Chapter – 2

FTIR Spectra of magnetic charge transfer Complexes of 2, 2, 6, 6 - tetramethyl piporidinyloxy free radical.

[Vishal R. Jain/Ph.D.Thesis (Physics)/S.P.U.-2012]
1. Introduction:

TEMPO (2, 2, 6, 6-tetramethyl piporidinyloxy) radical and its substituted derivatives are found to form charge transfer complexes (CTCS) with organic acceptors. These CTCS are magnetic, either ferromagnetic or anti-ferromagnetic, at very low temperatures (1–7). The CTCS of TEMPO radical with TCNQ, TCNE, DDQ, chloranil and iodine are prepared in the present work and studied with FTIR spectroscopy. TEMPO acts as a donor because of electron-accepting four methyl groups.

2. Experimental:

TEMPO free radical was obtained in powder form from Sigma-Aldrich chemical company, USA. It was mixed with acceptors such as TCNQ (7, 7, 8, 8-tetracyano-p-quinop-dimethane), TCNE (tetracyano-p-ethylene), DDQ (2, 3-dichloro-5, 6-dicyano-p-benzoquinone), Chloranil and iodine also obtained as analytical reagents. After mixing, the mixtures were grinded in an agate mortar with a pastel till the colours changed. These homogeneously fine powders were mixed and grinded with spectrograde dry (anhydrous) KBr powder. Semitransparent pallets were prepared using a manually operated compressing machine and a die. The circular discs prepared in this manner were placed in the dark chamber of an FTIR spectrophotometer.
The spectra in the range 400 – 4000 cm\(^{-1}\) were recorded using a GXFTIR single beam spectrophotometer manufactured by Perkin Elmer Comnpy, USA. It was having a resolution of 0.15 cm\(^{-1}\), a scan range of 15,000 – 30 cm\(^{-1}\), a scan time of 20 scan/sec, an OPD velocity of 0.20 cm/sec and MIRTGS and FIRTGS detectors. A beam splitter of opt KBr type was used having a range of 7800 – 370 cm\(^{-1}\). The spectra were recorded in purge mode.

3. Results and Discussion:

The molecular structure of TEMPO free radical is shown (figure 1). The Fourier – transformed infrared (FTIR) spectrum of TEMPO free radical is shown (figure 2). TEMPO radical itself is an insulator having a transmitting range in IR spectrum. An asymmetric gaussian in the range 1000 cm\(^{-1}\) – 1700 cm\(^{-1}\) is observed in this spectrum. It is related with optical properties in small polarons model. There is hopping of polarons giving rise to this gaussian band of about 10% absorption below a transmitting range. Also there is a semicircular distribution, marked as SCD, in the low frequency range and centered about 700 cm\(^{-1}\). This distribution is related with random orientations of TEMPO molecules in three dimensions in a crystal. The disorder is isotropic giving rise to a semicircular distribution.

The FTIR spectrum of TEMPO – TCNQ charge transfer complex is shown (figure 3). This spectrum covers a range of nature of transition for absorption function and a range of symmetric gaussian
distribution, the latter being marked as $G$. The absorption $A$ is fitted as $A_h = B (h\gamma - E_g)$ - a nature of allowed indirect or a forbidden direct transition in two-dimensional conductors (figure 4a). Thus TEMPO – TCNQ is a two-dimensional semiconductor having $E_g = 0.325$ eV. It is a non-universal Hubbard semiconductor. The gaussian distribution is also fitted by plotting $\ln A$ vs $(K-K_0)^2$ (figure 4b). The symmetric gaussian distribution is related with electronic delocalization.

The FTIR spectrum of TEMPO – TCNQ is shown (figure 5). The range of nature of transition is marked as NT and asymmetric gaussian background is marked as AG. The transition is found to be forbidden indirect type by fitting $A_h = B (h\gamma - E_g)^3$ (figure 6a). The asymmetric gaussian is fitted by plotting $(\ln A)^{1/2}$ vs $K-K_0$ on one side of the gaussian band (figure 6b). Both sides of gaussian have different full – widths at half – maxima. The slopes of $(\ln A)^{1/2}$ vs $K-K_0$ lines are different for $K > K_0$ and $K < K_0$. The asymmetric gaussian band can be explained with optical properties in small polaron model (12). Band gap is about 0.272 eV.

The FTIR spectrum of TEMPO – DDQ is shown (figure 7). The spectrum reveals a range of nature of transition and two gaussian bands. The nature of transition shows a forbidden direct type transition fitting $A_h = B(h\gamma - E_g)^3$ with $E_g = 0.225$ eV. It is a universal Peierls gap. Thus an asymmetric molecule like DDQ rather than symmetric TCNQ and TCNE shows a Peierls gap. Both two -
dimensional character with electronic delocalization. The nature of transition and one of the gaussian bands are analyzed (figure 8a and 8b).

The FTIR spectrum of TEMPO – chloranil is shown (figure 9). It shows a nature of transition and oscillations in the density of states. The transition is found to be allowed direct type by fitting $A \hbar \gamma = B (h \nu - E_g)^{1/2}$ with $E_g = 0.25 \text{ eV}$ – again a Hubbard gaps. The direct transition is usually found for small donor and small acceptor in a charge transfer complex (figure 10). Below 1800 cm$^{-1}$, three repeated structures of square – root singularities along a homomolecular sub lattice of chloranil stacks are found. This is observed usually in chloranil complexes.

Finally, the FTIR spectrum of TEMPO – I$_2$ is shown (figure 11). The range of nature of transition and asymmetric gaussian band are fitted (figure 12a and 12b). The transition is allowed direct type by fitting $A \hbar \gamma = B (h \gamma - E_g)^{1/2}$ with $E_g = 0.225 \text{ eV}$ again attributed to a universal Peierls gap. Thus asymmetric I$_3^-$ ion also leads to a Peierls gap. Asymmetric gaussian is fitted by plotting $(\ln A)^{1/2}$ vs $K-K_0$, on one side of the gaussian band. Other side shows a different slope giving a different full width at half – maximum. This can be explained with optical properties in small polaron model (12). The CTCS of TEMPO radical are found to be ferromagnetic of anti – ferromagnetic at very low temperature (11). However, semiconducting nature having either a Hubbard gap or a Peierls gap
found in the present work shows that these CTCS are diamagnetic at room temperature. These small band gap semiconductors cannot undergo a transition to a magnetic state without being converted into a Pauli paramagnetic phase. Thus a semiconductor to metal type transition should occur at intermediate temperature. This can happen due to thermal contraction of the semiconductors upon cooling. The intermolecular distance decreases due to thermal contraction at low temperature which can lead to more orbital overlap. Thus p – p overlap leads to a transition to metallic state which has Pauli paramagnetism. At still low temperatures these CTCS can become magnetic by increase in exchange interaction. Thermal contraction leads to more exchange interaction at low temperature.

4. Conclusions:

The symmetric acceptor molecules such as TCNQ, TCNE and Chloranil gives rise to Hubbard semiconducting nature of CTCS of TEMPO radical while asymmetric molecules like DDQ and I$_3^-$ ion give rise to Peierls semiconducting nature. Symmetric gaussians and asymmetric gaussans are related with electronic delocalization and small Polaron hopping. Direct and indirect transitions are related with one or two – dimensionalities of CTCS of TEMPO radical. Magnetic nature at low temperature can be explained with a transition to a Pauli paramagnrtic phase with orbital overlap due to thermal contraction.

[Vishal R. Jain/Ph.D.Thesis (Physics)/S.P.U.-2012]
Figure 1 Molecular structure of TEMPO and other organic accepters
Chapter – 2

Figure 2 FTIR spectra of TEMPO

Figure 3 FTIR spectra of TEMPO – TCNQ

[Vishal R. Jain/Ph.D.Thesis (Physics)/S.P.U.-2012]
Figure 4a NT of TEMPO –TCNQ

Figure 4b Symmetric Gaussian of TEMPO –TCNQ
Figure 5 FTIR spectrum of TEMPO – TCNE
Figure 6a NT of TEMPO – TCNE

Figure 6b Asymmetric Gaussian of TEMPO – TCNE

[Vishal R. Jain/Ph.D.Thesis (Physics)/S.P.U.-2012]
Figure 7 FTIR spectrum of TEMPO – TCNE

Figure 8a NT of TEMPO – DDQ

[Vishal R. Jain/Ph.D.Thesis (Physics)/S.P.U.-2012]
Figure 8b  Symmetric Gaussian of TEMPO – DDQ
Figure 9 FTIR spectrum of TEMPO – Chloranil

Figure 10 NT of TEMPO – Chloranil

[Vishal R. Jain/Ph.D.Thesis (Physics)/S.P.U.-2012]
Figure 11 FTIR spectrum of TEMPO – Iodine

Figure 12a NT of TEMPO – IODINE

[Vishal R. Jain/Ph.D.Thesis (Physics)/S.P.U.-2012]
Figure 12b  Asymmetric Gaussian of TEMPO – IODINE
References


[Vishal R. Jain/Ph.D.Thesis (Physics)/S.P.U.-2012]