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2	Insight into miscibility behaviour of cellulose ester blends with N-vinyl pyrrolidone
3	copolymers in terms of viscometric interaction parameters
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14 Abstract: We previously offered miscibility maps for blend systems of cellulose esters (CEs) including cellulose acetate (CA), propionate (CP), and butyrate (CB) with vinyl 15 16 copolymers containing *N*-vinyl pyrrolidone (VP) unit, an i.e., poly(*N*-vinyl 17 pyrrolidone-co-vinyl acetate) (P(VP-co-VAc)) and poly(N-vinyl pyrrolidone-co-methyl 18 methacrylate) (P(VP-co-MMA)); the maps were constructed based on data of thermal analysis 19 as a function of the degree of ester substitution (DS) of the CE component and the VP fraction 20 in the copolymer component. The blend system using CP among the three CEs imparted the 21 largest region of miscible pairings with the vinyl copolymers, and both of the maps for the 22 CP/P(VP-co-VAc) and CP/P(VP-co-MMA) systems comprised a "miscibility window" 23 associated with the respective copolymer compositions at high DSs of >2.65. The present work was made to interpret the expansion of the miscible markings for the CP/copolymer 24 systems in comparison with the cases using CA and CB, in terms of a Krigbaum-Wall 25 26 interaction parameter (μ) obtained by solution viscometry for selective polymer pairs involved in the respective CE/copolymer blends. The results of μ measurements were in good 27 28 accordance with the earlier miscibility estimations. The assessment of very small negative μ 29 values (i.e., extremely weak repulsion) for CP/PVAc and CP/PMMA combinations and that of considerably larger negative μ values for PVP/PVAc and PVP/PMMA combinations enabled 30 31 us to give a rational explanation for the CP systems. The strongly repellent character of the 32 two different monomer units constituting the copolymers permits accession of the CP component (DS > 2.65) to them, which would be responsible for the advent of the miscibility 33 34 window. Further expansion of the window observed when cellulose acetate propionate (CAP) was adopted instead of CP as the CE component was also well explained on the basis 35 36 of a μ data indicative of additional intramolecular repulsion in the CAP side.

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38 Keywords: Blend miscibility; Cellulose ester; Interaction parameter; Miscibility window;
 39 *N*-Vinyl pyrrolidone

43 Organic esters of cellulose (CEs) are commercially important polymers over nearly a century. 44 They are widely prevailing in application fields such as coating, drug delivery (excipients), 45 molded plastics including biodegradable ones, fibers, optical films, and membranes and other separation media (Edgar et al. 2001; Rustemeyer 2004). For improvement in physical 46 47 properties of CEs toward their further applications, the designing of high-functional 48 multicomponent materials based on the cellulosics via graft copolymerization or polymer 49 blending is a significant approach (Edgar et al. 2001; Nishio 2006; Yamaguchi 2010; 50 Suginura et al. in press). In the field of optical materials such as regulator or modulator of 51 polarized light in modern displays, great attention of researchers has been focused on the 52 delicate control of orientation birefringence and its wavelength dependence for CE-based 53 films (Ohno and Nishio 2007a; Yamaguchi 2010; Yamaguchi et al. 2012; Yamanaka et al. 54 2013; Sugimura et al. 2013b; Hayakawa and Ueda 2015; Sugimura et al. in press). 55 Especially, miscible polymer blending is practically useful to manipulate the physical 56 properties and functions of CEs readily at the lowest cost possible. Therefore, there have 57 been a number of fundamental and practical blend studies of CEs; the counter components to 58 CEs are categorized into mainly two sorts of polymers, biodegradable aliphatic polyesters 59 such as poly(3-hydroxybutyrate) and poly(ε -caprolactone) (Nishio et al. 1997; Edgar et al. 2001; Nishio 2006; Kusumi et al. 2008; Higeshiro et al. 2009), and synthetic vinyl polymers 60 61 (Miyashita et al. 2002; Ohno et al. 2005; Nishio 2006; Ohno and Nishio 2006; Ohno and 62 Nishio 2007a; Ohno and Nishio 2007b; Yamaguchi 2010; Yamaguchi et al. 2012; Yoshitake et 63 al. 2013; Sugimura et al. 2013a; Sugimura et al. 2013b; Sugimura et al. in press).

64 Against the background stated above, the authors' group has recently performed basic 65 characterization of miscibility and intermolecular interaction on binary blends of CEs with 66 non-crystalline vinyl polymers, particularly poly(*N*-vinyl pyrrolidone) (PVP) and its random

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67 copolymers (Miyashita et al. 2002; Ohno et al. 2005; Ohno and Nishio 2006; Ohno and Nishio 2007b; Sugimura et al. 2013a; Sugimura et al. 2013b). The CE component mainly 68 69 used in the previous studies was cellulose acetate (CA), propionate (CP), or butyrate (CB) 70 (Fig. 1a), and poly(N-vinyl pyrrolidone-co-vinyl acetate) (P(VP-co-VAc)) (Fig. 1b) or 71 poly(N-vinyl pyrrolidone-co-methyl methacrylate) (P(VP-co-MMA)) (Fig. 1c) was the 72 counter polymer component. Fig. 2a–c survey miscibility estimations for the blend systems 73 of CA, CP, and CB, each combined with P(VP-co-VAc) (designated as CE/P(VP-co-VAc)) 74 (Miyashita et al. 2002; Ohno and Nishio 2006; Sugimura et al. 2013a), by offering the 75 miscibility map constructed as a function of the degree of ester substitution (DS) of CE and 76 the copolymer composition of P(VP-co-VAc). The mappings were made based on thermal 77 analysis (T_g detection) by differential scanning calorimetry (DSC). As can readily be seen 78 by comparison of the three maps, the miscibility behaviour of CE/P(VP-co-VAc) blends is 79 seriously affected by a small difference in alkyl chain-length (carbon number) of the acyl 80 substituent in the employed CE. The CP system produced the largest miscible region.

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<< Figure 1 (a) & (b) & (c) >>

82 << Figure 2 (a) & (b) & (c) >>

Similar representations of miscibility estimations are given in Fig. 3 for two systems in which P(VP-*co*-MMA) was combined with either CA (Ohno and Nishio 2007b) or CP (Sugimura et al. 2013b); however, the mapping for CB/P(VP-*co*-MMA) blends is not made in this figure (see later discussion). Again interestingly, the miscible pairing region for the CP/P(VP-*co*-MMA) system is much larger than that for the CA/P(VP-*co*-MMA) system, with spreading to the upper right side of higher DS of CP and lower VP fraction of P(VP-*co*-MMA) in the map.

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<< Figure 3 (a) & (b) >>

Using supplementary data from Fourier transform infrared (FT-IR) and solid-state NMR
 measurements, we have tentatively concluded that the CE/VP-containing copolymer

93 combinations assume miscible or immiscible behaviour according to the balance in effectiveness of the following four factors (Sugimura et al. 2013a; Sugimura et al. 2013b): (1) 94 95 hydrogen-bonding attraction between residual hydroxyls of CE and VP-carbonyl groups of 96 the vinyl (co)polymer; (2) steric hindrance of bulky side-groups to the interaction specified in 97 (1); (3) indirect attraction via intramolecular repulsion between the comonomer units in the 98 copolymer; and (4) weak interaction due to structural affinity (e.g., dipole-dipole antiparallel 99 alignment) between the ester side-group of CE (such as CH₃-CH₂-CO-O-C-) and the VAc 100 (-(CH₂-CH(-O-CO-CH₃))-) or MMA (-(CH₂-(CH₃)C(-CO-O-CH₃))-) unit. To explain the 101 factor 3 more lucidly, when two monomer species having mutually repellent characters are 102 randomly combined by covalent bonding, the copolymers tend to form a miscible phase with 103 the CE component in the binary blends, rather than self-associate with the strong 104 intramolecular repulsion. Unfortunately, however, the factors 3 and 4 could not be directly 105 detected by the spectroscopic measurements.

106 In the present comparative study of the CE/vinyl polymer blends, we aim to clarify the 107 contributions of the copolymer effect and structural affinity to the miscibility attainment, by 108 another method besides thermal and spectroscopic techniques. In a previous work (Ohno 109 and Nishio 2007b), we preliminarily estimated the attractive or repulsive action between 110 chain segments of the polymer ingredients participating in the three systems, 111 CA/P(VP-co-VAc), CA/P(VP-co-MMA), and CB/P(VP-co-VAc), in terms of Krigbaum-Wall 112 polymer-polymer interaction parameters (Δb and μ) determinable by dilute solution 113 viscometry. Particularly μ data gave a satisfactory account of the difference in the 114 miscibility behaviour between the three blend systems (see later discussion). In this context, 115 the present paper covers complementary assessments of μ parameters for various ingredient 116 polymer pairs involved with the CP/P(VP-co-VAc) and CP/P(VP-co-MMA) systems. 117 Through comprehensive comparison of the results with the μ data formerly obtained for the CA and CB systems, some profound insights are provided into the positive effect of propionyl 118

- substitution leading to expansion of the miscible paring region in the maps of the CP systems.
- Additional attention is turned to miscibility behaviour of CB/P(VP-*co*-MMA) and cellulose
 acetate propionate (CAP)/P(VP-*co*-MMA) blends.

123 Experimental

124

- 125 Materials
- 126

127 CA was kindly provided from Daicel Corporation, and CAP was purchased from Eastman 128 Chemical Co. CP and CB samples were synthesized with acid chloride/base catalyst from 129 cotton cellulose via a homogeneous reaction in our laboratory, as has been described in the 130 preceding papers (Nishio et al. 1997; Ohno and Nishio 2006; Kusumi et al. 2008). Table 1 131 summarizes the characterization data including DS, molecular weight, and glass transition 132 temperature (T_g) determined by DSC (see below) for all the CE samples used in this study. 133 Codes "CE_x" and "CA_yP_z" denote CE of ester DS = x and CAP of acetyl DS = y and propionyl 134 DS = z, respectively.

135

<< Table 1 >>

136 The vinyl polymers employed as a mixing partner for the CEs were PVP, PVAc, 137 poly(methyl methacrylate) (PMMA), P(VP-co-VAc), and P(VP-co-MMA). Data of 138 characterization for all the vinyl polymers are also listed in Table 1. As shown in the table, 139 any of the copolymer samples exhibited a single $T_{\rm g}$, and the $T_{\rm g}$ -copolymer composition 140 relationships were in good obedience to a well-known Fox equation (Fox and Flory 1954), 141 with a possible extent of scattering due to the difference in molecular weight; thus they were 142 all regarded as essentially random copolymer. Hereafter, a P(VP-co-VAc) copolymer of 143 VP:VAc = m:n (in molar ratio) is encoded as P(VP_m-co-VAc_n), and the same encoding rule is 144 also applied for P(VP-co-MMA) samples.

146 Preparation of blend samples

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148 Powder materials of CEs and vinyl polymers were individually dissolved in 149 N,N-dimethylformamide (DMF) at room temperature (~25 °C), at a polymer concentration of 1.00 g dL^{-1} . Blend solutions for viscometric measurements were prepared by mixing equal 150 151 amounts of two solutions of the component polymers. For DSC measurements, two 152 solutions of the required pairing polymers were mixed at the desired weight proportions. The mixed polymer solutions (transparent) were then poured into a Teflon[®] tray and film 153 154 samples were made by evaporation of DMF at 50 °C under reduced pressure (< 10 mmHg). The as-cast films were further dried at 50 °C in vacuo for 3 days, before supplying to the 155 156 thermal analysis.

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158 Measurements

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160 Viscosity measurements were performed for dilute polymer solutions in DMF with an 161 Ubbelohde capillary viscometer, which was placed in a thermo-regulated water bath (30 °C). 162 The temperature of the water bath was controlled within an accuracy range of ± 0.1 °C. The polymer concentration of the starting sample was adjusted to $1.00 \text{ g } \text{dL}^{-1}$, and dilutions of the 163 solutions were made to yield at least 4 lower concentrations by adding appropriate doses of 164 165 DMF. The measurements following the respective dilutions were done after elapsing of an 166 equilibrium time of 15 min. As for the polymer solutions containing $CP_{2.72}$ or $CB_{2.67}$, however, the viscometric data were actually collected in a polymer concentration range below 167 ~0.30 g dL⁻¹, because the solutions of 1.00 g dL⁻¹ were appreciably viscous due to 168 169 comparatively high molecular weights of the cellulosics (see Table 1). The elution time of each solution from the set gauge of the viscometer was determined as the average of five 170

171 readings.

DSC thermal analysis was carried out with a Seiko DSC 6200/EXSTAR 6000 apparatus. 172 173 The temperature readings were calibrated with an indium standard. The calorimetry 174 measurements were conducted on ca. 5-mg film samples packed in an aluminum pan under a 175 nitrogen atmosphere. Each sample was first heated from ambient temperature (~25 °C) to ~220 °C at a scanning rate of 20 °C min⁻¹, and then immediately guenched to -50 °C at a rate 176 of 80 °C min⁻¹. Following this, the second heating scan was run from -50 °C to 230 °C at a 177 rate of 20 °C min⁻¹ to record stable thermograms. Thermograms presented in this paper 178 179 were all obtained in the second heating scan, and the $T_{\rm g}$ was taken as a temperature at the 180 midpoint of a baseline shift in heat flow characterizing the glass transition.

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182 **Results and discussion**

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184 Quantification of interaction parameters

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Following the preceding work (Ohno and Nishio 2007b), we applied a viscometric method developed by Krigbaum and Wall (Krigbaum and Wall 1950) and other groups (Cragg and Bigelow 1955; Chee 1990), to assess the attractive or repulsive interactivity between the CE-vinyl polymer constituents focused so far in this series of blend studies. The result was greatly useful to understand the difference in miscibility behaviour between the blend systems, as embodied in a later discussion.

A viscometric interaction parameter, *b*, for a non-electrolyte dilute polymer solution (usually, in the concentration range lower than ~1.0 g dL⁻¹) is defined to fulfill a liner relationship given by the Huggins equation (Huggins 1942):

195 $\eta_{\rm sp}/c = [\eta] + bc \tag{1}$

196 where c is the solute concentration, and η_{sp} and $[\eta]$ are the so-called specific and intrinsic

197 viscosities, respectively. The *b* is assumed to reflect an interaction between chain molecules 198 of the considered polymer and determined from a slope of the plot of η_{sp}/c vs. *c*. The 199 parameter *b* is also related to the Huggins coefficient *k* by

$$200 \qquad b = k[\eta]^2 \tag{2}$$

201 The *k* value generally ranges from 0.3 (in good solvents) to ~0.7 (in the Θ state) (Bohdanecký 202 and Kovář 1982).

With regard to a blend solution of two different polymers in a common solvent, Equation(1) is applicable in a rewritten fashion:

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$$\left(\eta_{\rm sp}\right)_{\rm m} / c_{\rm m} = \left[\eta\right]_{\rm m} + b_{\rm m} c_{\rm m}$$
(3)

where the subscript m denotes "mixture", and b_m is a comprehensive viscometric interaction parameter that reflects an overall interaction involving three possible combinations of polymer chains of the same species (1-1 and 2-2) or not (1-2).

In this viscometric treatment, the polymer-polymer miscibility is estimated by comparison between an experimentally obtained value and an ideally calculated one of $b_{\rm m}$. The former value, $b_{\rm m}^{\rm ex}$, is determined from the plot of $(\eta_{\rm sp})_{\rm m}/c_{\rm m}$ vs. $c_{\rm m}$ for blend solutions of a given polymer pair. The latter ideal value, $b_{\rm m}^{\rm id}$, is calculated by the following equation (Krigbaum and Wall 1950):

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$$b_{\rm m}^{\rm id} = w_1^2 b_{11} + w_2^2 b_{22} + 2w_1 w_2 b_{12}$$
 (4)

where w_i is the weight fraction of component *i* in the polymer mixture, and b_{ij} is an interaction parameter between the molecular chain of polymer *i* and that of polymer *j*, and thereby a potential value of b_{12} may be given by

218
$$b_{12} = \sqrt{b_{11} \times b_{22}}$$
 (5)

- 219 Here, a Krigbaum-Wall interaction parameter, Δb , is defined as
- $220 \qquad \Delta b = b_{\rm m}^{\rm ex} b_{\rm m}^{\rm id} \tag{6}$

If Δb is positive, the polymer 1 and polymer 2 are mutually attractive and therefore the pair is taken as miscible. Contrarily, if Δb is negative, the repulsive pair is considered to be immiscible. When there is a large difference between $[\eta]$ values of both polymers ($[\eta]_1$ and $[\eta]_2$), the following alternative parameter μ as a standard in non-dimensional unit may be more useful to predict the miscibility between the two components (Chee 1990).

226
$$\mu = \frac{\Delta b}{([\eta]_2 - [\eta]_1)^2}$$
(7)

227 The absolute value of μ , i.e., $|\mu|$, should represent the relative strength of attractive or 228 repulsive interaction between the two component polymer molecules.

Table 2 summarizes data of $[\eta]$ and b parameters (b_m^{ex} and b_m^{id}) obtained by the 229 230 viscometry for DMF solutions of CEs, vinyl polymers, and selected blending pairs of 50/50 231 composition, together with the polymer-polymer interaction parameters Δb and μ determined for the blends. The values of $[\eta]$ and $b_{\rm m}^{\rm ex}$ were obtained directly from the reduced viscosity 232 (η_{sp}/c) versus concentration plots, and those of b_m^{id} , Δb , and μ were calculated by the relevant 233 234 equations ((4), (6), and (7)) given above. For comprehensive purposes, some data were 235 quoted from the previous paper (Ohno and Nishio 2007b). As can be seen in the table, the 236 $[\eta]$ values of the cellulosics and those of the vinyl (co)polymers are fairly far apart, and hence 237 the standardized parameter μ is mainly used below for discussion on the interaction and 238 miscibility between the blend constituents.

239

<< Table 2 >>

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241 Overview of μ records for CA and CB blends

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First, we briefly review the preceding results of μ assessment for CA/P(VP-*co*-VAc), CB/P(VP-*co*-VAc), and CA/P(VP-*co*-MMA) blends (Ohno and Nishio 2007b). Figs. 4a, 4c, and 5a summarize simplified miscibility maps of the three blend systems, with addition of the illustrations in terms of μ data obtained for selected polymer combinations (DS of CEs, ~2.7; VP:VAc or MMA of copolymers, ~0.5:0.5) critical to the respective systems. The individual μ evaluations were in consistency with the respective miscibility mappings based on DSC 249 thermal analysis; viz., a positive μ value was obtained for miscible pairs of 250 cellulosic/synthetic polymers, while immiscible blends all provided a negative μ value.

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- 252

<< Figure 4 (a) & (b) & (c) >>

<< Figure 5 (a) & (b) >>

253 As exemplified for a highly butyrated CB/P(VP-co-VAc) series (Fig. 4c, right), the μ data 254 concerned with the "three" constituting polymer ingredients made an order with respective to the degree of "immiscibility": PVP/PVAc (-4.23×10⁻²) > CB_{2.67}/PVP (-1.43×10⁻²) \geq 255 CB_{2.67}/PVAc (-1.07×10^{-2}). The mutually repellent character of the PVP/PVAc pair is 256 257 considerably stronger than the corresponding ones of the other pairs CB_{2.67}/PVP and 258 CB_{2.67}/PVAc. Then it can be taken for the $CB_{2.67}/P(VP_{0.52}-co-VAc_{0.48})$ blend that the 259 P(VP-co-VAc) component was intimately mixed with the CB component showing less 260 repulsion to both the comonomer units, as a result of avoidance of the intense repulsion 261 between VP and VAc segments inevitable in the copolymer-copolymer association; the 262 blending pair of $CB_{2.67}/P(VP_{0.52}-co-VAc_{0.48})$ is surely attractive to each other, giving a positive μ value, +3.69×10⁻³. This reasoning would satisfy us about the appearance of the miscibility 263 264 window (Fig. 4c, left), as amplified in the following sections. On the other hand, such an 265 explicit window never appeared in the map of the CA/P(VP-co-VAc) system (see Fig. 2a and 266 4a), although there should have arisen the intra-copolymer effect improving the miscibility in 267 the blends of relatively high-acetylated CAs. The absence of the window may be interpreted 268 as due to an inhibiting factor, i.e., the strong self-association ability of highly substituted CAs 269 of DS > 2.7; the CAs are rather easily crystallizable as cellulose triacetate II form. Differing 270 from this, no crystallizing habit was detected even for a CB synthesized at DS = 2.94 (Ohno 271 and Nishio 2006). The lesser self-association nature of CB should be advantageous to that 272 attractive interaction with the P(VP-co-VAc) component.

273 Meanwhile, another vinyl polymer combination of PVP and PMMA provided a μ value 274 of -1.87×10^{-2} , from which the binary system is suggested to be immiscible. In fact, the blend samples showed a common behaviour of essentially double T_{gs} in DSC measurements (Ohno and Nishio 2007b). However, the $|\mu|$ value for the PVP/PMMA pair is smaller than that ($|\mu| = 4.23 \times 10^{-2}$) for the PVP/PVAc pair. Thus it is deduced that the constituents VP and MMA in P(VP-*co*-MMA) show a somewhat weaker repulsive interaction than the VP and VAc units in P(VP-*co*-VAc). Presumably, this deterioration of the latent copolymer effect is responsible for the observation of a narrower miscible region in the CA/P(VP-*co*-MMA) map (Fig. 5a) relative to that in the CA/P(VP-*co*-VAc) map (Fig. 4a).

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283 Inspection of miscibility maps for CP blends in μ terms

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285 *CP/P(VP-co-VAc) system*

As shown in Fig. 4b (right), a negative μ value -1.02×10^{-2} was obtained for the combination 286 287 of CP_{2.72} and PVP homopolymer, while μ of the CP_{2.72}/P(VP_{0.52}-co-VAc_{0.48}) pair was positive, 288 +1.50×10⁻². From these assessments, PVP and P(VP_{0.52}-co-VAc_{0.48}) are taken as immiscible 289 and miscible, respectively, with the highly esterified CP. The judgment is actually in 290 accordance with the result of miscibility estimation by thermal analysis for the blends (see Fig. 4b, left). For another essential pair, $CP_{2.72}/PVAc$, we obtained a negative μ of -7.19×10^{-5} , 291 292 but the absolute value is much smaller than that for the CP_{2.72}/PVP pair by more than two 293 orders of magnitude. The former pair was previously marked to be partially miscible by observation of two T_{gs} approaching each other to an appreciable extent, and the low 294 295 magnitude of μ reflects such a "better compatibility" of highly substituted CP with PVAc 296 homopolymer.

297 Despite no presence of strong intermolecular attraction between $CP_{2.72}$ and the two 298 homopolymers (PVP and PVAc), the CP component was able to be miscible with the 299 copolymer comprising VP and VAc units. This phenomenon is explicable as being due to 300 the more intense repulsive action between the VP and VAc segments in the P(VP-*co*-VAc) 301 copolymer component, as in the case of the CB/P(VP-co-VAc) system. We find for sure in Fig. 4b (right) that the PVP/PVAc pair shows the largest negative μ value (-4.23×10⁻²) in the 302 three polymer pairs participating in the $CP_{2.72}/P(VP-co-VAc)$ system. In general, when two 303 304 monomer species repelling each other are randomly combined by covalent bonding, the 305 resulting copolymer tends to intimately mix with the other polymer of less self-associating 306 nature, so as to reduce the strong repulsion between the comonomer units (ten Brinke et al. 307 1983; Paul and Barlow 1984). This is the reason why the high-esterified CP and CB can be 308 miscible with P(VP-co-VAc) in a restricted range of the copolymer composition, even though 309 there is a scarcity of specific attractive force (i.e. proton donor-acceptor interaction) between 310 the two mixing components.

As is obvious in Fig. 4, the miscible region in the CP/P(VP-*co*-VAc) map is larger than the corresponding ones in the other maps of CA/P(VP-*co*-VAc) and CB/P(VP-*co*-VAc). In perspective comparison, the region involved in the CP system expands particularly to the side of VAc-richer compositions. This improvement virtually comes from the better compatibility of CP with PVAc supported above by the μ data of -7.19×10^{-5} for CP_{2.72}/PVAc. This value in $|\mu|$ is overwhelmingly small, compared with $\mu = -2.12 \times 10^{-2}$ for CA_{2.70}/PVAc (Fig. 4a, right) and $\mu = -1.07 \times 10^{-2}$ for CB_{2.67}/PVAc (Fig. 4c, right).

318 For three pairs of P(VP_{0.52}-co-VAc_{0.48}) with the CEs of DS \approx 2.7, we can rank them $CA_{2.70}/P(VP_{0.52}-co-VAc_{0.48})$ (+7.12×10⁻²) > 319 according to μ data, as follows: $CP_{2.72}/P(VP_{0.52}-co-VAc_{0.48})$ (+1.50×10⁻²) > $CB_{2.67}/P(VP_{0.52}-co-VAc_{0.48})$ (+3.69×10⁻³), all 320 showing miscibility. The CA_{2.70}/P(VP_{0.52}-co-VAc_{0.48}) pair exhibited the highest μ value, 321 322 which is attributable to the direct interaction based on the actually detected hydrogen bonding 323 between CA-hydroxyl and VP-carbonyl groups (Miyashita et al. 2002; Ohno et al. 2005); however, the increase of μ relative to that for CA_{2.70}/PVP (+4.53×10⁻²) suggests a secondary 324 325 contribution of the intra-copolymer effect to the miscibility attainment. The hydrogen 326 bonding effect seriously declines in the other two systems adopting propionyl and butyryl

327 substitutions for the CE component. Consequently, the miscibility of $CB_{2.67}$ with 328 $P(VP_{0.52}-co-VAc_{0.48})$ is realized only through the intra-copolymer repulsion as an indirect 329 driving force. As to the $CP_{2.72}/P(VP_{0.52}-co-VAc_{0.48})$ pair, besides the copolymer effect, a 330 weak interaction due to structural affinity between the propionyl ester group and VAc unit 331 also acts as a factor contributory to the miscibility attainment.

332

333 CP/P(VP-co-MMA) system

334 Fig. 5b (left) displays a simplified diagram of the miscibility mapping conducted for 335 CP/P(VP-co-MMA) blends (Fig. 3b). In the right side of Fig. 5b, μ data are collected for 336 four combinations of CP2.72 with P(VP0.50-co-MMA0.50), P(VP0.22-co-MMA0.78), PVP, and PMMA, the values being $+9.33 \times 10^{-4}$, $+4.27 \times 10^{-3}$, -1.02×10^{-2} , and -3.23×10^{-4} , respectively. 337 338 Judging from the positive or negative sign of μ , the P(VP-co-MMA) copolymers are taken as 339 potentially miscible with CP_{2.72}, whereas both the homopolymers are not. These judgments 340 entirely agree with the actual markings for the CP_{2.72}/P(VP-co-MMA) series in the miscibility 341 map. In addition, PVP/PMMA blends are immiscible and this polymer pair provides a larger 342 negative μ (-1.87×10⁻²) than the CP_{2.72}/PVP and CP_{2.72}/PMMA pairs. The relationship in 343 repulsion (immiscibility) between the three ingredient polymer pairs participating in the 344 CP_{2.72}/P(VP-*co*-MMA) series is basically similar to that found for the CP_{2.72}/P(VP-*co*-VAc) 345 series (see Fig. 4b, right). Accordingly, it is reasonable to assume that the intramolecular 346 repulsive effect of the VP-MMA copolymer gave rise to the miscibility window in the map for 347 the CP/P(VP-co-MMA) system. However, the window region observed for this system is 348 obviously narrower than that for the CP/P(VP-co-VAc) system (see Fig. 4b, left). This 349 narrowing of the window may be ascribed to the weaker repulsion in the VP-MMA copolymer relative to that in the VP-VAc copolymer ($\mu = -4.23 \times 10^{-2}$), as has been applied to 350 351 the comparative discussion of the two maps for the corresponding blends of CA. The location of the window in the side of MMA-rich compositions owes to the better affinity 352

between CP and MMA segments, as supported by the lower order (10⁻⁴) of μ obtained for the

 $354 \quad CP_{2.72}/PMMA pair.$

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356 Complementary mapping for CB/P(VP-co-MMA) system by application of μ assessment

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358 In the miscibility characterization of CE/vinyl copolymer blends, we have not yet 359 accomplished the total mapping for the CB/P(VP-co-MMA) system by thermal analysis. A 360 main reason is that $T_{\rm g}$ s (ca. 110–120 °C) of CBs of DS $\approx 2.5-2.9$ are fairly close to those (ca. 361 100–115 °C) of P(VP-co-MMA)s of VP < 50 mol%. However, we previously acquired the 362 following data for the system concerned: (i) CB and PVP homopolymer formed miscible 363 blends of hydrogen-bonding type unless the butyryl DS exceeded ~2.5 (see Fig. 2c) (Ohno 364 and Nishio 2006); (ii) a polymer pair of CB (DS = 2.94) with $P(VP_{0.50}-co-MMA_{0.50})$ was 365 judged to be immiscible (double T_{gs}) (Ohno and Nishio 2007b).

To depict the miscibility map of the CB/P(VP-*co*-MMA) system more closely, we newly examined the blend miscibility of relatively low-substituted CBs (DS < 2.5) with P(VP-*co*-MMA)s by DSC and also quantified μ for additional pairs of CB (DS \geq 2.6) with MMA-rich P(VP-*co*-MMA)s by viscometry. A major concern is whether the miscibility window emerges or not in the CB/P(VP-*co*-MMA) map.

371 Fig. 6a illustrates DSC thermograms measured for blend samples of CB_{2.01}/PMMA 372 homopolymer; the binary cast films were mostly cloudy to the naked eye. As can be seen 373 from the data, two independent glass transitions originating from the two components were 374 detected for the 40/60-80/20 compositions (in wt% ratio), signalizing immiscibility of the 375 CB_{2.01}/PMMA pair. The same behaviour of double T_{gs} was also observed for CB_{2.41}/PMMA 376 In contrast, Fig. 6b and c offer a typical miscible evidence in DSC (i.e. blends. composition-dependent 377 single $T_{\rm g}$) for $CB_{2.01}/P(VP_{0.22}-co-MMA_{0.78})$ and 378 CB_{2.01}/P(VP_{0.50}-*co*-MMA_{0.50}) blends, respectively. Similar miscible behaviour was

379 confirmed for other polymer combinations using CB_{2.41} and/or P(VP-co-MMA)s of VP \geq 9 380 mol% (MMA \leq 91 mol%). Additionally, as-cast films of the CB blends with the 381 P(VP-co-MMA)s were all highly transparent in the visual inspection. Thus it turns out that 382 the lower limit in VP fraction of P(VP-co-MMA) that can be miscible with CB (DS $< \sim 2.5$) is 383 ~10 mol%, which is almost the same limit as that found when CP was the CE component (see 384 Fig. 5b). In a reasoning similar to that applied to interpret the CP/P(VP-co-MMA) map, the 385 miscibility of CB with P(VP-co-MMA)s so rich in MMA residues (e.g. MMA = 87 and 91 386 mol%) would be invited by a good compatibility between the butyl ester side-group and the MMA unit. This may be supported by μ assessment of an extremely small negative value 387 388 (-8.35×10^{-5}) for a polymer pair CB_{2.67}/PMMA (see Table 2).

389

390 In the present viscometric μ measurements, we found a definitely positive data such as μ $= +2.12 \times 10^{-3}$ for CB_{2.67}/P(VP_{0.22}-co-MMA_{0.78}). This indicates that even CB of DS > 2.5 is 391 392 potentially miscible with the vinyl copolymer rich in MMA. Fig. 7 (left) summarizes a 393 miscibility map constructed for the total system of CB/P(VP-co-MMA) by the combined use 394 of the DSC and μ -assessment results. In the map, solid lines separate the miscible and 395 immiscible regions connected with DS of CB and VP fraction of P(VP-co-MMA), to provide a 396 miscibility window in the upper right portion. As illustrated in the right side in Fig. 7. μ 397 parameters for three combinations of the ingredient polymers pertinent to the CB_{2.67}/P(VP-co-MMA) series are all negative, but the PVP/PMMA pair gives the largest 398 absolute value (1.87×10^{-2}) . 399 This situation again supports the contribution of the 400 intramolecular repulsion inherent in the P(VP-co-MMA) copolymer to the appearance of the 401 miscibility window. However, the region is diminished to some extent, compared to the 402 window in the CB/P(VP-co-VAc) map (Fig. 4c), because the repulsion between VP and MMA 403 units is weaker than that between VP and VAc units, as already mentioned above.

Here we should further note that $CB_{2.67}$ of $DS \approx 2.7$ is estimated to be immiscible with P(VP_{0.50}-*co*-MMA_{0.50}) of VP:MMA = 50:50 from the μ data of -3.39×10^{-3} . In contrast, a comparable pair using CP, i.e., $CP_{2.72}/P(VP_{0.50}$ -*co*-MMA_{0.50}), was miscible, which was decisive from both T_g and μ determinations (see Figs. 3b and 5b). It follows, therefore, that the miscible pairing region (mainly associated with the window) in the CB/P(VP-*co*-MMA) map is a little narrower than that of the CP/P(VP-*co*-MMA) map. This comparison is made clearer in Fig. 7 (left), as guided by solid lines and broken ones inserted therein.

412 As indicated above, intimate mixing of two polymer components through the copolymer 413 repulsion effect is unrealized on blending $CB_{2.67}$ with $P(VP_{0.50}-co-MMA_{0.50})$. In 414 interpretation of this, the following data should be recalled: $\mu = -1.87 \times 10^{-2}$ for PVP/PMMA and -1.43×10^{-2} for CB_{2.67}/PVP (see Fig. 7, right), the two values being close to each other. 415 416 In the employment of the copolymer of VP = 50 mol%, probably, the relatively strong 417 repulsion would still work between the CB component and the VP residue and inhibit the 418 mutual approach of the two polymer components. Consequently, the intramolecular 419 copolymer effect to attain miscible CB/P(VP-co-MMA) blends is active only at restricted 420 copolymer compositions considerably rich in MMA. On the other hand, the repulsion between CP_{2.72} and PVP ($\mu = -1.02 \times 10^{-2}$) is evidently weaker than that between PVP and 421 422 PMMA (see Fig. 5b, right), and the copolymer effect would be significant even at the composition of VP = 50 mol%, resulting in the miscible blending of the 423 424 In addition, a low frequency of intermolecular $CP_{2.72}/P(VP_{0.50}-co-MMA_{0.50})$ pair. 425 hydrogen-bondings might contribute to this miscibility attainment as a secondary effect. 426 This inference took into consideration the DS boundary of ~2.7 partitioning the mixing states 427 of CP/P(VP-co-MMA) blends (VP \ge 60 mol%) (Fig. 5b, left).

428

Inspection of estimation results of miscibility for CAP/P(VP-*co*-MMA) blends in μ terms

Finally, we refer to miscibility behaviour of CAP blends with P(VP-*co*-MMA). To make a comparison with the result for the CP_{2.72}/P(VP-*co*-MMA) series, a partially acetylated cellulose propionate sample, CA_{0.16}P_{2.52} (acetyl DS = 0.16; propionyl DS = 2.52), was selected as the mixed ester component.

435 Fig. 8 (left) collects the miscibility data (Sugimura et al. 2013b) based on thermal analysis for the target CA_{0.16}P_{2.52}/P(VP-co-MMA) blends, together with the corresponding 436 437 data in the uses of $CP_{2,72}$ and $CA_{2,70}$. In the right side, an additional illustration is given in 438 terms of μ assessment. The combination of CA_{0.16}P_{2.52} and P(VP_{0.50}-co-MMA_{0.50}) imparted a positive μ value of +5.79×10⁻³, while negative μ data of -1.01×10⁻² and -2.08×10⁻⁴ were 439 440 assigned to CA_{0.16}P_{2.52}/PVP and CA_{0.16}P_{2.52}/PMMA pairs, respectively. Therefore, the $P(VP_{0.50}-co-MMA_{0.50})$ copolymer is potentially miscible with the mixed ester CA_{0.16}P_{2.52}, 441 whereas both the homopolymers are not. These judgments are consistent with the results of 442 443 miscibility estimation by DSC for the respective blends, also supporting that the 444 $CA_{0.16}P_{2.52}/P(VP-co-MMA)$ series offers a miscibility window, as did the blend series using 445 CP_{2.72}. Furthermore, the immiscible polymer pair of PVP/PMMA provides a larger negative μ (-1.87×10⁻²) than the other immiscible pairs of CA_{0.16}P_{2.52}/PVP and CA_{0.16}P_{2.52}/PMMA. 446 From this triangular relationship, the intramolecular repulsive effect of the VP-MMA 447 448 copolymer may be regarded as being responsible for the emergence of the miscibility window 449 in the map for the CAP/P(VP-co-MMA) blends.

450

<< Figure 8 >>

However, it is astonishing that the VP:MMA range involved in the window became more expanded in the $CA_{0.16}P_{2.52}/P(VP-co-MMA)$ series, when compared with the situation in the CP_{2.72}/P(VP-co-MMA) series. In order to explain this expansion, we directed attention to another intramolecular repulsive interaction that might have arisen in the mixed ester component *per se*. Thereupon, a cellulose ester pair $CA_{2.70}/CP_{2.72}$ was explored by thermal analysis and viscometry for evaluations of the miscibility and interaction parameter; the 457 residual hydroxyl contents of the monoester derivatives (CA_{2.70} and CP_{2.72}) are equalized to 458 that of CA_{0.16}P_{2.52}. DSC measurements confirmed that CA_{2.70}/CP_{2.72} blends exhibited dual T_{g} signals corresponding to those of the two constituents at any blending proportion. The 459 460 Krigbaum-Wall interaction parameter of this polymer pair was estimated to be negative, as μ $= -8.12 \times 10^{-3}$ (see Fig. 8, right), in conformity with the immiscible behaviour of the blends. 461 The absolute value of this μ is appreciably large, although it is below $|\mu| = 1.87 \times 10^{-2}$ for the 462 463 PVP/PMMA pair. The present result suggests that a relatively strong repulsive interactivity can work between the two cellulosic ester components. 464

In view of the above context, it is deduced that the cellulose mixed ester would also behave as a kind of copolymer dangling two different ester groups along the carbohydrate backbone; thus, the CAP/P(VP-*co*-MMA) blends are taken as a copolymer/copolymer system where the miscibility should be affected by the duplicated, intramolecular copolymer effect. The expansion of the window in the mapping of the CA_{0.16}P_{2.52}/P(VP-*co*-MMA) blends can be ascribed to such an additional repulsion effect originating in the CAP side.

471

472 **Conclusions**

473

474 The blend miscibility of CP with the VP-containing vinyl copolymers P(VP-co-VAc) and 475 P(VP-co-MMA) is improved in respect of the miscible pairing number, compared with the 476 cases using CA and CB. This behaviour was satisfactorily explained by comparing the 477 attractive or repulsive interactivities between related polymer ingredients in terms of the 478 Krigbaum-Wall interaction parameter μ that was determined by solution viscometry. 479 Especially, great contributions of both the intra-copolymer effect and the structural affinity 480 effect to the miscibility attainment were made clear by the μ assessments. The former effect 481 is explicitly responsible for the miscibility window appearing in the maps constructed for the CP/vinyl copolymer systems, and this is also applicable to the maps for the CB systems. 482

The comparatively narrower window observed when the counter component to CP or CB was P(VP-*co*-MMA) is interpretable as due to the lesser strength in repulsion of the VP-MMA copolymer relative to that of the VP-VAc copolymer. The structural affinity effect is concretely connected with a good compatibility of the propionyl group of CP with the VAc or MMA unit of the partner copolymer in the CP-based two systems, and, in the employment of CB, this effect is active between the butyryl and MMA moieties in the CB/P(VP-*co*-MMA) system only.

Such a useful μ measurement was also applied to the inspection of miscibility mapping for CAP blends with P(VP-*co*-MMA). The observed expansion of the miscibility window relative to that for the comparable CP blends was explicable in terms of the μ data, which indicated additional repulsion in the side of the cellulose mixed ester component; therefore, the CAP/P(VP-*co*-MMA) blends should be taken as a copolymer/copolymer system where the duplicated copolymer effect works.

From a practical standpoint, the present results will be so useful for related researchers to expand the opportunities of material design based on the CE family including cellulose mixed esters. Delicate characterization and even prediction of the miscibility may be possible for many other series of CE/synthetic copolymer blends by examining the viscometric interaction parameters of the targeted constituent polymer pairs, in addition to the orthodox thermal and spectroscopic estimations.

502

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508 **Compliance with Ethical Standards**

The authors declare no conflict of interest.

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585 Figure Captions

586

587 Fig. 1 Structural formulae of (a) CEs (i.e., CA, CP, and CB), (b) P(VP-co-VAc), and (c)
588 P(VP-co-MMA).

589

Fig. 2 Miscibility maps for three blend systems (a) CA/P(VP-*co*-VAc) (Miyashita et al. 2002), (b) CP/P(VP-*co*-VAc) (Sugimura et al. 2013a), and (c) CB/P(VP-*co*-VAc) (Ohno and Nishio 2006), depicted as a function of DS of CE and VP fraction of the copolymer in a rearranged fashion with additional data. Symbols indicate that a given pair of CE/vinyl polymer is miscible (\bigcirc , single T_g), immiscible (\times , dual T_g s), or partially miscible (\triangle , dual T_g s approaching each other to an appreciable degree).

596

Fig. 3 Miscibility maps for two blend systems (a) CA/P(VP-*co*-MMA) (Ohno and Nishio 2007b) and (b) CP/P(VP-*co*-MMA) (Sugimura et al. 2013b), depicted as a function of DS of CE and VP fraction of the copolymer in a rearranged fashion with additional data. The meanings of two symbols \bigcirc and \times are the same as defined in Fig. 2.

601

602 **Fig. 4** Miscibility maps (left) with additional illustrations using μ data (right) for (a) 603 CA/P(VP-*co*-VAc), (b) CP/P(VP-*co*-VAc), and (c) CB/P(VP-*co*-VAc) systems. The meanings 604 of three symbols \bigcirc , \times , and \triangle are the same as used in Fig. 2. The miscibility maps are 605 represented in a simplified style retaining the essence of the data shown in Fig. 2.

606

607 **Fig. 5** Miscibility maps (left) with additional illustrations using μ data (right) for (a) 608 CA/P(VP-*co*-MMA) and (b) CP/P(VP-*co*-MMA) systems. The meanings of two symbols \bigcirc 609 and × are the same as used in Fig. 2. The miscibility maps are represented in a simplified 610 style retaining the essence of the data shown in Fig. 3.

612 **Fig. 6** DSC thermograms obtained for blends of CB_{2.01} with (a) PMMA, (b) 613 P(VP_{0.22}-*co*-MMA_{0.78}), and (c) P(VP_{0.50}-*co*-MMA_{0.50}). Arrows indicate a T_g position taken 614 as the midpoint of a baseline shift in heat flow.

615

616 **Fig. 7** Miscibility map (left) and additional illustration (right) using μ data for 617 CB/P(VP-*co*-MMA) blends. The meanings of two symbols \bigcirc and \times are the same as used in 618 Fig. 2. Solid lines in the map represent a boundary partitioning the miscible and immiscible 619 regions for the CB/P(VP-*co*-MMA) system, and, for comparison, the corresponding boundary 620 for the CP/P(VP-*co*-MMA) system (Fig. 5b) is drawn by broken lines.

621

Fig. 8 Mapping of miscibility data (Sugimura et al. 2013b) (left) and additional illustration in μ terms (right) for CA_{0.16}P_{2.52}/P(VP-*co*-MMA) blends. For comparison, miscibility data for the corresponding blends using CA_{2.07} and CP_{2.72} (see Fig. 3) are also mapped in the left figure. The meanings of two symbols \bigcirc and \times are the same as used in Fig. 2.

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⁶²⁸ In addition to the eight figures, there are two tables. See annexed sheets.

$M_{\mathrm{w}}{}^{d}$ M_n^{d} Sample code a $M_{\rm w}/M_{\rm n}$ d $T_{\rm g}/^{\rm o}{\rm C}$ Source $CP_{2.72}$ 1,070,000 367,000 2.92 Synthesized 134 CA0.16P2.52 258,000 73,400 3.51 143 Eastman Chemical Co. 237,000 73,000 186 Daicel Co. CA_{2.70} 3.25 114 998,000 285,000 3.50 CB_{2.67} Synthesized CB_{2.41} 952,000 218,000 4.37 132 Synthesized CB_{2.01} 651,000 294,000 Synthesized 2.21 139 $M_{\rm w}^{e}$ Sample code M_n^{e} $M_{\rm w}/M_{\rm n}^{\ e}$ $T_{\rm g}/^{\rm o}{\rm C}$ Source **PVP** 24,500^f Nacalai Tesque, Inc. 162 _ 90,000^{*f*} **PVAc** 41 Polyscience, Inc. _ $P(VP_{0.52}-co-VAc_{0.48})^{b}$ 28,000 89 5,120 5.47 Polyscience, Inc. **PMMA** 88,400 35,000 2.53 100 Aldrich Chemical Co. $P(VP_{0,22}-co-MMA_{0,78})^{c}$ 189,000 70,800 2.66 111 Synthesized^g $P(VP_{0.50}-co-MMA_{0.50})^{c}$ 61,300 3.00 119 Synthesized^g 184,000

632 **Table 1** Characterization of CEs and synthetic vinyl polymers used in the present study

^a The DS values were determined by ¹H NMR.

^b The VP content was determined by ¹H NMR.

^c The VP contents were determined by FT-IR in a way described by Liu et al. (1994).

^d Determined by gel permeation chromatography (mobile phase, tetrahydrofuran at 40 °C) with polystyrene standards.

^{*e*} Determined by gel permeation chromatography (mobile phase, 10 mM L^{-1} lithium bromide/DMF at 40 °C) with polystyrene standards.

^f Nominal value.

631

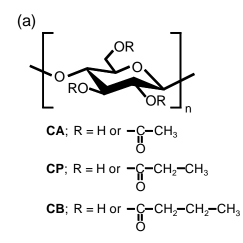
^{*g*} Synthesized in the authors' laboratory by radical polymerization of two distilled monomers, VP (Nacalai Tesque, Inc.) and MMA (Nacalai Tesque, Inc.), in the same way as that described in a previous paper (Ohno and Nishio 2007b).

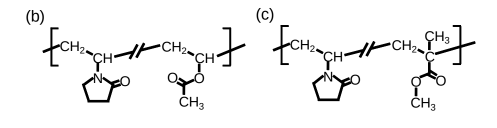
Samples	$[\eta]/dL \cdot g^{-1}$	$b_{\rm m}^{\rm ex}/{\rm dL^2 \cdot g^{-2}}$	$b_{\mathrm{m}}^{\mathrm{id}}/\mathrm{d}\mathrm{L}^2$ ·g ⁻²	$\Delta b/dL^2 \cdot g^{-2}$	μ
CP _{2.72}	6.27	1.84×10 ¹	_	_	_
$CA_{0.16}P_{2.52}$	1.85	1.84	_	_	_
CA _{2.70} ^{<i>a</i>}	2.28	1.86	_	_	_
$\operatorname{CB}_{2.67}{}^a$	5.61	1.13×10 ¹	_	_	_
PVP ^a	1.46×10 ⁻¹	1.18×10^{-2}	_	_	_
PVAc ^{<i>a</i>}	6.10×10 ⁻¹	1.32×10^{-1}	_	_	_
$P(VP_{0.52}$ -co-VAc $_{0.48})^{a}$	1.67×10^{-1}	1.21×10^{-2}	_	_	_
PMMA ^{<i>a</i>}	2.92×10^{-1}	3.01×10^{-2}	_	_	_
P(VP _{0.22} -co-MMA _{0.78})	3.64×10 ⁻¹	3.45×10^{-2}	_	_	_
$P(VP_{0.50}$ -co-MMA _{0.50}) ^a	5.54×10^{-1}	9.47×10 ⁻²	_	_	_
CP _{2.72} /PVP	3.77	4.46	4.85	-3.84×10^{-1}	-1.02×10^{-2}
CP _{2.72} /PVAc	3.42	5.42	5.42	-2.30×10 ⁻³	-7.19×10^{-5}
CP _{2.72} /P(VP _{0.52} -co-VAc _{0.48})	3.20	5.41	4.85	$+5.59 \times 10^{-1}$	$+1.50 \times 10^{-2}$
CP _{2.72} /PMMA	3.67	4.98	4.99	-1.15×10^{-2}	-3.23×10^{-4}
CP _{2.72} /P(VP _{0.22} -co-MMA _{0.78})	3.09	5.17	5.02	$+1.49 \times 10^{-1}$	$+4.27 \times 10^{-3}$
CP _{2.72} /P(VP _{0.50} -co-MMA _{0.50})	3.12	5.32	5.29	$+3.05 \times 10^{-2}$	+9.33×10 ⁻⁴
CA _{0.16} P _{2.52} /PVP	9.80×10 ⁻¹	5.06×10 ⁻¹	5.35×10^{-1}	-2.91×10^{-2}	-1.01×10^{-2}
CA0.16P2.52/PMMA	1.07	5.83×10^{-1}	5.84×10^{-1}	-5.03×10^{-4}	-2.08×10^{-4}
CA _{0.16} P _{2.52} /P(VP _{0.50} -co-MMA _{0.50})	1.13	7.01×10^{-1}	6.91×10 ⁻¹	$+9.71 \times 10^{-3}$	+5.79×10 ⁻³
CA _{2.70} /PVP ^{<i>a</i>}	1.38	7.50×10^{-1}	5.43×10 ⁻¹	$+2.07 \times 10^{-1}$	+4.53×10 ⁻²
CA _{2.70} /PVAc ^a	1.47	6.87×10^{-1}	7.47×10^{-1}	-5.95×10^{-2}	-2.12×10^{-2}
CA _{2.70} /P(VP _{0.52} - <i>co</i> -VAc _{0.48}) ^{<i>a</i>}	1.61	8.63×10 ⁻¹	5.44×10^{-1}	$+3.20 \times 10^{-1}$	+7.12×10 ⁻²
CA _{2.70} /PMMA ^a	1.28	5.85×10^{-1}	5.92×10 ⁻¹	-6.64×10 ⁻³	-1.67×10^{-3}
CA _{2.70} /P(VP _{0.50} -co-MMA _{0.50}) ^a	1.40	6.78×10^{-1}	6.99×10 ⁻¹	-2.12×10 ⁻²	-7.06×10^{-3}
CB _{2.67} /PVP ^{<i>a</i>}	2.97	2.59	3.01	-4.27×10^{-1}	-1.43×10^{-2}
CB _{2.67} /PVAc ^{<i>a</i>}	3.14	3.20	3.47	-2.69×10^{-1}	-1.07×10^{-2}
CB _{2.67} /P(VP _{0.52} -co-VAc _{0.48}) ^a	2.82	3.13	3.02	$+1.10 \times 10^{-1}$	+3.69×10 ⁻³
CB _{2.67} /PMMA ^a	3.04	3.13	3.13	-2.37×10^{-3}	-8.35×10^{-5}
CB _{2.67} /P(VP _{0.22} -co-MMA _{0.78})	2.42	3.21	3.15	$+5.84 \times 10^{-2}$	$+2.12 \times 10^{-3}$
$CB_{2.67}/P(VP_{0.50}-co-MMA_{0.50})^{a}$	3.16	3.28	3.37	-8.67×10^{-2}	-3.39×10 ⁻³
PVP/PVAc ^{<i>a</i>}	3.90×10 ⁻¹	4.66×10 ⁻²	5.57×10 ⁻²	-9.13×10 ⁻³	-4.23×10 ⁻²
PVP/PMMA	2.40×10^{-1}	1.95×10^{-2}	1.99×10 ⁻²	-4.02×10^{-4}	-1.87×10^{-2}
CA _{2.70} /CP _{2.72}	4.34	7.88	8.01	-1.29×10^{-1}	-8.12×10 ⁻³

634 CEs, synthetic vinyl polymers, and their respective 50/50 blends

^{*a*} Data were quoted from a previous paper (Ohno and Nishio 2007b).







640 Fig. 1 Structural formulae of (a) CEs (i.e., CA, CP, and CB), (b) P(VP-co-VAc), and (c)

- 641 P(VP-*co*-MMA).

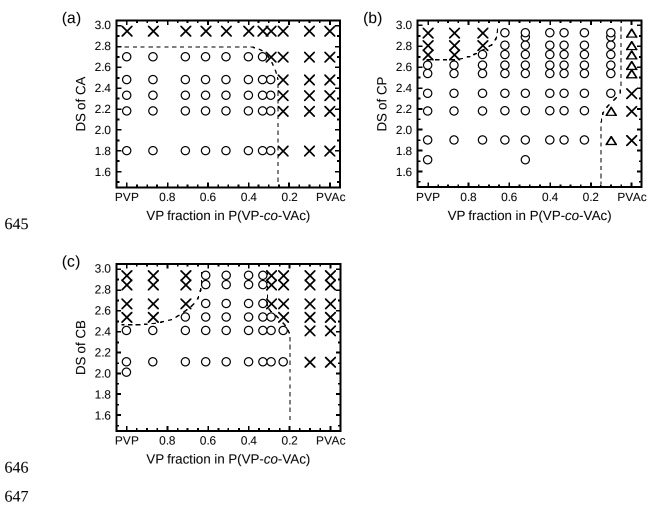


Fig. 2 Miscibility maps for three blend systems (a) CA/P(VP-*co*-VAc) (Miyashita et al. 2002), (b) CP/P(VP-*co*-VAc) (Sugimura et al. 2013a), and (c) CB/P(VP-*co*-VAc) (Ohno and Nishio 2006), depicted as a function of DS of CE and VP fraction of the copolymer in a rearranged fashion with additional data. Symbols indicate that a given pair of CE/vinyl polymer is miscible (\bigcirc , single T_g), immiscible (\times , dual T_g s), or partially miscible (\triangle , dual T_g s approaching each other to an appreciable degree).

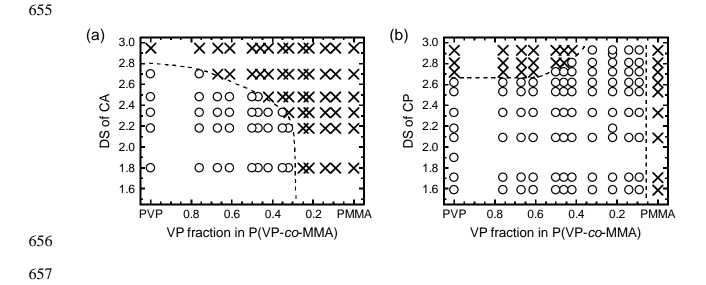


Fig. 3 Miscibility maps for two blend systems (a) CA/P(VP-*co*-MMA) (Ohno and Nishio 2007b) and (b) CP/P(VP-*co*-MMA) (Sugimura et al. 2013b), depicted as a function of DS of CE and VP fraction of the copolymer in a rearranged fashion with additional data. The meanings of two symbols \bigcirc and \times are the same as defined in Fig. 2.

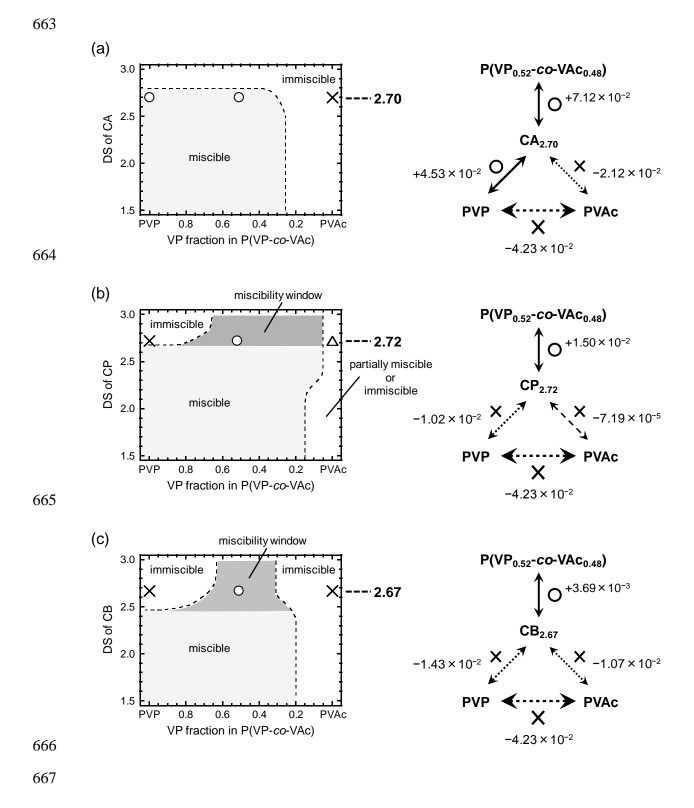


Fig. 4 Miscibility maps (left) with additional illustrations using μ data (right) for (a) CA/P(VP-*co*-VAc), (b) CP/P(VP-*co*-VAc), and (c) CB/P(VP-*co*-VAc) systems. The meanings of three symbols \bigcirc , \times , and \triangle are the same as used in Fig. 2. The miscibility maps are represented in a simplified style retaining the essence of the data shown in Fig. 2.

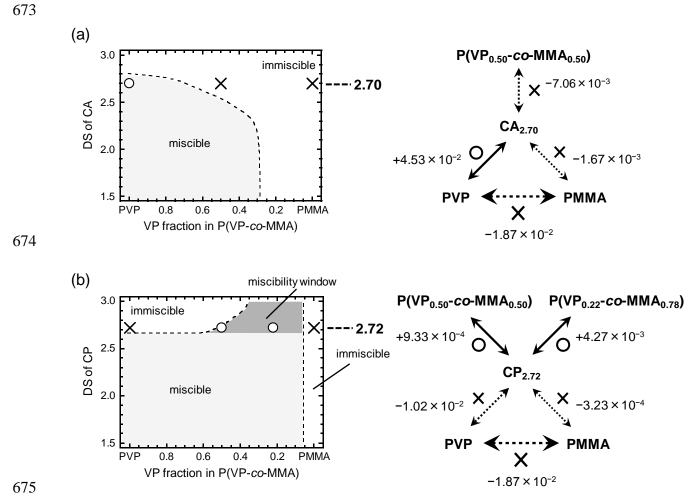


Fig. 5 Miscibility maps (left) with additional illustrations using μ data (right) for (a) CA/P(VP-*co*-MMA) and (b) CP/P(VP-*co*-MMA) systems. The meanings of two symbols \bigcirc and \times are the same as used in Fig. 2. The miscibility maps are represented in a simplified style retaining the essence of the data shown in Fig. 3.

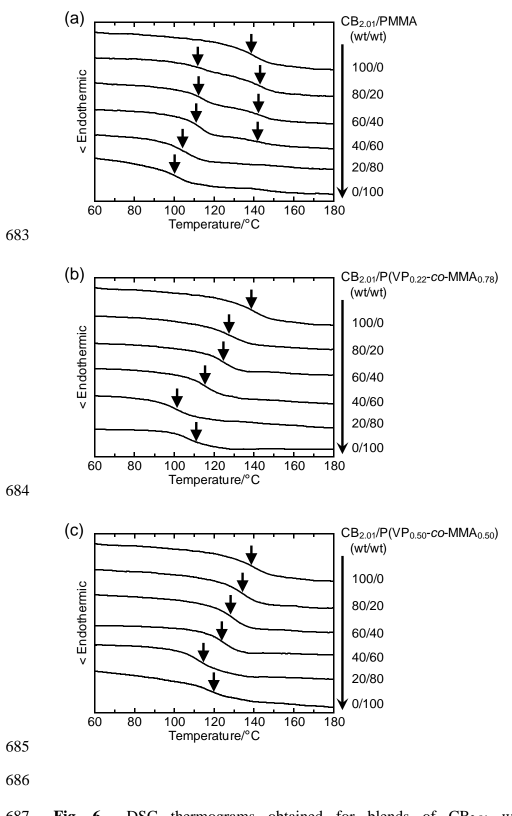


Fig. 6 DSC thermograms obtained for blends of CB_{2.01} with (a) PMMA, (b) 688 $P(VP_{0.22}\text{-}co\text{-}MMA_{0.78})$, and (c) $P(VP_{0.50}\text{-}co\text{-}MMA_{0.50})$. Arrows indicate a T_g position taken 689 as the midpoint of a baseline shift in heat flow.

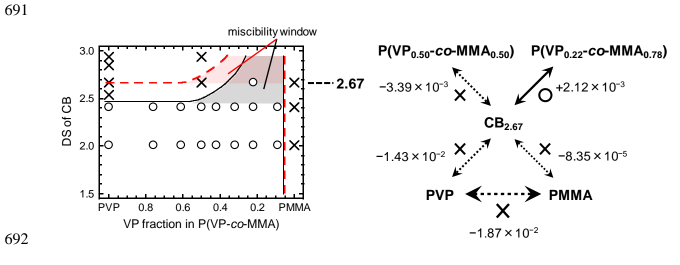
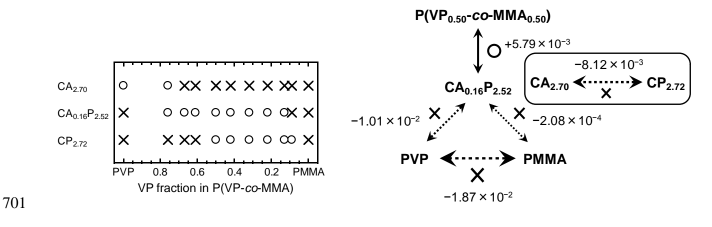


Fig. 7 Miscibility map (left) and additional illustration (right) using μ data for CB/P(VP-*co*-MMA) blends. The meanings of two symbols \bigcirc and \times are the same as used in Fig. 2. Solid lines in the map represent a boundary partitioning the miscible and immiscible regions for the CB/P(VP-*co*-MMA) system, and, for comparison, the corresponding boundary for the CP/P(VP-*co*-MMA) system (Fig. 5b) is drawn by broken lines.



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Fig. 8 Mapping of miscibility data (Sugimura et al. 2013b) (left) and additional illustration in μ terms (right) for CA_{0.16}P_{2.52}/P(VP-*co*-MMA) blends. For comparison, miscibility data for the corresponding blends using CA_{2.07} and CP_{2.72} (see Fig. 3) are also mapped in the left figure. The meanings of two symbols \bigcirc and \times are the same as used in Fig. 2.