# Determination of Monomer Reactivity Ratios of Poly (Acrylonitrile – co – vinylacetate)

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(Received n 15/11/2006) (Accepted for publication on 22/8/2007)

# Abstract :

Copolymers of Acrylonitrile (AN) and Vinyl Acetate (VAc) were synthesized in dimethyl formamide (DMF) using benzoyl peroxide (BPO) as initiator at 75°C. The copolymers were analyzed by IR spectroscopy. The monomer reactivity ratios were determined by the methods of Fineman – Ross (F-R) and Kelen – TűdÖs (K-T) giving the results shown in the following table:-

	$\mathbf{r}_1(A)$	AN)			r <sub>2</sub> (	VAc)	
F-	·R	K	-T	F-	R	K	-T
1 <sup>st</sup>	2 <sup>nd</sup>	1 <sup>st</sup>	2 <sup>nd</sup>	1 <sup>st</sup>	2 <sup>nd</sup>	1 <sup>st</sup>	2 <sup>nd</sup>
2.2347	2.5341	2.3889	2.7087	0.0303	0.002	0.1038	0.0719

#### الخلاصة

تم تحضير البوليمر المشترك للاكريلونايتريل (AN) و الفنيل استيت (V<sub>Ac</sub>) في مذيب ثنائي مثيل فورمأيد (DMF) باستخدام بيروكسيد البنزويل كبادئ وبدرجة حرارة 75°c . تم تحليل البوليمر المشترك بواسطة مطيافية الاشعة تحت الحمراء .

الفعالية النسبية للمونمرات تم تحديدها بواسطة طريقة فنمان و روس (F-R) وطريقة كلين-ثيودوس (K-T) و النتائج موضحة في الجدول التالي :

	$r_1(A$	AN)			r <sub>2</sub> (	VAc)	
F-	R	K	-T	F-	R	K	-T
1 <sup>st</sup>	2 <sup>nd</sup>	1 <sup>st</sup>	2 <sup>nd</sup>	1 <sup>st</sup>	2 <sup>nd</sup>	1 <sup>st</sup>	2 <sup>nd</sup>
2.2347	2.5341	2.3889	2.7087	0.0303	0.002	0.1038	0.0719

#### Introduction:

Japan has recently been the leading contributor in the diversification of the production of modified acrylic fibers(1) More than 85% of acrylonitrile monomer must be incorporated into copolymer in order for it to be termed an acrylic fiber(2). Copolymerization of acrylonitrile with various comonomers produces specialty fibers for different applications. Methyl acrylate, methyl methacrylate and vinyl acetate are the commonly used "neutral" comonomers. They increase solubillity and change the morphology of the fiber(3). The theory of radical copolymerization leads to the conclusion that the composition of the copolymer is determined by the reactivity ratios (r1 and r2) which reflect the inherent tendencies of a radical to react with its own monomer relative to the comonomer, copolymerizations are classified according to values of the product  $r_1r_2$ . When  $r_1r_2=0$ , an alternating copolymer is produced, when  $r_1r_2=1$ , the copolymerization is said to be ideal, since  $r_1 = \frac{1}{r_2}$ , the copolymer is random.

Many methods have been used to estimate reactivity ratios of a large number of comonomers but the method of increasing application are those of Fineman- Ross and Kelen – TÜdÖs (4). 

<sup>1</sup>H–NMR, 

<sup>13</sup>C–NMR and FTIR methodologies are used to determine the copolymer composition(5).

The aim of the present work is to evaluate the reactivity ratios of comonomers using two methods, both of them depend on the absorption and absorptivity of certain group in the monomers and copolymers.

## Experimental:

## 1- Materials and purification

Acrylouitrile (AN) (Aldrich) was washed with (10% NaOH) to remove inhibitors and then washed with distilled water to remove any trace of NaOH and finally distilled at reduced pressure. Vinyl acetate (VAc) was distilled before use. Dibenzoyl peroxide BPO (Aldrich ) was purified by dissolving in small amounts of chloroform followed by precipitation with n - hexane.

#### 2- Synthesis

For copolymerization, the AN and VAc in various proportions but with the total weight maintained at 4g and the initiator BPO being kept at concentration of (3×10<sup>-4</sup> mol/L) were dissolved in 25 ml of DMF. The content of the flask was degassed with argon for at least 2 minutes and polymerization was carried out at 75°C for a period of 30 minutes. Each copolymer was

## Results and Discussion

# 3-Determination of Molar Fractions (F)

The molar fractions of (AN) and (VAc) in poly (AN-co-VAc) were determined by two methods:-

$$F_{1} = \frac{ACN / M_{1}}{ACN / M_{1} + Aco / M_{2}}.....(1) \qquad F_{2} = \frac{Aco / M_{2}}{Aco / M_{2} + ACN / M_{1}}.....(2)$$

Where A is the absorbance of and CO at 2240,1720 cm-1 respectively. F1 and F2 are the molar fractions of (AN) and (VAc) in the

#### b) The second method

This method depends on the absorbance of the recorded analytical absorption bands for CN and CO for

$$F_1 / F_2 = (A_{CN} / A_{CO}) (\epsilon_{CO} / \epsilon_{CN}) ....(3)$$

precipitated by ethanol, washed repeatedly in methanol and dried. For the determination of reactivity ratio, the polymer conversions were always less than 5%. Homopolymers of AN and VAc were also prepared and their absorbances were measured using liquid cell with path length of 1cm. The IR spectra copolymers of homopolymers were recorded Pye Unicam (SP3-100 IR) as films KBr window, figure (1).on

## a) The first method

This method depends on the absorbance of the recorded analytical absorption bands that is CN and CO for (AN) and (VAc) respectively, besides the molecular weights of the (AN) and (VAc) according to the following equations(6):-

$$F_2 = \frac{Aco / M_2}{Aco / M_2 + ACN / M_1}.....(2)$$

copolymer respectively, M1 and M2 are the molecular weights of (AN) and (VAc) respectively.

(AN) and (VAc) respectively, besides the molar absorptivities of the CN and CO groups according to the following equation(7).

Where F1 and F2 are the molar fractions of (AN) and (VAc) in the copolymer, while  $\epsilon$ CO and  $\epsilon$ CN are the molar absorptivity of CN and CO respectively. The  $\epsilon$ CN and  $\epsilon$ CO were found to be 1.7543 and 3.2166 L.cm-

1.mol-1 respectively. From the monomer feed ratio and the resultant copolymer composition, the reactivity ratios of (AN) and (VAc) were evaluated by Fineman – Ross and Kelen – TüdŐs equations that are:-

$$f(F-1)/F = r_1 \frac{f^2}{F} - r_2 \dots (4)$$
 Fineman - Ross   
  $\eta = (r_1 + r_2 / \alpha) \zeta - \frac{r_2}{\alpha} \dots (5)$  Kelen - TüdŐs

Where:

$$\eta = \frac{f(F-1)/F}{f^2/F + \alpha}$$
  $\zeta = \frac{f^2/F}{f^2/F + \alpha}$   $\alpha = \sqrt{(f^2/F) \max(f^2/F) \min}$ 

A plot of f (F-1)/F against  $f^2$ /F gives straight line, with slope =  $r_1$ , and intercept=  $-r_2$ .

A plot of  $\eta$  against  $\zeta$  gives straight line the slope =  $r_1+r_2/\alpha$  while the intercept =  $-r_2/\alpha$  the results are presented in the following tables:-

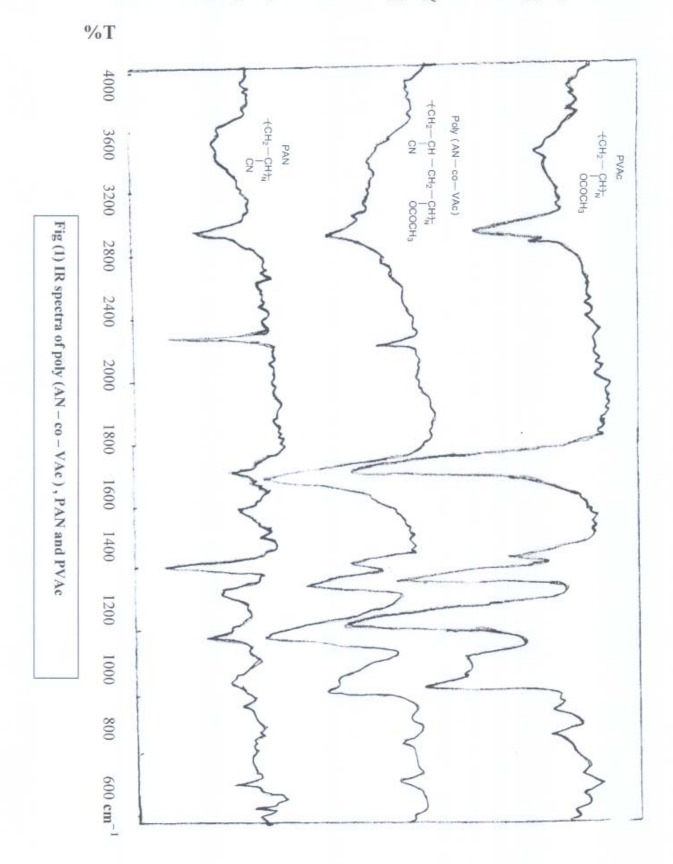


Table (1) IR analysis data for determining the composition of poly (AN - co-VAc ) prepared from various initial monomer mixture

Copolymer		raction of er in Feed	IR Abs	orbance		on of Monomer polymer
Copolymer	f <sub>1</sub> for AN	f <sub>2</sub> for VAc	AN CN	VAc CO	- F <sub>1</sub> for AN	F <sub>2</sub> for VAc
$A_1$	0.8295	0.1704	0.94	0.13	0.9214	0.0785
$A_2$	0.73	0.2699	1.33	0.31	0.87439	0.1256
$A_3$	0.6187	0.38129	0.46	0.16	0.8234	0.1765
$A_4$	0.4933	0.50669	0.378	0.16	0.7931	0.2068
A <sub>5</sub>	0.351	0.6489	0.183	0.14	0.67959	0.3204
$A_6$	0.1881	0.8118	0.108	0.24	0.422	0.5779

The following table shows the values of r1 and r2: -

Table (2): copolymerization constants r<sub>1</sub> and r<sub>2</sub> for poly (AN-co-VAc)

	(F-R) e	quation			(K-T) 6	equation	
1	st	2r	nd	1	st	21	nd
r1	r2	r1	r2	r1	r2	r1	r2
2.2347	0.0303	2.5341	0.002	2.3889	0.1038	2.7087	0.0719

From the values of r<sub>1</sub> and r<sub>2</sub> it seems that (AN) is more reactive than prefers to add (AN) more than (VAc).

(VAc), where the macroradical

Table (3) Fineman - Ross and Kelen - TüdŐs equations parameters of poly (AN - co- VAc) in DMF at 75°C when  $\alpha_{1st} = 0.3852 \ \alpha 2^{nd} = 0.3408$ 

		F=F	F=F1/F2	F-F	F-R Equation parameters	parameter	80	*	<-T Equati	K-T Equation parameters	ers
Copoly- mer	f=f <sub>1</sub> /f <sub>2</sub>			f2/F	[+:	f(F-	f(F-1)/F	-	n		πλ
		181	2 <sup>nd</sup>	18t	2 <sup>nd</sup>	$1^{st}$	2 <sup>nd</sup>	1 <sup>st</sup>	2 <sup>nd</sup>	1 <sup>st</sup>	2 <sup>nd</sup>
Aı	4.8679	11.7375	13.2576	2.0188	1.7873	4.4531	4.5007	1.8523	2.11489	0.8397	0.8398
A2	2.7047	6.9617	7.8663	1.0508	0.9299	2.3161	2.3608	1.6128	1.8578	0.7317	0.7318
A3	1.6226	4.6651	5.2713	0.5643	0.4994	1.2747	1.3147	1.34249	1.5647	0.5943	0.5943
A4	0.9735	3.8351	4.3316	0.2471	0.2187	0.7196	0.7487	1.138	1.3381	0.39079	0.3908
As	0.5409	2.121	2.3966	0.1379	0.122	0.2858	0.3152	0.5463	0.681	0.2636	0.2636
A <sub>6</sub>	0.2317	0.7302	0.825	0.0735	0.065	-0.0856	-0.0491	-0.1866	-0.1209	0.1602	0.1601

The following figures show the Plots of Fineman - Ross and Kelen - TüdŐs

equations for first and second method:

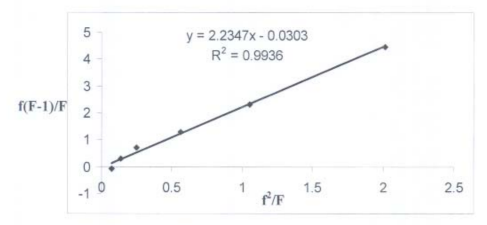


Fig (2) A plot of (F-R) (  $1^{st}$  method ) for the copolymer (AN-co-VAc)

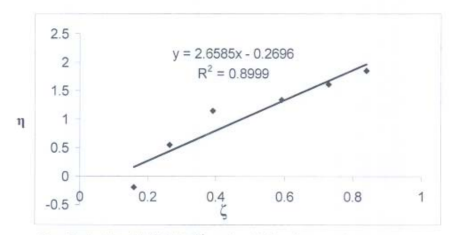


Fig (3) A plot of (K-T) (  $1^{st}$  method ) for the copolymer (AN-co-VAc)

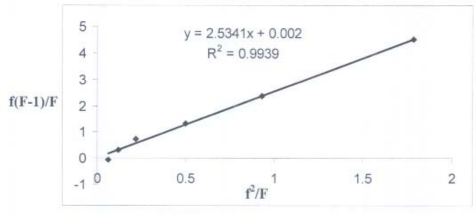


Fig (4) A plot of (F-R) (  $2^{nd}$  method ) for the copolymer (AN-co-VAc)

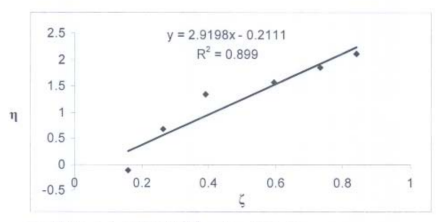


Fig (5) A plot of (K-T) (  $2^{nd}$  method ) for the copolymer (AN - co - VAc)

## 3-2 copolymerization behavior

We can predict the copolymer type (alternate, random or block) from

the product  $(r_1r_2)$  The values of this product are summarized in table (4):-

Table (4) the values of (r<sub>1</sub>r<sub>2</sub>) for poly(AN-co-VAc)

	$\mathbf{r}_1\mathbf{r}_2$		
(F-R) e	quation	(K-T) e	quation
1 <sup>st</sup>	2 <sup>nd</sup>	1 <sup>st</sup>	2 <sup>nd</sup>
0.0677	0.005	0.2479	0.1947

According to these values, the nature of the copolymer sequence of poly (AN-co-VAc) system is tending to alternate since the product r<sub>1</sub>r<sub>2</sub> closer to zero, so there is a greater tendency for (AN) and (VAc) to alternate in the copolymer poly (AN-co-VAc) due to big differences in polarity between the two monomers which leads to a transition state stabilized by resonance

which prefer to add either of the two monomers by the same preference.

Depending on the experimental values of  $r_{AN}$  and  $r_{VA}$ , the values of Q and e for each monomer were calculated and found as follows: for AN, Q= 0.539, e = 1.67, for VA, Q= 0.0339, e= -0.189 these values are in good agreement with values in literature(8). The copolymer

composition  $(F_1)$  vs initial composition  $(f_1)$  (fig 6), shows no

resemblance between the feed and the copolymer compositions. From the values of  $r_{AN}$  and  $r_{VA}$  one can deduce that AN is more reactive than VAc to go into copolymer.

This behavior can be explained on the basis that AN radical is more stable than VAc radical so the growing macroradical prefer to add to AN more than VAc.

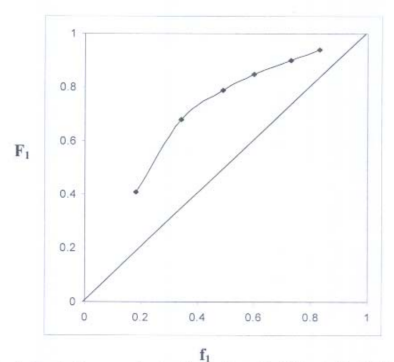


Fig (6) The relation between mole fraction of acrylonitrile in feed  $(f_1)$  and in copolymer  $(F_1)$ .

Finally, we can conclude that the two methods for reactivity ratio measurement used in the present work

are good methods for prediction of copolymer composition and reactivity ratios.

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