In this study, the effect of a liquid stabilizer based on organotins on PVC heat stability is investigated. Control and films with tin soap heat stabilizer Sn500K were prepared by solvent casting and they were heated at 160 and 180°C for different time periods. X-ray fluorescence spectroscopy analysis showed that tin was present in Sn500K and it was lead-free. DSC analysis showed that glass-transition temperature of films with Sn500K increased from 67 to 75–85°C by heating for 15 min at 180°C. IR spectroscopy indicated that concentration of carboxylate groups of Sn500K decreased with heating. Films with Sn500K did not degrade for short heating periods; however, degradation occurred in long heating times. Low concentration of Sn500K was recommended, since at high levels incompatibility with PVC and acceleration of degradation was observed. In the high temperature range, control and stabilized samples degraded in the same manner in two steps. © 2002 John Wiley & Sons, Inc. Adv Polym Techn 21: 65–73, 2002; DOI 10.1002/adv.10008

KEY WORDS: poly(vinylchloride), stabilization

Introduction

PVC degrades at operating temperatures by the removal of hydrogen chloride. During this degradation, color changes from yellow to orange,
brown, and black according to the double bond formation. Discoloration starts when the mass loss reaches to 0.1%. Degradation occurs, as shown in Eq. (1), by the formation of conjugated double bonds.

\[ \text{CH}_2=\text{CH}_2 \longrightarrow \text{CH}==\text{CH}_2 \]

or in a shorter notation

\[ \text{PVC} \longrightarrow \text{dePVC} + n\text{HCl} \]

where dePVC is dehydrochlorinated PVC which has conjugated double bonds.

Assuming the reaction as first-order with respect to potential double bonds to be formed in PVC, rate of double bond formation

\[ \frac{dC}{dt} = -kC, \]

where \( C \) represents concentration of potential double bonds, i.e. mol dm\(^{-3}\) HCl in PVC, and \( k \) is the first-order reaction rate constant in min\(^{-1}\).

Taking the integral between limits \( C = C_0 \) at \( t = 0 \) and \( C = C_t \) at \( t = t \), we have

\[ \ln\left(\frac{C_0}{C_t}\right) = kt \] (2)

Thus, conversions at any time can be found by this integrated rate equation.

Knumann and Bockhorne\(^2\) compared PVC and other polymers by TGA. They investigated the kinetics of pyrolysis of PVC and found a dehydrochlorination reaction order to be 1.5. Gupta and Viswanath\(^3\) studied the role of metal oxides in PVC degradation by TGA and proposed that the reaction was in two steps, and the rate-controlling step was the formation of metal chloride. Wu et al.\(^4\) studied the pyrolysis of PVC in the temperature range of 400–500 K by TGA. The entire process of pyrolysis was in two stages; thermal dehydrochlorination and the breakdown of the intermediate products produced after the dehydrochlorination stage. The pyrolysis behavior of PVC at high conversion was explained by a competitive reaction model with the production of volatiles and char.\(^5\)

According to the Frye–Horst mechanism,\(^6\) dehydrochlorination of PVC is prevented by metal soaps, with the substitution of labile chlorine atoms of PVC by carboxylate groups of the metal-soap-forming ester groups.

Garrigues et al.\(^7\) compared the thermal dehydrochlorination of PVC in the presence of dialkyl tin maleates and laurates in 1,2,4-trichlorobenzene solution by IR analysis, and they proved the superiority of tin maleates over tin laurates in the substitution reaction. Their attempts to trace intermediate monochlorotin derivatives in the case of maleates by polarography and Mössbauer spectroscopy were not conclusive. Rujian et al.\(^8\) reported that dibutyltin laurate (DBTDL) prevented the formation of longer polyene lengths, and coloration of PVC was retarded effectively in this way. DBTDL cannot prevent the formation of shorter polyene length. Thermal dehydrochlorination still occurred but its rate was decreased and longer polyenes were formed.\(^8\) There was no change in color of the PVC films with a commercial organo tin stabilizer with trade name Sn500K at 160°C for 105 min.\(^9\) Sterically hindered phenolic antioxidants were used together for long-time stabilization of PVC.\(^10\)

PVC heat stabilizers are sold on the market under trade names and no information about their exact composition are given by their producers. Lead-free stabilizers gained importance in recent years because of the harmful effects of lead on health. Thermal stabilization of the PVC under this study by using lead-free stabilizers, epoxidized soybean oil, and zinc stearate was investigated before.\(^11\) Low transparency, good thermal stability at 140°C, but accelerated degradation at higher temperature and at long heating times because of accumulation of ZnCl\(_2\) acting as a catalyst for dehydrochlorination were observed. The solid state of zinc stearate in PVC films at the temperatures studied prevented formation of a homogeneous mixture at the molecular level, causing inefficient thermal stabilization. Thus, a liquid-lead-free heat stabilizer having an organotin compound with trade name Sn500K was chosen as a representative example to show how a heat stabilizer can be characterized both in composition and in efficiency. Higher thermal stabilization efficiency was expected since better mixing of the liquid stabilizer with PVC would be obtained.

**Experimental**

In this study, emulsion PVC (PETVINYL P38/74 from Petkim, Turkey with number average molecular weight 50,000) was used. PVC was produced
by spray-drying of the PVC latex from the emulsion polymerization, and it contained all the emulsifying ingredients. Liquid organotin stabilizer (Sn500K from Farstab stabilizer company, Turkey) and tetrahydrofuran (THF), C4H8O, stabilized with antioxidant and butylated hydroxytoluene (BHT) (from Merck) were used in the film preparation, together with PVC.

The organotin stabilizer was analyzed by x-ray fluorescence analysis using the Jeol X-ray Fluorescence Spectrometer. The ash content was also determined after carbonizing in air and heating at 900°C till constant weight.

In the studied samples, 2.5 and 5% Sn500K were added to PVC. Sample preparation was done using solvent casting method. After mixing 6 g of PVC with 0.15 and 0.30 g of Sn500K in 125 cm³ of THF, they were allotted to 10 petri dishes. THF has high vapor pressure at 25°C and it boils at 65–66°C.12 After allowing for evaporation of THF for one day, PVC film samples with tin stabilizers and BHT were obtained. All films contained 0.49% antioxidant, BHT introduced with THF, and a small amount of THF. Since BHT melts at 69–70°C and boils at 265°C, it was present in PVC films at 25°C in solid state.12 Films were exposed to heat by static oven method at 160 and 180°C for 105 min.

A Shimadzu IR Spectrophotometer and a Jasco 750 UV Spectrophotometer were used to obtain transmission IR and UV spectra of the films. IR spectrum of a liquid film of Sn500K on KBr disc was also taken. A Setaram 92 DSC Colorimeter and a DuPont TGA Analyzer were used for thermal analysis. The samples were heated at 10°C min, heating rate in the ranges of 25–200°C and 30–500°C for DSC and TGA respectively.

The yellowness index of the films was determined using X–Y–Z parameters of the films placed on MgO white standard of Muruyama Color Machine.

Results

Sn500K CHARACTERIZATION

By x-ray fluorescence spectroscopic analysis, tin was detected in Sn500K. There was a reflection showing the presence of tin at 2θ = 14.04 with LiF crystal. No Pb peak was found at 2θ = 33.93 with LiF crystal.13,14 Thus, Sn500K was a stabilizer containing tin without lead. It contained 8.1% tin as indicated by its ash content.

The IR spectrum of Sn500K in Fig. 1 indicated the presence of C=O groups at 1720 cm⁻¹, double bonds at 1600 and 1640 cm⁻¹, and COO⁻ groups at 1540 cm⁻¹. Although the spectrum of Sn500K was very similar to that of dibutyltinbis(isooctyl maleate) (DBTIOM) given by Wypich15, Sn500K contained a smaller amount of tin, 8.1% as compared to the 11.5% of butyltinbis(isooctyl maleate) had. Thus, there should also be a diluting solvent in Sn500K.

EFFECT OF Sn500K ON PVC DEGRADATION

Yellowness Index and Transparency

The films having Sn500K had lower yellowness index than that of the control film heated at different periods at 160 and 180°C, as seen in Fig. 2. The higher the Sn500K concentration and the heating temperature were, the higher was the yellowness index. The color of the film with 2.5% Sn500K had a very small yellowness index up to 30 min time both at 160 and 180°C, indicating that a safe induction period for dehydrochlorination was present. The heat stability temperature range of the PVC under study was higher for Sn500K than for zinc stearate and epoxidized soybean oil stabilizers.11

While the control and 2.5% Sn500K-containing films were transparent, films with 5.0% Sn500K were opaque white, indicating limited solubility of Sn500K in PVC.

DSC Analysis

From DSC curves seen in Fig. 3, the glass-transition temperature (Tg) of unheated PVC was
FIGURE 2. Yellowness index of films versus heating time.

67°C and the glass-transition temperature of PVC heated at 180°C for 60 min was 92°C. By heating at 180°C for 60 min, the THF acting as a plasticizer was removed from the films and the glass-transition temperature was increased. As seen in DSC curves of films with Sn500K, the glass-transition temperature for unheated film increased from 67 to 75–85°C by heating the film at 180°C for 105 min (Fig. 4). A small endotherm observed near the glass-transition temperature could be attributed to the heat of evaporation of THF from the films. Sn500K also had a plasticizing action on PVC since \( T_g \) points of Sn500K-containing films as seen in Fig. 4 were not as sharp as the \( T_g \) points of control films as seen in Fig. 3.

**IR Analysis**

In the IR spectrum of PVC in Fig. 5a, 2965 cm\(^{-1}\) as CH stretch, 2910 cm\(^{-1}\) as CH\(_2\) stretch, 1425 cm\(^{-1}\) as CH\(_2\) bending, 1098 cm\(^{-1}\) as C–C vibration, 970 cm\(^{-1}\) as CH\(_2\) rocking, 690 cm\(^{-1}\) as CCI isotactic stretch, and 603–698 cm\(^{-1}\) as CCI syndiotactic stretch vibrations were observed. The decrease of C–O–C band for THF at 1060 cm\(^{-1}\) (Fig. 5b) by heating of PVC film indicated the removal of THF from PVC.

With increasing time and temperature of heating, the band intensity for COO\(^-\) groups at 1540 cm\(^{-1}\) decreased for the films with Sn500K, as seen in Figs. 6 and 7 and in Table I. Reaction of stabilizer with PVC can be shown as follows:

\[
\text{CH}_2-\text{CHCl}-\text{CHCl}-\text{CHClCH}_2\text{CHCl}-X- + (\text{R}_2\text{COO})_2\text{Sn(R}_1)_2
\]
PVC STABILIZATION BY ORGANO TIN SOAPS

FIGURE 5a. IR spectrum of unheated PVC.

\[
\begin{align*}
\text{CH}_2\text{-CHCl\text{-CHCl\text{-CHClCH}_2\text{CH-X-}}}
\end{align*}
\]

\[
\text{+Cl(R}_2\text{COO)Sn(R}_1\text{)_2 (3)}
\]

where \( (R_2\text{COO})_2\text{Sn(R}_1\text{)_2} \) is the suggested formula of the active ingredient.

Further reaction of stabilizer with PVC was expected to end with \( \text{Cl}_2\text{Sn(R}_1\text{)_2} \).

The labile chlorine atoms of PVC were substituted by carboxylate groups of organotin soap forming stable ester groups, according to the Frye–Horst mechanism.\(^6\)

Decrease of intensity of 1720 cm\(^{-1}\) and 1640 cm\(^{-1}\) bands as reported in Table I indicated the removal of carbonyl groups and groups having double bonds by evaporation of Sn500K from films on heating. Sn500K was volatile similar to DBTDL, which evaporates in the range of 445–523 K.\(^15\) The characteristic vibration of \(-\text{C=C-}\) bonds in polyene sequences at 1630 cm\(^{-1}\) was observed to be increasing with heating time after the induction period for PVC stabilized with DBTDL.\(^7\) Since the band intensity at 1640 cm\(^{-1}\), which was very close to the 1630 cm\(^{-1}\) band, was decreased with heating time, the polyene sequences formed by heating Sn500K-stabilized PVC was not at a concentration level that could be detected by IR spectroscopy.

UV Analysis

It is possible to calculate percent conversions related to a number of double bonds that may be formed by dehydrochlorination from the UV spectra of the films. Total concentration of 6–10 conjugated double bonds were found using the absorbance and molar absorption coefficients of different polyene types.\(^16\)

The concentration of polyene sequences is given by

\[
C_n = \frac{A}{(\epsilon_\lambda d)}
\]

where \( A \) is the absorbance at wavelength \( \lambda \), \( \epsilon_\lambda \) is the extinction coefficient, \( n \) is the number of conjugated double bonds, and \( d \) is the path length of the cell.\(^17\) \( \epsilon_\lambda \) values given by Shindo and Hirai\(^16\) were used in \( C_n \) calculations.

FIGURE 6. IR spectra of samples with 2.5% Sn500K for the 1500–1800 cm\(^{-1}\) range: (1) unheated, (2) heated at 160°C for 15 min, (3) heated at 160°C for 105 min, (4) heated at 180°C for 15 min, and (5) heated at 180°C for 105 min.

FIGURE 5b. IR spectrum of PVC heated at 180°C for 60 min.
The mole fraction of polyene sequences containing \( n \) conjugated double bonds can be determined from \( C_n \) as

\[
N_n = \frac{C_n}{C} \quad (5)
\]

where \( C \) is the concentration of vinyl repeat units in mol/dm\(^3\). Mole fraction of double bonds

\[
N_D = \sum nN_n \quad (6)
\]

For tin-stabilizer-containing films, UV absorption of the stabilizer also overlapped with conjugated double bond peaks up to 300 nm.\(^{16} \) For solid PVC having 1.4 g/cm\(^3\) density, \( C \) was 22.2 mol/dm\(^3\).

Since there was a peak of BHT at 275 nm, which overlaps the peaks of up to six conjugated -enes and Sn500K also absorbed UV light in the same range, the total number of double bonds having six and more than six double bonds were calculated using UV-spectra and Eqs. (4)–(6). Percent conversion of available repeat units to double bonds by dehydrochlorination equals to 100 \( N_D \).

Representative UV spectra of the films are given in Figs. 8 and 9. Percent conversions found for PVC, PVC with 2.5% Sn500K, and PVC with 5% Sn500K from UV spectra and their colors are given in Table II. Smaller conversion of PVC with 2.5% Sn500K than of PVC alone was observed. PVC without any additive became brown on heating at 180°C for 105 min.
indicating it was degraded with the formation of long polyene sequences. On the other hand, the white color of the unheated films with 5% Sn500K in Table II indicated that PVC was not degraded. The opaque and white color of the unheated films appeared to have superficially high % conversion [1.8% found from Eq. (6) as reported in Table II]. Thus, the conversion determination from Eq. (6) should be correct only for transparent samples. Since films were not transparent the UV analysis did not give reliable results.

**TGA**

The thermograms of the film samples shown in Fig. 10 indicate that there is a step mass loss around the glass-transition temperature of the films because of THF evaporation. This also confirmed that endotherm observed around the glass-transition point belonged to THF evaporation. The stabilizing effect of BHT and Sn500K on PVC above 175°C was observed, since the mass loss from PVC powder is much faster than the mass loss from cast PVC films.

Activation energy $E_a$, Arrhenius constant $A$, and degradation rate coefficient $k$ can be found from the following equation:

$$k = A \exp\left(-\frac{E_a}{RT}\right)$$

$$\ln\left[\frac{(d\alpha/dT)/(1-\alpha)^n}{(1-\alpha)^n}\right] = \ln\left(A/\beta - \frac{E_a}{RT}\right)$$

taking the reaction order to be 1 and by taking temperatures and mass loss % ($\alpha$) from TGA curves. $\alpha$ values, conversions, were taken as (100 $- a/58$)\%/100, where 58 is the weight % of HCl ($C_0$) in PVC and $a$ is the percentage of remaining mass of the sample at temperature $T$. $\beta$ is the heating rate expressed in degrees per minute. The slope and the intercept of the $\ln\left[\frac{(d\alpha/dT)/(1-\alpha)^n}{(1-\alpha)^n}\right]$ vs. $1/T$ line is $-E_a/R$ and $\ln(A/\beta)$, respectively. $k$ values at 180°C were found using $E$ and $A$ values found from TGA.

Conversions found from UV analysis were lower than conversions from TGA, using Eq. (3), because of the evaporation of THF, and Sn500K present in the films occurred as well as PVC degradation (Table III).

As seen in Table IV, the PVC film and samples with Sn500K have lower $k$ values than PVC powder. Antioxidant BHT acted as heat stabilizer since it was a radical acceptor. Only the rate of formation of polyene sequences longer than 6 units was included in UV analysis, excluding shorter polyene sequences because of the overlapping of the BHT and Sn500K spectra in their range. Higher conversion found form TGA represented all HCl molecules splitted from PVC, including short and long polyene sequences. The present study also confirmed the formation of short polyene sequences with organotin stabilizers, as reported by previous workers.

It was observed that the sample with 2.5% Sn500K was less degraded than the sample with 5% Sn500K. The level of Sn500K in the films should be kept low to have transparent films. At high Sn500K...
concentrations, films become opaque white because of the incompatibility of Sn500K and PVC.

In the high temperature range, Sn500K did not effect degradation of PVC. As seen in Fig. 10, degradation occurred in two steps, as observed by previous workers.4,5 At 425°C the remaining mass of PVC was in the range of 17–25% for control and stabilized films.

**Discussion and Conclusion**

The exact composition of PVC stabilizers was not revealed by their producers. Using x-ray fluorescence analysis, it is possible to find out which metals are present in any stabilizer. IR spectroscopy can be used to find out the functional groups that are effective in PVC stabilization. Thus, Sn500K was chosen as a representative liquid heat stabilizer and was analyzed to find out that it was a lead free, and was containing tin element. It is most probably DBTOM dissolved in a solvent.

The heat stability tests of films with Sn500K showed an induction period up to 30 min in terms of yellowing. Actually, shorter polyene sequences,8 which did not discolor PVC, were formed as indicated by UV spectroscopy.

At low temperatures (160°C) and short heating times (15 min) films having 2.5% Sn500K were more stable than PVC powder and PVC film without additive with respect to color, double bond formation, and rate of mass loss. For longer heating periods, Sn500K has not enough effectiveness. The effects of THF on glass-transition temperature and BHT on thermal stability were also observed. DSC and TGA showed that THF was evaporated from the films by heating at glass-transition temperature of the films. In film preparation, transparent coatings of PVC stable to heat for short periods could be obtained by using low Sn500K content. In the high temperature range of heating, Sn500K had no effect on the degradation behavior of PVC.

**TABLE III**

Percent Conversion of PVC to dePVC for Heating at 180°C for 15 min as Determined by UV Spectroscopy and TGA

<table>
<thead>
<tr>
<th>Sample</th>
<th>TGA</th>
<th>UV</th>
</tr>
</thead>
<tbody>
<tr>
<td>PVC film</td>
<td>4.2</td>
<td>0.354</td>
</tr>
<tr>
<td>2.5% Sn</td>
<td>4.7</td>
<td>0.219</td>
</tr>
<tr>
<td>5% Sn</td>
<td>9.9</td>
<td>1.130</td>
</tr>
</tbody>
</table>

**FIGURE 10.** TGA curves for PVC powder (----), PVC film (---), PVC film with 2.5% stabilizer (----), and PVC film with 5% stabilizer (——).
TABLE IV

<table>
<thead>
<tr>
<th>Sample</th>
<th>$E_a$ (J/mol)</th>
<th>$A$</th>
<th>$k$ (1/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PVC powder</td>
<td>91750.71</td>
<td>$1.04 \times 10^9$</td>
<td>0.020597</td>
</tr>
<tr>
<td>PVC film</td>
<td>82790.87</td>
<td>$1.26 \times 10^7$</td>
<td>0.002769</td>
</tr>
<tr>
<td>2.5% Sn500K</td>
<td>118151.6</td>
<td>$2.29 \times 10^{11}$</td>
<td>0.003774</td>
</tr>
<tr>
<td>5% Sn500K</td>
<td>83573.67</td>
<td>$4.02 \times 10^7$</td>
<td>0.007160</td>
</tr>
</tbody>
</table>

Higher thermal stabilization efficiency was obtained with Sn500K compared to the zinc stearate stabilizer, since better mixing of the liquid stabilizer with PVC had been possible.

References