Research Article

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Equibiaxially stretchable colorless and transparent polyimides for flexible display substrates

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Abstract: 4,4'- (4,4'-isopropylidenediphenoxy)bispthalic anhydride (BPADA) was reacted with three structurally different diamines to produce poly(amic acid)s, which were then imidized to produce colorless and transparent polyimide (CPI) films through stepwise thermal cyclization. The three amines used to synthesize CPI based on BPADA are: bis(3-aminophenyl)sulfone (APS), p-xylyenediamine (p-XDA), and bis[4-(3-aminophenoxy)-phenyl] sulfone (m-BAPS). The obtained CPI films were almost colorless and exhibited excellent optical transparencies. The solubility of the CPI films in various solvents was investigated, and all the CPI films were found to be soluble in common solvents such as chloroform, dichloromethane, N,N'-dimethyl acetamide, and pyridine. The thermo-optical properties and oxygen transmission rates (O\textsubscript{2}TRs) of the CPI films were examined for various biaxial stretching ratios in the range of 100–150%, and their properties were compared. When the stretching ratio changed from 100 to 150%, the glass transition temperature and yellow index did not show any significant change; however, the O\textsubscript{2}TR decreased for all CPI films.

Keywords: colorless and transparent polyimide; organosoluble polyimide; equibiaxial stretching; thermo-optical property; gas permeation

1 Introduction

Recently, various techniques have been developed to synthesize colorless and transparent polyimides (CPIs). For example, a -CF\textsubscript{3} group, which is a strong electron-withdrawing group, is often used as a substituent, or bent monomer structures are used to prevent charge-transfer complexes (CT complexes) from being formed in linear structures [1–3]. The resulting CPIs exhibit superior solubility, gas-barrier property, and optical properties, compared to other commercialized PIs [4–6]. CPIs with such outstanding characteristics find applications in several fields, as flexible electronic polymeric materials [5, 6].

CPIs can replace the glass in display devices, and have many applications in electronic devices. They can provide great advantages when used in electronic devices because they can be synthesized easily and mass-produced. Optically, CPI films are great alternatives for the glasses that are widely used in flat-panel displays such as plasma discharge panels (PDPs) and liquid crystal displays (LCDs) [7, 8]. Over the past few years, the use of flexible and transparent materials with a conducting oxide layer such as indium tin oxide (ITO) has increased in the fields of display substrates and microelectronics [8-10]. However, ITOs have some drawbacks because indium is a rare and expensive element, which requires processing at high temperatures to achieve good purity when applied on to glass plates. Moreover, glass cannot be used as a display material because it is fragile and not flexible [8]. Therefore, CPI is one of the best alternative materials to overcome the drawbacks of glass.

In the meantime, many researchers have used 4,4'- (4,4'-isopropylidenediphenoxy) bis(phthalic anhydride) (BPADA) anhydride for PI synthesis [11–13]. BPADA has a bended structure as a whole and does not have a -CF\textsubscript{3} substituent on its structure. Thus, if PI is synthesized using BPADA monomer, PI shows colorless and transparent characteristics. Some groups [12, 13] used BPADA to synthesize new PIs, and others [14, 15] used BPADA monomers to make nanocomposites. In particular, our group [16] and Huang et al. [15] used BPADA to synthesize PI nanocomposites using functionalized graphenes. However, studies on CPI films using biaxial stretching have not yet been published.

Many works have reported the effects of uni- or biaxial orientation on the morphology and crystalline behaviors of polymer films [17, 18]. Biaxial stretching means stretching in one direction and stretching perpendicularly to that direction, or stretching it in both directions at the same
time. Stretching makes the polymer structure of the film directional, increases the gas barrier, and improves the tensile strength and modulus. These methods are also commonly used in the film industry to improve the physical properties and introduce birefringence in films.

Many engineering polymer films such as poly(ethylene terephthalate) (PET), poly(lactic acid) (PLA), and polyamide (PA) are used in biaxial stretching processes. Most of these films are used for packaging food and snacks such as chips, breads, biscuits, etc.; they are also used as electronic encapsulating materials [19, 20]. However, in most cases, stretching of the PI film has been reported to cause whitening and/or damage to the film. The whitening phenomenon ultimately leads to deterioration of the optical transparency and physical properties of the PI film [21, 22]. Studies on stretching of the PI films have been quite rare. However, if the monomer structures in PIs are designed well, the PI films can be stretched without any whitening, regardless of the transparency.

The gas permeabilities are closely related to the sizes of the side groups and substituents and the chemical structure of the main chain. For example, bulky side groups or substituents disrupt molecular packing in the polymer chains, thereby increasing the free volume, which in turn, decreases the barrier property [23, 24]. The stretching of the film also has a great effect on the gas barrier property. The gas molecules transmitted through the polymer film are affected by the degree to which they are oriented by the stretching of the film. In fact, chemical structural packing resulting from stretching reduces the diffusion coefficient of the gases passing through the polymer film. This reduction in diffusion coefficient is due to the dense molecular packing caused by the stretching of the molecular structure [23–26].

In this work, we synthesized CPI films through the thermal cyclization of a precursor CPI obtained by the reaction of 4,4′-(4,4′-isopropylidenediphenoxy)bis(phthalic anhydride) (BPADA) with three different diamine monomers: bis(3-aminophenyl) sulfone (APS), p-xylenediamine (p-XDA), and bis[4-(3-aminophenoxy) phenyl] sulfone (m-BAPS). To clarify the relationship between the structure and properties of the CPI films, this paper explains the properties of the CPIs synthesized from the dianhydride BPADA and three different aromatic diamines, as mentioned previously.

This paper also describes the thermal properties, optical transparency, solubility, and oxygen permeability of the CPI films, as functions of the different diamine monomers. We also examine and compare the effects of varying the equibiaxial stretching ratio in the range 100–150% on the thermal properties, optical transparency, and oxygen permeability of the CPI films.

2 Experimental

2.1 Materials

The dianhydride and diamine monomers were purchased from Aldrich Chemical Co. (Yongin, Korea) and TCI (Tokyo, Japan), and were used without purification. N,N′-dimethylacetamide (DMAC) was purified and stored in molecular sieves (4 Å).

2.2 Preparation of CPI films

Three different CPIs (designated as samples I–III) were synthesized from BPADA and three diamines, namely, APS (sample I), p-XDA (sample II), and m-BAPS (sample III). The chemical structures of the synthetic route are shown in Figure 1.

As the synthesis methods of the three CPI films are almost the same, we will describe the synthesis process of BPADA/APS (sample I) only, representatively. Poly(amic acid) (PAA) was synthesized by reacting BPADA and APS dissolved in DMAC. BPADA (6.8 g; 1.3 × 10⁻² mol) and DMAC (40 mL) were placed in a beaker and reacted at 0°C for 30 min under nitrogen atmosphere. 20 mL of DMAC was added to another beaker with APS solution, and was mixed

![Figure 1: Synthetic routes for CPIs based on BPADA.](image-url)
Table 1: Synthetic conditions of CPI films based on BPADA.

| Sample | Temperature (°C) / time (hr) / pressure (Torr) |
|--------|-----------------------------------------------|
| PAA    | 0/1/760 → 25/14/760 → 50/2/760 → 80/1/1     |
| CPI    | 110/0.5/760 → 140/0.5/760 → 170/0.5/760 → 200/0.5/760 → 230/0.5/760 → 250/0.5/760 |

Table 2: Thermal properties of CPIs based on BPADA.

| Monomer | I.V. \(^a\) | \(T_g\) (°C) | \(T_{\text{ib}}\) (°C) | wt\(^{500\circ\text{C}}\) (%) | CTE\(^d\) (ppm/°C) |
|---------|-------------|---------------|----------------------|-----------------|-----------------|
| I       | 0.87        | 204           | 503                  | 57              | 43.94          |
| II      | 0.94        | 182           | 509                  | 34              | 58.02          |
| III     | 0.92        | 196           | 527                  | 59              | 52.43          |

\(^a\)Inherent viscosities were measured at 0.1 g/dL concentration in N,N\(^'-\)dimethylacetamide at 30°C.
\(^b\)2% initial decomposition temperature.
\(^c\)Weight % of the residue at 600°C.
\(^d\)Coefficient of thermal expansion range was 50–150 °C.

The films were equibiaxially stretched using a biaxial stretching machine, with stretching ratios in the range 100–150% at a strain rate of 1 mm/s. The stretching temperatures were 260°C for APS (sample I), 240°C for p-XDA (sample II), and 255°C for m-BAPS (sample III). The obtained films were warped and torn into wave patterns when the stretching temperatures exceeded the optimum temperature.

The biaxial stretching of all CPI films proceeded in the same manner. No whitening was observed even when the biaxial stretching ratio was 150%. When all the CPI films were stretched biaxially by more than 150%, the surface became nonuniform and tore, failing to produce a good film.

### 2.4 Characterization

Differential scanning calorimetry (DSC) and thermogravimetric analysis (TGA) were performed under N\(_2\) conditions using the DuPont 910 equipment. The samples were heated and cooled at a rate of 20°C/min. The coefficients of thermal expansion (CTEs) of the films were evaluated with a macro-expansion probe (TMA-2940), which was used to apply an expansion force of 0.1 N to the films at a heating rate of 5°C/min in the temperature range of 50–150°C.

The ultraviolet-visible (UV-vis) spectra of the CPI films were recorded on a Shimadzu UV-3600 instrument. The color intensity of the CPI film was measured using a Minolta spectrophotometer. The film was usually 71–76 µm thick. A CIE-D light source was used, and the CIELAB color difference equation was used to obtain the value of yellow index (YI). The O\(_2\) permeability of the sample was obtained by the ASTM E-96 method using Mocon DL 100. The O\(_2\) transmission rate (O\(_2\)TR) was measured at a temperature of 23°C, relative humidity of 0%, and pressure of 760 Torr. The CPI film was stretched biaxially, with a 4 × 4 stretching magnification, using a biaxial stretching tester (X6H-S, Toyo Seiki-Sho Ltd).

### 3 Results and discussion

#### 3.1 FT-IR analysis

The formation of PAA and the completion of imide formation from amic acid were confirmed by examining FT-IR spectra; the spectra of PAA and the PIs are shown in Figure 2. Only representative IR results for sample I (BPADA/APS) is discussed here. The C=O stretching peaks

![Figure 2: FT-IR spectra of PAA and PI (sample I).](image-url)
at 1700 and 1602 cm\(^{-1}\) are due to the acid and amide groups of PAA, and shift to higher frequencies in the imides, specifically to approx. 1782 cm\(^{-1}\) (C=O, in phase) and 1712 cm\(^{-1}\) (C=O, out of phase). In addition, the presence of a feature at 1378 cm\(^{-1}\) corresponding to C-N-C stretching confirms the formation of the imides [27].

### 3.2 Thermal behavior

The thermal behaviors of the CPIs fabricated using three different diamines are listed in Table 2. The glass transition temperatures \(T_g\) were in the range of 182–204°C, depending on the structure of the diamine. The molecular chains containing the sulfonyl (SO\(_2\)) group (APS, sample I) were much stiffer and rigid than those containing methylene (p-XDA, sample II) or ether (m-BAPS, sample III) linkages. Thus, the \(T_g\) value for sample I (204°C) was higher than that of the CPIs containing p-XDA (182°C) or m-BAPS (196°C) (see Table 2). The factors affecting the increase in \(T_g\) value were: (1) the effects of the rigid rod-like monomer structures on the free volume of the polymer, and (2) the restriction of inter-reacting polymer chains within the polymers, which prevented the segmental motion of the chains [28, 29]. The DSC curves of the CPIs with the three different diamine monomers are shown in Figure 3.

The initial decomposition temperatures \(T_{D1}\) of samples I–III were in the range of 503–527°C (see Table 2). In sample I, because of the difficulty in rotating freely about its own axis, the rigid SO\(_2\) group caused high torsional deformation. In addition, SO\(_2\) could be released easily because of the steric hindrance and conformational energy produced by the high torsion under high temperatures. As a result, SO\(_2\) separated easily into radicals [30]. Hence, the CPI containing SO\(_2\) groups showed the lowest \(T_{D1}\) value. The \(T_{D1}\) of sample II was lower than that of sample III because of the presence of flexible alkyl (–CH\(_2\)–) linkages, which degraded easily upon heating. However, sample III exhibited a higher \(T_{D1}\) than sample I because of the presence of a larger number of phenyl groups and the flexible ether linkage between the phthalimide units. These ether linkages allowed the free rotation of the SO\(_2\) group, inhibiting the dissociation of SO\(_2\) radicals in m-BAPS. The wt\(_{600}^R\) of samples I–III were in the range of 34–59%. Sample III exhibited a higher wt\(_{600}^R\) (59%) than sample I (57%) because of the presence of a larger number of phenyl groups per monomer unit in the entire molecular structure. Sample II showed the lowest weight residue (34%) because of the low thermal stability of the alkyl structure in the main chain. The TGA thermograms of samples I–III are shown in Figure 4.

The CTE depends on molecular rearrangement. High molecular rearrangement produces low CTE values and, eventually, low residual stress. When the temperature increases, the in-plane-oriented CPI molecules tend to relax in a direction normal to their original direction and, eventually, expand mainly in the out-of-plane direction [31, 32]. In the case of samples I–III, the rigid rod-type molecular structure present in the main chain made the movement of molecules difficult, and this led to low CTE values. The CTE values of the CPIs were 43.94–58.02 ppm/°C, depending on the molecular structure, and the values are summarized in Table 2. Sample I showed the lowest CTE of 43.94 ppm/°C because it was easy to build up close molecular packing because of the rigid rod-type SO\(_2\) structure and high intermolecular attraction between the molecular chains and chains, resulting in lower CTE [33, 34]. On the other hand, sample II showed the highest CTE of 58.02 ppm/°C because of the flexible (–CH\(_2\)–) linkages, which reduced the close packing. The CTE results obtained for various stretching ratios are shown in Figure 5.

![Figure 3: Differential scanning calorimetry (DSC) thermograms of CPIs based on BPADA.](image)

![Figure 4: Thermogravimetric analysis (TGA) thermograms of CPIs based on BPADA.](image)
3.3 Optical transparency

The optical properties of all solvent-cast CPI films were measured using a UV-vis. spectrometer, and the results are shown in Figure 6. The optical transparencies of the CPIs were clarified from the cutoff wavelengths ($\lambda_0$), transmittance at 550 nm, and yellow index (YI), as listed in Table 3. $\lambda_0$ is the first wavelength at which the transmittance occurs. YI is an index value indicating the degree of yellow.

Table 3: Optical properties of CPIs based on BPADA.

| Monomer | Film thickness ($\mu$m) | $\lambda_0$ (nm) | 550 nm trans (%) | YI $^b$ |
|---------|------------------------|-----------------|-----------------|-----|
| I       | 76                     | 371             | 98              | 1.27 |
| II      | 73                     | 362             | 92              | 2.15 |
| III     | 71                     | 358             | 92              | 2.62 |

$^a$ Cutoff wavelength. $^b$ Yellow index.

All CPI films showed values of $\lambda_0 < 400$ nm and 92–98% transmittance at 550 nm. CPI films containing SO$_2$ (sample I) showed the lowest YI value (YI = 1.27), and sample I was almost colorless. In general, it can be said that it is colorless when the YI value is 5 or less. The value of YI was almost similar to that of poly(methyl methacrylate) (PMMA) (YI = 1.50) [35].

Sample I, which contained SO$_2$, showed lower YI values than samples II and III with methylene and ether linkages, respectively. The electron-withdrawing and bulky SO$_2$ group in the diamine monomer probably contributed to the decrease of CT complexes between the main chains through steric hindrance and inductive effects [36]. On the contrary, the electron-donating ether group (Sample III) increased the intermolecular CT complexes by supplying electrons to the main chain phthalimide moiety, resulting in a higher YI value (YI = 2.62), as shown in Table 3. However, all CPI films containing different diamine monomers were almost colorless and transparent; the letters could be read easily through the films (see Figure 7). The transparency of the sample I was observed with varying stretching ratios of 100% to 150% in order to maintain transparency even after biaxial stretching, as shown in Figure 8. Regardless of the biaxial stretching %, all films were clear and transparent and no whitening was observed.

The colorlessness and transparency of the CPI films based on BPADA could be described by the reduction of intermolecular interaction. The bent structure and ether linkages in BPADA were effective in interrupting the CT complex formation between the polymer chains through steric hindrance and inductive effect [36, 37].
### Table 4: Solubility of CPIs based on BPADA.

| Monomer | Acetone | CHCl<sub>3</sub> | CH<sub>2</sub>Cl<sub>2</sub> | DMAc | DMF | DMSO | CH<sub>3</sub>OH | NMP | Pyridine | THF | Toluene |
|---------|---------|------------------|------------------|------|-----|------|-------------|-----|----------|-----|---------|
| I       | x       | o                | o                | o    | x   | o    | o           | o   | o        | x   |         |
| II      | x       | o                | o                | o    | x   | x    | o           | o   | o        | o   | x       |
| III     | x       | o                | o                | o    | o   | o    | x           | x   | o        | o   | △       |

⊚: Excellent, O: Good, △: Poor, ×: Very poor.

DMAc: N,N'-dimethylacetamide, DMF: N,N'-dimethylformamide, DMSO: dimethyl sulfoxide, NMP: N'-methyl-2-pyrrolidone, THF: tetrahydrofuran.

### Table 5: Thermo-optical properties of CPIs based on BPADA for various equibiaxial stretching ratios.

| Biaxial stretching ratio (%) | I | II | III |
|-----------------------------|---|----|-----|
| 100 (unstretched)           | 204 | 182 | 196 |
| 120                         | 207 | 184 | 196 |
| 130                         | 206 | 184 | 197 |
| 150                         | 205 | 184 | 196 |

a: Yellow index.

### 3.4 Solubility

CPI is a very special polymer material that is very strong and has heat and chemical resistance [38, 39]. However, because of the fully cyclized structure, it is often not melted or dissolved, which limits its use as an engineering material. Hence, solubility will influence its processability for polymeric applications [40, 41].

The solubilities of the CPIs were measured in various solvents and the results are summarized in Table 4. Samples I–III exhibited somewhat limited solubility. All the films were soluble in chloroform, dichloromethane, N,N'-dimethyl acetamide (DMAc), and pyridine; however, they were insoluble in acetone, dimethyl sulfoxide (DMSO), methanol, and toluene.

The solubility of sample II was better than that of samples I and III in NMP and THF. The difference in solubility depends on the monomer structure present in the main chain. Sample II with a flexible methylene structure had weak molecular chain interactions and could not provide effective close packing, resulting in increased solubility [1, 36, 42]. However, the solubility level was not affected significantly by the CPI composition in terms of the molecular structure of the diamine.

### 3.5 Thermo-optical properties of equibiaxially stretched films

In general, polymer films such as polyethylene (PE), polypropylene (PP), poly (vinyl chloride) (PVC), and poly (ethylene terephthalate) (PET), which are used as packaging materials, are uniaxial- and biaxial-stretched. However, most polymer films used in electronic- and flexible display-materials are used after biaxial stretching in consideration of optical properties [18, 20, 22].

Table 5 lists the $T_g$ and YI of the unstretched CPIs and the CPIs with various equibiaxial stretching ratios. Compared to the unstretched CPIs, the $T_g$ of the CPIs with various biaxial stretching ratios appeared to be virtually unchanged, regardless of the stretching ratio varying from 120 to 150%. As a result, biaxial stretching has little effect on free volume or segmental motion, and thus does not significantly affect $T_g$.

The YI values obtained for various stretching ratios are also summarized in Table 5. Even though the stretching ratio increased from 120% to 150%, the YI values of all samples were almost constant and all the films were colorless and transparent. In addition, there was no difference when compared with the unstretched film (100%). From these results, it was found that stretching ratios had no effect on the clarity and optical properties of the CPI films. The transparencies of the films stretched to various percentages are not shown here because they are the same as the results in Table 5.

### 3.6 Oxygen permeability of the equibiaxially stretched films

There are two important factors necessary to obtain a polymer film with increased gas barrier properties [43, 44]: (1)
the main chains in the polymer must be stiffened by hindering intra-rotational movements, and (2) the intermolecular close packing of the main chains can suppress good permeability.

In this work, our results are explained in terms of the relative permeability $P_{st}/P_p$, where $P_p$ is the permeability of the unstretched polymer and $P_{st}$ is the permeability of the stretched polymer. The $O_2$TR of CPI films with various equibiaxial stretching ratios ranging from 100 to 150% are summarized in Table 6. Compared to samples II and III, the molecular chains containing $SO_2$ groups in sample I were straighter and harder than those containing methylene and ether linkages; thus, the $O_2$TR of sample I ($0.64 \text{ cc/m}^2/\text{day}$) could be expected to be lower than that of sample II ($2.03 \text{ cc/m}^2/\text{day}$) or sample III ($3.62 \text{ cc/m}^2/\text{day}$).

Table 6: Oxygen permeabilities of CPIs based on BPADA for various equibiaxial stretching ratios.

| Biaxial stretching ratio (%) | I $O_2$TR $^a$ | $P_{st}/P_p$ $^b$ | II $O_2$TR | $P_{st}/P_p$ $^b$ | III $O_2$TR | $P_{st}/P_p$ $^b$ |
|-----------------------------|---------------|----------------|-----------|----------------|-----------|----------------|
| 100 (unstretched)          | 0.64          | 1            | 2.03      | 1            | 3.62      | 1            |
| 120                         | 0.53          | 0.83         | 1.04      | 0.51         | 0.05      | 0.01         |
| 130                         | 0.43          | 0.67         | 0.34      | 0.17         | 0.07      | 0.02         |
| 150                         | 0.37          | 0.58         | 0.37      | 0.18         | 0.06      | 0.02         |

$^a$ Oxygen transmission rate (cc/m$^2$/day). $^b$ Stretched polymer permeability/unstretched polymer permeability (i.e., relative permeability rate).

For equibiaxial stretching ratios between 100 and 130%, the $O_2$TR of sample I was found to decrease linearly from 0.64 to 0.43 cc/m$^2$/day (33% reduction). When the stretching ratio increased up to 150%, the $O_2$TR decreased to 0.37 cc/m$^2$/day. This tendency could be attributed to the orientation of the CPI film molecules by stretching, which showed a greater effect on gas barrier properties as the stretching ratio increased [45–47].

The $O_2$TRs for samples II and III with various equibiaxial stretching ratios were measured, and are summarized in Table 6. The $O_2$TR of sample II decreased from 2.03 to 0.34 cc/m$^2$/day (83% reduction) as the biaxial stretching ratio increased to 130%, and then remained constant for stretching ratios up to 150% (0.37 cc/m$^2$/day). Similar behavior was observed in the case of sample III. The $O_2$TR of sample III decreased significantly when the biaxial stretching ratio changed from 100 to 120%, and then leveled off upon stretching to more than 120%. For example, the biaxial stretching of only 120% for sample III resulted in 99% reduction of $O_2$TR ($0.05 \text{ cc/m}^2/\text{day}$), with respect to that of the unstretched PI film (3.62 cc/m$^2$/day). As the biaxial stretching ratio changed from 120 to 150%, the $O_2$TR of sample III remained fairly constant ($0.05–0.07 \text{ cc/m}^2/\text{day}$) and close to zero. This reduction in $O_2$TR is closely related not only to diffusion by changes in polymer chains during biaxial stretching but also to solubility between film and gas. The permeation is explained by the following equation [47]

$$P = D \times S$$

Where, $P$ = permeation, $D$ = diffusion, and $S$ = solubility.

The $D$ value can be explained by the movement of the polymer chains changed by biaxial stretching. As the crystallization of the polymer chain increases with the increase of the biaxial stretching ratio, the $P$ value decreases. However, the $S$ value obtained between film and gas is very difficult to explain. Since our sample showed no change in thermal and optical properties even if the stretching ratio increased from 100 to 150%, our results can be assumed to be affected more by $S$ than the $D$ effect.

4 Conclusions

The $T_g$ of CPI films containing three different amine monomers were in the range of 182–204°C, and thermal degradation was not observed at temperatures below 500°C, indicating excellent thermal stability. The CTE values of the three CPI films were in the range of 43.94–58.02 ppm/°C. Optical transparencies showed 92–98% transmittance at 550 nm and the YI values were in the range of 1.27–2.62; i.e., almost colorless. The $O_2$TR values were slightly different, depending on the amine structure, but they were usually in the range of 0.64–3.62 cc/m$^2$/day.

The CPI films were equibiaxially stretched with stretching ratios ranging from 120 to 150%, and the $T_g$ and $O_2$TR of the stretched films were investigated in detail. The $T_g$ of the films appeared virtually unchanged in the DSC thermograms, regardless of the biaxial stretching ratios up to 150%. The values of $O_2$TR, however, decreased upon increasing the stretching ratio.

In conclusion, it appears that the structure of the di-amine monomers influences the thermal properties, optical transparency, solubility, and gas permeation of the CPI films. We strongly claim that the present CPI films are vi-
able candidates for flexible display substrates in electrophotonic applications even though their long-term reliability needs to be tested further.

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