

Structure role of cesium bromide in calcium silicate glasses and glass ceramics

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Abstract: To make the interpretation of the complex $x\text{CsBr} \cdot (100 - x)(\text{CaO}:\text{SiO}_2)$ glasses easier to study, the structure of glasses in a binary composition $\text{CaO}-\text{SiO}_2$ is being initially investigated. The changes in the crystallization behavior of glasses and the local environment surrounding silicon atoms could be easily followed using X-ray diffraction (XRD) and nuclear magnetic resonance (NMR) spectroscopy. The XRD pattern of the binary $50\text{CaO}-50\text{SiO}_2$ glass showed that its structure was amorphous. But when $(\text{CaO}:\text{SiO}_2)$ is replaced with CsBr, some sharp diffraction lines appear in system of the ternary $x\text{CsBr} \cdot (100 - x)(\text{CaO}:\text{SiO}_2)$ glass composition. A Polycrystalline $\text{Cs}_2\text{Ca}(\text{SiO}_3)_2$ structure is the primary phase in CsBr rich glasses. The results based on transmission electron microscopy (TEM-EDP) and X-ray diffraction pattern (XRD) are in excellent agreement, indicating that crystalline-clustered species develop in glasses enriched with CsBr. Both the NMR and FTIR spectra are clearly defined, and they contain different features that distinguish between different silicate structural subunits. One and two bridging oxygen atoms (BO) can be found in the main SiO_4 structural units. Such units become less shielded due to increasing of nonbridging oxygen atoms (NBO) in the silicate network by increasing CsBr at the expense of both SiO_2 and CaO. There is a good correlation between the data obtained from FTIR and NMR spectroscopy. Both techniques could differentiate between BO and NBO involved in the silicate structural units.

Keywords: Ion exchange; CsBr; Invert structure; Silicate species

Introduction

The effect of thermal heat treatments (THT) can result in the formation of the crystalline Ca_2SiO_4 and CaSiO_3 polymorphs in the amorphous silicate glasses containing calcium oxide [1, 2]. The cement chemistry community is quite interested in producing materials under the application of a thermal heat treatment process [2–4]. Inorganic stabilizers such as metal halides, in addition to the (THT) methods, have a direct impact on the stability and crystallinity of the prepared materials. The well-formed structural units that make up the host glass network can be controlled through addition of specific concentrations from halides in the host glass matrix. Additionally, the stability

and crystallization of a glass network are affected by the indirect manner connected to the ion exchange process. For instance, when the metal halide salts and a metal oxide are introduced to the primary glass formers, such as P_2O_5 , TeO_2 , SiO_2 , B_2O_3 , etc., an exchange reaction between the two types in the primary glass forming network may occur. The ion exchange interaction (IXI) between the halogen atom from the metal halide and the oxygen atom from the modifier oxide changes the physical properties and structure of the produced glass samples [4].

An inversion structure with crystalline phases can be created as a result of IXI process [2–4]. In this situation, X-ray diffraction analyses showed that the base glass that is free from halogens, is entirely amorphous [5]. The amorphous glass network becomes polycrystalline with the addition of metal halides. For instance, more open spaces might be formed when CdO is increased at the expense of Na_2O and P_2O_5 [5] using invert phosphate glasses. This

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may contribute to structural weakness, which is manifested in an increase in glass-free volume (V_f). The concentration of (NBO) should probably grow with increasing CdO at the expense of a concentration of similar magnitude from P_2O_5 and Na_2O , according to increasing the latter in the glass network. The formation of the cadmium invert phosphate glasses results in the remaining of the amorphous structure even at high ion exchange rates.

The main aim of this study is obtaining new cesium invert crystalline silicate glasses, as a first trial, through ion exchange process between oxygen atoms from CaO and bromine ions from CsBr. In this case, opened spaces in the host glass matrix might be increased, to play the main role of increasing both the number and mobility of charge carriers [4, 6]. Increasing the latter recommend the studied material to be used as row material shared in the manufacturing of solid state batteries.

Experimental details

Glass samples in the system $x\text{CsBr} \cdot (100 - x)(\text{CaO}:\text{SiO}_2)$ were prepared using analytical grade materials of purity around 99.9% of SiO_2 , CaCO_3 , and CsBr from Aldrich company. Glasses were formed and obtained using the molar formula $x\text{CsBr} \cdot (100 - x)(\text{CaO}:\text{SiO}_2)$, where $x = 0, 5, 15, 30,$ and 40 mol%, Table 1. The melting process was carried out in crucibles made of silica at melting temperature between 1250 and 1560°C in an electric furnace. To ensure homogeneity, the melts were shacked multiple times. In an electric furnace, the resulting glasses were annealed at 400°C for 4 h to reduce the internal stresses and strains formed in the bulk of as prepared glasses.

To measure X-ray diffraction, a Shimadzu X-ray diffractometric is employed (the apparatus type Dx-30, Metallurgy institute, El Tebbin, Cairo, Egypt). By comparing the maximum peak and intensity values to patterns in the international powder diffraction file (PDF) database of the joint committee for powder diffraction standards (JCPDS), the material type can be identified.

Table 1 Glass composition in mol%

Composition mol%		
CsBr	CaO	SiO ₂
0	50	50
5	47.5	47.5
15	42.5	42.5
30	35	35
40	30	30

The powdered samples' NMR spectra were captured using a Joel NMR 500 (11.74T, Mansoura University) spectrometer. To solve the issue of the relaxation time, one plus detection techniques were used. Pulses with a duration of $2\text{--}3$ s, an angle of around 45° , and substantial delays (25 s) between them were gathered. Up to 1400 scans were accumulated depending on the specific sample. The zirconia rotating sample was capable of rapid rotation at the magic angle at a rate of 7.5 kHz due to two different double-bearing systems.

The FTIR absorbance spectra are measured in the $400\text{--}4000\text{ cm}^{-1}$ region with a spectral resolution of 2 cm^{-1} using a Mattson 5000 FTIR spectrometer. The resulting spectra were normalized to that of a blank KBr pellet after background and dark currents were removed using a two-point baseline. To minimize the effect of the powder sample's concentration on the KBr disc, normalization is required. At least three samples of each material were tested. The spectrum of every sample is collected through a collection of 20 scans.

Results and discussion

XRD results

XRD pattern of glass samples of different CsBr concentrations are shown in Fig. 1. The spectra of glasses of 0 and 5 mol% CsBr confirmed the amorphous structure, since there is no any sharp diffraction peaks but just a broad

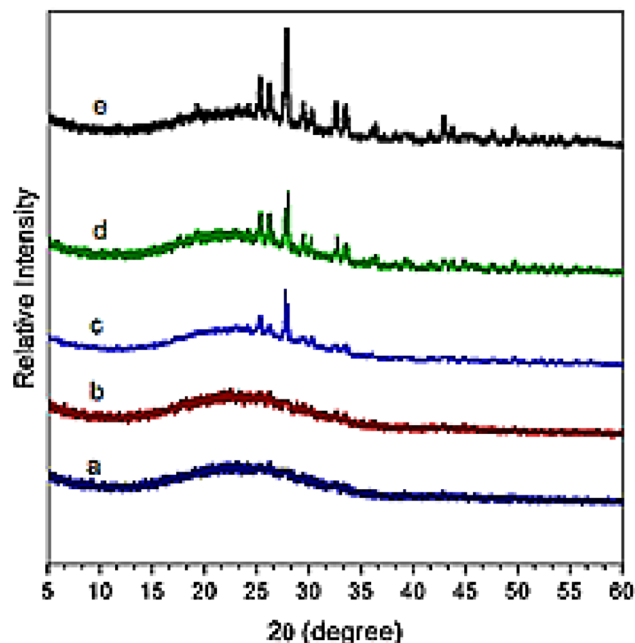


Fig. 1 XRD spectra for glasses of $0, 5, 15, 30$ and 40 mol% CsBr (a, b, c, d and e) respectively, all samples were annealed at 400°C for 4 h

Fig. 2 ^{29}Si NMR of the measured spectra (a) and of deconvoluted spectra (b) of sample free from CsBr

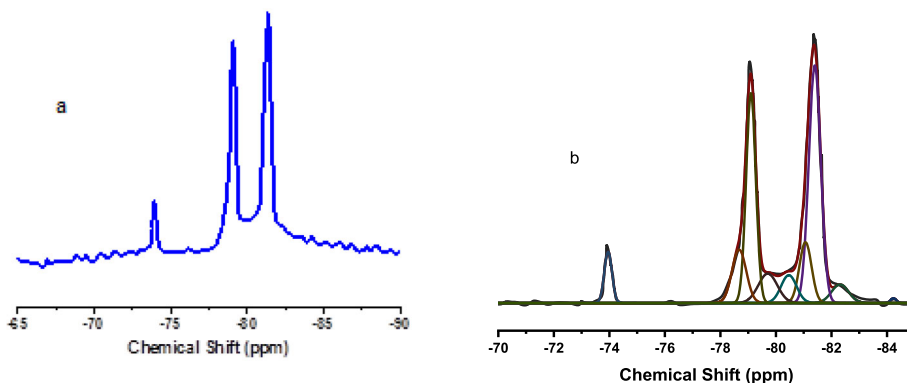


Fig. 3 ^{29}Si NMR of the measured spectra (a) and of deconvoluted spectra (b) of sample containing 40 mol% CsBr

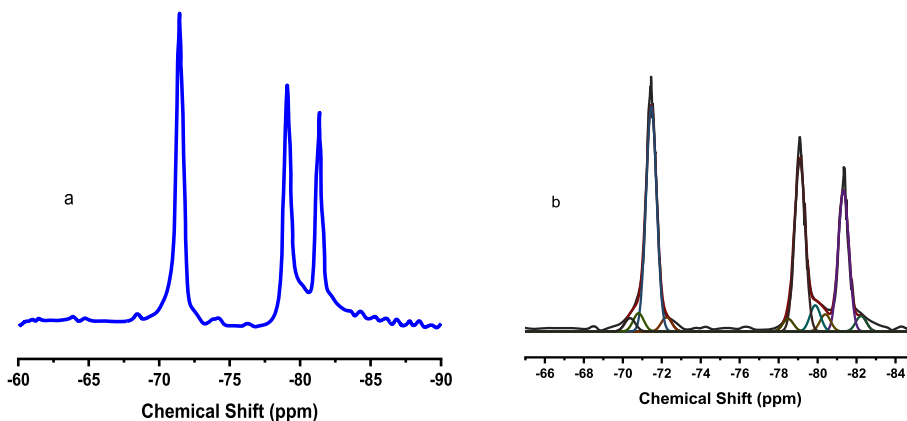


Table 2 Changes of relative area of the resonance peaks located around -74 ppm, and between -80 and -83 and -78 and -80 ppm

Position	74 area	78–80 area	80–83 area
Sample 1	0.048	0.413	0.537
Sample 2	0.413	0.347	0.238

hump lies between $2\theta \sim 12^\circ\text{--}37^\circ$ is appeared. On the other hand, the spectra of glasses containing 15, 30 and 40 mol% CsBr (c, d, e) contain some sharp diffraction peaks superimposed on the wide halo. The intensity of the sharp diffraction lines are gradually increased with increasing CsBr content at expense of both CaO and SiO_2 which leads to an increase in the percentage of crystalline species to become maximum at 40 mol% CsBr. Therefore, the samples containing 30 and 40 mol% CsBr are considered as the suitable composition presenting a glass ceramic type, since they contain a specific crystalline Cs_2SiO_3 and $\text{Cs}_2\text{Ca}(\text{SiO}_3)_2$ phases [7, 8]. The latter has special interest as a crystalline host for shielding materials against ionized radiations, due to its high chemical stability.

The formation of crystalline structure is considered as a direct result of ion exchange process between Br ions from CsBr and oxygen atoms from CaO. The ion exchange process increases the total modifier contents from Cs_2O , CaO and CaBr_2 to create more NBO which facilitate the mobility of ions to form a collective crystalline species. This aspect can be clearly clarified from the results based on NMR spectroscopy, which discussed below.

NMR spectroscopy

^{29}Si NMR and FTIR spectroscopy have been used to investigate the structure of glasses in the system $x\text{CsBr}$ ($100 - x$) ($\text{CaO}:\text{SiO}_2$) with $x = 0, 5, 15, 30$ and 40 mol%. Addition of CsBr at expense of both CaO and SiO_2 results in the decondensation of the calcium metasilicate chains, which is defined as silicate unit containing two bridging oxygen atoms (BO) named Q^2 [9–11]. The fraction of Q^2 sites of chemical shift lies at about (-82 ppm), Fig. 2a, b, decreases along with the formation of Q^1 and few Q^0 structural units of the average chemical shift of value between (-78 and -73 ppm), Fig. 3a, b. It can be detected from Figs. 2 and 3 and Table 2 that there is a clear change in both peak position and intensity of Q^n due to change of

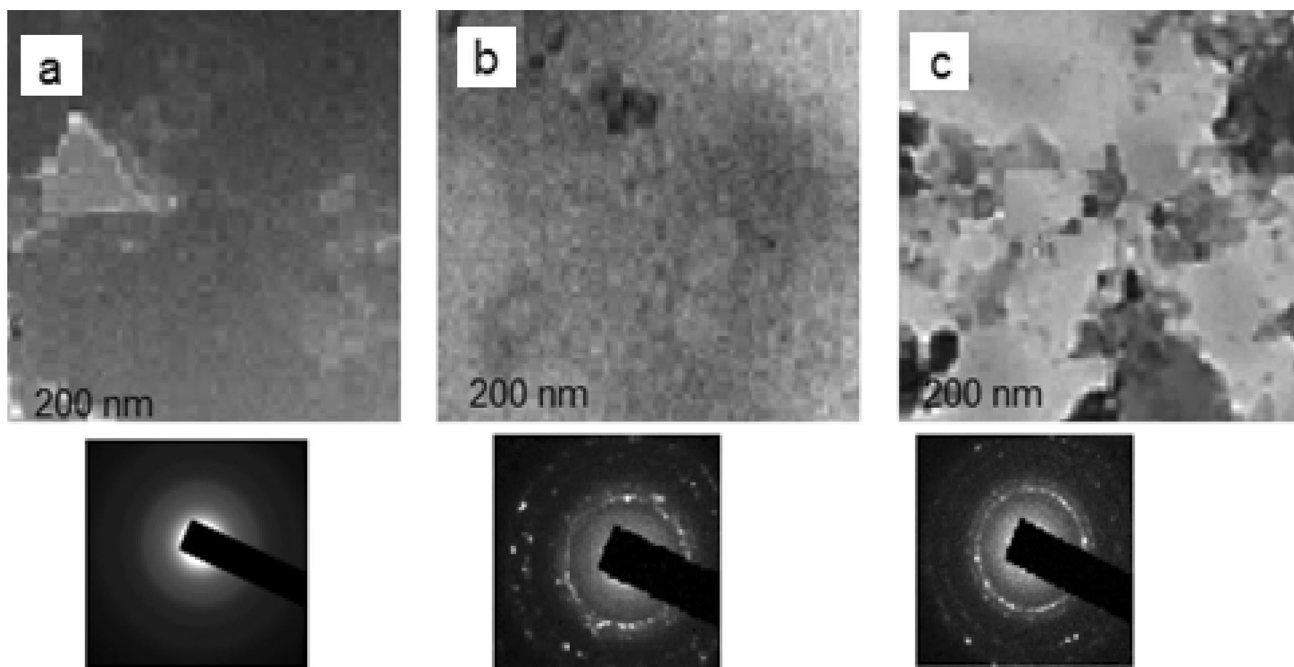
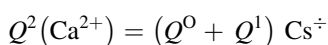


Fig. 4 TEM-EDP micrographs for sample free from CsBr (micrograph a) and of samples containing 30 and 40 mol% CsBr

CsBr from 0 to 40 mol%. The shift of peak to reach -73 ppm which represents a mixture from Q^1 and Q^0 sites leads to the formation of orthosilicate groups [12, 13] which are clearly formed upon exchange of both CaO and SiO₂ with CsBr. The changes in both spectral feature and chemical shift values of ²⁹Si NMR spectra (Figs. 2 and 3) leads to the conclusion that Cs⁺ ions are bonded mainly to the less shielded silicate units (Q^1 and Q^0 sites). Some Q^0 [12, 13] species which are present in the glass containing 30 and 40 mol% CsBr are formed by the following disproportionation reaction,



The increase in (Q^1 and Q^0) at the expense of Q^2 leads to conclusion that there is ion exchange between some of oxygen atom from CaO and bromine ion from CsBr which results in formation of Cs₂O and CaBr₂. This process was also confirmed in PbI₂ and Ag₂O telluride glasses [14]. The ion exchange process increases the total contents from (CaO and Cs₂O) as modifiers which leads to formation of extra nonbridging oxygen atoms (NBO) in the silicate network [12, 13]. Moreover, the bromine ions are inserted interstitially [14] in the silicate network resulting in elongation and weaken Si–O bonds which has a direct effect on NBO formation. In addition, the decreasing SiO₂ concentration upon CsBr exchange leads to formation of additional NBO atom which can be considered as agent for enhancing the crystallization process in silicate glasses [13, 14]. Presence of NBO ions and Cs⁺ with an extremely high concentration leads to formation of electron–hole

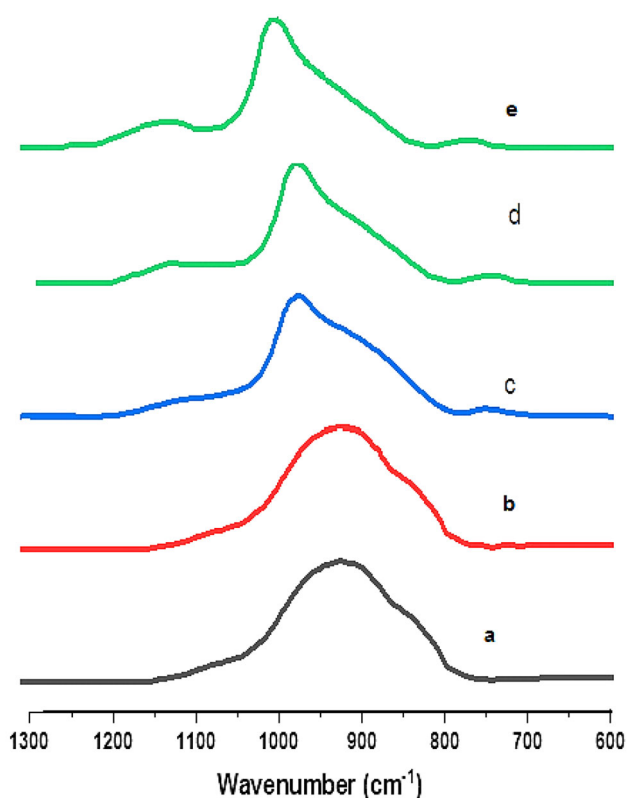


Fig. 5 FTIR absorbance spectra of glasses containing 0, 5, 15, 30 and 40 mol% CsBr, spectra a, b, c, d and e respectively

centers which makes the crystalline glasses useful as a shielding materials against ionized radiation [15].

From ^{29}Si NMR spectroscopy, both the first and second coordination neighbors of Si nuclei can be determined. Accordingly BO and NBO associated with SiO_4 tetrahedral units can be simply followed. From Figs. 2 and 3, it can be documented that the average coordination number of bridged oxygen atoms (BO) around Si atom decreases as CsBr increases which means increasing concentration of (NBO). The samples of 40 mol% CsBr contains the highest number of NBO founds in orthosilicate groups ($Q^1 + Q^0$) of chemical shift at about -73 ppm. Accumulation of NBO around Cs^+ cations leads to the formation of cluster species from Cs^+ and Si which is called alamosite [16] crystalline phase Cs_2SiO_3 . In such a case the silicate groups should be formed in orthosilicate crystalline type, i.e., highest number of NBO [12, 13, 16–20].

TEM-EDP investigation

It clear from TEM micrographs, Fig. 4a–c that the most accumulated and precipitated species are appeared in micrograph (c) which represents TEM-EDP of sample containing 40 mol% CsBr. The precipitation of Cs_2SiO_3 phase increases with increasing CsBr contents. The associated diffraction patterns confirm the enhanced precipitation process with increasing CsBr contents which has been also evidenced from XRD spectra of the same compositions. Generally increasing CsBr content at expense of CaO and SiO_2 in silicate glasses will change the feature of the well-formed crystalline units through formation of Cs_2SiO_3 phase [16, 20]. The nature of glass matrix was highly influenced by the formation of crystalline Cs_2SiO_3 species,

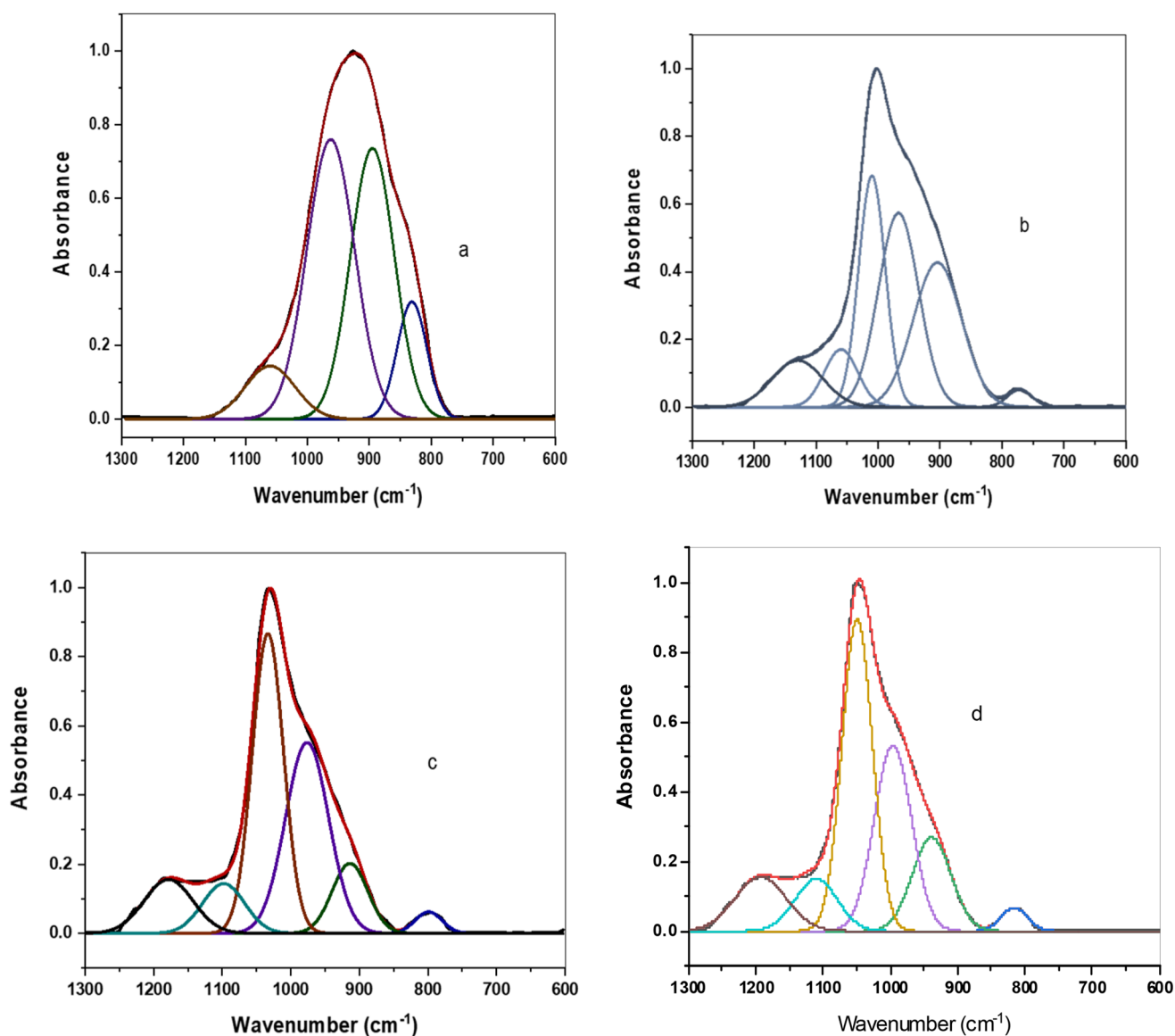
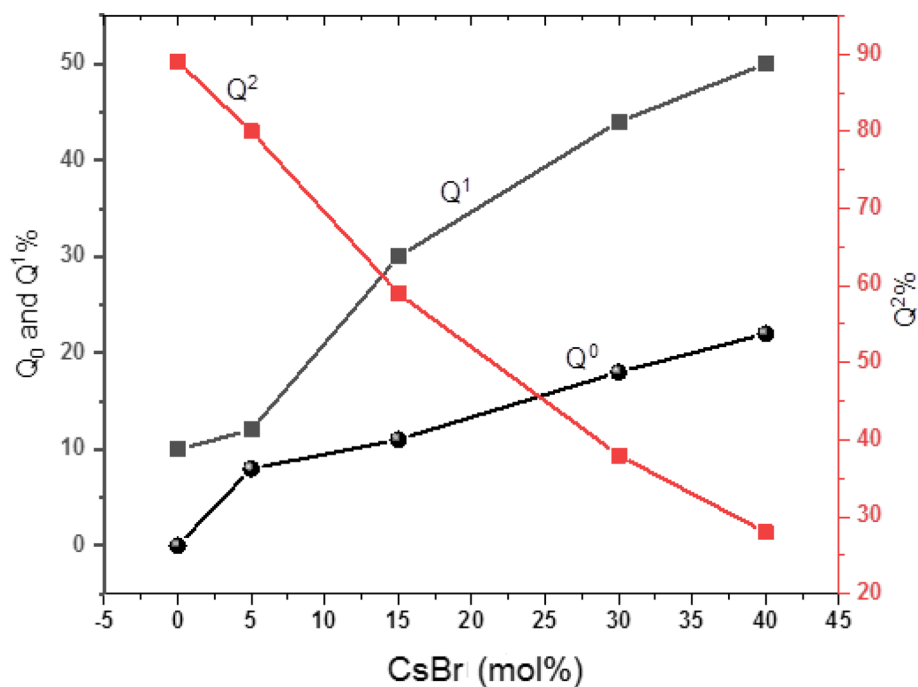


Fig. 6 FTIR analyzed spectra of glass containing 0, 15, 30 and 40 mol% CsBr

Fig. 7 Change of different silicate structural units Q^2 , Q^1 and Q^0 with CsBr content



since more ordered network is the main product as was evident from both X-ray and TEM-EDP results. Separated species of larger size are found in glass containing 40 mol% CsBr presented from TEM micrograph, Fig. 4c.

FTIR analysis

The FTIR spectra of the investigated glasses are displayed in Fig. 5. In the composition of 0 and 5 mol% CsBr, the CaO–SiO₂ spectra exhibited a broad absorption band with a peak located at roughly 950 cm⁻¹ which assigned to Si–O vibration in the middle unit, which contains two bridging oxygen atoms, Q^2 [16, 17]. The Q^1 sites [18, 21, 22], known as chain-end groups, are responsible for the well-defined, high peak (1050 cm⁻¹) in the FTIR spectra of glasses containing 15, 30, and 40% mol% CsBr. It can be assumed that these Q^1 sites are connected to mixed Ca²⁺ and Cs⁺ ions, since (CaO + Cs)/SiO₂ ratio in the glass is extremely high enough to form Q^1 species.

The decrease in peak intensity at 940 cm⁻¹ and increasing one at 980 cm⁻¹ confirm that most of cesium and calcium entered the phosphate glass network as a modifier. In some cases Cs₂O can be entered with two different coordination's which are tetrahedral CsO₄ and octahedral CsO₆. The electrostatic field strength of octahedral cesium is smaller than that of tetrahedral one. Then, some of Q units were subjected to more deshielding as a result of the octahedral or modifier cesium oxide. This hypothesis is thought to be the primary cause of the peak shift from 950 to 1050 cm⁻¹. This variation may possibly be a result of the Q^1 and Q^0 groups' interaction with the

octahedral cesium bond. The FTIR spectra of the glass with 30 and 40% CsBr can then be interpreted as showing extraordinarily high concentrations of Q^1 and Q^0 while glasses with 0 and 5 mol% CsBr had Q^2 as a significant species. Other minor peaks may be seen at 1150 and 750 cm⁻¹, respectively, which are attributed to Si–O vibrations in the middle groups of short chains and cycles [18, 21] metasilicate units.

As shown in Fig. 6a–e, line shape simulations or deconvolution techniques can be used to quantitatively calculate the proportion of different Q^n species in the silicate network [18–23]. The three silicate units Q^0 , Q^1 , and Q^2 in the ²⁹Si MAS NMR studied spectra, are the average main units. Each Q^n fraction can be determined and shown in Fig. 7 after peak deconvolution. The fractions of Q^0 and Q^1 both increase with increasing CsBr content, the fraction of Q^2 units decreases. This behavior leads to the conclusion that the CsBr–CaO exchange characteristic's tendency to decrease ($Q^2/Q^1 + Q^0$) ratio which is primarily caused by the local environment's distortion. The formation of additional NBO in conjunction with the addition of CsBr is what ultimately leads to variations in the percentages of Q^2 and ($Q^1 + Q^0$) with increasing CsBr content. The reversal variations in the peak intensity of the resonance lines of ²⁹Si NMR spectra provide strong support for this result (Fig. 2). Then from the results of NMR and FTIR spectroscopy, it can be confirmed that there are a good agreement between the data obtained from the different techniques. Both confirmed that NBO atom increased upon addition of CsBr at expense of both CaO and SiO₂.

Figure 7 summarizes the quantitative data obtained from the analysis of FTIR spectra. As can be seen from the figure, the percentage of Q^2 decreases and the values of both Q^1 and Q^0 increase with increasing CsBr content from 0 to 40 mol%. Such changes lead to confirm that less shielded silicate structural units are formed due to increasing of nonbridging oxygen atoms (NBO) in the silicate network by increasing CsBr at the expense of both SiO_2 and CaO . This argument was supported from NMR and FTIR techniques that could differentiate between BO and NBO involved in the silicate structural units.

Conclusions

The glasses prepared in mol% in the $x\text{CsBr} \cdot (100 - x)$ ($\text{CaO}:\text{SiO}_2$) system have been studied for the first time by several techniques. The XRD pattern of the binary $50\text{CaO}-50\text{SiO}_2$ glass showed that its structure was amorphous. Some crystalline phases appear in the ternary $x\text{CsBr} \cdot (100 - x)$ ($\text{CaO}:\text{SiO}_2$) glasses. Data based on (TEM) and X-ray diffraction patterns were in excellent agreement, indicating that the crystalline-clustered species can be developed in glasses enriched with CsBr. Both the NMR and FTIR spectra contain specific different features that can distinguish between different silicate structural subunits. Such units become less shielded and richer with (NBO) in the form of Q^0 that are electrically balanced with cesium cations by increasing CsBr at the expense of both SiO_2 and CaO . The analysis of FTIR and NMR spectra by peak separating method has a powerful action to distinguish different types of oxygen atoms.

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Availability of data and materials As authors, we are increasingly make our research data available and Data will be made available on request.

Declarations

Conflict of interest The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Ethical Statement Authors declare that we have no conflict of interest. We are agreed upon all the Ethical Rules applicable for this journal.

Consent for Publication Authors agreed to publish this work. This declaration ensures that the Publisher has the Author's permission to publish this work.

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