1 ml. acetone (containing .2 g. oxalic acid)

add  $0.2$  to  $2.4$  m.e. HCl to the final alcoholic extract

2 ml. curcumin solution  $(1 g.$  curcumin/100 ml. acetone)

Absorbance measurements were carried out at wavelengths of  $\mu$ 50 through 700 millimicrons. Ethyl alcohol was used as the standard.

### TABLE IV



### TABLE V







In run 1 with no oxalic acid and HCl added no maximum is reached and also no appreciable increase in absorbance is obtained at a wavelength of  $525$  to 575 millimicron indicating that no chelation seemed to occur when these two reactants are both absent.

In run 2 with HCl but no oxalic no maximum is reached either. There ia however an increase in absorbance indicating that there is some complexation occurring between the boric acid• and eurcumin in the presence of the mineral acid. In run 3 with HCl and oxalic acid both present chelation definitely occurs with the chelate absorbing at its maximum at <sup>525</sup> to 550 millimicrons.

In run  $\mu$  the water effect is evident from the addition of 0. $\mu$  m.e. to 1.0 m.e. of acid. The quantity of acid in the range of 0.2 to 1.0 m.e. was pipetted from a lN HCl stock bottle and therefore as the quantity of acid added was increased from  $0.4$  m.e. to 1.0 m.e. the amount of water containing this acid also increased.

### TABLE VII



### TABLE VIII



From the results of Tables VII and VIII, it is evident from the absorbance readings that  $Ca(OH)_2$  is definitely necessary in order to prevent loss of boron for the final alcoholic extracts of the colored substance formed in the modified Naftel reaction. When Ca(OH)<sub>2</sub> is not added, a great deal of the boron present as boric acid is probably lost during the hot plate evaporation. Contrary, with addition of  $Ca(OH)_2$ , boric acid is stabilized with  $Ca(OH)_2$  to form a calcium borate precipitate which is stable during the evaporation process.

D. The Effect of Hydrochloric Acid

In the preceding study of the effect of  $Ca(OH)_2$  addition in the modified Naftel reaction, there was evidence of an increase in absorbance as l•l HCl additions were increased from .001 to .1 m.e. In order to determine what quantity of HCl would give maximum absorbance when used in conjunction with 1.59 millimoles of oxalic acid (.2  $g$ ./ml. acetone), the quantity of HCl added to various runs was varied from .001 to 3.6 m.e. (in .2 ml.  $H_2O$ ).

Runs were made with and without addition of 1.59 millimoles of oxalic. When oxalic was not used, 1 ml. of acetone was added nevertheless. Samples of 2:5 microgram boron were used throughout and the modified Naftel procedure followed as usual. Absorbances were measured at 550 millimicrons with ethyl alcohol as reference.

determined. Although this kinetic investigation did not prove to be as successful as hoped, several important generalizations were deduced *tram* the Various rate studies made.

The rate studies were carried out during the water bath evaporation stage of the reaction when the actual complexation seems to occur.

As a preliminary step several samples containing 2 micrograms or boron were placed in indiVidual porcelain dishes and treated in the normal way, that is 1 ml. of  $Ca(OH)_2$  was first added to the individual dishes and evaporated to dryness on a hot plate at low heat. To the cooled individual residues was then added 1 ml. acetone containing .2 g. oxalic acid, 0.2 ml. 1-1 HCl and 2.ml. curcumin solution (.l  $g.$ /100 ml. acetone). The resulting solutions were then placed on a water bath (55<sup> $\bullet$ </sup>  $\pm$  3<sup> $\bullet$ </sup>C.). The samples were then removed from the bath at 5 minute intervals and cooled on ice far below the water bath temperature so as to stop any further reaction from occurring to any appreciable extent. The cooled samples were then extracted with *25* ml. *of* acetone instead of *95•* ethyl alcohol so as not to complicate things with another solvent. Absorbance of each sample was then measured as soon as possible at a wavelength of 550 millimiorons using acetone as reference.

 $-35 -$ 



TABLE I

0.20 0.36 0.62 *0.56*  0.60 0.61 0.60

O.dO 0.62

From the above absorbance data it is seen that the major portion of the color forming reaction is complete after 20 minutes. This time also corresponds approximately to the time required for the solutions to reach dryness on the water bath at *55•.* It can be observed from the above results that the additional 30 minutes standing period at this temperature is not strictly essential since the absorbance readings do not change appreciably during this additional stand-by period. From a standpoint of speed, the drying periods could be shortened by at least 20 minutes with the loss or very little sensitivity. Where accuracy is more desirable than speed, the additional standing period is desirable.

As a further step a kinetic otudy was oartied out using the same reagents except tor oxalic acid which was eliminated completely. In the place of oxalic acid l ml. or acetone containing no oxalic acid was added to the residue of the hot plate drying. Absorbanoes were found to be muoh lower than in the case where oxalic acid sensitizes the method considerably. Additional standing of the residues before extraction seems advisable when

oxalic acid is not used. Absorbanoe readings increased from a value of 0.157 after 20 minutes to 0.200 after approximately 50 minutes.

Next a study was made on the rate of the reaction for 3 different boron concentrations, *0.5,* 1.0, 2.0 micrograms. The reagents used were the same and oxalic acid was again added to the procedure.

No correlation between concentration and time of reaction could be deduced from the absorbance readings obtained.

The effect of increasing the amount of oxalic acid added to the dry residue of the preliminary drying (hot pl ate) was also studied. Quantities of 50 micromoles, 100 micromoles and 1.59 millimoles of oxalic acid ware added to the residues in three separate runs respectively. From absorbance values obtained, evidence again pointed to the fact that the sensitivity was greatly increased by addition of larger and larger quantities of oxalic acid. The optimum value of oxalic acid was not detennined having previously bem found to be *1.59* millimoles by Powell and coworkers (18). As a final step to the kinetic studies, the effect of curcumin on the reaction was studied. Three runs ware again made using 2 micrograms of boron and varying the curcumin added (100 micrograms, 500 micrograms, 2 milligrams). The other reagents were used in their usual quantities. Absorbance readings were noticed to increase with the use of larger amounts of curcumin but not in the same order of magnitude as in the case of larger additions or oxalic acid. Unfortunately a simple relationship between concentration and time was not evident in any of the rate studiea made. All evidence, however, seems to indicate that the reaction is not a simple order reaction but is complicated due to several independent variables.

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V. Determination of Mechanism of Reaotion

As the final stage of this project, the molar ratios of boron to ourcumin and boron to oxalic acid were investigated in order to supplement the 1-l·l boron-curoumin-oxalic acid data obtained by Powell and coworkers (18). Finally along with data obtained from a pH study involving boron and oxalic acid in conjunction with a non-aqueous potentiometrlc titration investigation of the boron-oxalic acid system, a possible explanation or the mechanism of the modified Naftel reaction was attempted.

An explanation regarding the structure of the colored substance formed during the reaction was also sought.

A. Molar Ratio of Boron to Curcumin

A stwy was made to determine the molar ratio in which boron and curcumin combined (in the absence or oxalic acid) using Job• s method or continuous variation  $(5)$ . Job's method is a most important investigation tool in order to gain a greater insight of the molar ratio of the color substance formed in the boron-curcumin reaction. Powell and coworkers  $(18)$  using this method, found that boron and curoumin in the presence of oxalic acid combined in a lilil molar ratio.

It was recognized early that a characteristic color is one of the most important and distinguishing features of chelate compounds. In recent years a simple colorimetric method has been uaed to determine the composition *ot* chelate compounds in solution. This technique is called Job's method of continuous variation (5).

 $- 38-$ 

Consider the formation of a chelate compound AB<sub>n</sub> where A is a metallic ion and B is a chelating agent.

$$
\mathbf{A} + \mathbf{B}^{\circ}_{n} = \mathbf{A}\mathbf{B}_{n}
$$

$$
K = \frac{(AB_n)}{(A) (B)^n}
$$

where  $n$   $\bullet$  number of chelating groups

where the quantities in brackets represent molar activities which may be taken approximately equal to molar concentrations.. Ir the restrictions  $(A)$  + (B) = C is imposed where C is a constant, it can be shown readily that when concentration  $AB_n$  is a maximum then

$$
\frac{B}{A} = n
$$

or the first derivative of  $AB_n$  with respect to A is equal to zero.

$$
\frac{d (AB_n)}{d (A)} \bullet \circ
$$

In other words, tor a constant total concentration of metal A and chelating agent, the concentration of chelate is greatest when the metal and chelating agent are brought together in the same molar ratios in which they exist in the ehelate. If the chelate is the only colored substance present, the optical density of absorbance of the solution is proportional to the chelate concentration. Hence a plot of absorbance vs. composition of the solution

under the restrictions imposed would give a curve with a maximum at the composition corresponding to the formula of the chelate. The method is not generally applicable to systems in which more than one compound is formed.

In Powell's work the total number of moles was held constant at 0.926 micromoles land the mole percent boron varied from 20 to 80%. Since the absorbance of the colored species in the absence of  $\alpha$  alic acid is much less than when oxalic acid ispreeent, it was decided to use a larger total molar quantity. A total molar quantity of  $\mu$ .63 mioromoles was chosen after several trials as an ideal quantity to work with.

Twelve samples were then run in duplicate with the boron content ranging from 20 to 80% of the total moles. To the required amount *of boron in porcelain dishes was added 1 ml. of*  $0.1N Ca(OH)_2$ *. The re*sulting solution was then evaporated to dryness on a hot plate. To the cooled residues was then added 1 ml. acetone, 0.2 ml. lsl HCl and the required curcumin made up in 2 ml. of acetone. The resulting solutions were then evaporated to dryness in a water bath at  $55^{\circ}$   $\pm$  3<sup>°</sup>C. letting the dried residues stand an additional 30 minutes after complete dryness was obtained. The residues were then extracted with 25 ml. ethyl alcohol, the solutions centrifuged and absorbance readings obtained at 550 millimicrons using ethyl alcohol as a reference solution. The data are shown in Table XI and Figure 2.

- 40 -

### TABLE XI



#### Job's Method of Continuous Variation

A stock curcumin solution was prepared by dissolving 0.1853 g. curcumin in 50 ml. of acetone. Each 2 ml. of solution contained  $\mu$ .63 micromoles of curcumin. The curcumin required for the above runs was pipetted from the stock bottle and diluted with acetone so as to use a total of 2ml. curcumin solution.

When total moles boron was plotted vs. absorbance it was readily seen that a maximum occurred at an absorbance reading of 2.00 which corresponded to a 1:1 boron-curcumin ratio.

Thus we concluded that as in the case of boron-curcumin-oxalic system so also in the boronecurcumin system in which no oxalic acid was used a 1:1 molar ratio is obtained for boron and curcumin.

### B. Determination of the Boron-Oxalic Acid Molar Ratio

The boron-oxalic acid molar ratio was next determined by varying the total quantity of oxalic acid while maintaining boron constant in several runs. As a preliminary step the total volume of acetone giving



maximum absorbance without oxalic aoid was determined. This amount of acetone was then used to contain the amount of oxalic acid being investigated. The following scheme was followed: To 2 micrograms of boron in porcelain dishes was added 1 ml. O.1N  $Ca(OH)_2$  and evaporated to dryness on a hot plate. To the dried and cooled residues was added acetone in varying amounts from l ml. to 12 ml. to each of the individual dishes. This was done in order to determine what volume of acetone would give maximum absorbance. Hext wrre added 0.2 ml. lrl HCl and 2 ml. of 0.1% curcumin solution in acetone and the resulting solutions were evaporated to dryness on a water bath held at a constant temperature (55  $\pm$  3<sup>°</sup>C.).<br>The residues were then left on the water bath an additional 30 minutes after complete dryness was reached and then extracted with *25* ml. *of* ethyl alcohol. After the resulting solutions were centrifuged, absorbance readings were measured. When the volume at which maximum absorbance was determined, it was then used in subsequent runs to dissolve oxalic acid which was then varied while the other reagents were all held constant. Absorbance readings were again taken and maximum absorbance indicated at what molar ratio boron and oxalic acid combined.



 $\frac{2.0 \times 10^{-6} \text{ g}}{10.8 \text{ g}}$ . Foron **\*** 0.18 micromoles boron

$$
\frac{20 \times 10^{-6} \text{ g. Oxalto}}{90 \text{ g./mole Oxalio}}
$$
 = 0.22 micromoles oxide

### C. Potentiometric Titration Study of the Boric Acid-OXalic Acid Reaction. in a Non-Aqueous Medium

A potentiometric titration study of the reaction between boric acid and oxalic acid was next carried out. *A* O.Q981N NaOH alcoholic solution was prepared by dissolving 2 grams of NaOH pellets in *50* ml. ethyl alcohol. The resulting alkaline solution waa then standardized against 0.8976 g. or acid potassium phthalate using phenolphthalein aa indicator. The standardized base was then used as titrant in the potentiometric titration of solutions or oxalic acid and mixtures of oxalic and boric acids. From the titration it was hoped that the extent of reaction, if any, between oxalic and boric acid could be determined. Also from pH measurements during the t1trations a knowledge of the type of compound formed between oxalic and boric acids was hoped to be gained.

Initially *50* ml. of ethyl alcohol containing *0.5* millimoles of oxalic acid was placed in a porcelain casserole and titrated potentiometrioally with the 0.0981N NaDH. pH readings were then recorded after additiom of increments of 0.5 or 1 ml. of the standard base. A plot was then made of the pH vs. ml. of NaOH added. To another casserole was added the same amount of oxalic acid *as* above plus *5* millimoles of boron in ethyl alcohol. The initial

 $- 44 -$ 

volume was held constant as above at 50 ml. with the addition of ethyl alcohol. This solution was then titrated in a similar manner and again pH measurements were plotted vs. ml. NaOH added.

Finally 1 millimole of oxalic acid and then a solution of 1 millimole oxalic acid plus 0.5 millimoles of bornn were titrated in an analogous manner as above with NaOH.

### TABLE XIV

### Potentiometric Titration Data







From the preceding results and from a study of figures  $\mu$  and  $5$ we may postulate that the compound formed between boric acid and oxalic acid is a neutral species and not a positively charged compound. ٧Ź



is formed since from an observation of figures  $\mu$  and  $5$  we can see that initially upon addition of boric acid to oxalic acid the pH is greater than when oxalic acid is present alone at the start of the titrations. If compound B) were formed the initial pH of the solution containing the borin and oxalic acid mixture should be much lower (more acidic) than the pH of the oxalic acid solution alone since compound  $B$ ) is a stronger acid

than oxalic acid alone.

TABLE XV

rotentiometric: Titration Data					
		1 Millimole Oxalic Acid vs. 0.0981NNaOH	1 Millimole Oxalic 5 Millimoles boron	0.0981N VS. NaOH	
	ml. NaOH	<u>pH</u>	ml. NaOH	<u>pH</u>	
	0.0 1.0 2.0 3.0 4.0 5.0 6.0 7.0 7.5 8.0 8.5 9.0	3.05 3.31 3.48 3.55 3.69 3.88 4.05 4.31 4.48 4.61 4.89 5.28	0.0 1.0 2.0 3.0 4.0 5.0 5.5 6.0 7.0 7.5 8.0 8.5	3.30 3.35 3.57 3.70 3.82 4.02 4.20 4.32 5.09 8.30 9.02 9.30	
	9.5 10.0 11.0 12.0 13.0 18.0	7.60 9.70 11.02 11.29 11.31 11.33	9.0 10.0 11.0	9.35 9.39 9.41	

From the preceding results and from a study of figures  $\mu$  and 5 we may postulate that the compound formed between boric acid and oxalic acid is a neutral species and not a positively charged compound.

A)

is formed since from an observation of figures  $\mu$  and  $5$  we can see that initially upon addition of boric acid to oxalic acid the pH is greater than when oxalic acid is present alone at the start of the titrations. If compound B) were formed the initial pH of the solution containing the borin and oxalic acid mixture should be much lower (more acidic) than the pH of the oxalic acid solution alone since compound  $B$ ) is a stronger acid

than oxalic acid alone.



Figure 4  $\mathbb{R}^2$ 

## POTENTIOMETRIC TITRATION DATA



This is also another proof that boron and oxalic acid combine in a  $l-1$  ratio before complexation with curcumin. If compound B) were formed this would involve a  $1-2$  boron-oxalic complexation prior to the main reaction with curcumin.

From the two curves of oxalic acid titrated with NaOH there is evidence that the monocster of oxalic acid is formed when oxalic acid is dissolved in ethyl alcohol. Esterification between oxalic acid and ethyl alcohol is not complete however and therefore the titration with NaOH is actually a titration of the monoester. This explains why there is only one break in the titrating curves when we would normally have expected two if we were titrating oxalic acid as such.

Certainly evident from the titration curves involving the mixture of boric and oxalic acids, is a definite shift in curve breaks with respect to the titration curves for the solutions containing oxalic acid aloge. This is conclusive evidence of a definite reaction occuring between boric and oxalic acid (or ester). The following are the possible mechanisms that could occur in the reaction between boric and oxalic acid (or monoester of oxalic acid) .

1. Molecule of Oxalic Acid + Molecule of Boric Acid  $0 = 0 - 0$  $0 = C - OH$  HO,<br> $\uparrow$   $\uparrow$   $\uparrow$   $\uparrow$   $\uparrow$  $\rightarrow$   $\rightarrow$   $\rightarrow$  $\angle$ B-OH +  $2H_2$ OH  $0 \bullet 0 \bullet 0$  $0 \div 0 \div 0H$ 

 $- 47 -$ 

2. Monoester of Oxalic Acid + One Molecule of Boric Aoid

$$
0 = C - OC_2H_5 + HOC_3 + HOC_4 + O = C - O
$$
  
\n
$$
0 = C - O
$$
  
\n
$$
0 = C - O
$$
  
\n
$$
0 = C - O
$$
  
\n
$$
B = OH + HOH + C_2H_5OH
$$

3. Two Molecules of Oxalic Acid + One Molecule of Boric Acid

$$
0 = C - OH \n0 = C - OH \n0 = C - OH \nHO - C = 0 \nHO - C = 0 \nHO - C = 0 \nH0 - C = 0 \n0 = C - 0 \nB0 = C = 0
$$

3. has been ruled out as improbable from the potentiometric study carried out in this section.

The following is the probable mechanism occurring during the modified Naftel reaction for Boron determination which can now be deduced.

$$
0 = 0 - 0
$$
\n
$$
0 = 0 - 0
$$

The hydrogen ion from the 1-1 HCl acts as catalyst serving to initiate the chelation process. The hydrogen ion goess to one of the oxygens of the carbonyl groups of eurcumin. This gives the carbonyl carbon a residual plus charge. In order to recompensate for this electron unbalance, a hydrogen from the carbon between the two carbonyls or curcumin is lost leaving its pair of electrons to aid in rebalancing the electron unbalance created by the catalytic action of the hydrogen ion initiating the reaction. A double bond is thus formed between the oarbon and the electron deficient carbon of the carbonyl.

### **SUMMARY**

In the presant research project an attempt has been made to determine the mechanism of the reaction followed in the curcumin spectrophotometric method for boron determination using a modification of Naftel's method.

The reagents taking part in the reaction were studied by varying their concentrations and often omitting some entirely.

The use of  $Ca(OH)_2$  was proved to be essential in the method. Without it some of the boron as borio acid escapes during the preliminary hot plate drying. The Ca(OH)<sub>2</sub> tends to form a non-volatile calcium borate residue during the hot plate drying.

Several other organic acids were studied as possible substitutes for oxalic, but none was found to be more suitable than oxalic acid *as* a sensitizer. Either with or without oxalic acid Beer's Law seems to be obeyed in the miororange of boron concentration studied (0 to3 micrograms boron).

The use of different solvents for curcumin does not seem to have any noticeable effect on the sensitivity of the method. Acetone and ethyl alcohol were both used as solvents, both proving satisfactory. A possibility in the use of ethyl alcohol is the possible esterification with oxalic acid. That this esterification does occur, was evident when a potentiometric titration study was carried out for a solution containing oxalic acid dissolved in ethyl alcohol. Upon titration of this solution with an alcoholic sodium hydroxide solution only one break in the titration curve of oxalic acid could be observed when two breaks would have been expected since oxalic is a diearboxylic acid.

 $-19 -$ 

A potentiometric titration study was also carried out on a solution containing boric acid and oxalic acid in ethyl alcohol. It was at this point postulated that the compound formed between borio and oxalic acids is a neutral compound and not an anionic complex, since no hydrogen ions were found to be released in the process. It was further proved from this study that boric and cx alic acids combined in a  $l-1$  ratio. If a  $l-2$ boric-oxalic acid reaction occured, then release of hydrogen ions would have been expected.

From a kinetic study of the modified Naftel method, evidence obtained from absorbance data seemed to indicate that not much colored complex seems to form after the main reaction reaches dryness at  $55 + 3$ °C. on the water bath. If the additional 30 minute drying period is eliminated after complete dryness is reached, no great error seems to be introduced. This is quite an advantage since time of analysis may be shortened somewhat, making the method more appealing when time is a major factor in choice of a method.

The actual structure of the chelate formed between boron and curcumin was investigated. It was found that when boron and curcumin com bined under normal experimental conditions, a l:1 complex formed in the absence or oxalic acid. Powell and coworkers (18), using oxalic acid, found that boron and curcumin combined in a 1-1 ratio also. A 1-1 boron-oxalic compound was also found to be formed in another study carried out in this project. Maintaining the boron concentration constant, oxalic acid was varied until a maximum of absorbance was obtained. This was found to correspond to a  $l-1$  boron-oxalic compound.

Summarizing, boron-oxalic and curcumin definitely seem to form a 1-1-1 chelate type compound in the presence of a mineral acid. The mineral acid (HCl) seems to initiate the final complexation process between a 1-1 boro-oxalate compound and curcumin by catalytic action.

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I, Frank Xavier Pollio, was born on June 4, 1928 in Weehawken, New Jersey. I attended elementary school at Sarah H. Gilmore School, Union City, N.J.

From September 1939 to October 1945, I resided in Sorrento, Italy. There I graduated from Nino Bixio Technical High School and later attended the Nino Bixio Merchant Marine Academy for two years.

In June 1947 I was graduated from Emerson High School, Union City, N .J. which I attended from October 1945. In September 1948 I enrolled at St. Peter's College (N.J.) obtaining a B.S. in Chemistry in June 1952. From June 1952 to October 1957 I was enployed by Nitrogen Di vision, Allied Chemical & Dye, Hopewell, Va. as a control chemist.

In November 1957, I accepted employment on the research staff of the Rohm & Haas Co., Philadelphia, Pa. where I am presently employed.