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<td><strong>Author(s)</strong></td>
<td>Joshi, Sunil Chandrakant; Lam, Yee Cheong</td>
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Modelling heat and degree of gelation for methyl cellulose hydrogels with NaCl additives

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Abstract

Methyl cellulose (MC) hydrogels are thermo reversible physical hydrogels and their gelation is endothermic. A model consisting of a general mathematical expression (for bell-shape curves) is formulated to describe and capture the enthalpy changes that take place during gelation of an aqueous solution of MC, SM 4000, in the presence of different concentrations of sodium chloride, NaCl. The constants for the mathematical expression were obtained based on calorimetric measurements. The model could describe the salt-out effects of NaCl very well.

It was noticed that more than one bell-shape curves as defined by the expression, one major and one minor, were necessary to depict the enthalpy changes appropriately. It was hypothesized that the major curve is associated with the major enthalpy change related to hydrophobic association of MC groups leading to the development of a gel network and its swelling. The minor curve describes the minor enthalpy change, which include primarily the ionic interaction of salts with water, water-to-water and water-to-polymer hydrogen bonding and energy associated in stabilizing the gel network. Subsequently, an analytical model for the degree of gelation was introduced and its development discussed to study further the effects of various salt concentrations on the gelation of MC.
1. Introduction

Hydrogels are cross-linked networks of hydrophilic/ hydrophobic polymers which are formed either by chemical or physical bonds in water. They possess the ability to absorb large amounts of water and swell while maintaining their three-dimensional structure.

Hydrogels are increasingly being used in biomedical, food and pharmaceutical industries. Most hydrogels undergo transformation from solution to gel form (commonly termed as gelation process) upon either cooling or heating. Such thermo-sensitive hydrogels are gaining more importance and being explored as media for controlled drug release [1-4].

Methyl cellulose (MC), which is extensively used as a thickener or binder in pharmaceutical, ceramic processing and food applications, undergoes gelation in aqueous solution upon heating [5-7]. Gelation of MC has been a key area of research and a clear and complete picture of the gelation mechanism has not yet been established. The sol-gel transformation process for MC, as illustrated in Figure 1, is based on various explanations provided by other researchers [5-8] and our earlier studies [9-10]. According to Kobayashi et al. [8], at low temperatures, water molecules surround the hydrophobic methaxyl groups of MC, so that MC becomes water-soluble (Figure 1a). However, upon heating, these surrounding water structures deform and break to expose the hydrophobic regions of MC (Figure 1b). Because of hydrophobic nature of these side groups, the MC chains in the aqueous solution move towards each other, bringing side groups together to associate, and form aggregates or inter-chain clusters (Figure 1c). As temperature rises further, more and more hydrophobic aggregates are formed. Eventually these aggregates join together through the hydrophobic groups developing into a three-dimensional
physical network of MC chains (Figures 1d-1f). Even when the network is fully developed, the MC mers are still in a relaxed state rendering the network cells flexible. Gradually, as temperature increases, water uptake and entrapment of water molecules into these network cells increases resulting in enlargement of these cells to full-size reducing their flexibility (Figures 1g-1h). The continued swelling of the network of MC chains is due to osmotic forces, which are opposed by the elastic retraction forces of the network. Eventually an equilibrium swelling level is attained [11] resulting into a fully developed and stabilized MC gel (Figure 1i).

As such, the sol-gel transformation of MC in an aqueous solution is distinctively marked by two processes namely, formation of MC chain network and swelling or expansion of the various cells in the network resulting into a fully developed gel. Both these transformations are linked to the absorption and utilization of heat energy, generally referred to as the heat of gelation. Differential scanning calorimeter (DSC) measurements have been successfully used to obtain quantitative information about the heat of gelation [6, 12].

Kundu and Kundu [13] investigated the effect of sodium chloride, NaCl, when added to aqueous solutions of MC, with the MC content below 1%, on the DSC measurements. It was observed that the temperature at which absorption of heat was at the peak decreased with increasing salt concentration. This is an important salt-out phenomenon that can be used to alter the sol-gel transition process to suit particular application. Xu et al. [14-15] also carried out DSC measurements on MC solutions in the presence of NaCl in order to understand the salt-out mechanism further. Newer inventions are seen where MC is modified to match their salt-out effects at body temperature [16].
The current work focuses on the modeling of the changes in enthalpy associated with the gel formation process for MC with NaCl additives at different concentration. Unless such models are available, various applications of MC hydrogels, including drug delivery and drug release, can not be systematically analyzed, understood, and improved. In our earlier work [9-10] a generalized formulation was developed and validated for modeling MC gelation. The same formulation is now modified and extended further to capture the salt effects. The necessary parameters in the formulation were determined such that close agreement was established with the DSC heating thermograms. During the model formation, it was noticed that more than one bell-shape curves, one major and one minor, were necessary to depict the enthalpy changes adequately. It was hypothesized that the major curve was related to major enthalpy change associated with the movement of MC chains, formation of MC aggregates, and hydrophobic association of MC groups finally leading to the development of a three-dimensional gel network and its subsequent swelling. The term swelling is used to highlight the enlargement of MC network cells due to uptake and entrapment of water molecules into them. The minor curve was associated with minor enthalpy change primarily involving the ionic interaction of the salts with water, water-to-water and water-to-polymer hydrogen bonding and energy associated in stabilizing the swollen gel network. Subsequently, an analytical model for the degree of gelation was introduced and its development discussed to study the effects of various concentrations of NaCl on the gelation of MC further.
2. Materials and Experimental Data

The DSC data, i.e. the relative thermal capacity, $C_p$, was acquired [14-15] using a micro differential scanning calorimeter (VP-µDSC, MicroCal Inc.) for the aqueous solutions of 0.03mM (0.93wt%) of MC with different % of NaCl, namely 0.2M, 0.4M, 0.6M, and 0.8M. De-ionized water was taken as a reference during the heating process at the rate of 1 °C/min. Methylcellulose, with the trade name, SM4000, was provided by Shinetsu Chemical Co. Ltd. It has a weight-average molecular weight ($M_w$) of 310,000 g/mol and an average degree of substitution (DS) of 1.8. NaCl was purchased from Sino Chemical Co. (Pte) Ltd.

The DSC results for the MC-NaCl solutions as documented and discussed in [14-15] were used in the present work to demonstrate the development and validity of the models. It was observed that the baseline $C_p$ values in all cases showed small fluctuations around zero due to the initial unsteady state of the solutions and the temperature conditions before the DSC measurements attained a steady trend. As the magnitude of the base $C_p$ value was insignificant, its small fluctuation would have little effect. As such, for convenience, a single base $C_p$ value, set at $1 \times 10^{-4}$ KJ/mol/°C, was used to normalize the experimental data as depicted in Figure 2. The plots indicated that the addition of salt had an effect on the onset ($T_{on}$), peak ($T_p$), and offset ($T_{off}$) temperatures for the sol-gel transformation.
3. Model Development and Implementation

It was observed that most of the heating thermograms for gelation of MC hydrogels were un-symmetrical and bell-shaped (Figure 2), which required a generalized mathematical formulation to represent them. One such formulation (equation 1) was established earlier by the authors [9-10]:

\[
C_p(T - T_p) = \frac{NR}{DR_1 + DR_2} = \frac{(\lambda_1 + \lambda_2)C_p}{\lambda_1 e^{(T - T_p)C_1} + \lambda_2 e^{-(T - T_p)C_2}}
\]

where \( C_p(T - T_p) \) refers to the thermal function of the specific heat capacity, NR signifies the numerator term, and DR1 and DR2 designate part 1 and part 2 of the denominator. \( \lambda_1, \lambda_2, C_1, \) and \( C_2 \) are the empirical constants which are to be determined by minimizing the discrepancy between the experimental data (DSC thermogram) and the mathematical formulation (equation 1).

There are a few similarities in the thermal behavior of hydrogels [9-10] and other polymers [17-18]. Crystallization of thermoplastics, curing of thermosetting resins and gelation of hydrogels are all governed by thermodynamic considerations. Crystallization and curing are exothermic reactions whereas gelation, depending on the type of hydrogel, can be either endothermic or exothermic process. Both crystallization and physical gelation are thermo-reversible processes. Thermosetting resins and hydrogels both have three-dimensional networks. Therefore, the term ‘degree of gelation’ (\( \alpha_g \)), similar to the degree of cure or the degree of crystallization, was introduced for quantitative assessment of the state of gelation [9-10]. Once the DSC data is mapped, \( \alpha_g \) at temperature T can be calculated as:
\[ \alpha_g = \frac{\text{Heat absorbed up to a stage for complete gelation}}{\text{Heat for complete gelation}} = \frac{\int_{T_{on}}^{T} f(T) dT}{\int_{T_{on}}^{T_{off}} f(T) dT} \]

Note that \( \alpha_g \) is a fraction that varies from 0 to 1 and is dimensionless. \( \alpha_g = 0 \) at \( T = T_{on} \) and when \( T = T_{off}, \alpha_g = 1 \). Equations 1 and 2 were successfully used to describe the DSC heating thermograms and to obtain the heat of gelation, the degree of gelation and the swell initiation points for MC solutions of different concentrations without any additives [10]. The significance of the various constants was yet to be elaborated and will now be discussed.

From equation 1, NR is a multiplier that is determined based on the scaling requirement for \( 1/(DR_1 + DR_2) \). The denominator terms control the spread and the shape of the \( C_p \) curve. \( DR_1 \) describes the decaying arm from \( T_p \) and \( T_{off} \) whereas \( DR_2 \) maps the growing arm from \( T_{on} \) to \( T_p \). Both, \( DR_1 \) and \( DR_2 \), have a form similar to the Arrhenius equation [19] describing the progress of a reaction as a function of temperature with time. Thus the associated constants could have similar significance. \( \lambda_1 \) and \( \lambda_2 \) are the frequency parameters with the unit \( 1/\degree C \). The terms \( C_1 \) and \( C_2 \) resemble \( R/\Delta E \), where \( \Delta E \) is the activation energy (J/mol) and \( R \) the universal gas constant (8.314 J/mol.K). Since gelation takes place only within a certain temperature window when the temperature is allowed to increase at a certain rate, the temperature difference holds the key for the reaction. Under isothermal conditions or at slow rate of heating, the gelation process appears to attain equilibrium rapidly with no further change in the state of gelation. This suggests that for a reasonably slow scan rate of 1 \( \degree C \)/min the temperature difference and not the duration of
the temperature change is the important parameter for gelation modeling. It was therefore possible to establish a model based on the usual temperature unit °C. The units of $C_1$ and $C_2$ in the current case thus remain as $1/°C$, which cancel out with the temperature units for $(T-T_p)$ terms. $DRI$ and $DR2$ will have the same unit as the $\lambda$ terms. Finally, $C_p(T-T_p)$ would have the same unit as that of $C_{p_{\text{max}}}$ (J/mol °C) as required.

Equations 1 and 2 were first used to describe the $C_p$ values and $\alpha_g$ for MC solutions with 0.2M and 0.8M NaCl. The comparison between the model and the experimental data are shown in Figure 3. It is observed from these plots that the basic model, namely equation 1, could not describe the process well around the shoulders of the $C_p$ curves. Rather large discrepancies between the model and the experimental data manifested themselves at the beginning and towards the end of gelation. These differences were reflected for the degree of gelation curves also. It is observed that the discrepancies increased with increasing NaCl concentration. Thus the model, which reasonably predicted the gelation process for MC without salt, was unable to describe the heat absorption mechanism successfully for sol-gel transformations for aqueous solution of MC in the presence of NaCl.

These discrepancies suggest that there could be additional mechanisms taking place due to the addition of NaCl. By hypothesizing that these additional mechanisms follow a similar trend for the rate of energy absorption, and that the effects of more than one binding process and mechanism can be superimposed, an expression similar to equation 1 can be added to describe these mechanisms. Thus, in order to account for the effects of
the salt additive, a model that consisted of two expressions, with each similar to that of equation 1, was employed; see equation 3.

\[ C_p(T - T_p) = \frac{(\lambda_1 + \lambda_2)C_{p1}}{(T - T_{p1})^C1} + \frac{(\lambda_3 + \lambda_4)C_{p2}}{(T - T_{p2})^C2} \]

Where \( \lambda_1, \lambda_2, \lambda_3, \lambda_4, C_1, C_2, C_3 \) and \( C_4 \) are the empirical constants. Note that \( C_{p\text{max}} \) and \( T_p \) used in equation 1 are replaced by \( C_{p1}, T_{p1}, C_{p2} \) and \( T_{p2} \) in equation 3. This two-expression model was expected to provide two bell-shape curves, which when superimposed, would closely follow the experimental DSC curves.

The constants for the equation could be determined by minimizing the discrepancies between the model (equation 3) and experimental data. The least square technique was adopted in which the values of the constants were obtained by iteratively minimizing the sum of square of errors in \( C_p \) values calculated as:

\[ \sum_{i=1}^{N} [C_{p(\text{expr})} - (C_{p1(\text{model})} + C_{p2(\text{model})})]^2 \rightarrow 0 \]  \hspace{1cm} (4)

where, \( i = 1 \) at \( T = T_{on} \) and \( i = N \) when \( T = T_{off} \). This generated two bell-shape curves with \( C_{p1} \) and \( C_{p2} \) as their respective peak \( C_p \) values.

### 4.0 Results and Discussions

Figure 4 shows the heat capacity curves obtained using the model (equation 3) and its comparison with the DSC thermogram data for different concentrations of NaCl. The empirical constants for the model are listed in Table 1.
As NaCl is an electrolyte, the Cl⁻ ions compete with the MC chains for hydrogen (H) bonding with water molecules, which promotes early exposure of hydrophobic groups and rapid formation of MC aggregates at lower temperature. This, in turn, results in a lower $T_p$ that decreases monotonically with an increase in salt concentration [14].

All the curves exhibited the trend that the dominant or major bell-shape curves was always able to map most of the experimental data and was designated as the primary curve. The secondary curve, which always appeared small, however, was necessary to provide good fit to the left- and right-hand shoulders of the curve. As mentioned earlier, the low temperature activities are dominated by water-to-water and water-to-MC hydrogen bonding. The low-magnitude, initial portion of the primary curve describes the energy consumption causing disruption of these bonding processes. In a similar manner and in small magnitude also, the ionic interaction of salts with water, which is also through H bonding, is represented by the initial portion of the secondary curve. This reflects that such H bonding does not involve substantial amount of energy.

Following the breakage of water-cages surrounding the hydrophobic side groups of MC chains, these chains tend to move towards each other for association. However, these macromolecules of MC are considerably long in comparison with the molecules of NaCl and H₂O. It is expected that a substantial amount of energy is required to facilitate their movement in the solution. This justifies the substantial and fast rise in $C_p$. As the MC molecules come closer, they start to form groups through a few hydrophobic bonds. These groups further join together and start to form a network. This initial phase of hydrophobic association is referred to as non-cooperative association and requires more
energy as compared to the subsequent bonding. The subsequent hydrophobic association occurs among the already connected MC chains thereby increasing the density of the network cells. This type of bonding requires much less energy and called as cooperative association. As long as the process of non-operative association goes on, the energy absorption continues at high rate, marked by $C_p$ reaching $C_{pv1}$. It is anticipated that the cooperative association will go on for some more time until the network develops into a full three-dimensional structure. However, this stage has a much subsided energy requirement and the $C_p$ value starts to recede. Gradually, as temperature increases, the network cells start to take in more water molecules and hold them within. Because of this expansion or swelling of the cells and reduction in their flexibility, the gel starts to exhibit higher and higher elastic properties.

The peak of the second curve shifted towards the right indicating increased energy utilization with salt additives during the swelling of the gel network and the gel stabilization phase. The gradual shift of the peak to the right and the increase in the magnitude of $C_{pv2}$ with higher NaCl concentration may be attributed to the increasing interaction between the salt ions and water molecules that delayed water molecules from entering the orderly state within the gel network. This is evident from Figure 4 for all NaCl concentrations. As the salt concentration increased, higher $C_{pv2}$ values were observed indicating a rise in the heat energy absorbed. With additional interactions between the salt ions and the water molecules, the total enthalpy change, $H_T$, was higher for higher NaCl concentrations. It is evident from Figure 5 that $H_T$ values increased linearly with NaCl concentration. The trend-line as depicted by equation 5 shows that an additional energy of 1067.64 kJ was required per 0.2M/mol of NaCl.
\[ H_f \text{ (kJ/mol)} = 3815.7 + 5338.2 \times (NaCl \text{ Concentration in M}) \]  

(5)

However, it could not be ascertained whether the original energy requirement for MC reduced in the presence of the salt.

The plots of the degree of gelation versus temperature for all NaCl concentrations (determined using equation 2) are shown in Figure 6. The \( \alpha_g \) curves approximately have the same slope between \( \alpha_g = 0.1 \) and 0.6. This implies that the rate of gelation was not affected by the presence of NaCl during this phase. This was the window when the MC macromolecules were moving closer and the development of the gel network was in progress.

It is evident that the formulation presented in equation 3 could describe the energy utilization for MC gelation in the presence of NaCl. However, no specific trend for the various constants listed in Table 1 was observed. One of the primary reasons might be the mechanism and the transformations instigated by a specific amount of NaCl salt added to MC solution. Although the total heat of gelation appeared to follow a linear growth with salt concentration (Figure 5), the internal mechanisms may not be sharing the energy in the same proportion.

**5.0 Sigmoidal Model for Degree of Gelation Curves**

Being a representation of a transformation process in a bio-polymeric material, the degree of gelation curves are rather similar to the degree of cure curves for thermosetting polymers [20] and degree of crystallization curves for thermoplastics [21]. The difference is that gelation of MC is driven predominantly by the change of temperature, and the time
parameter is of little significance. As such, one could map $\alpha_g$ curves directly instead of trying to model the rate of change in $\alpha_g$. A good candidate is the sigmoidal curve of the logistics curves family. It is known to describe ‘S’ curves well and has been used for biomedical and biomaterials related studies [22-23]. The following sigmoidal growth curve (equation 6) was found to describe $\alpha_g$ curves very well for gelation of MC with NaCl additive.

$$
\alpha_g(T - T_p) = \frac{(T - T_{on})}{(T_{off} - T_{on})} \left[ \psi_1 + \xi \frac{e^{(T - T_p)B_1}}{\psi_2 + \xi e^{(T - T_p)B_2}} \right]
$$

Note that six empirical constants were employed in an unconstrained manner deliberately to avoid an over-constrained equation that could result in poor data fitting. $\left( \frac{T - T_{on}}{T_{off} - T_{on}} \right)$ is a temperature range fraction to enforce the limiting conditions. Similar to the $C_p(T - T_p)$ function, empirical parameters in equation 6 are analogous to the Arrhenius terms. $\xi_1$ and $\xi_2$ are the frequency parameters and carry the unit $1/°C$. $\psi_1$ and $\psi_2$ also have the same unit $1/°C$. $B_1$ and $B_2$ are the activation energy parameters and their units are the same as $R/\Delta E$ terms. Finally, all the numerator and denominator terms in equation 6 bear the unit $1/°C$ rendering the function $\alpha_g(T - T_p)$ non-dimensionless as required.

The same least-square-error-fit procedure as discussed in Section 3 was adopted to map $\alpha_g$ values calculated using equation 2 based on $C_p$ curves modeled using equation 3. Error minimization was achieved by employing equation 7:
The results for all the cases are shown in Figure 7. Good agreement was established with the analytically derived values of $\alpha_g$ using the $C_p$ model, which was earlier found to agree well to the experimental data. The values of all the empirical constants are tabulated in Table 2. It can be observed from Table 2 that for the no-salt case, $\psi_1$ term was not essential. This implied that parameter $\psi_1$ was associated with the salt-out effects of NaCl. Thus, the same conclusion is reached as before that with the addition of salt, an additional mechanism is involved in the gelation process. Similar to the $C_p$ model (Table 1), no clear trend was observed for the sigmoid curves constants for different salt concentrations.

An attempt was made to estimate the sol-gel activation energy parameters for the degree of gelation ($\Delta E\alpha_g$) by correlating $B_1$ and $B_2$ with $R/\Delta E$ terms. The calculated $\Delta E\alpha_{g1}$ and $\Delta E\alpha_{g2}$ values, which are based on the normalized energy represented by $\alpha_g$, are shown in the last two rows of Table 2. These energy parameters are the process-specific remainders obtained after eliminating the effects of the process-independent universal gas constant. Thus, they relate more directly to the gelation process and the mechanism of sol-gel transformation.
6.0 Conclusions

A two-bell shaped model was developed for simulating the $C_p$ curves defining the heat absorption during gelation of MC in the presence of NaCl additives. The import of various constants in terms of gelation kinetics was identified. The formulation worked well and could describe the MC gelation with different % of NaCl. Out of the two bell-shaped curves, which upon superimposing described the $C_p$ variations with temperature, the dominant curve always mapped most of the experimental data and was designated as the primary curve. It appeared that the energy described by the curve was utilized mainly for the movement of MC macromolecules and their hydrophobic association leading to a gel network. The secondary curve, which always accounted for a small amount of energy, however, was necessary in order to describe accurately the left and right hand shoulders of the curve. This appeared to represent the energy utilized in the ionic interaction of salt with water and its effects during the stabilization of the gel.

A sigmoidal model with the appropriate units for each terms was formulated to describe the degree of gelation, $\alpha_g$, for all cases. The correlation between the values derived using the $C_p$ curves and the sigmoidal model was excellent. All $\alpha_g$ curves had approximately the same slope, except at the beginning and towards the end of gelation. This implied that the rate of gelation was not affected by the presence of NaCl during the phase when the movement of MC chains was a dominant activity.
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\[ y = 5338.2x + 3815.7 \]
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Table 1: The model constants for $C_p$ curves (Equations 1 and 3)

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Table 2: The sigmoidal model constants for $a_\phi$ curves (Equation 6) and the sol-gel activation energy parameters for MC gelation

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