OPTICAL AND THERMAL BLEACHING OF X-IRRADIATED BARIUM ALUMINOBORATE GLASSES

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New EPR studies of boron electron centers (BEC), boron-oxygen hole centers (BOHC) and interstitial atomic hydrogen centers (H\textsuperscript{0}) in aluminoborate glasses X-irradiated at 77K are reported. Optical bleaching experiment suggests that protons (H\textsuperscript{1}) are also present.

INTRODUCTION

Radiation damage in borate glasses induces several paramagnetic centers such as BEC, BOHC and H\textsuperscript{0}. The BEC was first studied by Griscom\textsuperscript{(1)} in potassium borate glasses. As \textsuperscript{11}B nucleus is most abundant in nature (80.29%), three of the four transitions, due to hyperfine interaction between the trapped electron and the nuclear spin I = 3/2, are observed and easily recognised. It was proposed\textsuperscript{(1)}, among other possibilities, that BEC is a center originated from the trapping of an electron at a vacant site of a non-bridging oxygen of a tetrahedral BO\textsubscript{4} unit, forming a \textsuperscript{sp}\textsuperscript{3} dangling bond.

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Almost all X-irradiated borate glasses show a characteristic EPR line of 40 gauss width attributed to \textsuperscript{57}Fe\textsuperscript{(2)}. After some controversy\textsuperscript{(3)}, it was generally accepted that this center is formed by the trapping of a hole on the \textit{\tau}_\text{g} level of an oxygen bonded to a boron atom. Studies of these centers in glassy and crystalline compounds containing alkali and boron oxides proportional to 1:4, 1:3, 1:2 and 1:1, respectively, showed several characteristic structural units containing planar BO\textsubscript{3} units and BO\textsubscript{4} tetrahedra. In 1:1 compound the EPR line attributed to the BOHC has a structure of four lines, showing that the BOHC is localized in the neighbourhood of a bridging oxygen with substantial delocalization in the \textit{p}_z orbital of the trigonal boron. In borate glasses containing 20% of alkaline oxides, the BOHC EPR line is structureless and it was assigned to a hole trapped a non-bridging oxygen.

The BOHC spectrum of 30at%BaO; 50%B\textsubscript{2}O\textsubscript{3}; 20%Al\textsubscript{2}O\textsubscript{3} X-irradiated at 77K, studied here, is shown in fig. 3.

Insert Figure 1

The kinetic studies of the EPR absorption lines of the atomic hydrogen center (H\textsuperscript{0}) in barium aluminoborate glasses X irradiated at 77K have supported the assumption that they are trapped in B\textsubscript{10} ring structures\textsuperscript{(4)}.

The growth curves of optical absorption in barium aluminoborate glasses doped with Ce \gamma-irradiated with \textsuperscript{60}Co showed to be a superposition of saturation exponentials\textsuperscript{(5)}. This observation showed that the process of formation of BEC and BOHC is somewhat complex.
EXPERIMENTAL PROCEDURES

The samples of barium aluminoborate glasses (30 at% BaO; 60% B₂O₃; 10% Al₂O₃) were prepared and kindly furnished by Prof. A. Bishay of the Cairo American University. The details about the sample preparation, EPR measurements and temperature control conditions are explained elsewhere (4). The X-irradiations were done using a Philips X-ray apparatus operating at 40 KV and 20mA. The samples were cooled by a cold finger within a cryostat at 10⁻⁵ torr. The quartz window of this cryostat was located in front of a tungsten target tube. Care was taken in order to avoid sample heating above 110K during the transfer from the cold finger to the EPR spectrometer. The UV-irradiation was done with a 400W mercury lamp and the exposure to visible light with a Jarrell-Ash monochromator directed on the sample assembled in the EPR cavity.

The growth curves of the paramagnetic centers with X-irradiation times of 2, 4, 6, 8 and 24 hours, respectively, were obtained from average values determined from several samples. The error bars include the fluctuations among samples and measurements. After each measurement the sample was heated by 8 to 10 hours at 400°C in order to bleach the centers induced in the previous X-irradiation.

RESULTS

Barium aluminoborate glasses, X-irradiated at 77K, exhibit the characteristic EPR spectrum shown in figures 2 and 3.

Insert Figures 2 and 3

In the figure 2 we see three of the four lines of BEC, a strong line of BOHC at the center of the spectrum and two additional lines of \( n_1^0 \) provenient from hydrogen impurity in the sample.

The detailed structure of the BOHC central line appears in figure 3. The nature of these lines is sensitive to the alkali content in the glass and a detailed description of this center is found in the literature (2). The BOHC lines of our sample consist of a superposition of a structure of four lines ascribed to a hole trapped at a bridging oxygen and a single line attributed to a hole trapped at a non-bridging oxygen. On heating, the four-structured line is bleached before the single one, showing that the BOHC at bridging oxygen is less stable than the BOHC at non-bridging oxygen.

In figure 4 there are shown the growth curves of BEC and \( n_1^0 \), averaged over four samples. The results were normalized in order to allow comparisons.

Insert Figure 4

As shown in figure 2, one of the four lines of BEC was obscured by the BOHC spectrum. This line can be observed by means of the suppression of the BOHC line introducing an impurity ion which is a good hole scavenger, such as Ag⁺ or Ca⁺⁺⁺. Both capture holes forming Ag⁺⁺ and Ca⁺⁺⁺ ions, respectively. The \( n_1 = \frac{3}{2} \) line of BEC can be seen in figure 5, at 3138 Gauss, in the spectrum of an X-irradiated barium aluminoborate glass containing
cerium impurity.

With the gradual increase of temperature of annealing the spectra (see figure 6) show clearly the corresponding suppression of the BEC spectrum, remaining the more stable component, attributed to BOHC.

The $m_1 = +\frac{1}{2}$ line is also visible in the spectrum of X-irradiated aluminoborate glass containing silver impurity (see figure 7).

The samples X-irradiated at 77K, when heated in air, showed a blue luminescence. The EPR measurements showed that the BEC and $H_1^0$ were bleached, remaining only the BOHC spectrum.

The annealing rates of BEC and $H_1^0$ are not linearly correlated, as seen in the graphs of the respective isochronal decays (see figure 8). First, we see that the decay of BEC starts before than one of $H_1^0$, but with lower rate. At -60°C all the $H_1^0$ centers are bleached but the BEC concentration is still considerable.

The most interesting observation was the proof that the decay of BEC increases the concentration of $H_1^0$. We studied this correlation inducing the decay of BEC by the exposure of the sample to visible light. In figure 9 it is shown the decay of BEC ($m_1 = -\frac{1}{2}$) line and the corresponding growth of $H_1^0$ by exposition to a mercury lamp at 113K. The practically linear growth above 60 minutes of exposition is due to the creation of $H_1^0$ centers by the effect of the UV component of this light. In the same figure we see the growth curve of $H_1^0$ with the exposition to UV component of the mercury lamp light of a non-irradiated sample. Subtracting the production of $H_1^0$ by UV light, we see the increase of $H_1^0$ due to the BEC bleaching (see figure 10).

The decay rate of BEC is faster than the corresponding rate of increase of $H_1^0$ due to the additional electron-hole recombinations.

We have examined the limit of light induced decay of BEC. We observed that even red light induces the decay of BEC, although at slower rate, with a measurable increase of $H_1^0$ line intensity. On the other hand, only the UV component of light of the mercury lamp produces $H_1^0$ in non-irradiated samples, in which case is followed by the production of BOHC.

The isothermal decay of BEC is plot in figure 11. The points are experimental data and the full line shows the results.
of an empirical adjustment using the empirical formula.

\[ I(t) = I_0 t^{-b} \]

The values of \( I_0 \) and \( b \) coefficients are shown in Table 1, together with the determination coefficient \( r^2 \).

\begin{center}
Insert Table 1
\end{center}

The Arrhenius plot from these values is shown in figure 12 and the activation energy was obtained using the expression

\[ E_{\text{BEC}} = \frac{\Delta \log b}{\Delta T} = (8.6 \pm 0.3) \times 10^{-21} \text{ J} \]

where \( E_{\text{BEC}} = \text{BEC activation energy (eV)} \) and \( k = \text{Boltzmann's constant} \).

\begin{center}
Insert Figure 12
\end{center}

We see that both \( I_0 \) and \( b \) are not constant with temperature of the isothermal decay, showing the need of a more elaborated theoretical approach.

In addition to the study of the growth rate of BEC and \( H_1^0 \) centers, we followed the evolution of the \( \text{Fe}^{3+} \) line at \( g = 4,1 \). The intensity of this line decays with the time of exposure of X-rays (see figure 13). Similar observation was reported earlier by Bishay et al \(^{11}\) and was assigned to the reduction of \( \text{Fe}^{3+} \) to \( \text{Fe}^{2+} \).

\begin{center}
Insert Figures 13a and 13b
\end{center}

\[ .7. \]

\[ .8. \]

The BOHC lines of the EPR spectra do not change at room temperature. These lines are bleached by heating at 400°C by 8 h. During the bleaching it was observed an emission of orange light.

\[ .8. \]

**DISCUSSION AND CONCLUSIONS**

The samples of barium aluminoborate glasses, when X-irradiated at 77 K, showed the formation of paramagnetic centers of the same kind of those observed in other borate glasses, i.e., BEC, BOHC, \( H_1^0 \), and the reduction of \( \text{Fe}^{3+} \) with irradiation dose.

The BEC EPR spectrum observed is essentially the same described by Griscom et al \(^{11}\) in potassium aluminoborate glasses. It consists of a hyperfine structure of four lines due to the interaction between a single electron trapped in a \( sp^3 \) dangling orbital of boron and the \( ^{11}\text{B} \) nucleus (80.2%) which has the nuclear spin \( I = 3/2 \). The BEC is stable at temperatures below 110 K.

The heating of BEC above this temperature produces strong blue luminescence.

The BOHC spectra observed in our samples have two components: (I) a structureless component of the line due to a hole trapped at a non-bridging oxygen \(^{2}\); (II) a quartet due to a hole trapped at a bridging oxygen \(^{2}\). Heating of BOHC at temperatures up to 400°C produces an orange luminescence.

In figure 14 we show a sketch of the mechanisms of

\begin{center}
Insert Figure 14
\end{center}
formation of paramagnetic centers by X-irradiation at liquid nitrogen temperature in barium aluminoborate glass. The X-ray absorption induces the formation of electron-hole pairs. The analysis of the short time behaviour and the microscopic structure of these pairs is beyond the scope of the present work. We propose that the electrons and holes only can migrate throughout the sample if the electrons are excited to the conduction band and the holes injected in the valence band, otherwise they are always localized. This condition, obviously, is only satisfied when the sample is exposed to an irradiating source having photon energy at least equal to the band gap. The migration of holes is supported experimentally by the scavenging of holes by Ag⁺ and Ca³⁺ impurities.

The observed sites for the holes in the sample are the following: (a) non-bridging oxygen bonded to a boron atom; (b) bridging oxygen between a planar BO₃ and a tetrahedral BO₄ structural unit.

The BEC is probably a sp³ dangling orbital having a non-paired electron captured likely at an oxygen vacancy.

The mechanisms of formation of H⁺ on the light of the present experimental data can be: (a) the photodissociation of hydroxyls bonded to boron atoms forming BO₃⁻ and H⁺ followed by the capture of an electron; (b) the capture of a photo-electron by hydroxyls bonded to boron atoms forming BO₃⁻ and H⁺; (c) the photodissociation of hydroxyls forming BO₃⁻ (BOHC) and H⁺.

The reduction of Fe³⁺ into Fe²⁺ is assigned to the trapping of photo-electrons.

Figure 15 shows a sketch of the mechanism of formation of paramagnetic centers under UV light irradiation at 77K. In this process BEC is not formed.

Figure 16 shows a sketch of the effect of visible light on X-irradiated sample at liquid nitrogen temperature. The BEC is bleached even with red light showing that the BEC trap is shallow. The release of electrons from the BEC trap increases the concentration of H⁺ center. The blue luminescence was assigned to the electron-hole recombination at the BOHC site. The EPR spectrum of aluminoborate glass X-irradiated at 77K, exposed to 400W mercury lamp for 1h at 113K is shown in figure 17.

The sketch shown in figure 18 represents the effects of heating to room temperature of X-irradiated samples. The BEC bleaching releases electrons which partially recombine with holes of the BOHC producing the emission of blue light. A fraction of these electrons react with Fe³⁺ forming Fe²⁺.

It is shown in figure 19 a sketch of the effects of further heating of the X-irradiated sample to 400°C. The bleaching of BOHC release holes which are trapped by Fe²⁺ ions, which are oxidated to Fe³⁺.
From the above discussions we conclude that Fe and OH impurities take part in the mechanisms of luminescence and irradiation effects in barium aluminoborate glasses.

REFERENCES


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TABLE I - Constants of the expression I_BEC = I₀_BEC t⁻b calculated from experimental values.
FIGURE CAPTIONS

Fig. 1 - Diborate-type structural unit in 30at% BaO:60% B₂O₃; 10% Al₂O₃ glass showing the likely sites for trapping holes. The proportion 1:2 corresponds to the local relative [alkaline oxide]:[boron oxide] molar concentrations.

Fig. 2 - EPR spectrum of 30at% BaO:60% B₂O₃:10% Al₂O₃ glass X-irradiated at 77K, showing the characteristic BEC, BOHC and H°1 centers.

Fig. 3 - BOHC EPR spectrum of aluminoborate glass X-irradiated at 77K.

Fig. 4 - Growth of H°1 and BEC EPR peak-to-peak amplitudes of the derivative of microwave absorption (I°1 and I_BEC, respectively) in function of the X-rays exposure time (40 KV; 20 mA). Each error bar represents the distribution of measurements done with 4 samples. The exposure of 24 h was taken with only one sample.

Fig. 5 - EPR spectrum of 30at% BaO:50% B₂O₃:20% Al₂O₃ glass containing 3.5at% of cerium, X-irradiated at 77K during 8 hours (40 K; 20 mA; tungsten target). The BOHC line is sufficiently weak in order to show clearly the m₁ = 1/2 BEC line, usually obscured in undoped aluminoborate glasses.

Fig. 6 - EPR spectra of 30at% BaO:50% B₂O₃:20% Al₂O₃ glass containing 3.5at% of cerium, X-irradiated at 77K during 8h (a) after annealing at 253K for 3h; (b) after further annealing at 295K for 3h.

Fig. 7 - EPR spectrum of 30at% BaO:65% B₂O₃:5% Al₂O₃ containing 5at% of silver. The absence of BOHC line allows the observation of the usually obscured m₁ = 1/2 BEC line.

Fig. 8 - Isochronal annealing of BEC and H°1 centers in 30at% BaO: 60% B₂O₃:10% Al₂O₃ glass, X-irradiated at 77K during 8h. The EPR signal amplitudes were obtained by measuring the height (cm peak-to-peak) of the derivative of absorption of BEC m₁ = -1/2 and H°1 (upper magnetic flux density line). The dashed lines were drawn in order to aid the reader's eye.

Fig. 9 - Effect of optical bleaching of 30at% BaO:60% B₂O₃; 10% Al₂O₃ glass, X-irradiated at 77K, on the BEC (m₁ = 1 line) and H°1 (upper magnetic flux density line) EPR line amplitudes in function of the time of exposure to the light source at 113K. The measurements were taken using (a) 250 W and (b) 400 W mercury lamps. The growth of H°1 at the bottom (●) was obtained from the UV-component of the 400 W mercury lamp, shielded on the same sample when it was non-irradiated. The solid lines were drawn to aid the reader's eye. In this experiment it was used an optical-access cavity.

Fig. 10 - Growth curve of H°1 center after subtracting the growth of H°1 produced by the UV-component of the 400W mercury lamp (see Fig. 9b), compared with the decay curve of BEC. After 75 min of exposure, the curve above (●) is nearly saturated whereas the BEC (●) is almost completely bleached.
Fig. 11 - Isothermal annealing of BEC in 30at% BaO; 60% B₂O₃; 10% Al₂O₃ glass X-irradiated at 77K. The solid lines represent the best fit to the empirical formula \( I(t) = I_0 t^{-b} \).

Fig. 12 - Arrhenius plot \( b \times 1/T \) of the angular coefficients obtained from Fig. 11. The calculated activation energy is \( E_{BEC} = (8.6 \pm 0.3) \times 10^{-21} J \).

Fig. 13 - a) EPR of g=4.1 line of Fe³⁺ in 30at% BaO; 60% B₂O₃; 10% Al₂O₃ glass.

b) Decay curve of the Fe³⁺ EPR peak-to-peak amplitude in function of time of exposure to X-irradiation (40 kV; 20 mA) at 77K.

Fig. 14 - Mechanisms of formation of paramagnetic centers in aluminoborate glasses X-irradiated at 77K. The Fe³⁺ ions were partially reduced to Fe²⁺.

Fig. 15 - UV-irradiation effects in barium aluminoborate glass at liquid nitrogen temperature.

Fig. 16 - Visible light exposition effect on X-irradiated barium aluminoborate glass at liquid nitrogen temperature.

Fig. 17 - EPR spectrum of 30at% BaO; 60% B₂O₃; 10% Al₂O₃ glass X-irradiated at 77K for 8h and bleached with the light of 400 W mercury lamp for 1h at 113K. The amplitude of \( \delta^0 \) lines has grown whereas the BEC spectrum was completely bleached. The central strong line is attributed to BOHC.

Fig. 18 - Effects of heating to room temperature of X-irradiated aluminoborate glasses.

Fig. 19 - Effects of heating X-irradiated aluminoborate glasses from room temperature to 400°C.
Fig. 1

- \( h^+ \) Boron
- \( h^+ \) Bridging Oxygen
- \( h^+ \) Trapped Hole

Fig. 2

Derivative of Absorption (Arbitrary Units)

Magnetic Flux Density \( B(T) \)

- \( H_1^0 \)
- \( \text{BEC} \) \( m_{1/2} \)
- \( \text{BEC} \) \( m_{3/2} \)
- \( \text{BOHC} \)
Fig. 3

Derivative of Absorption (Arbitrary Units)

Magnetic Flux Density B(T)

Fig. 4

Amplitude (V p-p)

Time of Exposure (H)
Derivative of Absorption (Arbitrary Units)

Magnetic Flux Density \( B(T) \)

(Fig. 5)

Derivative of Absorption (Arbitrary Units)

Magnetic Flux Density \( B(T) \)

(Fig. 6)