Statistical Thermal Stability of PVC

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ABSTRACT: Experimental design was used to optimize the processing parameters for the decomposition of poly(vinyl chloride). Factorial design and face centered composite design (FCC) were applied to determine the optimum conditions. A total of 10 g PVC powder was mixed with different amounts of zinc stearate (ZnSt2) and natural zeolite and tested for thermal stability. Factorial fitted model was explained by first order pattern due to the significant main effect regression constants, and FCC model was described by second order model owing to higher order polynomial coefficients. FCC design was superior to factorial design as FCC considers not only its pure quadratic effects contribution but also its higher overall desirability for thermal stability of PVC. For factorial design the optimum conditions were determined as 163.06 mg for ZnSt2, 399.99 mg for zeolite, and 140°C for temperature with desirability of 0.933. However, 400 mg for ZnSt2, 333.24 mg for zeolite, and 140°C for temperature with desirability of 0.956 were obtained as the optimum conditions by FCC design.

Key words: poly(vinyl chloride) (PVC); zeolite; degradation; kinetics; stabilization

INTRODUCTION

Poly(vinyl chloride) (PVC) is one of the most versatile polymers because it can accept a wide range of additives, which make PVC, among other properties, durable and difficult to break down. It can be applied to different fields from construction (e.g., in pipes and window frames) to packing (foods, plasma, and blood bags).1,2 PVC is rarely used alone and is mixed with various additives such as plasticizers and stabilizers.3 One of the most serious problems for PVC processing and handling is the rather low thermal stability.

The thermal degradation of PVC has two step processes. The first step mainly involves progressive unzipping of neighboring labile chlorine atoms along the polymer chain by heat, while PVC undergoes a partial carbonization with the release of benzene and other aromatics.4 The dehydrochlorination of PVC starts at about 100°C and is the reason for discoloration, owing to the formation of poly-ene sequences. It is catalyzed by HCl evolved (autocatalysis) and also be promoted by other strong acids. To prevent undesirable degradation processes, a number of chemicals can be used as stabilizers, such as lead compounds, organotin compounds, compounds of other metals, and inorganic stabilizers (zeolite and hydrotalcite). Even though thermal stabilizers based on tin and cadmium were of the practical interest as stabilizers for PVC, cadmium use in PVC was restricted as this metal was toxic, environmentally hazardous, and had negative health effect, such as lung cancer. Moreover, organotin stabilizers that are derived from tetravalent tin has unpleasant odor during processing and in the finished product of PVC. Besides that, semi volatile compounds can migrate into environment over time from the PVC materials stabilized with organotin compounds.5 To overcome the disadvantages of tin- and cadmium-based thermal stabilizers, these compounds are substituted by calcium and zinc stabilizers that are nontoxic. Zinc-based stabilizer, which has stronger Lewis acidity, is able to scavenge HCl and react with labile chlorine atoms of PVC; therefore, it is used as a primary stabilizer in PVC degradation processes. When zinc and calcium stearate stabilizers were compared in the stabilization of PVC, it was observed that the traces in the induction time of the samples stabilized with zinc stearate were more stable than the one stabilized with calcium stearate stabilizer, indicating the power of zinc soap to inhibit the dehydrochlorination of PVC.6,7 The main role of zinc stearate in the degradation of PVC is the substitution of allylic chlorine in PVC with carboxylate group of the zinc soap as indicated in eq. (1), and this stabilizer makes PVC more resistant to heat.

\[ (\text{RCOO}^-)_2\text{Zn} \rightarrow 2\text{R}_1 - \text{Cl} \rightarrow 2\text{RCOOR}_1 + \text{ZnCl}_2 \]

where \( R_1 \) represents the PVC polymer chain except an allylic chlorine. However, the formed ZnCl2 in...
The effect on the thermal stability of PVC was investigated as the interlayer counterions of this anion-exchanging clay initially react with HCl, and second, the layers react with HCl to form metal chlorides with loss of the layered structure. It can be seen that the thermal stability time of PVC-hydrotalcite composite resins is greater than that of pure PVC resin and increases as the weight fraction of the hydrotalcite increases. A natural zeolite, clinoptilolite tends to neutralize the aqueous medium acting either as proton acceptor or as a proton donor due to its amphoteric character. Bronsted and Lewis acidic/basic sites existing in the microporous structure of the zeolitic materials are responsible for their chemical behavior in aqueous solutions. The neutralization of the aqueous medium can be result of the removal of protons from surface Bronsted acidic sites or the binding of protons to the Lewis basic sites of the zeolite. Zeolite addition in PVC not only enhances the thermal stabilization of PVC but also effects the mechanical properties of PVC. Young modulus of PVC-zeolite foams increases with increasing the concentration of zeolite in PVC foam. The addition of 20% (by weight) zeolite increases Young Modulus value from 1.53 to 8.67 N/mm².

In the previous study, the stabilizing effect of natural zeolite (clinoptilolite) on the stabilization of PVC plastigel at 140 and 160°C was confirmed. Thermal stability of PVC can be determined by PVC Theromat instrument in which induction and stability time, initial and linear region rate constant of dehydrochlorination reaction can be measured. PVC powder was provided by Petkim (Petvinil P.38/74). The samples were prepared by mixing PVC powder with different amounts of zinc stearate (ZnSt₂) and clinoptilolite rich natural zeolite having 45 µm average particle size which was supplied from Gördes Region in Western Anatolia. The composition of clinoptilolite rich mineral was 66.36% SiO₂, 11.36% Al₂O₃, 1.227% Fe₂O₃, 3.844% K₂O, 0.998% Na₂O, 2.344% CaO, 0.420% MgO, 14.22% H₂O. The clinoptilolite content was estimated as 80%. All chemicals were of analytical grade and used as received.

**MATERIALS AND METHODS**

**Materials**

PVC powder was provided by Petkim (Petvinil P.38/74). The samples were prepared by mixing PVC powder with different amounts of zinc stearate (ZnSt₂) and clinoptilolite rich natural zeolite having 45 µm average particle size which was supplied from Gördes Region in Western Anatolia. The composition of clinoptilolite rich mineral was 66.36% SiO₂, 11.36% Al₂O₃, 1.227% Fe₂O₃, 3.844% K₂O, 0.998% Na₂O, 2.344% CaO, 0.420% MgO, 14.22% H₂O. The clinoptilolite content was estimated as 80%. All chemicals were of analytical grade and used as received.

**Method**

Different amount of stabilizers were added to 10 g PVC powder and mixed to form homogeneous blends. Thermal stability test was performed by 763 PVC Theromat, which was equipped with two heating blocks each with four measuring positions. The reaction vessels that contained 0.5 g. of PVC samples were placed into the heating blocks of PVC Theromat after they were heated to 140, 150, and 160°C. During the reaction, HCl formed was taken up by a nitrogen gas stream and transferred into the measuring vessels. The deionized water absorbed HCl gas and the change in conductivity of the solution was determined with respect to time by the conductimeter. Therefore, the decomposition process was monitored by measuring the conductivity of aqueous HCl solution.

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Experimental design

In dehydrochlorination of PVC, induction time, stability time, initial rate constant, and linear region rate constant were chosen as the responses, while the amount of ZnSt₂, zeolite, and temperature were considered as factors which were varied over two levels (low and high). The experimental factors and the levels investigated for thermal stability of PVC are given in Table I. The experiments were performed by considering a $2^3$ full factorial design with 4 center points, consisting of 20 experiments (two replicates for each experiment). The main purpose of the addition of center point was to provide protection against curvature from the second order effect as well as allow an independent estimate of error to be obtained. Also, FCC design was used to observe the polynomial of higher degree and to compare the results obtained by full factorial design. Both designs were carried out by using Design of Expert 6.0.11.

RESULTS AND DISCUSSION

Kinetic study of PVC

The kinetic study was carried out not only to optimize the amount of additives in PVC but also to expose the effect of zinc stearate and zeolite on the stabilization of the polymer at two different temperatures. Figures 1 and 2 indicate the change of conductivity of aqueous solution with respect to time for PVC, PVC stabilized with zinc stearate, and PVC stabilized with zeolite. By using the curves in these figures, kinetic rate constants of dehydrochlorination reaction were evaluated. The steps used in the calculation of rate constants of degradation reaction were described in the recent studies. In this model, it was assumed that the degradation of PVC was a first order reaction. The rate constant was related to the concentration of HCl in water ($C_{HCl}$), potential number of double bonds to be formed when 100% of HCl was eliminated per 1 g of PVC ($C_o$), moles of evolved HCl gas ($n_{HCl}$), mass of plasticizers in reaction vessel ($m$):  

$$k = \frac{dC_{HCl}}{dt} \cdot \frac{1}{C_o} = \frac{dn_{HCl}}{dt} \cdot \frac{1}{m \cdot CF} \cdot \frac{1}{C_o} \quad (2)$$

where CF is the correction factor based on the fraction of PVC amount in PVC mixtures. The dehydrochlorination curves indicated in Figures 1 and 2 were separated into two parts, initial region and linear region. Therefore, two rate constants were evaluated for each reaction. Initial rate constants were obtained from the slopes of initial region of the curves where the release of HCl gas started; whereas linear region rate constants were determined from the slopes of linear region where the release of HCl gas accelerated. The induction and stability periods of PVC at 140 and 160°C are higher than the PVC having additives as this PVC was synthesized by emulsion polymerization. Therefore, it has some organic compounds acting as heat stabilizers such as surface active agents, plasticizer, and anti oxidants, and in literature, it was found that PVC particles are covered by carbon-rich materials, and these additives are 1.6% of PVC as determined by ethanol extraction. The obtained induction and stability periods and the calculated rate constants are reported in Table II. The increase in rate constants for PVC stabilized with zinc stearate, in other words,
the increasing rate of dehydrochlorination after the consumption of zinc stearate indicated the autoacceleration effect of HCl and the produced ZnCl₂. Unlike zinc stearate, zeolite did not promote dehydrochlorination; therefore, it decreased the rate constants of the sample. The use of zeolite effect in PVC was considered to result from the absorption of HCl which catalyzed the degradation of PVC. To determine the optimum process parameters for the stabilization of PVC, the conductivity changes of PVC stabilized with different amounts of ZnSt₂ and zeolite with respect to time at 140, 150, 160°C are plotted in Figures 3–5. Table III and IV list the kinetic results corresponding to the factorial and face centered composite design. At 140°C, the sample stabilized with 400 mg ZnSt₂ and 400 mg zeolite has the induction value 18.23 h, while the lowest linear region rate constant was obtained for the sample only 100 mg ZnSt₂ and 100 mg zeolite which meant that small amount of ZnCl₂ are released during the dehydrochlorination. The enhancing effect of ZnSt₂ on the induction period is verified in Figure 4 as the highest induction time is observed for the sample having highest amount of ZnSt₂. However, PVC stabilized with 100 mg, ZnSt₂ and 400 mg, zeolite has the lowest linear region rate constant due to the absorption of HCl evolved at the initial stages of dehydrochlorination by zeolite. At higher temperatures (160°C), the addition of 400 mg zeolite into PVC not only increase the induction time of PVC but also decrease the linear region rate constant of the sample.

**Factorial design for PVC stability**

In factorial design, the interaction between the factors was postulated by the fitted regression model, which can used to obtain the predicted or fitted value of responses. The model representation of this design could be written as:

$$\hat{y} = \beta_0 + \beta_1X_1 + \beta_2X_2 + \beta_3X_3 + \beta_{12}X_1X_2 + \beta_{13}X_1X_3 + \beta_{23}X_2X_3$$

where \(\hat{y}\) is the response, \(\beta_0\) is the average value of the experimental responses, \(\beta_1, \beta_2, \beta_3\) are the main

### Table II

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<tr>
<th>Samples</th>
<th>Temperature (°C)</th>
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<th>Stability time (h)</th>
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<th>Linear region rate constant × 10⁵ (min⁻¹)</th>
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**Figure 3** The change of conductivity of aqueous solution with respect to time at 140°C for 10 g PVC stabilized with ZnSt₂ and zeolite, 1-PVC having 100 mg ZnSt₂ and 100 mg zeolite, 2-PVC having 100 mg ZnSt₂ and 400 mg zeolite, 3-PVC having 400 mg ZnSt₂ and 100 mg zeolite, and 4-PVC having 400 mg ZnSt₂ and 400 mg zeolite.

**Figure 4** The change of conductivity of aqueous solution with respect to time at 150°C for 10 g PVC stabilized with ZnSt₂ and zeolite, 1-PVC having 100 mg ZnSt₂ and 250 mg zeolite, 2-PVC having 250 mg ZnSt₂ and 100 mg zeolite, 3-PVC having 250 mg ZnSt₂ and 250 mg zeolite, 4-PVC having 250 mg ZnSt₂ and 400 mg zeolite, and 5-PVC having 400 mg ZnSt₂ and 250 mg zeolite.
Effect coefficients, \( b_{12}, b_{13}, b_{23} \) are the interaction coefficients, \( X_1, X_2, X_3 \) are coded variables that represent the ZnSt\(_2\), zeolite, and temperature, respectively.

The fitted regression models for each response are given in eqs. 4–7. The model results for induction time indicated that zeolite had positive effect, whereas temperature had negative effect on this response. Only temperature had negative effect on the stability time. However, all main and interaction effects are significant on the initial rate constant. Zeolite, temperature and zeolite-temperature effects are considerable on the linear region rate constant.

\[ \hat{y} = 9.06 + 0.92X_2 - 5.9X_3 \]  

(4)

Stability time
\[ \hat{y} = 10.72 - 7.38X_3 \]  

(5)

Initial rate constant
\[ \hat{y} = 7.742 \times 10^{-6} + 1.741 \times 10^{-6}X_1 - 3.648 \times 10^{-6}X_2 + 5.972 \times 10^{-6}X_3 + 1.427 \times 10^{-6}X_1X_3 + 3.501 \times 10^{-6}X_2X_3 \]  

(6)

Linear region rate constant
\[ \hat{y} = 1.266 \times 10^{-4} + 1.025 \times 10^{-4}X_2 + 1.134 \times 10^{-4}X_3 + 9.644 \times 10^{-5}X_2X_3 \]  

(7)

However, it is always necessary to examine any interaction that are important as the main effect do not have much meaning when they are involved in significant interaction. Zeolite and temperature; ZnSt\(_2\) and temperature effects on initial rate constant and zeolite and temperature effect on linear region rate constant are shown in Figure 6(a,b) respectively.

Figure 6(a) indicated that zeolite had not a significant effect at low level of temperature although it had a prominent effect at high level of temperature for both initial rate constant. This result implied that the change in the amount of zeolite in PVC did not significantly affect the degradation behavior of the sample at low temperature. On the other hand, the initial rate constant of degradation reaction increased by increasing the amount of zeolite in PVC at high temperature.

![Figure 5](image_url)  

Figure 5: The change of conductivity of aqueous solution with respect to time at 160°C for 10 g PVC stabilized with ZnSt\(_2\) and zeolite, 1-PVC having 100 mg ZnSt\(_2\) and 100 mg zeolite, 2-PVC having 100 mg ZnSt\(_2\) and 400 mg zeolite, 3-PVC having 400 mg ZnSt\(_2\) and 100 mg zeolite, and 4-PVC having 400 mg ZnSt\(_2\) and 400 mg zeolite.

### TABLE III

Full Factorial Experiment Design 2\(^3\) with 4 Center Point

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<th>Std</th>
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<th>Induction time (h)</th>
<th>Stability time (h)</th>
<th>Initial rate constant ( \times 10^6 ) (min(^{-1}))</th>
<th>Linear region rate constant ( \times 10^5 ) (min(^{-1}))</th>
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temperature. Same degradation behavior was observed for the sample stabilized with zinc stearate at 140 and 160°C. Linear region rate constant was increased when PVC stabilized with 400 mg zeolite at high level of temperature as displayed in Figure 6(b).

The response surface and contour plots for all responses are given in Figures 7–10. Figures 7 and 8 demonstrated that the response surface plots were planar, and the contour plots contained parallel straight lines. The result plots confirmed with the fitted regression model, which was explained only by the main effect instead of interaction. In that cases, the induction and stability results were well modeled by first order model.

Figures 9 and 10 show the response surface and contour plots of initial and linear region rate constants, respectively. In these figures, the significant interaction effects twisted the plane resulting curved contour lines. Thus, interaction was a form of curvature in the underlying response surface model for these responses. Initial and linear region rate constants were modeled by second order model. Temperature-zeolite and temperature-ZnSt₂ contributed curvature effect on initial rate constant, whereas temperature-zeolite gave the curvature effect on linear region rate constant with the higher regression coefficient.

The optimum conditions for dehydrochlorination of PVC were determined as 163.06 mg ZnSt₂, 399.99 mg zeolite, and 140°C with a desirability of 0.932.

### Table IV

Face Center Composite (FCC) Design

<table>
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<th>Std number</th>
<th>Additive (mg/10 g PVC)</th>
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<td>2.40</td>
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<td>3.47</td>
<td>33.40</td>
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<td>13.27</td>
<td>18.56</td>
<td>1.20</td>
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<td>7.28</td>
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<td>17.64</td>
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<td>16.04</td>
<td>18.43</td>
<td>1.12</td>
<td>0.48</td>
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</table>

**Face centered composite (FCC) design for PVC stability**

FCC model was fitted by the polynomial of higher degree which could be shown as:

\[
y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_3 x_3 + \beta_{12} x_1 x_2 + \beta_{13} x_1 x_3 \\
+ \beta_{23} x_2 x_3 + \beta_{11} x_1^2 + \beta_{22} x_2^2 + \beta_{33} x_3^2
\]

where \(\hat{y}\) is the response, \(\beta_i\) is the average value of the experimental responses, \(\beta_{11}, \beta_{22}, \beta_{33}\) are the main effect coefficients, \(\beta_{12}, \beta_{13}, \beta_{23}\) are the interaction coefficients, and \(\beta_{11}, \beta_{22}, \beta_{33}\) are the coefficients of pure quadratic terms, \(X_1, X_2, X_3\) are coded variables that represent the ZnSt₂, zeolite, and temperature, respectively.

The fitted regression models for each response are given in eqs. 9–12. These equations revealed that the main, interaction, and pure quadratic terms were significant for all responses.

**Induction time**

\[
y = 6.83 + 0.48 X_1 + 0.95 X_2 - 6.32 X_3 - 0.044 X_1^2 \\
- 0.34 X_2^2 + 3.18 X_3^2 - 0.03 X_1 X_2 - 0.52 X_1 X_3 \\
- 0.37 X_2 X_3
\]

**Stability time**

\[
y = 7.21 - 0.32 X_1 + 0.30 X_2 - 7.03 X_3 + 0.51 X_1^2 \\
- 0.077 X_2^2 + 2.83 X_3^2 + 0.34 X_1 X_2 + 0.46 X_1 X_3 - 0.21 X_2 X_3
\]
Figure 6  Interaction effect plot for (a) initial rate constant and (b) linear region rate constant. [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]

Figure 7  Response surface and contour plot of induction time.
Initial rate constant
\[ \dot{y} = 2.699 \times 10^{-6} + 1.562 \times 10^{-6}X_1 + 4.892 \times 10^{-6}X_2 \\
+ 7.705 \times 10^{-6}X_3 + 2.434 \times 10^{-6}X_1^2 + 1.422 \times 10^{-7}X_2^2 \\
+ 4.792 \times 10^{-6}X_3^2 + 9.875 \times 10^{-3}X_1X_2 + 7.437 \times 10^{-7}X_1X_3 + 6.009 \times 10^{-6}X_2X_3 \]  
(11)

Linear region rate constant
\[ \dot{y} = 3.939 \times 10^{-5} - 9.829 \times 10^{-6}X_1 + 1.058 \times 10^{-4}X_2 \\
+ 1.614 \times 10^{-4}X_3 - 3.484 \times 10^{-5}X_1^2 - 3.051 \times 10^{-5}X_2^2 \\
+ 1.894 \times 10^{-4}X_3^2 - 2.852 \times 10^{-3}X_1X_2 - 2.621 \times 10^{-3}X_1X_3 + 1.266 \times 10^{-4}X_2X_3 \]  
(12)

The main and interaction effect plots for each response are shown in Figures 11 and 12, respectively. For induction time, ZnSt2 had a linear positive effect, whereas for initial rate constant it pointed out positive curvature effect. On the other hand, ZnSt2 showed negative curvature effect for stability time and linear region rate constant. Zeolite indicated linear positive effect for stability time and initial rate constant, although it had positive curvature effect on responses, induction time and linear region rate constant. Moreover, temperature demonstrated negative curvature effect on induction and stability periods, but reverse behavior was observed for the initial and linear region rate constant.

At low level of temperature, ZnSt2 positively affected the induction time, whereas at high level of temperature, it had no significant effect on induction time as shown in Figure 12(a). Varying the amount of zeolite affected the induction time at low level of temperature as indicated in Figure 12(a). The effect of zeolite and zinc stearate on induction time could be assumed negligible when compared to the other interaction parameters. However, increasing the temperature pointed out the curvature effect on this response due to its high pure quadratic interaction.
Figure 10  Response surface and contour plot of linear region rate constant.

Figure 11  Main effect plot for (a) induction time, (b) stability time, (c) initial rate constant, and (d) linear region rate constant.
value as shown in eq. 9. In Figure 12(b), there was significant interaction between zeolite and ZnSt2 on stability time because of the intersection of the interaction lines. Therefore, it was found that the use of zeolite and zinc stearate in PVC increased the stability time of the sample. The stability time was not affected by the increasing the amount of zeolite at two different temperatures, whereas stability period of the sample decreased when 400 mg. zeolite was used at 140°C. This result confirmed the autoacceleration effect of ZnCl2 which was produced by consuming zinc stearate during the degradation of PVC. Figure 12(c), represents the interaction effects on initial region rate constant. Both at low and high levels of zeolite and temperature, the addition of ZnSt2 increased the initial rate constant, showing a curvature effect. Although the addition of zeolite had negative effect on initial rate constant at low level of temperature, it showed positive effect on the same response at high level of temperature. This result revealed that the use of high level of zeolite in PVC decreased the degradation of the polymer. As shown in Figure 12(d), at low level of zeolite and temperature, the addition of ZnSt2 increased the linear region rate constant, whereas at high level of zeolite and temperature, this response showed a negative

Figure 12 Interaction effect plot for (a) induction time, (b) stability time, (c) initial rate constant, and (d) linear region rate constant. [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]
effect by increasing ZnSt$_2$ amount indicating curvature behavior. By increasing zeolite amount, linear region rate constant slightly decreases at low level of temperature, whereas it showed positive effect on the same response at high level of temperature.

Figure 13 shows the response surface and contour plots for all responses. Twisted planes were obtained due to their pure quadratic and interaction effects for the responses resulting in curved contour lines. Among all the responses, linear region rate constant implied the most significant curvature effect with higher pure quadratic interaction coefficient.

The optimum conditions for dehydrochlorination of PVC were determined as 400 mg ZnSt$_2$, 333.24 mg zeolite, and 140°C with a desirability of 0.956 which was higher than the desirability obtained by factorial design.

CONCLUSIONS

Two factorial and FCC designs were used to investigate the dehydrochlorination of PVC with four responses (induction time, stability time, initial rate constant, and linear region rate constant) and three factors (ZnSt$_2$, zeolite, temperature).

The regression model representation of the experiment was fitted to the first order model including the main and interaction effect of the responses. The results of main effects revealed that the temperature had a positive effect on initial and linear region rate constants, although it had a negative effect on other responses. ZnSt$_2$ only attributed towards to the initial rate constant. Zeolite had a positive effect on all responses except stability time. The stabilizing effect of zeolite on the other responses was deliberated the results from the absorption of HCl which was thought to reduce the autocatalytic effect of this gas evolved at initial stage of the dehydrochlorination. The interaction effects should have been considered regardless the main effect, as the main effects did not have much contribution in the presence of significant interaction. Zeolite-temperature interaction indicated that there was no significant effect on the initial and linear region rate constant at low level, whereas a positive effect was observed on these responses at high level. ZnSt$_2$-temperature interaction was effective only for initial rate constant at high level of temperature indicating the power of ZnSt$_2$ to inhibit the dehydrochlorination but as soon as this was finished HCl gas evolved instantaneously. The fitted model was described by first order pattern for induction and stability time because of the fact that the response surface and contour plots were composed of parallel straight lines. As the shape of these plots gave a clue which parameters were significant, it can be concluded that the model
was well explained only by main effects for these responses. In other responses (initial and linear region rate constant), the significant interaction effects, which was a form of curvature in the underlying response surface model, twisted the plane resulting curved contour lines. Hence, these responses were modeled by second order model.

FCC model including the main, interaction, and pure quadratic term stated that the second order model was an appropriate approximation. In the main effects plots, ZnS\textsubscript{2} had a linear and curvature positive effect on induction time and initial rate constant, whereas it had a negative curvature effect on stability time and linear region rate constant. Zeolite had positive effect on all responses, linear for stability time and initial rate constant and curvature for induction time and linear region rate constant. Temperature had a negative curvature effect on induction and stability time, whereas it had a positive curvature effect on initial and linear region rate constants. At low level of temperature, ZnS\textsubscript{2} positively affected the induction time. Increasing the amount of zeolite indicated the curvature effect on induction time due to its high pure quadratic interaction value. There was significant interaction between zeolite and ZnS\textsubscript{2}, whereas ZnS\textsubscript{2} and temperature effect implied a curvature behavior on stability time. Both at low and high levels of zeolite and temperature, the addition of ZnS\textsubscript{2} increased the induction rate constant, showing a curvature effect. It was observed that the addition of zeolite had negative effect on initial rate constant at low level of temperature while, it showed positive effect on the same response at high level of temperature. The results implied that at low level of zeolite and temperature, the addition of ZnS\textsubscript{2} increased the linear region rate constant, whereas at high level this response showed a negative effect by increasing ZnS\textsubscript{2} amount indicating curvature behavior. Response surface and contour plots imparted that pure quadratic and interaction effects were dominant in all responses. Contour lines were curved resulting the plane to twist. The high polynomial order model was fitted for FCC model.

The optimum process parameters for factorial and FCC designs were determined as 163.06 mg ZnS\textsubscript{2}, 399.99 mg zeolite, and 140°C and 400 mg ZnS\textsubscript{2}, 333.24 mg zeolite and 140°C, respectively. The evaluated values were confirmed by PVC Thermomat after the design had been completed. All the results pointed out that FCC design was superior to factorial design because of its pure quadratic effects contribution and its higher overall desirability for thermal stability of PVC.

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References

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