Optical absorption and fluorescent behaviour of titanium ions in silicate glasses

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MS received 23 May 2002; revised 16 December 2002

Abstract. Titanium in normal melting conditions in air atmosphere present as Ti^{4+} ion in basic silicate glasses exhibited an ultra-violet cut-off in silicate glasses, viz. soda-magnesia-silica, soda-magnesia-lime-silica and soda-lime-silica glasses. This indicates that Ti^{4+} ion can be a good replacement for Ce^{4+} ion in producing UV-absorbing silicate glasses for commercial applications. The wavelength maxima at which the infinite absorption takes place in glasses was found to be around 310 nm against Ti-free blank glass in UV-region. The mechanism of electronic transition from O^{2-} ligands to Ti^{4+} ion was suggested as $L \rightarrow M$ charge transfer. The low energy tails of the ultra-violet cut-off were found to obey Urbach's rule in the optical range 360–500 nm. The fluorescence spectra of these glasses were also studied and based on the radiative fluorescent properties it was suggested that the soda-lime-silica glass containing Ti^{4+} ion with greater emission crosssection would emit a better fluorescence than the corresponding soda-magnesia-lime-silica and soda-magnesia-silica glasses. The shift of emission wavelengths maxima towards longer wavelength in titania introduced silicate glasses was observed on replacement of MgO by CaO which may be attributed due to an increase in basicity of the glass system.

Keywords. Optical; fluorescent characteristic; titania; silicate glasses; UV-visible range.

1. Introduction

In normal melting conditions titanium can be present in basic silicate glasses in air as furnace atmosphere in the form of Ti⁴⁺ ion whereas under strongly reducing conditions it can exist as Ti³⁺ ion (Kumar 1959; Bamford 1962, 1977; Bates 1962; Weyl 1967; Lee and Bruckner 1982; Paul 1982; Singh and Singh 2001). Several other systematic investigations have been carried out on optical absorption characteristics of glasses containing transition metal ions as well as their role as redox with particular reference to 3d transition metal ions viz. Cu⁺/Cu²⁺, Fe²⁺/Fe³⁺, Mn²⁺/ $Mn^{3+},\,V^{3+}\!/V^{5+}$ and $Cr^{3+}\!/Cr^{6+}$ pairs, but information on the mechanism of charge transfer process resulting in the absorption bands due to Ti⁴⁺ ion in commercial type of silicate glasses had been lacking over past several years (Johnston 1964; Nath and Douglas 1965; Paul and Douglas 1965; Banerjee and Paul 1974; Singh et al 1978a,b; Duram and Fernandes Navarro 1985; Cable and Ziang 1989a,b; Bae and Weinberg 1991; Singh and Kumar 1992, 1995; Morinaga et al 1994; Singh and Singh 1998, 2000, 2001). However, Singh and Singh (2001) had studied the spectrochemical behaviour of charge transfer bands due to d^0 , d^5 and d^{10} ions in a sodium silicate glass

and reported the intensity of the bands due to Ti⁴⁺ ion as 1.65×10^3 and 1.13×10^3 g mol/l/cm of glass at their wavelengths maxima of 250 nm and 290 nm in the glass, respectively. These authors suggested the mechanism of charge transfer from ligand to central metal ion in the binary sodium silicate glass based on the values of intensity of the bands. Further, they had also reported the appearance of UV-bands in the glass due to Ce⁴⁺, V⁵⁺ and Cu⁺ ions and attributed the bands to ligand to metal (L \rightarrow M) charge transfer in V⁵⁺ ion and Ce⁴⁺ ion and metal to ligand to metal (M \rightarrow L \rightarrow M) cooperative charge transfer in the case of Cu⁺ ions (Cu⁺ \rightarrow O \rightarrow Cu²⁺), respectively in the glass (Singh *et al* 1986; Singh and Singh 2001).

Steele and Douglas (1965) suggested that in silicate glasses on the one hand ferrous ion is responsible for charge transfer band in UV region in between 200 and 230 nm with an intensity of 3×10^3 g mol/l/cm, while on the other hand, in silicate and borate glasses ferric ion has strong charge transfer band with molar extinction coefficient of 7×10^3 g mol/l/cm centred at its wavelength maxima at around 230 nm in ultraviolet region. Nath *et al* (1965) studied the ultraviolet absorption characteristics of hexavalent chromium in alkali silicate glasses as well as aqueous alkaline chromate solution and in the light of identical absorption spectra of glasses and solutions containing chromium, they identified the presence

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of Cr^{6+} ions as chromate complex ions (CrO_4^{-}) . They reported the intensity of the bands due to Cr^{6+} ions as $4 \cdot 2 \times 10^3$ g mol/l/cm in a sodium silicate glass at its wavelength maxima of 370 nm.

In the case of iron present in the silicate glass, the mechanism of charge transfer has been suggested as exchange of electrons in between $M \rightarrow L \rightarrow M$ cooperative charge transfer in the glass (Steele and Douglas 1965; Wood and Remeika 1966). Whereas in the case of sodium silicate glass containing hexavalent chromium the mechanism of charge transfer has been reported as transfer of electron from oxygen ligand to central metal ion as $L \rightarrow M$ charge transfer (Nath *et al* 1965; Singh and Singh 2001). Paul (1982) had also studied the mechanism of charge transfer with mixed cerium-titanium redox in a sodium boroalumino silicate glass and suggested that the broad charge transfer band had arisen at about 345 nm due to cooperative charge transfer within $Ce^{3+} - O - Ti^{4+}$ coloured chromophore. He also observed the optical absorption behaviour of 12Na2O·20B2O3·3Al2O3·65SiO2 glass melted with 1.5 wt.% TiO₂ in the wavelength range 290-450 nm and found that the low energy tail of the UV cutoff for Ti⁴⁺ ion obeyed the Urbach's rule for relation between log (absorbance/cm) vs frequency of radiation in the range of around 22000-27000 cm⁻¹ in the glass. However, in the higher frequency range beyond 27000 cm⁻¹ (370 nm) the deviation of the curve from linearity had indicated its tendency for an approach towards an UVband due to Ti⁴⁺ ion as marked by a sharp cut-off at around 300 nm also in the sodium alumino borosilicate glass even for higher concentration of titania than 0.1% by weight in the glass.

In the light of this the present study was undertaken with a view to investigate the optical absorption characteristics of titanium ions in soda-magnesia-silica, sodalime-silica and soda-magnesia-lime-silica glasses in the UV-visible region as well as to suggest the mechanism of transfer of electrons which result in the UV bands in the aforesaid optical range.

Since the phenomena of fluorescence in the visible region due to a characteristic species occur only when its absorption behaviour is ceased as such, studies on fluorescent behaviour of titanium ions was also undertaken in the proposed glasses here. Although several investigations had been carried out in the past to dictate the excitation and emission properties of rare earth ions in glasses but the studies on optical fluorescent characteristics of transition metal ions in glasses are still very small. However, in order to explore the possibilities for the presence of even traces of titanium in trivalent state due to its redox property in glasses (Johnston 1965), it was also planned to investigate the fluorescence spectra of the proposed silicate glasses doped with titania along with its optical transmission spectra as Ti³⁺ ion had been known to exhibit optical absorption as well as fluorescence characteristics in glasses.

2. Experimental

Silicate glasses having molar compositions: 20Na₂O·10MgO· 70SiO₂, 20Na₂O·10CaO·70SiO₂ and 20Na₂O·4MgO· 6CaO·70SiO₂, were selected as the base glass. For preparing the glass batches, analytical reagent grade 99.9% pure BDH sodium carbonate, calcium carbonate, Thomas and Baker 4MgCO₃·Mg(OH)₂·nH₂O as well as acid washed natural quartz up to less than 0.001% Fe₂O₃ impurities were used. Desired concentration of titanium as 500-1000 ppm TiO_2 was introduced in the glass batches as analytical reagent grade titania. Silicate glasses doped with TiO₂ were melted in a 100 ml capacity platinum-2% rhodium crucible. The crucible containing the glass batch was kept in a Globar rod electric furnace in air atmosphere at 1425°C. The temperature was controlled within $\pm 10^{\circ}$ C with an automatic R-type thermocouple temperature indicator-cum-controller. The glasses were melted for ~ 6 h and then were cast and poured onto an aluminium plate. After cooling they were crushed and remelted in the furnace for another 4 h to ensure homogeneity. They were taken out of the furnace and poured into a rectangular mould on an aluminium plate and annealed at 500°C for 1 h and cooled to room temperature slowly by controlling the rate of cooling. After grinding and polishing the transmission spectra of the glass samples were recorded on a Varian-Cary 2390 recording spectrophotometer in the UV-visible region against a blank glass prepared without titania. The thicknesses of the glass samples were measured and their densities were also determined by Archimedes principle at room temperature. The fluorescence spectra of the glass samples were recorded on a Perkin-Elmer LS-45 fluorescence spectrometer.

3. Results and discussion

Results presented in figure 1 show the optical transmission spectra of titanium ions present in soda–magnesia–silica, soda–lime–silica and soda–magnesia–lime–silica glasses. The relation between log (absorbance/cm) and wave number (cm⁻¹) indicate the verification of Urbach's rule as given in (1) where the low energy tail of the UV cut-off follows a linear behaviour in the optical range of around 20000–28000 cm⁻¹:

$$\log (\text{absorbance/cm}) = A + B\nu, \tag{1}$$

where A and B are constants and v the frequency of radiation. The optical transmission spectra of glasses containing titania show a sharp ultraviolet cut-off at around 310 nm for all these silicate glasses, viz. $Na_2O - MgO - SiO_2$, $Na_2O - CaO - MgO - SiO_2$ and $Na_2O - CaO - SiO_2$, with reference to their blank glass samples which is attributed due to the presence of Ti⁴⁺ ion in the glass. Although the ultraviolet absorption of Ti⁴⁺ ion dependent upon the glass composition originates from different points of ordinate in figure 1 but it shows an UV cut-off at around ~ 310 nm showing infinite ultraviolet absorption at zero transmission. The transmission spectra of glasses were recorded in UV-visible region (200–800 nm) in order to explore the possibilities for the presence of the Ti^{3+} ion but the optical transmission curves do not show sign of minimum transmission (%) due to trivalent titanium in the visible range at any particular wavelength maxima. At this stage it is ensured that Ti^{3+} ion is not at all present in the glass to show any absorption band in the visible region at a I_{max} . So the entire amount of titanium is present in the glass in its tetravalent state.

In order to marking the UV-bands for Ti⁴⁺ ion having high intensities as found earlier to be of the order of around 10³ g-mol/l/cm of a 3Na₂O·7SiO₂ in ultraviolet region (Singh and Singh 2001), the concentration of TiO_2 was kept quite low (0.05-0.10 wt%), but the absorption of the UV-radiation even in thinner glass samples at $I_{\rm max}$ at ~ 310 nm was found to be infinite showing zero transmission in the curve (figure 1). At this stage it is mentioned that Ti⁴⁺ ion would be a good substitute for Ce⁴⁺ ion for producing UV absorbing glasses for commercial applications as titania being cheaper than ceria. The deviation of the curves from linearity indicate that the UVband for Ti⁴⁺ ion is being approached in the glass (figure 2). However, it would be interesting to refer to the work of Steele and Douglas (1965, 1966) who studied using Lambert-Beer's law the effect of absorption of a blank sodium silicate glass contaminated with very small proportion of ferric iron on the UV-absorption characteris-



Figure 1. Optical transmission spectra of titania doped silicate glasses.

tics of the silicate glasses doped with iron in the range of 200-350 nm (figure 3). The authors reported that the background absorption of the blank glass was found to decrease with increasing wavelength of the UV-radiations and showing an almost negligible effect in the range of around 310-330 nm as presented in figure 3. Further, they mentioned that the non-bridging oxygen apart from ferric iron in the glass probably might be contributing to the UV cut-off at around 210 nm causing an exciton charge transfer to the neighbouring positive ion in the glass. The slight levelling over the range of 230-280 nm was probably due to ferric iron impurity. The intensity of the background absorption due to ferric iron impurities was more significant in this range and also greater at around 230 nm in the silicate glass. Whereas it did not affect the iron spectra as Fe³⁺ ion had no absorption in the longer range of UV-radiations (~ 310-330 nm). In view of this it is clearly mentioned herewith that the UV cut-off at ~ 310 nm is definitely due to charge transfer from oxygen ligands to Ti⁴⁺ ion in the present silicate glasses as the effect of the background impurities due to contamination of ferric iron in the glass was almost negligible and insignificant at around this wavelength. Hence the use of the semiconductor grade silica was not so required at this stage irrespective of whether the chemically bonded traces of iron impurities were present in the acid washed quartz.

The comparison of the energy of a redox pair with another has been already observed and practically become a productive process with reference to mixed redox interaction in glasses at high temperature like redox titrations in aqueous acidic or alkaline solutions in air atmosphere at room temperature. Although the melting conditions in glasses such as high temperature and base glass compositions are different for redox reactions from aqueous media at room temperature in atmospheric



Figure 2. Verification of Urbach's rule by relation between log (absorbance/cm) and frequency of UV-radiation for Ti^{4+} ion in silicate glasses.

air. The production of ferro-manganese amber glass, physico-chemical decolourization of intense ferrous blue colour by manganic ion, preparation of heat absorbing glass by thermo reduction of ferric iron to ferrous ion with stannous tin, development of emerald green colour due to trivalent chromium by reducing Cr^{6+} ion with Sn^{2+} ion and chemical decolorization of ferrous blue colour by oxidizing it into ferric straw yellow with As^{5+} ion in glasses as well as preparation of $Mn^{3+} + Cr^{3+}$ black colour are some of the common examples of redox reactions where their redox potentials have played a vital role as per the following reactions in glass melts.

$$\operatorname{Fe}^{2+} + \operatorname{Mn}^{3+} \rightleftharpoons \operatorname{Fe}^{3+} + \operatorname{Mn}^{2+},$$
 (2)

$$2\mathrm{Fe}^{3+} + \mathrm{Sn}^{2+} \rightleftharpoons 2\mathrm{Fe}^{2+} + \mathrm{Sn}^{4+}, \qquad (3)$$

$$2Cr^{6+} + 3Sn^{2+} \rightleftharpoons 2Cr^{3+} + 3Sn^{4+}, \qquad (4)$$

$$2Fe^{2+} + As^{5+} \rightleftharpoons 2Fe^{3+} + As^{3+}, \qquad (5)$$

$$3\mathrm{Mn}^{2+} + \mathrm{Cr}^{6+} \rightleftharpoons 3\mathrm{Mn}^{3+} + \mathrm{Cr}^{3+}.$$
 (6)

The development of photosensitive copper ruby, silver yellow and gold ruby colloidal colours in glasses in presence of CeO₂ as sensitizer in the glass melts where the Ce³⁺/Ce⁴⁺ redox species played an important part, had



Figure 3. Relation between background absorption vs wavelength for a $30 \text{ Na}_2\text{O} \cdot 70 \text{ SiO}_2$ (A) blank glass and (B) glass doped with iron (Steele and Douglas 1966).

been done by mixed $(Cu^+/Cu) + (Ce^{3+}/Ce^{4+})$; $(Ag^+/Ag) + (Ce^{3+}/Ce^{4+})$ and $(Au^+/Au) + (Ce^{3+}/Ce^{4+})$ redox interactions (Dwivedi and Nath 1977, 1978, 1980). So was the case of Ti^{3+}/Ti^{4+} and Ce^{3+}/Ce^{4+} redox interactions as per the reaction

$$Ce^{3+} + Ti^{4+} \rightleftharpoons Ce^{4+} + Ti^{3+}, \qquad (7)$$

for preparation of cerium–titanium intense yellow colour in glass. Kuhl *et al* (1938) tabulated redox species commonly used in the glass in order of increasing their redox potentials as $Cr^{3+}/Cr^{6+} > Mn^{2+}/Mn^{3+} > Ce^{3+}/Ce^{4+} > Cu/$ $Cu^+ > As^{3+}/As^{5+} > Fe^{2+}/Fe^{3+} > Sn^2/Sn^{4+} > Ti^{3+}/Ti^{4+}$ which had dictated that any reduced state of a redox having lower potential would be oxidized by the oxidized state of another with higher potential which was placed above it in the electrochemical series. Similar table prepared on the basis of standard thermodynamic data for pure oxides was also reported by Tress (1960).

In silicate glasses because of the random network model of [SiO₄] tetrahedra Ti⁴⁺ ion even having greater ionic radii could form tetrahedra and enter into the network resulting in a random packing of tetrahedra. The analysis of an infra-red spectra of a titania silica glass prepared with 10 wt% TiO2 by Smith and Condrate (1975) had shown earlier that the IR band at around 735 cm^{-1} out of the four bands marked at 940, 735, 360 and 240 cm⁻¹ could be assigned due to vibrational mode of the tetrahedral [TiO₄] group in silica glass. The tetrahedral coordination of Ti⁴⁺ ion deduced as above was in accordance with the previous findings of Evans (1970) that Ti⁴⁺ ion was present as four-fold coordinated Si⁴⁺ ion in silicate glasses. There was no fundamental mode with characteristic frequencies to indicate octahedral coordination of titanium in the glass.

Since the glasses melted with titania had contained only tetravalent titanium as an ultraviolet absorbing species at ~ 310 nm, in the light of the present justification, therefore, the probable mechanism of charge transfer is suggested herewith as the transfer of an electron from oxygen (O^{2-}) ligands to fill up the vacant d^0 -orbital of Ti^{4+} ion (L \rightarrow M) present in the glass as $[TiO_4]^{4-}$ complex ion which is based on the facts. The presence of similar charge transfer species such as $[CrO_4]^{2-}$, $[CeO_4]^{4-}$ and $\left[VO_4 \right]^{3-}$ had already been reported in the silicate glass by previous workers (Nath et al 1965; Singh et al 1986; Singh and Singh 2001). Although the $I_{\rm max}$ at around 310 nm is the same in the curve (figure 1) but the intensity of Ti⁴⁺ bands is expected to be dependent upon the glass composition and mixed effect of these alkaline earth oxides as well at the wavelength maxima. The applicability of Urbach's rule as tested and verified for Ti⁴⁺ ion in glasses (figure 2) follows similar patterns as it was found in the case of other charge transfer species like Cu⁺, V⁵⁺, Ce⁴⁺, Fe³⁺ and Cr⁶⁺ ions in glasses (Nath et al 1965; Singh et al 1986; Singh and Singh 2001). The present results with lower concentration of titania are in conformity to silicate glass containing higher concentration of 1.5 wt% TiO₂ as referred to in figure 4 after Paul (1982). They are also in close analogy to the results reported in the fluorophosphate glass melted with titanium for an UV-band at ~ 300 nm due to Ti⁴⁺ – F⁻charge transfer process (Fuxi 1992).

The ability to generate spectrophotometrically active reduced species under non-equilibrium condition is of vital importance for determining the coordinations and symmetries of the newly formed ions such as Ti^{3+} in glasses. The formation of Ti^{3+} ions was observed earlier as a product of irradiation of Ti^{4+} ion present in the glass and presumably located in the same oxygen environment possessed by the original Ti^{4+} ion before the occurrence of electron capture (Smith and Cohen 1963; Wong and Angell 1976). The trivalent titanium being $3d^1$ ion should exhibit a ligand field spectrum the form of which would identify directly the oxygen coordination, but Ti^{3+} ion was not observed presently in glasses at all.

The excitation and emission spectra of Ti⁴⁺ ions present in the glasses have been given in figures 5 and 6, respectively. The relative intensity of emitted radiations was found to be of the order of around $1{\cdot}02 \rightarrow$ 1.03×10^3 . The excitation spectra in the present ternary silicate glasses as in figure 5 show that an excitation at around 310 nm which corresponds to the $Ti^{4+}(3d^0)$ $\leftarrow O^{2-}(2s^2 3p^6)$ charge transfer band might impart an intense fluorescence radiative emission transition from an excited energy level to the ground level due to $M \rightarrow L$ charge transfer. This process of transition from excited Ti⁴⁺ ion to O²⁻ ligands exhibits the fluorescent behaviour of Ti⁴⁺ ion in glasses which might be attributed due to energy transfer by ion-phonon coupling. Similar observations in form of $Ti^{4+} - F^- - Ti^{3+}$ charge transfer were also reported earlier in reduced fluorophosphate glass which contained both the Ti^{3+} and Ti^{4+} ions (Fuxi 1992). Park and Webb (1972) as well as Annapurna et al (2000)



Figure 4. Optical absorption characteristics of a sodium boro–alumino silicate glass containing 1.5 wt\% TiO_2 (Paul 1982).

studied the UV-excitation of Cu⁺ ion from $3d^{10}$ configuration to $3d^9 4s^1$ and reported the fluorescent transition as the phonon assisted greenish blue emission the broader visible bands for which at around 500 nm were attributed due to spin forbidden transition from excited ³Eg $(3d^9 4s^1) \rightarrow {}^{1}Ag (3d^{10})$ ground level in phosphate and zinc borosilicate glasses.

When the light radiation interacts with an electronic system in an ionic state the radiative absorption or emission transition between two energy levels y_a and y_b is induced. The spontaneous emission transition probability, $A(y_b \rightarrow y_a)$ from the excited ion level y_b to a level y_a can be approximately given by the following equation (Paul 1982; Fuxi 1992)

$$A(\mathbf{y}_b \to \mathbf{y}_a) = \frac{8\mathbf{n}^2 \times 10^6}{e^2}, \qquad (8)$$

where **n** is the frequency of emission transition and *e* the charge of the electron. The stimulated emission cross-section for each emission band (s_{se}) in the present silicate glasses for the same excitation wavelength was calculated using the equation (Fuxi 1992)

$$\boldsymbol{s}_{se} = \left[\frac{\boldsymbol{l}^4}{8\boldsymbol{p}cn^2\Delta\boldsymbol{l}}\right] \mathbf{A} \left(\boldsymbol{y}_b \to \boldsymbol{y}_a\right),\tag{9}$$

where DI is the effective half band width and I the emission band position as well as n is the refractive index of



Figure 5. Excitation spectra of silicate glasses doped with TiO_2 .

Physical properties	$\begin{array}{c} 20Na_2O{\cdot}10MgO{\cdot}\\ 70SiO_2 \end{array}$	$\begin{array}{c} 20Na_2O{\cdot}4MgO{\cdot}\\ 6CaO{\cdot}70SiO_2 \end{array}$	$\begin{array}{c} 20Na_2O{\cdot}10CaO{\cdot}\\ 70SiO_2 \end{array}$
Concentration of TiO ₂ (wt.%)	0.100	0.100	0.100
Refractive index (<i>n</i>)	1.511	1.520	1.532
Density $(d, mg/cm^3)$	2.474	2.478	2.488
Reflection loss (<i>R</i> %) [$R = (n - 1/n + 1)^2$]	4.120	4.250	4.410
Excitation wavelength, I_{ex} (nm)	310	310	310
Emission peak position, \hat{I} (nm)	326	418	425
Emission band width, Dl (nm)	10.0	12.5	12.5
Stimulated emission cross-section,			
$\boldsymbol{s}_{se} \times 10^{-32} \text{ (cm}^2)$	5.30	6.91	7.03

Table 1. Physical properties and stimulated emission cross sections (s_{se}) of Ti⁴⁺-doped alkali–alkaline earth–silicate glasses.



Figure 6. Emission spectra of silicate glasses doped with TiO_2 .

the glass. The values of stimulated emission cross-section $(s_{se} cm^2)$ presented in table 1 was found to increase from 5.30×10^{32} to 7.03×10^{32} with increasing basicity or oxygen ion activity of the ternary silicate glass on replacing MgO by CaO. The other physical parameters have also been given in table 1. The order of magnitude of these results follow the similar pattern reported earlier in ternary silicate, phosphate and alkali-magnesia-alumino silicate glasses containing rare earth Nd³⁺ and Sm³⁺ ions (Fuxi 1992; Annapurna et al 1999). Further, it is concluded that a soda-lime-silica glass containing Ti⁴⁺ ion with higher s_{se} (cm²) value would emit a better radiative fluorescence than the corresponding soda-magnesialime-silica and soda-magnesia-silica glasses with their lower values. The present results presented in table 1 are also in conformity with the results obtained by Sreenivasulu and Rao (2001) in fluoroborophosphate glasses

prepared with Pr^{3+} ion who mentioned that an increase in the value of emission cross-section would lead to a better fluorescence emissivity in glasses at room temperature.

In oxide glasses the shifting of emission bands obtained for active sites is not only dependent upon temperature but also it depends upon the glass composition which can be marked from the peak position and the band width as reported previously for other ions and evident from figure 6 for Ti⁴⁺ ion at room temperature. In case of glasses particularly at least at room temperature the coupling between ions and phonons becomes stronger for cross relaxation and the interaction between them would result in emission by energy transfer. The same may not take place probably at lower temperature. The nature and intensity of the fluorescent Ti⁴⁺ centre in the glass is its specific characteristics like other species which depends upon its band position, temperature and the glass composition, its emission spectra would not be generally influenced due to the presence of background small iron impurities in the glass till it is overlapped by them which was not observed during present investigation.

Acknowledgements

The authors gratefully dedicate this work to the Department of Ceramic Engineering, Institute of Technology, and they are also thankful to the Director, Institute of Technology, Banaras Hindu University, Varanasi, for providing the necessary facilities for the present work.

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