# The Optimization of Copper Sulfate and Tincalconite Molar Ratios on the Hydrothermal Synthesis of Copper Borates

E. Moroydor Derun, N. Tugrul, F. T. Senberber, A. S. Kipcak, S. Piskin

**Abstract**—In this research, copper borates are synthesized by the reaction of copper sulfate pentahydrate (CuSO<sub>4</sub>.5H<sub>2</sub>O) and tincalconite (Na<sub>2</sub>O<sub>4</sub>B<sub>7</sub>.10H<sub>2</sub>O). The experimental parameters are selected as 80°C reaction temperature and 60 of reaction time. The effect of mole ratio of CuSO<sub>4</sub>.5H<sub>2</sub>O to Na<sub>2</sub>O<sub>4</sub>B<sub>7</sub>.5H<sub>2</sub>O is studied. For the identification analyses X-Ray Diffraction (XRD) and Fourier Transform Infrared Spectroscopy (FT-IR) techniques are used. At the end of the experiments, synthesized copper borate is matched with the powder diffraction file of "00-001-0472" [Cu(BO<sub>2</sub>)<sub>2</sub>] and characteristic vibrations between B and O atoms are seen. The proper crystals are obtained at the mole ratio of 3:1. This study showed that simplified synthesis process is suitable for the production of copper borate minerals.

*Keywords*—Hydrothermal synthesis, copper borates, copper sulfate, tincalconite.

## I. INTRODUCTION

**B**ORON is the chemical element with atomic number 5 and the chemical symbol B. The atomic mass is 10.81. It is a low-abundance element in both the solar system and the Earth's crust. Borate is the common name of the boroncontaining minerals. General boron reserves of world can be classified as sodium borate, calcium borate, sodium-calcium borates and magnesium borates. Copper borates can be thought as special boron compounds due its low percentage of reserves in world. Natural copper borates reserves of world are the forms of Jacquesdietrichite  $(Cu_2(H_2BO_3)(OH)_3)$  in Morocco, Santarosaite  $(CuB_2O_4)$  and Bandylite  $(Cu(B(OH)_4)Cl)$  in Chile [1]-[4].

There are more than 150 types of boron minerals in nature. Tincalconite ( $Na_2B_4O_7.5H_2O$ ) is the kind of boron mineral that has the  $B_2O_3$  content of 47.8%. It can be used as both boron sources of industry and raw material of synthesis process for the specific boron compounds [1].

Copper sulfate is a kind of salt that exists as a series of compounds that differ in their degree of hydration. The

E. Moroydor Derun, F. T. Senberber, and S. Piskin are with the Yildiz Technical University, Department of Chemical Engineering, Davutpasa Campus, 34210 Esenler, Istanbul, Turkey (e-mail: moroydor@yildiz.edu.tr, tsenberber@gmail.com, piskin@yildiz.edu.tr).

is with the Yildiz Technical University, Department of Chemical Engineering, Davutpasa Campus, 34210 Esenler, Istanbul, Turkey (e-mail:).

anhydrous form is a pale green or gray-white powder, whereas the pentahydrate (CuSO<sub>4</sub>·5H<sub>2</sub>O), the most commonly encountered salt, is bright blue [5].

Copper borates are the member of both delafossite (CuMO<sub>2</sub>) family and the metal borate groups [1], [6], [7]. With the lattice structure of copper borates that can be explained with a two-dimensional spin system; they exhibit specific features of optical transparency, high electrical conductivity  $(1.65 \text{ S/cm}^2)$ and an indirect gap of 2.2 eV [8]. Copper metaborate CuBO<sub>2</sub> is considered a potential system. This is because of recent theoretical investigations, which predicts that the band gaps of  $CuMO_2$  should increase as the ionic radius of M decreases. According to these predictions, CuBO<sub>2</sub> should have the largest band gap and hence better transmission characteristics. So they applications have the potential in the areas of superconductivity, transparent conductive oxides (TCOs) and diluted magnetic semiconductors [6], [7]. Also there are experiments about usage copper borates in preservation of wood materials, in catalysts of the dehydrogenation of organic compounds [9], [10].

Copper borate mineral can be synthesized at different structures of unit cells with the changing of experiment conditions. One of the typical copper borates [11] is CuBO<sub>2</sub>, which can be seen in Fig 1. The synthesis procedures of copper borates generally involve thermal operations. The copper borate (Cu<sub>3</sub>B<sub>2</sub>O<sub>6</sub>) was prepared by a solid-state reaction of copper oxide (CuO) and boric acid (H<sub>3</sub>BO<sub>3</sub>) at the reaction temperature of 900°C and time of 24 hours [12]. A kind of copper aluminum borate (Cu<sub>2</sub>Al<sub>6</sub>B<sub>4</sub>O<sub>17</sub>) was obtained via thermal synthesis method using copper sulfate (CuSO<sub>4</sub>), aluminum sulfate octadecahydrate  $(Al_2(SO_4)_3 \cdot 18H_2O),$ potassium sulfate (K<sub>2</sub>SO<sub>4</sub>) and H<sub>3</sub>BO<sub>3</sub> [10]. Malavi et al., synthesized copper metaborate of CuB<sub>2</sub>O<sub>4</sub> using the starting materials of Cu(NO<sub>3</sub>)<sub>2</sub>.3H<sub>2</sub>O and B<sub>2</sub>O<sub>3</sub> at the reaction temperature of 500°C [13].

There also few studies about the preparation of copper borates via sol-gel method. Zheng et al, managed to synthesize the two different types of copper borates in nano-scale using sol-gel reactions of  $CuSO_4.5H_2O$  and  $Na_2O_4B_7.10H_2O$  between reaction temperatures of 10-70°C, although there are unknown impurities in structure [14]. In another sol-gel study, Santra et al., synthesized obtained the copper borates with reaction of CuO and  $B_2O_3$  in the medium of citric acid at 75°C and 10 h [15].

A. S. Kipcak is with the Yildiz Technical University, Department of Chemical Engineering, Davutpasa Campus, 34210 Esenler, Istanbul, Turkey (phone: 0090-212-3834751; fax: 0090-212-3834725; e-mail: skipcak@ yildiz.edu.tr).

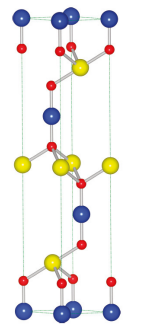


Fig. 1 Unit cell figure of rhombohedral CuBO<sub>2</sub>. The red, blue, and yellow balls are oxygen, copper, and boron respectively [11]

In this study, copper borate mineral has been synthesized without using gelation agent in hydrothermal conditions. Applied synthesis process in literature has been simplified. The effect of molar ratio of starting materials (CuSO<sub>4</sub>.5H<sub>2</sub>O to Na<sub>2</sub>O<sub>4</sub>B<sub>7</sub>.5H<sub>2</sub>O) to structure of obtained mineral has been investigated by identification and characterization methods of XRD and FT-IR.

### II. EXPERIMENTAL

#### A. Materials and Method

Raw materials used in synthesis experiments were copper (II) sulfate pentahydrate (CuSO<sub>4</sub>.5H<sub>2</sub>O) and tincalconite (Na<sub>2</sub>O<sub>4</sub>B<sub>7</sub>.5H<sub>2</sub>O). CuSO<sub>4</sub>.5H<sub>2</sub>O and Na<sub>2</sub>O<sub>4</sub>B<sub>7</sub>.5H<sub>2</sub>O were provided from Merck Chemicals and Kırka Boron Management Plant (ETIMINE Kırka Works) in Eskisehir-Turkey, respectively.

Preparation method of copper borate used in study was hydrothermal synthesis method. The liquid phase of hydrothermal conditions was used as ultra-pure water (18.3 m $\Omega$ .cm) that was produced from the equipment of Human Power I<sup>+</sup> Water Purification System. Reaction temperature and time were selected as 80°C, and 1 hour, respectively; to investigate the interaction between raw materials according to the mole ratio changes. Determined molar ratios of CuSO<sub>4</sub>.5H<sub>2</sub>O to Na<sub>2</sub>O<sub>4</sub>B<sub>7</sub>.5H<sub>2</sub>O were 1:1, 2:1, 3:1, 4:1 and 5:1.

After the reaction, the filtration process was used for the removal of excess copper sulfate. In this process, distilled water was used for the washing and dispersing the synthesized copper borates below the filter paper. After that the slurry content was dried in Ecocell model oven at 60°C. The dried content was triturated for the characterization operations.

#### B. Characterization

Raw materials were subjected to X-Ray Diffraction (XRD) analysis with Philips PANanalytical brand (Fig. 2 (a)) where in this equipment X-rays are produced from Cu-K $\alpha$  tube at the parameters of 45kV and 40mA [16].

Perkin Elmer Spectrum One (Fig. 2 (b)) Fourier Transform Infrared Spectroscopy (FT-IR) technique was used to determine which functional groups are presented in the samples. In the FT-IR technique Universal ATR sampling accessory – Diamond / ZnSe is used and measurement range is selected as 4000–650 cm<sup>-1</sup>, scan number is 4 and resolution set as 4 cm<sup>-1</sup> [16].



Fig. 2 (a) Philips PANanalytical XRD, (b) Perkin Elmer Spectrum One FT-IR

#### III. RESULTS AND DISCUSSION

## A. Raw Material Characterization Results

XRD patterns of the starting materials used in experiments are shown in Figs. 3 and 4 respectively. In the XRD pattern of copper sulfate pentahydrate (Fig. 3), the first three major peaks are seen in the 2 $\theta$  values of 18.746°, 16.146° and 48.476°, respectively. In the XRD pattern of tincalconite (Fig. 4), the first three major peaks are seen in the 2 $\theta$  values of 30.639°, 20.365° and 20.289°, respectively.

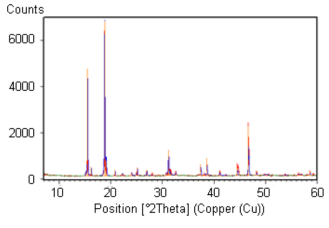


Fig. 3 XRD pattern of copper sulfate pentahydrate

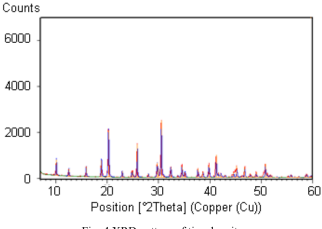


Fig. 4 XRD pattern of tincalconite

Results of XRD analyses are given in Table I. From the XRD results obtained it is seen that the raw materials used in experiments are identified as "01-077-1900" and "01-079-1529" pdf numbered "copper sulfate pentahydrate" and "tincalconite" minerals, respectively.

TABLE I XRD RESULTS OF THE RAW MATERIALS					
Mineral Name	Chemical Formula	Pdf #	Score		
Copper Sulfate Pentahydrate	CuSO <sub>4</sub> .5H <sub>2</sub> O	01-077-1900	42		
Tincalconite	Na2O4B7.10H2O	01-079-1529	74		

FT-IR spectra of the  $CuSO_4.5H_2O$  and  $Na_2O_4B_7.10H_2O$  are shown in Fig. 5.

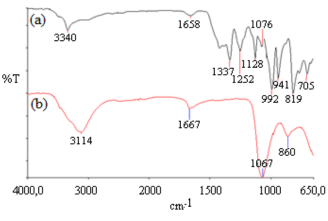


Fig. 5 FT-IR spectra of raw materials; (a) tincalconite, (b) copper sulfate pentahydrate

According to the FT-IR analysis of the tincalconite, characteristic peak are seen in the band values of 705, 819, 941, 992, 1076, 1128, 1252, 1337 and 1658 cm<sup>-1</sup>. In the CuSO<sub>4</sub>.5H<sub>2</sub>O spectrum have the characteristic peaks at the band values of 3114, 1667, 1063 and 860 cm<sup>-1</sup>. In these spectra, the peaks over 3000 cm<sup>-1</sup> may explain with the crystal water in structure. The peaks at lower band values can be explain with the vibrations between O and nonmetal atoms.

## B. Synthesized Copper Borate XRD Results

XRD results and patterns of the synthesized copper borates are shown in Table II and, Fig. 6 respectively.

TABLE II   XRD RESULTS OF THE SYNTHESIZED COPPER BORATES						
Mole Ratio	Pdf#	Mineral Name	Mineral Formula	Score		
1:1	00-001-0472	Copper Borate	$Cu(BO_2)_2$	41		
2:1	00-001-0472	Copper Borate	$Cu(BO_2)_2$	65		
3:1	00-001-0472	Copper Borate	$Cu(BO_2)_2$	70		
4:1	00-001-0472	Copper Borate	$Cu(BO_2)_2$	64		
5:1	00-001-0472	Copper Borate	$Cu(BO_2)_2$	68		

From the XRD results (Table II) obtained it is seen that "00-001-0472" pdf numbered "Copper Borate" mineral with the chemical formula of  $Cu(BO_2)_2$  is formed at all experiments varying crystal scores. The highest XRD score is seen at the mole ratio of "3:1".

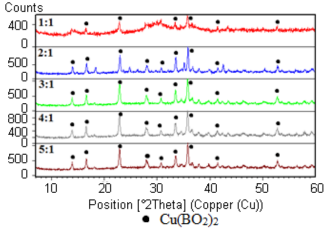


Fig. 6 XRD patterns of the synthesized copper borates

In Fig. 6, the first three major peaks of synthesized copper borate are seen in the 20 values of  $16.714^{\circ}$ ,  $22.902^{\circ}$  and  $35.744^{\circ}$ . Obtained peaks are not proper enough at the mole ratio of 1:1. With the increasing amount of copper sulfate in solution, proper peaks, which have similar band values of Copper Borate (00-001-0472 – Cu(BO<sub>2</sub>)<sub>2</sub>), are obtained. In the experiments for 4:1 and 5:1, minor changes in the structure of synthesized mineral are seen which may be due to excess copper sulfate pentahydrate in solution.

## C. Synthesized Copper Borate FT-IR Results

FT-IR spectrums of synthesized minerals are shown in Fig 7. When vibrations in each spectrum are compared, it is seen that percentage changes in transmissions of synthesized copper borate minerals are similar except the spectrum of 1:1 (Fig. 7 (a)). This difference can be explained by the formation of insoluble byproducts in hydrothermal media. Thus, it may not be separated by filtration stage. These peaks are seen between band values of 1404 and 1231 cm<sup>-1</sup> in the FT-IR spectra for mole ratio 1:1.

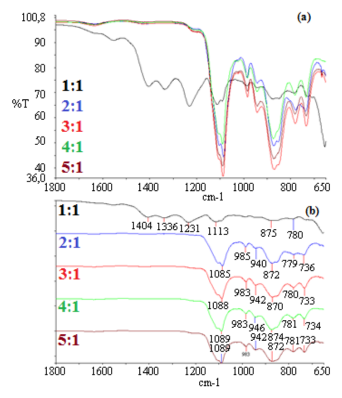


Fig. 7 FT-IR spectrum of synthesized minerals (a) overlaid, (b) split

In Fig. 7 (b), the peaks between 1120–980 cm<sup>-1</sup> might be the asymmetric stretching of tri-coordinate boron ( $B_{(4)}$ –O). Symmetric stretching of  $B_{(3)}$ –O can be seen between 980–870 cm<sup>-1</sup>. Symmetric stretching of tri-coordinate boron ( $B_{(4)}$ –O) can be seen in the band values of approximately 780 cm<sup>-1</sup>. Other peaks which have values lower than 750 cm<sup>-1</sup> explains the bending of  $B_{(3)}$ –O in the structure.

Obtained vibration band values are compatible with the previous studies in literature [17].

## IV. CONCLUSIONS

Boron found in nature in the combinations of other elements. Majority of boron reserves are found in Turkey (%73). Copper borates are a sub-group of boron minerals and have the usage fields of superconductivity, transparent conductive oxides (TCOs) and diluted magnetic semiconductors due to their optical and magnetic properties.

In literature copper borate synthesis have complex processes. The novelty of this research is the synthesis of copper borate using simplified processes. The effect of mole ratio to synthesis of copper borates is investigated and optimum mole ratio is determined in light of characterization analyses. These experiments can be seen as a preliminary step.

In future studies, copper borate synthesis at higher crystallinity and different structures are intended with the optimization of reaction temperature and time.

#### ACKNOWLEDGMENT

The author would like to thank S. Ucpinar and Y. Yarinineli for the contribution to our study.

#### REFERENCES

- [1] ETI MINE General Directorate, "Bor sector report", Istanbul, 2013.
- [2] J. Schlüter, D. Pohl, U. Golla Schindler, "Santarosaite, CuB<sub>2</sub>O<sub>4</sub>, a new mineral with disordered structure from the Santa Rosa mine, Atacama desert, Chile", *Neues Jahrbuch für Mineralogie-Abhandlungen*, vol. 185, 27-32, 2008.
- [3] A.R. Kampf, G. Favreau, "Jacquesdietrichite, Cu<sub>2</sub>[BO(OH)<sub>2</sub>](OH)<sub>3</sub>, a new mineral from the Tachgagalt mine, Morocco: Description and crystal structure", *European Journal of Mineralogy*, vol. 16, 361-366, 2004.
- [4] J.W. Anthony, R.A. Bideaux, K.W. Bladh, M.C. Nichols, "Handbook of Mineralogy, 1997, vol. 3, 35,.
- [5] P. A. Kokkoros, P. J. Rentzeperis, "The crystal structure of the anhydrous sulphates of copper and zinc", *Acta Crystallographica*, vol. 11, 361–364, 1958.
- [6] M. R. Snure, A. Tiwari, "CuBO<sub>2</sub>: A p-type transparent oxide", *Applied Physics Letters*, vol. 91, 1-4, 2007.
- [7] N. V. Kuratieva, D. Mikhailova, H. Ehrenberg, "A new polymorph of Cu<sub>3</sub>B<sub>2</sub>O<sub>6</sub>", Acta Crystallographica Section C - Crystal Structure Communications, vol. C65, i85-i86, 2009.
- [8] M. R. Snure, "Transparent conducting oxides and their applications", 2009, PhD Thesis, The University of Utah, Utah, USA.
- [9] S.S. Nair, "Effectiveness of Copper-Boron Diffusion Treatments for Wood", 2006, MSc. Thesis, University of Idaho, Moscow, USA.
- [10] C. Zhu, W. Li, X. Nai1, D. Zhu, F. Guo, S. Song, "Preparation of copper aluminum borate whiskers via flux method", *Crystal Research and Technology*, vol. 47, 73-78, 2012.
- [11] D. O. Scanlon, A. Walsh, G. W. Watson, "Understanding the p-Type Conduction Properties of the Transparent Conducting Oxide CuBO<sub>2</sub>: A Density Functional Theory Analysis", *Chemistry of Materials*, vol. 21, 4568–4576, 2009.
- [12] P.S. Malavi, S. Karmakar, S.M. Sharma, "High pressure structural investigations of copper metaborate (CuB<sub>2</sub>O<sub>4</sub>)", SOLID STATE PHYSICS: Proceedings of the 57th DAE Solid State Physics Symposium, vol. 1512, 88-89, 2013.
- [13] Y. Zheng, Z. Wang, Y. Tian, Y. Qu, S. Li, D. An, X. Chen, S. Guan, "Synthesis and performance of 1D and 2D copper borate nano/microstructures with different morphologies", *Colloids and Surfaces A: Physicochemical and Engineering Aspects*, vol. 349, 156– 161, 2009.
- [14] S. Santra, N.S. Das, K.K. Chattopadhyay, "Sol-gel synthesis and characterization of wide band gap p-type nanocrystalline CuBO<sub>2</sub>", *Materials Letters*, vol. 92, 198–201, 2013.
- [15] E. Moroydor Derun, A. S. Kipcak, "Characterization of some boron minerals against neutron shielding and 12 year performance of neutron permeability", *Journal of Radioanalytical and Nuclear Chemistry*, vol. 292, 871-878, 2012.
- [16] C.E. Weir, R.A. Schroer, "Infrared spectra of the crystalline inganic borates", *Journal of Research of The National Bureau of Standards- A. Physics ond Chemistry*, vol. 68A, 465–486, 1964.
- [17] J. Yongzhong, G. Shiyang, X. Shuping, L. Lun, "FT-IR spectroscopy of supersaturated aqueous solutions of magnesium borate", *Spectrochimica Acta Part A*, 56, 1291–1297, 2000.



**Emek Moroydor Derun** was born in Istanbul in 1976. Moroydor Derun graduated from B.Sc. in 1998, M.Sc. in 2000 and Ph. D. in 2005 from Chemical Engineering Department at Yildiz Technical University, Istanbul. Her research interest is in the area of waste management, lightweight concrete, semi conductive materials and She has many articles and studies in international and

boron technology. She has many articles and studies in international and national conference proceedings and articles.

#### World Academy of Science, Engineering and Technology International Journal of Chemical and Molecular Engineering Vol:8, No:10, 2014



Nurcan Tugrul was born in Gaziantep in 1973. Tugrul was graduated from B.Sc., M.Sc. and Ph.D. in Chemical Engineering Department at Yildiz Technical University, Istanbul. Her research interest is in the area of chemical technologies, evaluation of industrial wastes, food drying. She has many articles and studies in international and national conference proceedings and articles.



Fatma Tugce Senberber was graduated from B.Sc. at Yildiz Technical University in 2010. After she completed her M.Sc. studies at Yildiz Technical University in 2012, she started to Ph.D. studies at the same year and same department of university. She is interested in boron technologies such as alternative synthesis methods of boron minerals and evaluation of

industrial wastes in synthesis process. She also studied the characterization methods by instrumental analysis, kinetic studies of minerals and alternative application areas of synthesized minerals.



Azmi Seyhun Kipcak was graduated from Department of Chemical Engineering in Ege University in 2002. After completing the university studies he graduated from Bilgi University from the department of Master of Business Administration in 2004. He worked in Kultur University from 2003 to 2007 as a research assistant then he transferred to Yildiz Technical University at ted his MSc. studies about Chemical Engineering in

2008, where he started his M.Sc. studies about Chemical Engineering in 2006. He completed his M.Sc. studies at Yildiz Technical University in 2009 and Ph.D. studies in 2013. Now he is studying on different types of borate synthesis from different raw materials and wastes.



Sabriye Piskin graduated from Istanbul Technical University on Chemical Engineering with M.Sc. degree in 1974. She completed a Ph.D. degree at the same department in 1983. Her research interests include boron minerals and compounds, hydrogen storage technologies, fuel cell applications, materials characterization, coal, waste management, corrosion,

implants and synthetic materials production. She has more than fifty articles and eighty conference manuscripts pressed at the international area.