

3.1.1 Optical basicity

The optical basicity of an oxide glasses can be conveniently measured in terms of the ability of the glass to donate a negative charge to an acidic probe ion [7-10]. Duffer and Ingram [11] reported that the optical basicity can be predicted from the composition of the glass and the basicity moderating parameters of various cations present. The theoretical values of optical basicity (Λ_{th}) of glasses can be estimated using the formula,

$$(\Lambda_{th}) = \sum_{i=1}^n \frac{Z_i r_i}{2\gamma_i} \quad (1)$$

where 'n' is the total number of cations present. Z_i is the oxidation number of the i^{th} cation to the number of oxides present and ' γ_i ' is the basicity moderating parameter of the i^{th} cation. The basicity moderating parameter ' γ_i ' can be calculated from the following equation

$$\gamma_i = 1.36(x_i - 0.26) \quad (2)$$

where x_i is the Pauling electro negativity of the cation [12]. The theoretical optical basicity values computed of the glasses in the present study are given in **Table.2**.

Table 2

Physical parameters of Chromium ions doped in SLCB glasses.

Physical Parameters (Units)	Glass System			
	SLCB ₀	SLCB ₁	SLCB ₂	SLCB ₃
Average molecular weight(M)(g/mol)	67.69	64.04	67.73	71.42
Density (gr/cc)	2.65	2.54	2.64	2.77
Transition metal ion (N_i) Concentration ($10^{19}i$ ions/cc) (± 0.005)	2.38	2.36	2.34
Inter ionic distance (r_i)(\AA)(± 0.005)	34.71	34.90	34.94
Polaron radius (r_p)(\AA)(± 0.005)	13.98	14.06	14.05
Optical basicity (Λ_{th})	0.43	0.43	0.43	0.42

It is observed that the Λ_{th} values gradually increases from SLCB₁ to SLCB₃. It is observed that the optical basicity value is minimum for a glass with 15% SrO, 5% Li₂O and maximum for 10% SrO, 10% Li₂O contents.

3.2 Powder XRD

The powder XRD diffractograms of undoped and chromium ions doped glasses are shown in **Fig.1**. This reveals the amorphous nature the synthesized glasses.

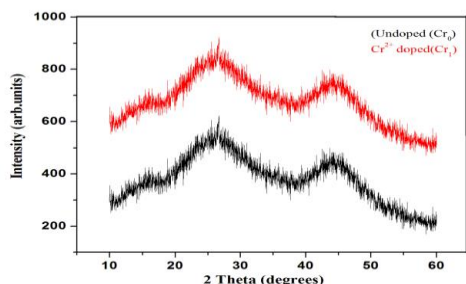


Figure 1: X-ray diffractograms of undoped and Chromium ions doped SLCB glasses.

3.3 FT-IR Studies

The IR spectra of SLCB glasses are shown in **Fig.2**.

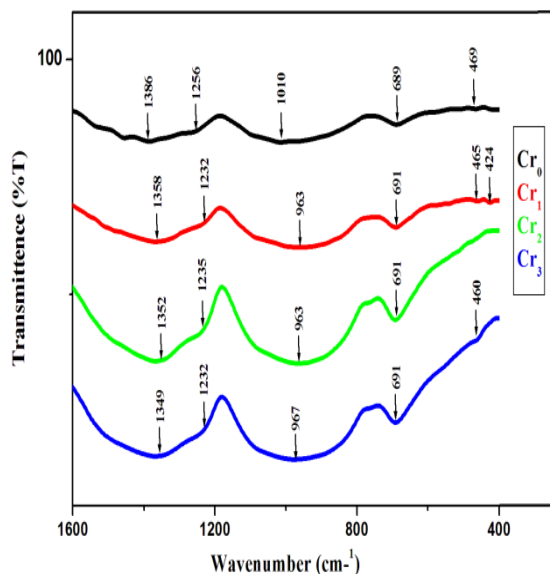


Figure 2: FT-IR spectrum of Cr³⁺ ions doped SLCB glasses.

The assignments of various bands are given in **Table 3**. The IR spectra can be alienated into three regions such as 600–800 cm⁻¹ which are due to B-O-B bending vibrations of various borate units. The second region 800 – 1200 cm⁻¹ is attributed to B-O symmetric vibrations of tetrahedral BO₄ units. The third region 1200–1600 cm⁻¹ is related to B-O stretching vibrations of BO₃ units [13-15]. The spectra contain three broad and intense absorption bands centered around 1386 cm⁻¹, 1010 cm⁻¹ and 686 cm⁻¹, with some shoulders and feeble bands are also observed in the undoped (Cr₀) sample. The broad band around 1386 cm⁻¹ and the and the shoulder around 1256 cm⁻¹ are due to symmetric stretching vibrations of B-O bonds in BO₃ units varied from types of pyro, ortho borate groups respectively. The intensity of the broad band and the shoulder increases with the increase of SrO content. The broad band around 1010 cm⁻¹ is due to boroxil rings [16]. This band is the characteristic of glass containing high B₂O₃ content. The intensity has increased with the increase of SrO content. The absorption band at 689 cm⁻¹ is due to B- O-B deformation vibrational mode [17, 18]. The intensity of the band increases with increase in SrO content. The shoulder at 469 cm⁻¹ is due to Ca-O₆ stretching mode of Borate units.

Table 3

Assignment of the bands in the infrared spectra of Chromium ions doped in SLCB glasses.

Cr ₀	Cr ₁	Cr ₂	Cr ₃	Assignment
-	424	-	-	Ca-O stretching mode of BO ₃ units
469	465	-	460	Ca-O ₆ stretching mode of BO ₃ units
689	691	691	691	B-O-B bending
1010	963	963	987	B-O stretching Vibrations
1256	1232	1235	1232	Stretching vibrations B-O

				bonds in BO ₃ Units from pyro-ortho borate groups
1386	1358	1352	1349	Stretching vibrations of B (III)-O-B (IV)- units

The intensity at this peak is maximum for 15 mol% of SrO and 5 mol% of Li₂O [19].

4. Conclusion

- 1) The density values are increasing with the increase of SrO mol%.
- 2) The optical basicity value is maximum when the mol% of SrO and Li₂O are equal i.e. 10%.
- 3) XRD has confirmed the amorphous nature of the
- 4) Present glasses under study.
- 5) The FT-IR spectra have shown the characteristic

Spectra of Borate glasses.

5. Acknowledgements

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Author Profile



Dr. Sandhya Cole M.Sc., M. Phil, Ph.D., She is working as Assistant professor in the department of Physics since 2006, Acharya Nagarjuna University, Nagarjuna Nagar, Guntur, A.P., India. Her research fields are Solid state Spectroscopy, Nanotechnology, Material Science, and Glass Materials. She is a life member of several scientific bodies like Indian Association of Physics Teachers, Indian Physics Association and associated fellow of A.P. Akademy of sciences, A.P, 2014. She has successfully completed UGC major research project. She also received best research paper award in sciences-2014. At present 15 research scholars including one CSIR-UGC (SRF) fellow and two BSR research fellows are working for their research degree under her guidance.



Mr. M. Ratna Raju, working as Lecturer in Physics, Andhra Christian College, Guntur, A.P, India. He is having 19 years of teaching experience. He is currently associated with research for his Ph.D., work under UGC (FIP) program under the guidance of Dr. Sandhya Cole.



M. Subba Rao had received his M.Sc., degree in Physics with specialization in condensed Matter physics from Department of Physics, Acharya

Nagarjuna University, Nagarjuna Nagar, Andhra Pradesh, India. He is currently Ph.D., Student in the field of Solid State Spectroscopy and Nanomaterials. He is CSIR-UGC (SRF) Fellow and he had attended International and national Conferences, workshops and International Symposium. Under the guidance of Dr. Sandhya Cole.

