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DRYING AND SHRINKAGE OF POLYMER GELS

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Abstract - The polymer hydrogel was synthesized by photo-polymerization process (UV light, 60 $^{\circ}$ C) in presence of Photo-initiator (Irgacure^R) and Cross-linker (NN'-methylene bisacrylamide; MBAM). In the present work, the drying of polymer hydrogel was carried out to study the effect of temperature, gel-sheet thickness, monomer ratio of acryl acid to acrylamide (AA/AM), concentration of MBAM and quantity of monomers. A correlation has been developed for modified sheet thickness as a function of contraction coefficient and degree of drying. Effective diffusivity was estimated from Fickian-diffusive model considering modified sheet thickness and was found to be in the range of 1.1×10^{-10} - 5.93×10^{-10} m²/s. The activation energy obtained using Arrhenius type equation was found to be in the range of 2979-10737 kJ/kmol H₂O. The drying behavior shows an initial shoot-up in drying rate followed by constant rate and two falling rate periods.

Keywords: Polymer gel; Drying characteristics; Diffusion coefficient; Modified gel thickness; Contraction coefficient.

INTRODUCTION

Hydrogels, which are crosslinked hydrophilic polymers, are used in the areas like biotechnology, medicine, pharmacy, agriculture, food industry and others. The hydrophilicity of hydrogels is attributed to the presence of hydrophilic functional groups such as alcohols, carboxylic acid, and amides. The understanding of the phenomena occurring in the drying of these hydrogels is necessary for the designing of the entire process. Drying of hygroscopic solids is generally known to be a complex process because of simultaneous heat and mass transfer and variable physical properties of the materials (Hawlader, 1996). Polymers and gels are highly hygroscopic and shrinking colloidal materials and the drying of these encompass many fields of technology. The quantity of bound water associated with polymer varies as per the internal structure of the macromolecule. Monomer and cross-linking agent proportions are responsible for both the porous structure and the pore size of the gel. An acrylamidederived gel is a cross-linked network of polymer whose molecular weight is fairly high; it can absorb solvent (water), but is itself insoluble.

During the water migration in drying process, shrinkage corresponds simply to the compacting of solid mass. Jomma et al. (1991) have reported a diffusive drying model for the drying of highly shrinking materials like polyacrylamide gel and cellulosic paste. Bonazzi and Ripoche (1996) have used a conventional diffusive model with variable effective diffusivity to evaluate the moisture diffusivity in a water-gelatin system as a model for non-porous, shrinking colloidal and thermoreversible materials. Okos et al. (1992) have reported many different functional forms of the relationship between moisture content and effective diffusivity.

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Recently approximate models have been used by Coumans (2000) to predict the drying kinetics for geometry. Diffusion coefficient slab in polymer/solvent system over a wide range of concentrations and temperatures can be obtained by using a "free volume theory" of Vrentas and Duda (1979). In the present work, drying of acrylic acidacrylamide copolymer was carried out under various operating conditions. The drying data has been analyzed using Bonazzi and Ripoche (1996) model with altercations in modified gel thickness which is a of gel moisture content. strong function Conventionally, the gel thickness is given in terms of dry gel thickness and average moisture content is related by contraction coefficient. This is given by Eq. (1) as, (Bonazzi and Ripoche, 1996; Gehrmann and Kast, 1978)

$$Z = Z_{\rm dry} \left(1 + \varepsilon X \right) \tag{1}$$

Where, Z, ε and X are the gel thickness at any time, coefficient of contraction and moisture content respectively. However, it is difficult to know Z_{dry} apriori and therefore, we propose the following Eq. (2) to relate the gel thickness with the initial gel thickness and degree of drying (which can be easily obtained during the course of experiments).

$$Z = Z_{o} \left(1 - \varepsilon D_{d} \right)$$
⁽²⁾

Where, D_d is the degree of drying (discussed later).

MATERIALS

Acrylic acid (AA), acrylamide (AM) and methyl methacrylate (MMA) (Wardhaman Polymers Ltd., Mumbai, India.) used were of technical grade. Irgacure^R (Ciba specialty chemicals Inc., Mumbai, India) was of analytical grade, while NN'-methylene bisacrylamide was prepared in the laboratory. Potassium hydroxide (s. d. fine chemicals, Mumbai, India) was of commercial grade. Distilled water was used for sample preparation.

EXPERIMENTALS

The copolymer gel was synthesized using photopolymerization process. Drying experiments were carried out in a vacuum tray dryer where the temperature and vacuum could be maintained independently. For drying experiments nine samples were prepared (Table 1) to study the effect of temperature (samples 1-3), initial sheet thickness (samples 2, 4, 5), concentration of cross-linker (samples 2, 6, 7) and the monomer ratio (AA/AM) (samples 2, 8 9). The experiments were carried out at 80-120 $^{\text{O}}$ C, 4-8 mm sheet thickness, 0.156-0.312% MBAM concentration based on AA and 30/70-70/30 (%) monomer molar ratio. The data of moisture removed with respect to time was used to evaluate the drying characteristics. The initial and final sheet thickness was measured to find out the contraction coefficient.

RESULTS AND DISCUSSION

Drying Characteristics

The drying characteristics of the gel were evaluated from the data of moisture removed with respect to time. Figure (1) shows the plot of moisture content versus time, from which the drying curve was plotted by using Eq. (3) as,

$$N = -\frac{w}{A}\frac{dX}{dt}$$
(3)

Where, N, w, A and t are the drying rate, bonedry gel weight, surface area for drying and drying time respectively. Figure (2) shows a typical drying curve (sample 2) at 20 mm Hg. It can be seen that the drying curve consists of an initial shoot-up followed by a wavy constant drying period and two falling rate periods. Since, the formation of crust after some time at the surface of the gel during the drying leads to decrease in the drying rate in the early stages. In the constant rate period, the concentration gradient, dX_{dZ} , required for the moisture removal remains constant and the decrease of moisture content takes place with the corresponding decrease in the sheet thickness. Also in gel drying, it can be seen that small vapor pockets were formed just below the drying surface and then migrate towards the drying surface where they break, release the vapors and form small cavities at the surface. As the formation and collapse of these vapor pockets were irregular, the drying rate also changes and therefore the wavy nature of the drying curve. As the drying proceeds, moisture level goes down and polymer sheet becomes harder, which results into decreased drying rate and hence the two falling rate periods.



Figure 2: Characteristic Drying Curve

Moisture Effective Diffusivity and Drying Rate

The drying rate was calculated from the experimental data of moisture removed versus time by using Eq. (3). The drying rate was then used to calculate the moisture effective diffusivity by applying Fick's law with modified sheet thickness (Z), (discussed later) as,

$$N = -\rho_s D_{eff} \frac{dX}{dZ}$$
(4)

Where, ρ_s and D_{eff} are the true polymer density and effective moisture diffusivity respectively. From Table 1, it can be seen that the moisture diffusion coefficient and hence the drying rate increases with an increase in the drying temperature from 80 $^{\rm O}$ C to 120 $^{\rm O}$ C. The relationship between the effective diffusivity and the drying temperature can be given by the Arrhenius type equation. From Table 1 it can be said that effective diffusivity and drying rate decreases with increase in the concentration of cross-linker from 0.156 to 0.624 % (mole) based on acrylic acid. As the higher concentrations of crosslinker leads to hardening of the gel due to increased cross-linkage between the polymeric chains, the escape of the moisture through the gel becomes difficult. Also from Table 1, it can be said that the effective diffusivity and the drying rate decreases with an increase in AA/AM ratio from 30/70 to 70/30. The probable reason for the decrease in the effective diffusivity with an increase in AA/AM ratio may be attributed to the increase in the number of carboxylic groups within the polymer gel, which are having the tendency to retain more water than the amides.

Sample No.	T (OC)	Z _o (mm)	AA/AM (%)	MBAM (%)	$N_c \times 10^4$ kg/m2s	$D_{eff} \times 10^{10}$ m2/s	CMC kg/kg	EMC kg/kg	E kJ/kmol
1	80	4	30/70	0.312	1.818	0.484-2.47	1.03	0.15	4783
2	100	4	30/70	0.312	3.10	1.24-3.71	1.25	0.143	3398
3	120	4	30/70	0.312	4.849	3.54-5.93	1.36	0.064	1698
2	100	4	30/70	0.312	3.10	1.24-3.71	1.25	0.143	3398
4	100	6	30/70	0.312	3.03	-	1.15	0.252	-
5	100	8	30/70	0.312	2.22	-	0.85	0.333	-
6	100	4	30/70	0.156	3.30	2.12-4.0	1.34	0.125	1969
2	100	4	30/70	0.312	3.10	1.24-3.71	1.25	0.143	3398
7	100	4	30/70	0.624	2.87	0.49-3.62	1.19	0.225	6202
2	100	4	30/70	0.312	3.10	1.24-3.71	1.25	0.143	3398
8	100	4	50/50	0.312	3.14	0.82-3.58	1.30	0.1	4578
9	100	4	70/30	0.312	3.27	0.685-3.5	1.61	0.033	5058

Table 1: Effect of Temperature (T), initial sheet thickness (Z₀), monomer ratio of AA/AM and %crosslinker (MBAM) on Nc, Deff, CMC, EMC and activation energy.

Critical Moisture Content (CMC) and Equilibrium Moisture Content (EMC)

CMC can be obtained from the drying rate curve. An empirical approximation (Gehrmann and Kast, 1978) of the drying rate curve was used to confirm the obtained values of CMC. This approximation states that if the product of drying rate and slab thickness (which depends on moisture content) referred to the data of the constant rate period is plotted versus average moisture content (X) normalized with CMC (X_c), the results is a relationship with two nearly linear portions, divided by a slight break in the curve at $\frac{X}{X_c} = 1$. The first

linear part of the curve is due to the proportionality of the sample thickness to average moisture content X while the drying rate during the first drying period is constant. Figure (3) shows this approximation for sample 4.



Figure 3: Identification of Critical moisture content (CMC)

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Table 1 shows the effect of drying temperature on the CMC and EMC. It was observed that CMC increases and EMC decreases with an increase in drying temperature. High constant drying rate at higher temperature develops a steep concentration profile within the solid. During low-intensive evaporation of moisture (corresponding to low drying temperature) from the surface of the material, major part of moisture will migrate to the surface layer of evaporation before reaching the level of EMC. In this case, the moisture content profile in the material appears more equalized and therefore CMC is lower. The effect of cross-linker concentration on CMC and EMC is given in Table 1. It can be seen that CMC decreases and EMC increases with an increase in concentration of cross-linker. The probable reason for this observation is that increased concentration of MBAM harden the gel material which results into decrease in the drying rate and hence the CMC. Also because of the increase in cross-linking density of copolymer, most of the moisture trapped inside the cross-linked polymer chain inhibits lower vapour pressure, which results into an increase in EMC. Table 1 also shows that both CMC increases and EMC decreases with an increase in AA/AM form 30/70 to 70/30. It is also seen that CMC decreases and EMC increases with an increase in the gel thickness from 4 to 8mm. As the thickness of the gel material in the tray increases, there exists a temperature profile inside the material. The temperature of material surface is same as the heating medium and decreases with distance from the surface towards the center of the material. Also the moisture at the bottom of the tray has lot of diffusional resistances and it increases with an increase in the thickness of the gel material.

Modified Gel Thickness (Z)

As copolymer gel shrinks during drying, the thickness through which the moisture diffused also changes with moisture content. Therefore, in order to calculate the correct values of the diffusion coefficient, the variation of gel thickness with moisture content and hence the degree of drying must be taken into account. The extent of contraction of gel depends on the drying temperature, monomer ratio, cross-linker concentration and initial gel thickness. For all the samples, the extent of contraction (hereafter, contraction coefficient, ε) was measured and a correlation was developed. Eq. (5). For the given experimental conditions (drying temperature, T; MBAM concentration, C; monomer ratio, R and initial gel sheet thickness, Zo etc.), the contraction coefficient varied in the range of 0.5-0.77. Figure (4) shows the parity plot between the correlation and the experimental results.

$$\varepsilon = 0.0107 \times T^{0.8196} R^{0.175} C^{-0.243} Z_0^{0.075}$$
(5)

The modified length (Z) has been calculated by using contraction coefficient (ε), degree of drying (D_d) and initial gel sheet thickness (Z_o) as in Eq. (2). Where the degree of drying (D_d) is given as,

$$D_{d} = \frac{X}{X_{o}}$$
(6)

Where, X_o is the initial moisture content.



Figure 4: Parity plot for coefficient of contraction

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Activation Energy (E)

The activation energy can be calculated with the help of Arrhenius type equation:

$$D_{(X,T)} = D_{o(X_1)} e^{\frac{-E(X_1)}{RT}}$$
(7)

Where, D_o is the moisture diffusivity at constant drying period and E is the activation energy for moisture diffusion. Figure (5) shows the plot of activation energy versus average moisture content. The values have been tabulated in Table 1. The activation energy of diffusion for all the samples obtained was found to be in the range of 2979-10737 kJ/kmol H₂O.

Porosity (ϕ) of the Dried Material Sheet

Calculation of the porosity is based on the assumptions that the volume of solid phase is constant and that the volume of the solid phase (V_s), the liquid phase (V_m) and the void (V_c) are additive quantities expressed as:

$$V_{\rm T} = V_{\rm s} + V_{\rm m} + V_{\rm c} \tag{8}$$

The creation of void volume and hence the porosity of the sheet depends on the drying conditions and system parameters. Figure (5) shows that the increase in the porosity of sheet as the drying proceeds. Figure (6) shows the porous structure of the dried sheet.



Figure 5: Activation energy and Porosity as a function of moisture content



Figure 6: Porous structure of the dried polymer sheet by using Image Analyzer

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True Density of Polymer Sheet

From Figure (7) it can be seen the true density of polymer sheet increases as drying proceeds. During the drying the solid mass compactness resulting into increase in the true density of the sheet provided that the assumption, the volumes of cavities, moisture and solid are additive is valid.

Identification of Shrinkage Behavior

A physical model describing the volume reduction factor (α) to identify the shrinkage behavior of the polymer gels is given as below,

1.8

$$\alpha = 1 + \frac{\Delta V_c}{\Delta V(\langle w \rangle)} \tag{9}$$

Where ΔV_c and $\Delta V(\langle w \rangle)$ are void volume and volume of the sheet at average moisture content $\langle w \rangle$ respectively. Figure (7) also shows the plot of reduction factor (α) versus average moisture content (X). It is observed that α increases with decrease in moisture content. This indicates that the volume of cavities formed increases as drying proceeds. It was observed that initially the value of α is close to one and as drying proceeds α deviates from unity and hence non-ideal shrinkage behavior of the gel.

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Figure 7: Reduction factor and true density of a co-polymer gel as a function of moisture content

CONCLUSIONS

The drying behavior of hydrogel has been characterized in terms of CMC, EMC, effective diffusion coefficient, activation energy and shrinkage behavior. It can be concluded that the shrinkage of the polymer gel is non-ideal. As poly (AA-co-AM) hydrogel with higher monomer ratio of AA/AM and MBAM concentration retain more water hence can be used as water retainer in various applications in pharmaceuticals, health care products. Effect of crosslinker has been found to be more significant with respect to drying kinetics.

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NOMENCLATURE

А	drying surface area	(m^2)
С	concentration of cross-linker	(wt %)
D _d	degree of drying	(-)
D	diffusion coefficient	(m^2/s)
E	activation energy	(J/mol)
Ν	drying rate	$(kg/m^2.s)$
R	monomer ratio	(AA/AM)
R	universal gas constant in Eq. (7)	(-)
Т	drying temperature	(K)
t	drying time	(s)
V	volume	(m^3)
W	bone-dry solid weight	(kg)
<w></w>	average moisture content	(kg/kg)

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Х	moisture content	(kg/kg)
Ζ	polymer gel thickness	(m)

Greek Letters

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α	reduction factor	(-)
3	contraction coefficient	(-)
ρ	density	(kg/m^3)

Subscripts

c	critical or constant	(-)
m	liquid phase	(-)
0	initial	(-)
S	solid phase	(-)
Т	total	(-)
W	water	(-)

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