

nal group is parallel for the oligosilylenes having even number of silylene unit, but nonparallel for the oligosilylenes having odd number of silylene unit, and the former is symmetric but the later is not. Thermal properties of pentamer and hexamer are not investigated yet because of the difficulty of the synthesis of them, but this assumption is possibly reasonable to explain the even-odd effect of chain length on the glass-forming properties. Although such even-odd effect is known for the effect of alkyl spacer length of liquid crystalline materials, there is no report of the effect on glass-forming property.

Table 1. Thermal properties of oligosilylenes^a

	$T_g/^\circ\text{C}^b$	$T_c/^\circ\text{C}^c$	$T_m/^\circ\text{C}^d$
S1	-89		
S2			142
S3	14 ^e		103 ^f
S4	(31) ^g	(86) ^g	215 ^f
S7	77 ^e		217 ^f

^aScan rate: 10 degree min⁻¹. ^bGlass-transition temperature. ^cCrystallization temperature. ^dMelting temperature. ^eSecond heating. ^fFirst heating. ^gSecond heating after the melt was cooled faster than 10 degree min⁻¹.

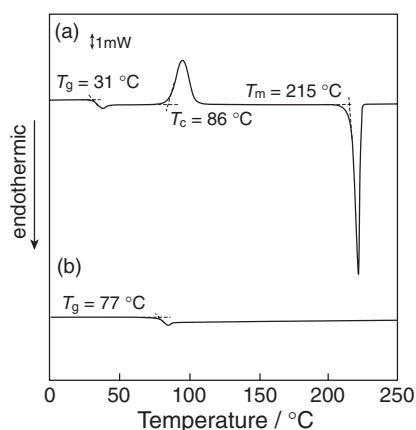


Figure 2. DSC thermograms of second heating process of (a) **S4** and (b) **S7**. Scan rate = 10 degree min⁻¹.

To obtain the information on the morphology of the glassy samples, X-ray diffraction (XRD) was investigated (Figure 3). While XRD of recrystallized **S7** showed typical sharp peaks due to the crystallinity of the sample, XRD of the glassy **S7** showed two broad peaks at $2\theta = 8.5^\circ$ ($d = 10.4 \text{ \AA}$) and 20.6° ($d = 4.31 \text{ \AA}$). It is reported that polysilylenes having two alkyl groups show a strong peak at $7-9^\circ$ indexed as superposition of d_{110} and d_{020} of an orthorhombic lattice with hexagonally packed chains, and a broad peak centered at around 20° indexed as d_{013} .⁸ Thus, peaks observed in glassy **S7** can be assigned similarly to the above-mentioned case of polysilylenes. From these data, the distance between two neighbor oligosilylenes can be estimated as $12.0 \text{ \AA}$, which is coincident with the diameter of the oligo-(diphenylsilylene) molecules having trans-zigzag structures. In polarized microscopy, the image of glassy samples in cross-nicol state is dark, and suggests to be isotropic. Thus, the arrangement of molecules in glassy **S7** might be mainly amorphous, with only two-dimensional order of hexagonally packed structure.

To our best knowledge, our result is the first report on the

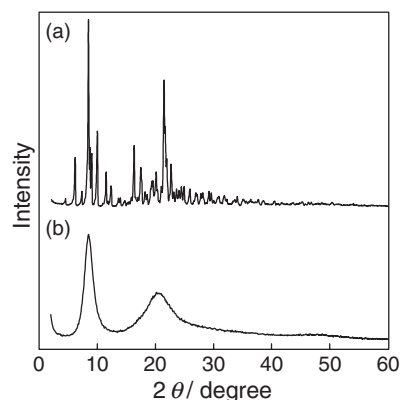


Figure 3. X-ray diffraction patterns of (a) crystalline and (b) glassy **S7**.

creation of glassy low-molecular weight materials based on silicon-containing compounds.

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