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Manika Varma-Nair, and Bernhard Wunderlich





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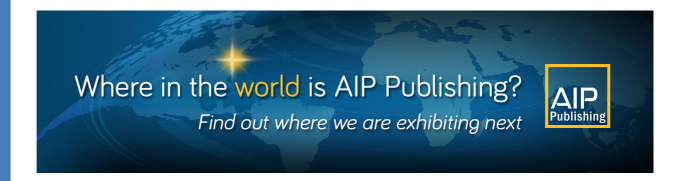
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Heat Capacity and Other Thermodynamic Properties of Linear Macromolecules

X. Update of the ATHAS 1980 Data Bank

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The prior published ATHAS 1980 Data Bank of experimental heat capacities of linear macromolecules that included critically reviewed material on almost 100 polymers is updated. In addition, the data bank has been computerized so that future updates can be made continuously, and new print-outs or computer files will be available from the authors from 1990 on.

Key words: data bank, enthalpy, entropy, Gibbs function, glass transition, heat capacity, linear macro-molecule, polymer.

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1. Introduction

In an effort to correlate the macroscopically observed heat capacity of linear macromolecules with the microscopic structure, a compilation and critical analysis of all available literature on experimental heat capacities was completed in 1980. In a series of nine publications heat were recommended for capacities selenium,1 polyethylene,² various types of polyoxides,³ polypropylene,4 polystyrene,5 acrylic polymers,6 other carbon backbone polymers,⁷ polyesters and polyamides,⁸ and various aromatic and inorganic polymers.9 The data bank of heat capacities was incorporated in ATHAS (Advanced THermal AnalysiS). Excerpts and reviews based on this data bank were contributed to the Polymer Handbook¹⁰ and the Encyclopedia of Polymer Science and Technology.¹¹ Since then new heat capacity measurements in our laboratory were concentrated in the areas where prior efforts to establish the data bank revealed shortcomings. In addition, a computer search from 1981 to the end of 1989 accumulated all new heat capacities measured in other laboratories. The ATHAS data bank of recommended experimental heat capacity data has thus been updated to the level of 1990 and contains now data on nearly 150 macromolecules. The data bank has been transferred to an IBM-compatible PC to be updated continually. This paper is the final publication in the printed literature displaying a review of experimental heat capacities of linear macromolecules. The data bank will in the future be updated every six months and is then available on software based on Lotus 1-2-3. In addition to the critically reviewed experimental data, descriptive text on individual polymers as presented here, and data tables on thermodynamic functions [enthalpy, entropy and free enthalpy (Gibbs function)] derived with the help of theory-based computed heat capacities from 0 to 1000 K, fitted to the here discussed experimental data over the range of measurement will also be available on request from the authors.

A wide variety of polymers are reviewed in this paper using the methods outlined before. 1.2 These include carbon-backbone polymers and polymers with hetero-atoms and aromatic groups in the main- and side-chains. In ad-

dition, polymers with long pendant side groups have been covered. In all, new recommended heat capacities are presented for 42 new polymers and four polymers on which recommendations had not been made before. The ATHAS Table of Thermal Properties given in the Appendix shows a summary of all updatings performed since 1980. A complete table containing data on nearly 150 macromolecules is available in the ATHAS data bank and is updated every six months.

2. Update on Polymers Already Present in ATHAS Data Bank 1980

Based on the standards of acceptability laid down in 1980,1 all the available literature heat capacities were critically reviewed for quality and reliability. For this purpose sample characterization, experimental technique used, error limits and accuracy of representation of data are considered. In case recommended heat capacities had already been included in the ATHAS data bank (1980), the new data were compared with it and if deviations were less than $\pm 1\%$, the new data were incorporated in the data bank without altering the recommended data. All acceptable data (smoothed, in case unsmoothed data are reported) are stored in the standard temperature intervals of the data bank. All the available and acceptable data are given in Tables A1-A57 which have been deposited with the Physics Auxillary Publication Service of the American Institute of Physics. The tables contain more significant figures than justified by both source of data and data treatment. The extra significant figures are included only for the purpose of smooth representation.

Additional heat capacities were found in the literature from 1980 onwards on polyethylene, polypropylene, poly(1-butenylene) (cis and trans), poly(vinyl fluoride), poly (vinylidene fluoride), polytrifluoroethylene, polytetrafluoroethylene, polystyrene, polyoxymethylene, polyoxyethylene, poly(ethylene terephthalate), polycarbonate, poly (oxy-2,6-dimethyl-1,4-phenylene), poly (diethyl siloxane) and Nylon 6.12. The recommended data (1990) of polyethylene are only insignificantly changed from those of 1980. This includes the change of glass transition temperature (temperature of half devitrification) from 252 K1 to a value of 237 K which is based on more extensive experiments done in our laboratory. The five investigations 12-16 published after 1980 that did not meet our standards and have, therefore, not been included in the update are listed in Table 1. Heat capacities of polypropylene are altered slightly in the glass transition region to include the glass transition temperature as 270 K (instead of the previous value at 260 K) as reported by Bair and Schilling¹⁷ on atatic polypropylene made by hydrogenation of poly (1,3-dimethyl-1-butylene). The change in heat capacity at $T_g(\Delta C_p)$ thus obtained is 22.93 J/(K mol) and close to the previous recommended value of 23.54 J/(K mol). Three additional investigations 18-20 did not meet our data standards as measurements were made on uncharacterized commercial samples over limited temperature ranges and have, therefore, not been included in the update (see Table 1).

Four additional investigations 15,18,21,22 have been reported for polystyrene for which the recommended heat capacity had earlier been arrived-at based on nineteen out of the forty-five investigations reported in the literature. Among these new publications the data reported by Chang et al. Pand Richardson et al. Pagreed to better than $\pm 1\%$ over the range of measurement, so that it was not considered necessary to alter the ATHAS recommended data 1980. The details of the measurements are given in Table 2 and the new acceptable data are given in Tables A1-A4. Two investigations which did not meet our standards of acceptability are listed in Table 1 together with the reasons for their exclusion.

On the basis of measurements on well characterized, semicrystalline poly (vinyl fluoride), poly (vinylidene fluoride) and poly (trifluoroethylene);²³ polyoxymethylene and polyoxyethylene;²⁴ it was possible to arrive at a preliminary, recommended set of heat capacities for the melt. In addition Feklina et al.²⁵ and Rekhleta et al.²⁶ have reported heat capacity of poly (vinylidene fluoride) (PVF2) and poly (trichlorofluoroethylene) (PCTFE), respectively. These measurements have been excluded as for PVF2 the data are on uncharacterized, semicrystalline polymers above the glass transition temperatures and for PCTFE data are presented over a narrow temperature range in the form of two small graphs and the main topic of this investigation was the effect of pressure on heat capacity.

The recommended heat capacities of fluorinated polymers above and below the melting temperature, but above glass transition temperature are given in Table 3 and can be described by two linear equations. The data for the melt between $T_{\rm g}$ and $T_{\rm m}$ were obtained by the authors by subtracting the fractional contribution of solid heat capacity (calculated from the vibrational spectrum)²³ from the semicrystalline data. The liquid heat capacities thus obtained were found to be non-linear. The liquid heat capacities of all polyfluoroethylenes could be established as folows.²³

```
N_{\rm F} (104.749 + 0.0568 T) + (1-N_{\rm F}) (36.840 + 0.0854 T) (1)
and below 480 K as: C_p =
N_{\rm F} (54.976 + 0.162 T) + (1-N_{\rm F}) (37.754 + 0.0829 T) (2)
```

above 480 K: $C_p =$

Where $N_{\rm F}$ is the mole fraction of fluorine atoms. In case of PVF2 and P3FE agreement between data calculated using these equations and the experiment was better than \pm 1%. The agreement was less good for PVF, but still less than 5%. This disagreement could, in addition, have been caused by the beginning of decomposition in PVF. The recommended melt data for these polymers was, therefore, obtained with these equations and is given in Table 3.

The recommended heat capacities for molten polyoxymethylene and polyoxyethylene extrapolated down to $T_{\rm g}$ (190 K for POM, and a new recommended value of 206 K for POE) are given in Table 4. They are based on the data listed in Table 2 and a critical review of all data on aliphatic polyoxides.²⁴

Additional data poly (4,4'-isopropylidene for diphenylene carbonate)27 and poly (oxy-2,6-dimethyl-1,4phenylene)²⁷⁻³⁰ are given in Tables A10-A12. The details of the measurements are given in Table 2. Recommended heat capacities (1990) of polycarbonate are only insignificantly changed in the glass transition region, from those of 1980. Based on an extensive investigation carried out in our laboratory, T_g is changed from 418 K to 424 K. The ΔC_p remains almost unchanged at about ~ 49 J/(K mol). Melt heat capacities (1980) were retrieved from graphs and were, therefore, not so accurate. These have also been replaced by the more reliable new data²⁷ which is slightly different from the previous recommended heat capacities.

In case of poly(oxy-2,6-dimethyl-1,4-phenylene) the previously recommended data (1980),³ were based on only one investigation. This has been updated to include Jauhianen's²⁸ and Cheng's³⁰ data. Later is based on a more extensive work carried out in our laboratory and agrees with Karasz's³ and Jauhianen's data in the region of overlap to a precision of better than 1%.

No recommendations were made earlier for heat capacities of polytetrafluoroethylene, poly(1-butenylene) (cis and trans), poly(ethylene sebacate) and poly(diethyl siloxane) as the reported data were found to show large uncertainties. In addition, heat capacity data for Nylon 6.12 were based on only one investigation. Since then, new measurements have been carried out on these polymers and a recommended set of data will be discussed in this paper together with the polymers that were not discussed before.

TABLE 1. Investigations not included

Polymer	Reference	Reasons for exclusion
Polyethylene	1. Mathot and Pÿpers (1983) [12].	Data could not be read accurately from to small graphs.
	2. Fritzsch (1983) [13].	Heat capacity of crosslinked thin layer samples of unknown degree of crystallinity.
	3. Finlayson and Mason (1985) [14].	Sample characterization not reported. Heat capacity data below 2 K reported for semicrystalline samples.
	4. O'Reilly and Connelly (1985) [15].	Semicrystalline data reported from 325 to 440 K are within the melting range and data could not be read accurately from the small graphs.
	 Shut, Musaelyan, Besklubenko, Karpovich and Kasperskii (1986) [16]. 	Heat capacity on low density polyethylene before and after γ -irradiation are reported. Data could not be read accurately from too small graphs.
Polypropylene	1. Zhongde, Xuexin (1983) [17]	Data retrieved from the graph was $\sim 3\%$ lowe below T_g and $\sim 3\%$ higher above T_g than the recommended data.
	2. Vargha-Butler and Neumann (1982) [18].	Commercial sample, characterization not reported. Deviations of nearly 5% from the recommended data.
	3. Mostafa and Gaber (1986) [19].	Commercial sample, characterization not reported. Decrease in heat capacity at 350 K has been reported.
	4. Cometto and Giulianelli (1986) [20].	Commercial samples, characterization not reported. Data could not be read accurately from too small graphs.
Polystyrene	1. Vargha-Butler and Neumann (1982) [18].	Sample characterization not reported. Large deviations from the recommended data.
	2. O'Reilly and Connelly (1985)	Small graphs and very few data points.

Table 2. Heat capacity measurements of various linear macromolecules

Polymer	Investigator Sar	nple num	ber, characterization	Temperature range (K)	Experimental technique (claimed uncertainty)	Source of data
Polystyrene	Richardson <i>et</i> <i>al</i> . (1979) [21]	1.	$\frac{\text{Quenched}}{M_{\text{w}}} = 36,000$	250-320	DSC (1%)	Graph
	Chang et al. (1984) [22]	2.	Amorphous ^a $ \underline{\overline{M}}_{n} = 35,800 $ $ \underline{\overline{M}}_{w}/\overline{M}_{n} = 1.045 $	5–360	Adiabatic (Above 50 K it is 0.02%)	Table
		3.	Quenched	80–360	Adiabatic (Above 50 K it is 0.02%)	Table
		4.	Annealed	40–360	Adiabatic (Above 50 K it is 0.02%)	Table
poly (Vinyl- fluoride)	Loufakis (1986) [23]	1.	$\frac{\text{Melt}^{\text{b}}}{M_{\text{w}}} = 126,000$	480–530	DSC-2 (0.1%)	480–530 K: Equation ^c
ooly (Vinyl- idene fluoride)	Loufakis (1986) [23]	1.	$\frac{\text{Se}_{\text{micrystalline}}^{\text{b}}}{M_{\text{w}}} = 120,000$	260–580	DSC-2 (0.1%)	260–310 K: Equation ^d
						450–580 K: Equation ^e
ooly (Tri- fluoro- ethylene)	Loufakis (1986) [23]	1.	Semicrystalline ^f	330–600	DSC-2 (0.1%)	330–410 K: Equation ⁸
cinyiche)						480–600 K: Equation ^h
Polyoxy- methylene	Wunderlich et al. (1985) [24]	1.	$\frac{\underline{M}\text{elt}^{i}}{\overline{M}_{n}} = 38,000$ $\overline{M}_{w} = 76,000$	430–540	DSC-2 (0.1%)	430–540 K: Equation ^j
Polyoxy- ethylene	Wunderlich et al. (1985) [24]	1.	Melt and $\underline{\underline{Semicrystalline}}^{k}$ $\overline{\underline{M}}_{w} = 15,000 - 20$	330–430	DSC-2 (0.1%)	330–430 K: Equation ¹
poly (4,4'- Isopropyli- denediphen- ylene carbonate)	Wunderlich et al. (1986) [27]	1.	Quenched ^m $ \underline{M}_{w} = 40,250 $ $ \underline{M}_{n} = 14,400 $	440–610	DSC-2 (0.1%)	440–610 K: Equation ⁿ
ooly (Oxy-2,6- dimethyl-1,4- phenylene)	Jauhiainen (1982) [28]	1.	Semicrystalline	160–280	Adiabatic	Table
	Bopp et al. (1982) [29]	2.	$\frac{\text{Amorphous}^{\circ}}{\overline{M}_{\text{n}}} = 16,800$ $\overline{M}_{\text{w}} = 57,400$	310–600	DSC (1%)	Table
	Wunderlich <i>et al</i> . (1987) [30]	3.	Amorphous and Semicrystalline $\overline{M}_{w} = 44,000$ $\overline{M}_{n} = 19,000$	220–460	DSC-2 (0.1%)	220–460 K: Equation ^p
		4.	Melt	480–580	DSC-2 (0.1%)	480–580 K: Equation ^q
poly (2- Methyl-1,3- pentadiene)	Zhongde et al. (1985) [17]	1.	$\frac{\text{Quenched}}{M_{\text{m}}} = 11.7 \times 10^4$ $M_{\text{w}} = 12.5 \times 10^4$	230–320	DSC-2 (1%)	Graph

Table 2. Heat capacity measurements of various linear macromolecules - Continued

Polymer	Investigator	Sample numb	ber, characterization	Temperature range (K)	Experimental technique (claimed uncertainty)	Source of data
poly (1-Buten- ylene)	Dainton et al. (1962) [7]	1.	trans poly (1-butenylene 96.2% pure 3.8% 1,2 addition	e) 22–340	Adiabatic (1%)	Table
		2.	cis poly (1-butenylene) 94% pure 3% trans isomer 3% 1,2 addition	22–310	Adiabatic (1%)	Table
	Wunderlich <i>et al</i> .(1986) [31]	3.	$\frac{100\% \ poly(1-\frac{\text{butenylene})^{\text{r}}}{M_{\text{w}}} = 2.3 \times 10^{5}$	220–450	DSC-2 (0.1%)	260–450 K: Equation ^s
		4.	100% trans poly(1-butenylene) ^t	220-450	DSC-2 (0.1%)	220-270 K: Graph
						410–450 K: Equation ^s
poly (1- Butenylene)	Wunderlich et al. (1986) [31]	5.	89% -trans 9% cis, 2% vinyl poly (1-butenylene) ^u $M_{\rm w} = 3.67 \times 10^5$	220–450	DSC-2 (0.1%)	360–450 K: Equation ^s
		6.	83%-trans, 12.4 cis, 4.6%-1,2-trans poly(1-butenylene) ^v	220–450	DSC-2 (0.1%)	310–450 K: Equation ^s
Polytetrafluoro- ethylene	Furukawa et al. (1952) [7]	1.	Teflon ^w Powder $\Delta H_{\rm f} = 1.90 \text{ kJ·mol}^{-1}$	5–365	Adiabatic	Table
		2.	Teflon ^w Average data for molded and quenched samples	5–365	Adiabatic	Table
			Molded: $\Delta H_{\rm f} = 1.744 \text{ kJ·mol}^{-1}$			
			Quenched: $\Delta H_f = 1.82 \text{ kJ} \cdot \text{mol}^{-1}$			
		3.	Teflon ^w Annealed	5-365	Adiabatic	Table
	Noer et al. (1959) [7]	4.	Teflon*	1.4-4.2	Heat pulse (5%)	Equation*
	Reese and Tucker (1965) [7]	5.	Teflon ^w $\rho = 2.160 \text{ Mg m}^{-3}$	1.0-4.5	Transient (10%)	Equation ^y
	Douglas and Harman (1965) [7]	6.	Teflon ^w Powder $w^c = 0.95$	340–560	Drop calorimeter (5%)	Equation ^z
		7.	Teflon ^w Quenched	330–720 (5%)	Drop calorimeter	330-560 K: Equation ^{a a} 660-720 K: Equation ^{a b}
Polytetra- fluoroethylene	Choy et al. (1979) [7]	8.	Teflon ^w $w^c = 0.9$	5–100	Adiabatic (2%)	Table

Table 2. Heat capacity measurements of various linear macromolecules - Continued

Polymer	Investigator	Sample num	per, characterization	Temperature range (K)	Experimental technique (claimed uncertainty)	Source of data
	Salinger and Cieloszyk (unpublished) [7]	9.	Teflon*	0.3-4.4	Heat pulse (2%)	Equation ^{a c}
	S. F. Lau, B. Wunderlich, <i>et al</i> . (1984) [32]	10.	Semicrystalline ^{a d} $\Delta H_{\rm f} = 1.36 \text{ kJ/mol}$ $w^{\rm c} = 0.332$	200–700	DSC-2 (0.5%)	Table
		11.	Semicrystalline ^{a d} \overline{M} w = $\sim 10^6$ $\Delta H_f = 1.76$ kJ/mol $w^c = 0.43$	390–480	DSC-2 (0.5%)	Table
		12.	Semicrystalline ^{a.d} $\overline{M}_{w} = 5 \times 10^{5}$ $\Delta H_{f} = 2.72 \text{ kJ/mol}$ $w^{c} = 0.662$	390–480	DSC-2 (0.5%)	Table
	S. F. Lau, B. Wunderlich, <i>et al</i> . (1984) [32]	13.	Semicrystalline ^{a d,a c} $\Delta H_{\rm f} = 0.703 \text{ kJ/mol}$ $w^{\rm c} = 0.171$	200–700	DSC-2 (0.5%)	Table
		14.	Semicrystalline ^{a d,a f} $w^c = 0.95-0.99$	220–570	DSC-2 (0.5%)	Table
Polytetra- fluoroethylene	S. F. Lau, B. Wunderlich, <i>et al</i> . (1984) [32]	15.	Branched PTFE ^{a d,a g}	220–570	DSC-2 (0.5%)	Table
Polyglyco- lide	Lebedev <i>et al</i> . (1978) [8]	1.	$\overline{M}_{v} = 38,000$ $w^{c} = 0.67$	13–550	Adiabatic (from 13.8 to 30 K 1%) (from 30 to 90 K 0.5%)	Table ^{a h}
					Adiabatic (from 90 to 370 0.25%) (from 370 to 550 K 1%)	Table ^{a h}
	Lebedev et al. (1984) [35]	1.	$\overline{M}_{\rm w} = 1.0 \times 10^6$ $\Delta H_{\rm f} = 10.519 \text{ kJ/mol}$	13.8-5\$0	"	Table ^{a h}
poly (β-Propiolac- lactone)	Lebedev <i>et al</i> . (1984) [35]	1.	$\overline{\underline{M}}_{w} = 3.0 \times 10^{4}$ $\overline{M}_{n} = 8.0 \times 10^{3}$ $\Delta H_{f} = 8.058 \text{ kJ/mol}$ $w^{c} = 0.79$	13.8-400	и	Table ^{a li}
poly (γ-Butyro- lactone)	Lebedev et al. (1984) [35]	1.	$\overline{M}_{n} = 3.0 \times 10^{3}$ $\Delta H_{f} = 8.844 \text{ kJ/mol}$ $w^{c} = 0.67$	13.8-350	n	Table ^{a h}
poly(δ-Valerolac- tone)	Lebedev et al. (1984) [35]	1.	$\overline{M}_{\rm w} = 3.6 \times 10^3$ $\overline{M}_{\rm n} = 1.4 \times 10^3$ $\Delta H_i = 12.74 \text{ kJ/mol}$ $w^c = 0.70$	13.8–350	a	Table ^{a h}
Polycaprolactone	Lebedev <i>et al</i> . (1974) [8] and	1.	Melt crystallized $\overline{M}_{w} = 29,000$ $\Delta H_{f} = 12.27 \text{ kJ/mol}$ $w^{c} = 0.76$	13.8–350	n	Table ^{a h}
	Lebedev <i>et al</i> . (1984) [35]	2.	$\overline{M}_{\rm w} = 29,000$ ${\rm w'}^{\rm c} - 0.63$	13.8 - 350		

Table 2. Heat capacity measurements of various linear macromolecules - Continued

Polymer	Investigator	Samplenum	ber, characterization	Temperature range (K)	Experimental technique (claimed uncertainty)	Source of data
Polyundecano- lactone	Lebedev <i>et al</i> . (1984) [35]	1.	$\overline{M}_{\rm w} = 3.4 \times 10^4$ $\overline{M}_{\rm n} = 1.0 \times 10^4$ $\Delta H_{\rm f} = 27.122 \text{ J/mol}$ $w^{\rm c} = 0.71$	13.8 - 400	н	Table ^{a h}
Polytridecano- lactone	Lebedev <i>et al</i> . (1984) [35]		$[\eta] = 0.82 \text{ d}\ell/\text{g}$ in CHCl ₃ at 298.15 K. $\Delta H_t = 34.5 \text{ kJ/mol}$ $w^c = 0.75$	13.8 – 370	"	Table ^{a h}
Polypropio- lactone	Lebedev et al. (1984) [35]		$[\eta] = 0.93 \text{ d}\ell/\text{g}$ in CHCl ₃ at 298.15 K. $\Delta H_{\rm f} = 42 \text{ kJ/mol}$ $w^{\rm c} = 0.75$	13.8 – 370	n	Table ^{a h}
poly (Ethylene oxalate)	Lebedev et al. (1982) [36]	1.	$\overline{M}_n = 2 \times 10^5$ Quenched sample (amorphous)	8 - 360	Adiabatic $(\pm 0.2\%)$	Table ^{a í}
poly (Butylene adipate)	Rabinonvich <i>et al.</i> (1984) [37]	1.	$\overline{M}_{n} = 1912$ $\overline{D}_{p} = 9.1$ $\Delta H_{f} = 20.83 \text{ kJ/mol}$ $w^{c} = 0.84$	80-470	Adiabatic (80-320 K 0.3%) (320-470 K 0.8%)	Table ^{a j}
poly (Trimethy- lene adipate)	Pan (1987) [38]	1.	Semicrystalline ^{a k}	310–360	DSC-2 (0.1%)	Equation ^{a 1}
poly (Hexa- methylene sebacate)	Pan (1987) [38]	1.	Semicrystalline ^{a k}	340-400	DSC-2 (0.1%)	Equation ^{a m}
poly (Trimethy- lene succinate)	Pan (1987) [38]	1.	Semicrystalline ^{a k}	310–360	DSC-2 (0.1%)	Equation ^{a n}
poly(Ethylene- sebacate)	Wunderlich an Dole (1958) [8]		Semicrystalline $\rho = 1.167 \text{ Mg m}^{-3}$ $\Delta H_f = 72.8 \text{ J/g}$ $w^c = 0.54$	250-410	Adiabatic (1%)	250–310 K: Equation ^{a o} 350–410 K: Equation ^{a p}
	S. Lim and B. Wunderlich (1987) [39]	2.	Semicrystalline, Five samples of different crystal- linities (w ^c unknown)	120–220	DSC (0.1%)	120–220 K: Equation ^{a q}
Nylon 6	Wunderlich <i>et (</i> (1989) [40]	al. 1.	Semicrystalline ^a ^r	230–600	DSC-2 (0.1%)	230–320 K: Table of smoothed solid data
						500-600 K: smoothed melt data
Nylon II	Wunderlich <i>et a</i> (1989) [40]	al. 1.	Semicrystalline ^{a r}	230–550	DSC-2 (0.1%)	Table of smoothed data below 320 K
						470–550 K: smoothed melt data
Nylon 12	Wunderlich et a (1989)	<i>al</i> . 1.	Semicrystalline ^{a r}	230–540	DSC-2 (0.1%)	Table of smoothed data below 320 K
						460-540 K: smoothed melt data

Table 2. Heat capacity measurements of various linear macromolecules - Continued

Polymer	Investigator Sam	plenum	ber, characterization	Temperature range (K)	Experimental technique (claimed uncertainty)	Source of data
Nylon 6.6	Wunderlich <i>et al</i> . [40] (1989)	1.	Semicrystalline ^{ar}	230–620	DSC-2 (0.1%)	Table of smoothed data below 320 K
						540-620 K smoothed melt data
Nylon 6.9	Wunderlich <i>et al</i> . (1989) [40]	1.	Semicrystalline ^{a r}	230–590	DSC-2 (0.1%)	Table of data smoothed data below 320 K
						500–590 K: smoothed melt data
Nylon 6.10	Wunderlich <i>et al</i> . (1989) [40]	1.	Semicrystalline ^{a r}	230590	DSC-2 (0.1%)	Table of smoothed data below 320 K:
						510-590 K smoothed melt data
Nylon 6.12	Wunderlich <i>et al</i> . (1989) [40]	1.	Semicrystalline ^{a r}	230–590	DSC-2 (0.1%)	Table of smoothed data below 320 K:
						500-590 K smoothed melt data
poly (Diethylsiloxane)	Beatty and Karasz (1975) [9]	1.	$\frac{\text{Semicrystalline}}{M_{\text{w}}} = 112,000$	50–300	Adiabatic (not reported)	Equation ^{a s}
	Turdakin <i>et al</i> . (1976) [9]	2.	$\frac{\text{Semicrystalline}}{\overline{M}_{\text{w}}} = 1,530$	55–330	Adiabatic (0.3%)	Table
	Lebedev <i>et al</i> . (1984) [41]	3.	Semicrystalline $T_d = 203 \text{ K}$ $\Delta H_d = 2.63 \text{ kJ/mol}^{\text{a u}}$ $T_i = 295 \text{ K}$ $\Delta H_i = 2.62 \text{ kJ/mol}^{\text{a u}}$ $\Delta C_p = 7.374 \text{ J/K mol}$ $w^c = 78.6\%$	10–330	Adiabatic (1%)	Graph and Table
	Wesson et al. (1988) [42]	4.	Semicrystalline ^{a t} Cooled at 1 K min ⁻¹ $w^c = 0.585$ $\Delta C_p = 13.67 \text{ J/(K mol)}$ $\Delta H_d = 0.989 \text{ (kJ/mol)}^a$ $\Delta H_i = 0.977 \text{ (kJ/mol)}^a$	u	DSC (1%)	Table
	Wesson <i>et al</i> . (1988) [42]	5.	Semicrystalline ^{a t} Cooled at 10 K/min $w^c = 0.469$ $\Delta C_p = 14.13 \text{ J/(K mol)}$ $\Delta H_d = 0.793 \text{ (kJ/mol)}^a \text{ u}$ $\Delta H_i = 0.85 \text{ (kJ/mol)}^a \text{ u}$		DSC (1%)	Table
	Wesson et al. (1988) [42]	6.	Annealed at 264.8 K $w^c = 0.759$ $\Delta C_p = 6.92 \text{ J/(K mol)}$ $\Delta H_d = 1.283 \text{ (kJ/mol)}^a$ $\Delta H_i = 1.223 \text{ (kJ/mol)}^a$		DSC (1%)	Table

Table 2. Heat capacity measurements of various linear macromolecules - Continued

Polymer	Investigator	Samplenum	ber, characterization	Temperature range (K)	Experimental technique (claimed uncertainty)	Source of data
	Wesson <i>et al</i> . (1988) [42]	7.	Semicrystalline, Cooled to 201 K, heated to 223 K and Quenched $w^c = 0.472$ $\Delta C_p = 13.91 \text{ J/(K mo)}$ $\Delta H_d = 0.798 \text{ (kJ/mol)}$		DSC (1%)	Table
	Wesson <i>et al</i> . (1988) [42]	8.	Quenched Cooled to 201 K, heated to 223 K Quenched $w^c = 0.431$ $\Delta C_p = 21.12 \text{ J/(K mo}$ $\Delta H_d = 0.729 \text{ (kJ/mol)}$	110–360° v	DSC (1%)	Table
	Wesson <i>et al</i> . (1988) [42]	9.	Quenched $\Delta C_p = 31.67 \text{ J/(K mo}$	110–350° v l)	DSC (1%)	Table
ooly-p-Fluoro- styrene	Wunderlich <i>et</i> (1986) [43]	al. 1.	Amorphous $\overline{M}_n = 363,400$ $\overline{M}_w/\overline{M}_n = 2.9$	130–350	DSC-2 (0.1%)	Equations *
ooly-p-Chloro- styrene	Wunderlich <i>et</i> (1986) [43]	al. 2.	$\frac{\underline{A}\underline{m}\underline{n} = 140,400}{\underline{M}_{\text{w}}/\underline{M}_{\text{n}} = 1.9}$	300-550	DSC-2 (0.1%)	Equations ^{a x}
ooly-p-Bromostyrene	Wunderlich <i>et</i> (1986) [43]	al. 1.	Amorphous $ \underline{\overline{M}}_{n} = 80,100 $ $ \underline{\overline{M}}_{w}/\overline{\overline{M}}_{n} = 6.3 $	300–550	DSC-2 (0.1%)	Equations ^{a x}
ooly-p-Iodo- styrene	Wunderlich <i>et e</i> (1986) [43]	al. 1.	Amorphous $\overline{\underline{M}}_{n} = 84,800$ $\overline{\underline{M}}_{w}/\overline{\underline{M}}_{n} = 4.3$	300–550	DSC-2 (0.1%)	Equations ^{a x}
oly-p-Methyl- styrene	Wunderlich et (1986) [43]	al. 1.	Amorphous $ \underline{M}_{n} = 43,900 $ $ \underline{M}_{w}/\underline{M}_{n} = 1.9 $	300–550	DSC-2 (0.1%)	Equations ^{a x}
oly (Dimethylitaconate)	Cowie <i>et al</i> . (1983) [44]	1.	Quenched from 470 K at $\frac{320 \text{ K min}^{-1}}{M_{\text{n}}} = 7.1 \times 10^4$	110-450	DSC-2 (not reported)	Graph
oly(di-n - Propyl- itaconate)	Cowie <i>et al</i> . (1983) [44]	1.	Quenched from 470 K at $\frac{320 \text{ K min}^{-1}}{M_{\text{n}}} = 8.3 \times 10^4$	110-410	DSC-2 (not reported)	Graph
ooly (di-n - ¡Heptyl- itaconate)	Cowie <i>et al</i> . (1983) [44,45]	1.	Quenched from 470 K at 320 K min^{-1} $\overline{M}_n = 9.7 \times 10^4$	110–170	DSC-2 (not reported)	Graph
oly (di-n - Octyl- ìtaconate)	Cowie et al. (1983) [44,45]	1.	Quenched from 470 K at 320 K min^{-1} $M_n = 11.7 \times 10^4$	110-170	DSC-2 (not reported)	Graph
oly (di-n - Nonyl- itaconate)	Cowie et al. (1983) [44,45]	1.	Quenched from 470 K at $\frac{320 \text{ K min}^{-1}}{M_n} = 12.2 \times 10^4$	110-180	DSC-2 (not reported)	Graph
oly (Dido- decyl- itaconate	Cowie <i>et al</i> . (1983) [45]	1.	Quenched from 470 K at 320 K min ⁻¹	110–200	DSC-2 (not reported)	Graph
oly (Dicyclo- octyl- itaconate	Cowie et al. (1982) [46]	1.	Quenched from 470 K at 320 K min^{-1} $\overline{M}_n = 1.8 \times 10^5$	110 -440	DSC-2	Graph

Table 2. Heat capacity measurements of various linear macromolecules - Continued

Polymer	Investigator Sa	mplenum	ber, characterization	Temperature range (K)	Experimental technique (claimed uncertainty)	Source of data
poly (Dicyclo- decyl- itaconate	Cowie et al. (1983) [47]	1.	Quenched from 470 K at 320 K min ⁻¹	110-280	DSC-2	Graph
poly (Dicyclo- dodecyl- itaconate)	Cowie <i>et al</i> . (1983) [47]	1.	Quenched from 470 K at 320 K min ⁻¹	110–280	DSC-2	Equation ^{b a}
poly (p - Metha- cryloloxy- benzoic acid)	Lebedev et al. (1984) [48]	1.	$\frac{\text{Amorphous}}{M_{\text{v}}} = 3 \times 10^{5}$	10–310	Adiabatic (~0.5%)	10–80 K: Table 90–310 K: Graph
<i>poly</i> (<i>p</i> -Pheny- lene)	Rabinovich et al. (1981) [49]	. 1.	Semicrystalline	80–300	Adiabatic	Table
poly (Thio-1,4- phenylene)	Wunderlich <i>et al</i> . (1987) [50]	1.	Semicrystalline ^{b b} $ \underline{w}^{c} = 0.43 - 0.59 $ $ \underline{M}_{n} = 0.91 \times 10^{4} $	220-600	DSC-2 (0.1%)	220-340 K: Equation ^{b c}
•		2.	Quenched ^b °	220-600	DSC-2 (0.1%) Equation ^{b e}	540-600 K:
<i>poly</i> (p - Xylylene)	Wunderlich et al. (1986) [51]	1.	Semicrystalline ^{b f} $w^c = \sim 90\%$	220–410	DSC-2 (0.1%)	Table ^{bg}
poly (Oxy-2,6- dimethyl-5- bromo-1,4- phenylene)	Wunderlich <i>et al</i> . (1982) [29]	2.	Amorphous $\overline{\underline{M}}_{n} = 16,800$ $\overline{M}_{w} = 57,400$	310–530	DSC-2 (0.1%)	Table ^{b h}
poly[Oxy-2,6- bis (1-methyl- ethyl)-1,4- phenylene]	Jauhiainen (1983) [52]	1.	State not reported $\overline{M}_n = 5630 \pm 60 \text{g/r}$	270–450 nol	DSC-2 (1%)	(i)270–410 K: Equation ^{b i} (ii) 430–450 K: Graph
poly (Ethylene terephtha- late)	Collocot (1987) [53]	1.	Amorphous, Quenched	0.4–15	Adiabatic heat pulse (1% above 1.5 K, 2% < 1.5 K)	Graph
		2.	Semicrystalline (i) Annealed at 100 for 100 min $w^c = 14.5\%$	°C "	,	"
			(ii) Annealed at 140 for 1560 min	°C "	n	,,
			w ^c = 33.6% (iii) Annealed at 210 20 h)°C ″	"	,,
			$w^{c} = 53.6\%$ (iv) Annealed at 160 for 26 h, 240 °C for $w^{c} = 63.6\%$		W	,
poly(Butylene- terephthalate)	Cheng et al. (1988) [54]	1.	Semicrystalline ^{b j} $\overline{M}_n = 46,000$ Isothermally crystallized at 480 K for 48 h, then cooled to 220 K at 0.31 K/m	i	DSC-2 (0.1%)	220–280 K: Equation ^{b k} 340–460 K: Graph 490–540 K: Equation ^{b 1}

Table 2. Heat capacity measurements of various linear macromolecules - Continued

Polymer	Investigator	Samplenum	ber, characterization	Temperature range (K)	Experimental technique (claimed uncertainty)	Source of data
		2.	Quenched ^{b j} in a glass bead cooling bath	220–540 g	DSC-2 (0.1%)	Graph
poly (Butylene- terephthalate)	Aleman (1985)	[55] 3.	Semicrystalline ^{b m} $\overline{M}_n = 19,600$ Crystallized at 473 K to 20 min followed by co to 143 K with liquid N	oling	DSC (0.1%)	Graph
	Menczel and Wunderlich (1986) [56]	4.	Semicrystalline ^{b n} $w^c = 0.47$	270–570	DSC-7	Graph
		5.	Semicrystalline ^{b n} $w^c = 0.31$	270–570	DSC-7	Graph
		6.	Semicrystalline ^{b n} $w^{\circ} = 0.37$	270–570	DSC-7	Graph
poly (4-Hydroxy- benzoic acid)	Wunderlich <i>et</i> (1985) [58]	al. 1.	Semicrystalline ^{b o} (i) Samples kept at 500 K (below T _d) for 2 min, cooled at 20 K/min.	170–215	Mettler DSC (not reported)	170–280 K: Equation ^{6 q}
			(ii) Samples kept at 630 K (above T_d) for 2 min, cooled at 20 K/ $T_d^{bp} = 616.5 \pm 4$ K $\Delta H_d = 3.8 \pm 1.1$ kJ/m		DSC-2	290–480 K: Table
poly (2,6-Hydrox naphthoic acid		al. 1.	Semicrystalline ^b (i) Sample kept at 500 K (below T_d) for 2 min, cooled at 20 K/	170–215 min.	Mettler DSC (not reported)	170–280 K : Equation ^{b (}
			(ii) Sample kept at 630 K (above T_d) for 2 min, cooled at 20 K/min. $T_d^{\text{bp}} = 614.5 \text{ K}$ $\Delta H_d = 0.4 \text{ kJ/mol} \pm 6$	215–480	DSC-2 (not reported)	290–480 K: Table ^{b u}
		2.	Melt	500-650		500–650 K:
poly (Ethylene- 2,6-naphtha- lene dicarbo- xylate)	Cheng and Wunderlich (1988) [59]	1.	Semicrystalline, a. Isothermally crystallized from melt. T_c and t_c varied	220–360 (below T_g) 330–450 (Glass	DSC~2 (0.1%)	Equation ^{b v} 220–300 K: Equation ^{b w} 400–450 K:
жушсу			between 450-520 K and 0.5-15 h respectively. $w^c = 0.3 - 0.51$	transition region)		Graph
			b. Isothermally crystallized from glass T_c and T_c varied betwe 450–520 K and 0.5–15 respectively. $w^c = 0.28 \ 0.50$			
		2.	Quenched with liquid N_2	220–600	DSC-2 (0.1%)	220–300 K: Equation ^{b w} 400–460 K: Graph

Table 2. Heat capacity measurements of various linear macromolecules - Continued

Polymer	Investigator	Samplenum	ber, characterization		erimental technique claimed uncertainty)	Source of data	
		3.	Melt	540–600	DSC-2 (0.1%)	540-600 K: Equation ^{b x}	
poly (Oxy-1,4- phenylene- oxy-1,4- phenylene- carbonyl- 1,4-phenylene)	Kemmish <i>et al</i> . (1985) [61]	1.	Amorphous ^{b y} (Quenched with ice/water) $\overline{M}_{\rm w} = 3.9 \times 10^4$ $\overline{M}_{\rm n} = 1.7 \times 10^4$	340–570 (not reported)	DSC-2 Equation ^{b z}	(i) 340–420 (ii) 440–570	K: K:
1,4-риспуюне)		2.	Melt	610–680	DSC-2 (not reported)	Table	
	Cheng and Wunderlich	3.	Semicrystalline ^{c a} Isothermally	130–570	DSC-2 (0.1%)	(i) 130–200 Equation ^{c h}	K:
	(1986) [62]		crystallized from melt and glass at			(ii) 200–240 Equation ^{c c}	K:
			various T _c (463.2–593.2 K)			(iii)340-410 Graph	K:
			followed by cooling at -0.31 K/min to b $w^c = 0.30 - 0.51$	pelow T_{g}		(iv)430-570 Graph	K:
poly (Oxy-1,4- phenylene-	Cheng and Wunderlich	4.	Amorphous (Quenched with	340–570	DSC-2 (0.1%)	(i) 130–200 Equation ^{c b}	K:
oxy-1,4-	(1986) [62]		liquid N ₂)		(not reported)	(ii) 200–240	K:
phenylene- carbonyl- 1,4-phenylene)						Equation ^{c c} (iii)340–410 Graph	K:
	Cheng and Wunderlich (1986) [63]	5.	Melt	430–680	DSC-2 (0.1%)	(i) 430–680 Equation ^{c d}	K:
poly[Oxy-1,4- phenylene- sulfonyl-1,4- phenylene-oxy- 1,4-phenylene- (1-methylidene)- 1,4-phenylene]	Richardson <i>et a</i> (1979) [21]	l. 1.	Amorphous	380–540	DSC (1%)	380–440 K: Equation ^{c c} 480–540 K: Equation ^{c f}	
	Novoselova et al. (1985) [64]	2.	Amorphous	10–480	10–480 K: Adiabatic 330–480 K: DSC (5%)	Table of smoothed data	

^aAtatic polystyrene - NBS samples - 1478

^bCommercial sample from Polysciences, Inc.

^cAverage of 6 runs. $C_p = 65.3320 + 0.06326T [\pm 2\%, J/(K mol)].$

^dAuthor's curve fitted equation: $C_p = 5.933 + 0.263 T [\pm 5\%, J/(K mol)]$. Equation is derived by fitting melt data obtained from subtracting contribution of solid heat capacity (calculated using ATHAS computation scheme) from semicrystalline data.

^{*}Author's curve fitted equation: $C_p = 77.3227 + 0.058517 T [\pm 2\%, J/(K mol)].$

fGift sample from IBM.

^gAuthor's curve fitted equation: $C_p = 43.296 + 0.170 T [\pm 2\%, J/(K mol)]$. Equation is derived by fitting melt data obtained from subtracting contribution of solid heat capacity (calculated using ATHAS computation scheme) from semicrystalline data.

^hAuthor's curve fitted equation: $C_p = 88.0236 + 0.06223 T [\pm 1.5\%, J/(K mol)].$

Commercial samples from Celanese (Celcon M90) and duPont (Delrin 500 ×). Later had a M.W 40,000.

ⁱAuthor's curve fitted equation: Average of 36 runs. $C_p = 46.51 + 0.0372 T [\pm 1.2\%, J/(K mol)]$.

^kCommercial sample from Polysciences, Inc.

^{&#}x27;Author's curve fitted equation from an average of 8 runs. $C_p = 67.85 + 0.0668 T [\pm 0.2\%, J/(K mol)]$.

^mGift sample and commercial sample (uncharacterized) from General Electric Company.

ⁿAuthor's curve fitted equation: Average of 10 runs. $C_p = 0.559 T + 249 [\pm 0.1\%, J/(K mol)]$.

[°]Commercial PPO sample from General Electric Company.

PAuthor's curve fitted equation: $C_p = \exp[-0.13222 (\ln T)^3 + 2.2301 (\ln T)^2 - 11.1681 (\ln T) + 23.614][\pm 1.6\%, J/(K mol)]$. Average of 30 runs.

```
<sup>q</sup>Author's curve fitted equation: C_n = 0.2282 T + 141.6 [\pm 1.5\%, J/(K mol)].
'Sample obtained from Polysciences Inc.
<sup>s</sup>Author's curve fitted equation: C_p = 52.63 + 0.178 [\pm 0.2\%, J/(K mol)].
'Sample obtained from Chemische Werke Hüls.
"Sample obtained from the General Tire and Rubber Co., Research Division, Akron, OH, USA.
<sup>v</sup>Sample obtained from Firestone, Central Research Laboratories, Akron, OH, USA.
"Teflon is the trade name for polytetrafluoroethylene manufactured by E. I. duPont de Nemours Co., Inc.
{}^{x}C_{p} = 10^{-5}T^{3} \text{ cal g}^{-1} \text{ K}^{-1}
{}^{y}C_{p} = 98 T^{3} \text{ erg K}^{-1} \text{ cm}^{-3}
{}^{z}C_{p} = 0.54268 + 1.345 \times 10^{-3} T + 208/(603.4 - T)^{2} \text{ J g}^{-1} \text{ K}^{-1}
{}^{a} {}^{c}C_{p} = 0.54921 + 1.45154 \times 10^{-3}T + 410.8/(608 - T)^{2} \text{ J g}^{-1} \text{ K}^{-1}
^{a\,b}C_p = 0.61488 + 1.949 \times 10^{-3} T J g^{-1} K^{-1}
^{a} ^{c}C_{n} = \exp[-0.174744 (\ln T)^{3} + 0.128498 (\ln T)^{2} + 3.13938 (\ln T) + 5.85181] erg g^{-1} K^{-1}
<sup>a</sup> <sup>d</sup>Gift samples from duPont Research Laboratories (Polymer Products and Central Research Departments)
<sup>a c</sup>PTFE contains 12 ± 2 wt% hexafluoropropylene or 4 CF<sub>3</sub> branches per 100 carbon backbone atoms.
a fX-ray, IR and NMR indicate complete crystallinity. A series of four samples analyzed to determine heat of fusion and heat capacity of the melt.
<sup>a g</sup>PTFE has a perfluoroether group of 2-5 carbon atoms per branch (1.4 branches per 100 backbone atoms).
<sup>a</sup> hTabulated smoothed data presented by the author from 10 to 200 K or 150 K was interpolated using the spline function technique to determine
   heat capacity. For PGL from 210 K and for all other polylactones from 160 K up to just below the glass transition temperature, the Cp data were
  curve fitted into the linear equations listed in Table 11.
a i Author has tabulated the smoothed data. In order to retrieve the data at the standard data bank intervals, interpolation was done using spline
  function technique and curve fitting the presented data into linear equations at temperatures close to the glass transition (see text).
a i Author has tabulated the smoothed data. In order to retrieve the data at the standard data bank intervals, the data was refitted into equations
   described in the text.
**Commercial samples obtained from Polysciences Inc., Sample no. 1, 2 and 3 are poly(trimethylene adipate), poly(hexamethylene sebacate) and
  poly (trimethylene succinate) respectively.
a Author's curve fitted equation: C_p = 0.397038 \ T + 225.3193 \ J/(K \ mol) a Author's curve fitted equation: C_p = 0.755081 \ T + 309.156 \ J/(K \ mol)
<sup>a n</sup>Author's curve fitted equation: C_p = 0.253009 T + 212.5816 \text{ J/(K mol)}

<sup>a o</sup>Author's curve fitted equation: C_p = 0.159 + 1.26 \times 10^{-8} T^3 \text{ cal/g.C}
<sup>a p</sup>Author's curve fitted equation: C_p = 0.444 + 5.81 \times 10^{-4} T \text{ cal/g.C}
<sup>a q</sup>Author's curve fitted equation: C_p = 52.11 + 0.8269 T \text{ J/(K mol)}
Commercial samples from Scientific Polymer Products, Inc.,
<sup>a s</sup>Authors tabulated data were curve fitted into the following equations:
  60–140 K: 1.0987 \times 10<sup>-4</sup> T^2 - 0.016778 T + 1.297 J g^{-1} K<sup>-1</sup> (RMS dev. 2.3%)
 150–190 K: 1.1864 × 10^{-4} T^2 – 0.033338 T + 3.505 J g<sup>-1</sup> K<sup>-1</sup> (RMS dev. 2.4%) 200–230 K: 3.2433 × 10^{-4} T^2 – 0.1382 T + 16.17 J g<sup>-1</sup> K<sup>-1</sup> (RMS dev. 1.6%)
 270-300 K: 2.8756 \times 10<sup>-3</sup> T + 0.7695 J g<sup>-1</sup> K<sup>-1</sup> (RMS dev. 2.1%)
at Heat capacity data of semicrystalline samples provided by Boyer (Midland Macromolecular Institute) show a rigid crystal to condis crystal transition
  at T_d 206 K and condis crystal to liquid crystal transition at T_i = 272 K and a final isotropization to melt at \sim 300 K. w^c is the crystallinity of
  rigid crystal.
^{a} ^{u}\Delta H_{d} = and \Delta H_{i} = are the heats of transitions of the two transitions.
a Author has curve fitted the data from 290 to 360 K into the equation: C_p = 93.75 + 0.2365 T which represents melt C_p data.
<sup>a</sup> "For the structure and abbreviations used see the Table given in Appendix.
a *Author's curve fitted equations are listed in Table 25.
<sup>a y</sup>The structure and abbreviations used are given in the Table in Appendix.
<sup>a z</sup>The structure and abbreviations are given in the table in Appendix.
<sup>b</sup> <sup>a</sup>Author's curve fitted equation: C_p = 103.3 + 0.8635 T + 0.003429 T^2 [\pm 2.6\%, J/(K mol)].
bbSample obtained from Polysciences Inc. Semicrystalline samples were prepared by the author by isothermal and non-isothermal crystallization from
  melt and glass.
<sup>b c</sup>Average of 15 runs on 10 samples. C_p = \exp \left[0.109038 \left(\ln T\right)^3 - 1.7846 \left(\ln T\right)^2 + 10.7735 \left(\ln T\right) - 18.9156\right] \left[\pm 1.5\%, J/(K \text{ mol})\right]
^{\text{b}} \, ^{\text{d}}C_p = 0.12574 \, T + 119.73
<sup>b e</sup>Samples had a crystallinity of about 5%.
b fAnnealed sample.
bg Table of smoothed data.
<sup>b</sup> Table of smoothed data.
<sup>b</sup> Author's curve fitted equation: C_p = 0.258337 + 0.003835 T in J/(K g).
<sup>b</sup> Commercial sample from Eastman Kodak (Valox, Lot 6 P6D).
<sup>b k</sup>Author's curve fitted equation: C_p = 52.16 + 0.5203 T + 0.000713 T^2 J/(K mol) from an average of 5 measurements.

<sup>b l</sup>Author's curve fitted equation: C_p = 219.6479 + 0.452209 T J/(K mol) from an average of 7 measurements.
<sup>b m</sup>Commercial sample from Ciba-Geigy AG (Germany).
<sup>b</sup> <sup>n</sup>Commercial sample from GAF Corporation: Gafite 1600.
<sup>b</sup> <sup>o</sup>Commercial samples supplied by Celanese Research Company.
<sup>b p</sup>Disordering transition peak temperature.
<sup>b q</sup>Author's curve fitted equation: C_p = 3.1526 + 0.3977 T [\pm 0.03\%, J/(K mol)].
<sup>b</sup> Commercial samples supplied by Celanese Research Company.
b sDisordering transition peak temperature.
<sup>b</sup> Author's curve fitted equation: C_p = 3.1514 + 0.5951 T [\pm 0.02\%, J/(K mol)].
<sup>b u</sup>Author's tabulated smoothed data used as such.
<sup>b</sup> Author's curve fitted equation: C_p = 0.340361 T + 138.336 [\pm 0.1 \%, J/(K mol)].
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TABLE 3: Recommended heat capacity data for molten fluorinated polymers

TABLE 4: Recommended heat capacity data for molten polyoxymethylene and polyoxyethylene

Temp (K)	Heat capacity [J/(K mol)] PVF	Heat capacity [J/(K mol)] PVF2	Heat capacity [J/(K mol)] P3FE	Temp (K)	Heat capacity [J/(K mol)] POM	Heat capacity [J/(K mol)] POE
220.00	NA	73.304	NA .	190.00 (T _g) POM	53.578	NA
230.00	NA	74.529	NA	200.00	53.950	NA
240.00	NA	75.753	NA	$206.00 (T_g) POE$	54.173	81.611
250.00	NA	76.978	NA	210.00	54.322	81.878
260.00	NA	78.202	NA	220.00	54.694	82.546
270.00	NA	79.427	NA	230.00	55.066	83.214
273.15	NA	79.812	NA	240.00	55.438	83.882
280.00	NA	80.651	90.494	250.00	55.810	84.550
290.00	NA	81.876	91.916	260.00	56.182	85.218
298.15	NA	82.873	93.075	270.00	56.554	85.886
300.00	NA	83.100	93.338	273.15	56.671	86.096
310.00	NA	84.325	94.760	280.00	56,926	86.554
320.00	74.916	85.549	96.183	290.00	57.298	87.222
330.00	75.942	86.774	97.605	298.15	57.601	87.766
340.00	76.969	87.998	99.027	300.00	57.670	87.890
350.00	77.996	89.223	100.449	310.00	58.042	88.558
360.00	79.023	90.447	101.872	320.00	58.414	89.226
370.00	80.049	91.672	103.294	330.00	58.786	89.894
380.00	81.076	92.896	104.716	340.00	59.158	90.562
390.00	82.103	94.120	106.138	350.00	59.530	91.230
400.00	83.130	95.345	107.561	360.00	59.902	91.898
410.00	84.156	96.570	108.983	370.00	60.274	92.566
420.00	85.183	97.794	110.405	380.00	60.646	93.234
430.00	86.210	99.019	111.827	390.00	61.018	93.902
440.00	87.236	100.243	113.250	400.00	61.390	94.570
450.00	88.263	101.468	114.672	410.00	61.762	95.238
460.00	89.290	102.692	116.094	420.00	62.134	95.906
470.00	90.317	103.917	117.516	430.00	62.506	96.574
480.00	91.344	105.141	118.939	440.00	62.878	97.242
490.00	92.160	105.634	119.107	450.00	63.250	97.910
500.00	92.942	106.345	119.747	460.00	63.622	98.578
510.00	93.725	107.056	120.386	470.00	63.994	99.246
520.00	94.507	107.767	121.026	480.00	64.366	99.914
530.00	95.290	108.478	121.665	490.00	64.738	100.582
540.00	96.072	109.189	122.305	500.00	65.110	101.250
550.00	96.855	109.900	122.944	510.00	65.482	101.918
560.00	97.637	110.611	123.584	520.00	65.854	102.586
570.00	98.420	111.322	124.223	530.00	66.226	103.254
580.00	99.202	112.033	124.863	540.00	66.598	103.922
590.00	99.985	112.744	125.502			
600.00	100.767	113.455	126.142			

 $^{^{}b}$ WAuthor's curve fitted equation: $C_p = 0.0001616$ $T^2 + 0.7544$ T + 29.256 J/(K mol), average of 18 measurements of semicrystalline PEN and 5 of amorphous PEN with RMS deviation being 0.8%.

^{b x}Author's curve fitted equation: $C_p = 0.40941 T + 268.35 J/(K mol)$ average of 15 measurements.

^{b y}Commercial sample from ICI Co.

 $^{^{}b}$ Author's curve fitted equation: $C_p = (0.1036 \pm 0.017) + (3.34 \pm 0.05) \times 10^{-3} T \text{ in J/(K g)}$

^c ^aCommercial sample from ICI Co.

^{cb}Author's curve fitted equation: $C_p = \exp[0.1695 (\ln T)^3 - 2.193 (\ln T)^2 + 10.08 (\ln T) - 11.63] [\pm 2.5\%, J/(K mol)].$

c°Author's curve fitted equation: $C_p = 0.00005455 \ T^2 + 1.057 \ T + 9.631 \ [\pm 1.2\%, J/(K mol)].$ c°Author's curve fitted equation: $C_p = 0.4955 \ T + 308.15 \ [\pm 0.1\%, J/(K mol)].$ c°Author's curve fitted equation: $C_p = 0.309 + 0.00288 \ T \ J/(K \ g).$ c°Author's curve fitted equation: $C_p = 0.926 + 0.00204 \ T \ J/(K \ g).$

3. Recommended Heat Capacities of Polymers Added to the Data Bank

3.1. Other Carbon Backbone Polymers

3.1.1. Introduction

In the following subsections all newly recommended heat capacities are discussed using the prior established criteria. Overall, only one new polymer is included in this listing together with recommendations on two polymers described earlier. A summary and quick reference to the location of the description of all heat capacities is contained in the appendix.

3.1.2. poly (2-Methyl-1.3-pentadiene)

Only one investigation is reported in the literature for heat capacities of poly (2-methyl-1,3-pentadiene) (PMP).¹⁷ The details of the measurements are given in Table 2. Heat capacity of PMP over the range over which data were recovered from the graph is given in Table Λ 13. The preliminary recommended heat capacities below and above the glass transition temperature at 277.8 K (T_g) are given in Table 5. The change in heat capacity at 277.8 K (T_g) thus obtained agrees within 1.5% with that reported by the authors.¹⁷ Below the glass transition temperature the data recovered from the graph from 220 to 250 K were curve-fit into the equation:

$$C_p = -0.8438 + 0.0088767 T - 0.0000166811 T^2$$

(± 0.3%) J/(K mol) (3)

and from 240 to 270 K the data were fitted into the equation:

$$C_p = 0.1450 + 0.0007507 T (\pm 0.35\%) \text{ J/(K mol)}$$
 (4)

Equation (4) was used to extrapolate the data up to T_g . Melt heat capacities were obtained from T_g up to 320 K by curve fitting the data from 280 to 320 K into the equation:

$$C_p = 0.2575 + 0.0007148 T (\pm 0.4\%) J/(K mol)$$
 (5)

3.1.3. poly (1-Butenylene)

In the ATHAS data bank (1983),⁷ the heat capacities on *cis* and *trans poly* (1-butenylene) also reffered to as 1,4-polybutadiene were based on only one investigation for semicrystalline samples of unknown crystallinity. Since then a detailed investigation was carried out by Wunderlich *et al.*³¹ on both isomers. The heat capacities of both isomers of polybutadiene are given in Table A14 and A15. The details of the measurement are given in Table 2. The glass transitions of *cis poly* (1-butenylene) and *trans poly* (1-butenylene) were established at 171 K and 190 K, respectively. The *trans*-isomer exhibits a condis phase (conformationally disordered and mobile)

above 356 K.³¹ The melting temperature was established at 284.7 K for the *cis*-isomer and 437 K for *trans*-isomer. Below the glass transition the data of semicrystalline samples (1 and 2) were used to represent the solid state. From 30 to 100 K, tabulated data⁷ for the *cis*-isomer were used as such. Equation (6) represents the heat capacity data of the *cis*-isomer from 110 to 171 K (T_g) and was obtained by curve fitting the experimental data reported earlier⁷ from 120 to 130 K.

$$C_p = 6.945004 + 0.2761 T (\pm 0.02\%) \text{ J/(K mol)}$$
 (6)

For the *trans*-isomer, C_p data were reported earlier.⁷ The data from 30 to 120 K were used as such, while the data from 130 to 170 K were curve-fit into the equation:

$$C_p = 7.856993 + 0.2743 T (\pm 0.04\%) J/(K mol)$$
 (7)

Equation (7) was used from 130 to 190 K (T_g).

Above the glass transition temperature the data on samples 1 and 2 were not considered, as the new data were based on a more extensive investigation. Heat capacities of melt of both, the *cis* and *trans*-isomers could be represented by the smoothed equation given by the authors³¹:

$$C_p = 52.63 + 0.178 T (\pm 0.02\%) J/(K mol)$$
 (8)

Since the condis phase of the *trans*-isomer was shown by the authors to have heat capacities close to that of the extrapolated melt, Eq. (8) was used to extend the heat capacities to 190 K for *trans poly* (1-butenylene) and up to 171 K for *cis*-butadiene. The recommended heat capacities of *cis* poly (1-butenylene) and *trans poly* (1-butenylene) are given in Tables 6 and 7 and are plotted in Fig. 1.

TABLE 5: Recommended heat capacity data for poly(2-methyl-1,3-pentadiene)

Temp	Heat capacity
(K)	[J/(K mol)]
230.00	105.565
240.00	109.034
250.00	111.387
260.00	113.878
270.00	116.391
273.15	117.182
$277.80 (T_g)$	118.350
$277.80 (T_{\rm g})$	152.649
280.00	153.176
290.00	155.568
298.15	157.518
300.00	157.961
310.00	160.353
320.00	162.746

TABLE 6a. Recommended heat capacity data for cis poly (1-butenylene) below the glass transition

TABLE 7a. Recommended heat capacity data for trans poly(1-butenylene) below the glass transition

Temp (K)	Heat capacity [J/(K mol)]	Temp (K)	Heat capacity [J/(K mol)]
30.00	11.790	30.00	9.704
40.00	16.150	40.00	14.390
50.00	19.980	50.00	18.740
60.00	23.180	60.00	22.560
70.00	26.160	70.00	25.890
80.00	29.070	80.00	29.000
90.00	31.870	90.00	32.020
100.00	34.630	100.00	34.980
110.00	37.316	110.00	37.860
120.00	40.077	120.00	40.700
130.00	42.838	130.00	43.516
140.00	45.599	140.00	46.259
150.00	48.360	150.00	49.002
160.00	51.121	160.00	51.745
170.00	53.882	170.00	54.488
$171.00 \ (T_{\rm g})$	54.158	180.00	57.231
		$190.00 \ (T_{\rm g})$	59.974

Table 6b. Recommended heat capacity data for molten cis poly(1-butenylene)

Table 7b. Recommended heat capacity data for molten trans poly(1-butenylene)

Temp (K)	Heat capacity [J/K mol)]	Temp (K)	Heat capacity [J/(K mol)]
171.00 (T _g)	83.068	190.00 (T _g)	86.450
180.00	84.670	200.00	88.230
190.00	86.450	210.00	90.010
200.00	88.230	220.00	91.790
210.00	90.010	230.00	93.570
220.00	91.790	240.00	95.350
230.00	93.570	250.00	97.130
240.00	95.350	260.00	98.910
250.00	97.130	270.00	100.690
260.00	98.910	273.15	101.251
270.00	100.690	280.00	102.470
273.15	101.251	290.00	104.250
280.00	102.470	298.15	105.701
290.00	104.250	300.00	106.030
298.15	105.701	310.00	107.810
300.00	106.030	320.00	109.590
310.00	107.810	330.00	111.370
320.00	109.590	340.00	113.150
330.00	111.370	350.00	114.930
340.00	113.150	360.00	116.710
350.00	114.930	370.00	118.490
360.00	116.710	380.00	120.270
370.00	118.490	390.00	122.050
380.00	120.270	400.00	123.830
390.00	122.050	410.00	125.610
400.00	123.830	420.00	127.390
410.00	125.610	430.00	129.170
420.00	127.390	440.00	130.950
430.00	129.170	450.00	132.730
440.00	130.950		
450.00	132.730		



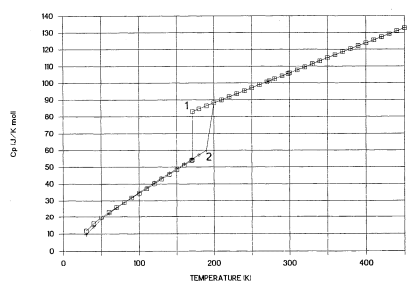


Fig. 1. Recommended heat capacities for: 1. cis poly (1-butenylene); 2. trans poly (1-butenylene).

3.1.4. Polytetrafluoroethylene

Eleven investigations have been reported in the literature for the heat capacity of PTFE. Eight investigations which meet our standards of acceptability are listed in Table 2. The investigations not included are given in Table 8. Heat capacity data on the first six investigations had already been deposited (Table A15, Ref., 7) with the Physics Auxillary Publication Service of the American Institute of Physics. The new data are given in Table A16. In the ATHAS data bank (1980), heat capacity of PTFE could not be recommended as at that time no definite information on the glass transition temperature and room temperature transitions was available. A detailed investigation on heat capacity of PTFE was carried out in our laboratory³² on samples of widely ranging crystallinities $(w^{\circ} = 0.17 - 0.95)$. This investigation permits us to recommend experimental heat capacities for completely crystalline and completely amorphous PTFE [obtained by the two phase model, assuming $C_p = w^c C_p^c + (1-w^c) C_p^a$].

Recommended heat capacities for crystalline PTFE were obtained as follows: From 0.3 to 4 K data on sample 9 was used as the uncertainty claimed by the authors is only 2%. Heat capacities of sample 4 were 14% higher at 1.4 K and 23% at 4 K than sample 9 and the data on sample 5 were higher by 24% and 34%, respectively. From 5 to 25 K the data on sample 8 (crystallinity 0.9) were smoothed by fitting into the equation:

$$C_p = \exp[-7.839547 + 5.168145 (\ln T) - 0.8187535 (\ln T)^2 + 0.03979258 (\ln T)^3] (\pm 0.0\%) J/(K mol)$$
 (9)

The heat capacities from 30 to 100 K of sample 8 and 1 were averaged in the region of overlap (the standard de-

viation is 0.7%) and together with data on sample 1 up to 170 K were fit into the equation.

$$C_p = 0.12087 + 0.2145618 T - 0.0002071 T^2$$

(± 0.43%) J/(K mol) (10)

Above 170 K the solid heat capacity in the ATHAS data bank (1980) is not considered since PTFE shows a glass transition in the 160–240 K region ($T_{\rm g}=200~{\rm K}$) and two room-temperature transitions at 292 and 303 K.³² Between 180 and 210 K the C_p was obtained using Eq. (10). From 220 to 280 K the heat capacity of sample 14 is taken as the recommended data. Equations (11) and (12) represent the smoothed C_p data from 220 to 240 K and 250 to 280 K, respectively.

$$C_p = 29.52984 - 0.06640669 T + 0.000466798 T^2$$
 $(\pm 0.04\%) \text{ J/(K mol)}$ (11)

$$C_p = 157.3948 - 1.099639 T - 0.002553868T^2$$

(± 0.13%) J/(K mol) (12)

At 292 K and 303 K PTFE undergoes two transitions (rigid crystal-condis crystal), hence the heat capacities above 270 K cannot be recommended at this stage.

The heat capacity determinations on practically completely crystalline sample 14 (average of two to four measurements) showed an internal standard deviation of only 0.3%. Between 390 and 480 K, the data on samples 11 and 12 showed a standard deviation of 0.1% (claimed by the authors). This agreed closely with the data on sample 1 (maximum discrepancy + 0.7%). From 340 to 560 K agreement with data on samples 6 and 7 was also quite good (maximum discrepancy -0.9%).

Heat capacity data for completely amorphous PTFE was obtained by using the crystallinity extrapolated data

as reported by Wunderlich et al.³² of the semicrystalline PTFE above the glass transition region (160–240 K). Between 240 K and 480 K the data on samples 10–14 were extrapolated to get 100% amorphous heat capacities. This was further smoothed by curve fitting into the equation:

$$C_{pa} = 29.51 + 0.06917 T + 0.00001292 T^2$$

(± 0.4%) J/(K mol) (13)

Between 560 and 700 K the C_p data of melt was smoothed by curve fitting into the equation:

$$C_p = 49.45 + 0.0332 T (\pm 0.2\%) J/(K mol)$$
 (14)

Equation (13) was used to obtain C_p data from 240 K to 470 K and from 480 to 700 K the heat capacities were obtained using Eq. (14). Between 180 K and 240 K the heat capacities were obtained by making a linear connection between amorphous and crystalline heat capacity. Between 180 K and perhaps 50 K the amorphous and crystalline heat capacities can be assumed to be the same, as in this region heat capacity is usually almost independent of crystallinity. Recommended experimental data of crystalline and amorphous PTFE are given in Tables 9 and 10, respectively.

TABLE 8. Investigations for polytetrafluoroethylene not included in this study

References	Reason(s) for exclusion
Terziiska et al. (1981) [33]	Sample characterization not reported. Data shows higher error limits.
Vargha-Butler <i>et al</i> . (1982) [18]	Sample characterization not reported for semicrystalline data.
Fritzsch (1983) [13]	Data could not be retrieved accurately from small graphs. Polymer was made by glow discharge polymerization and had uncertain chemical structure.
Boyer et al. (1983) [34]	Heat capacity of commercial grade PTFE has been measured by heat pulse technique. Very large deviations observed from recommended data. Up to 5 K data are within the experimental error of Choy's data and show deviations of nearly 10–15 % from Salinger's data discussed in ref. [7].

TABLE 9. Recommended heat capacity data for crystalline polytetrafluoroethylene

Heat capacity

[J/(K mol)]

Temp

(K)

(K)	[3/(12 11101)]
0.30	0.000065
0.40	0.000125
0.50	0.000223
0.60	0.000370
0.70	0.000582
0.80	0.000871
0.90	0.001252
1.00	0.001232
1.20	0.003094
1.40	0.005041
1.60	0.007687
1.80 2.00	0.011110 0.015380
3.00	0.051
4.00	0.109
5.00	0.228
6.00	0.376
7.00	0.554
8.00	0.759
9.00	0.985
10.00	1.228
15.00	2.565
20.00	3.915
25.00	5.155
30.00	6.371
40.00	8.372
50.00	10.331
60.00	12.249
70.00	14.125
80.00	15.960
90.00	17.754
100.00	19.506
110.00	21.217
120.00	22.886
130.00	24.514
140.00	26.100
150.00	27.645
160.00	29.149
170.00	30.611
180.00	32.032
190.00	33.411
200.00	34.749
210.00	36.046
220.00	37.513
230.00	38.950
240.00	40.480
250.00	42.102
260.00	44.130
270.00	46.669
273.15	47.575
280.00	49.719
200.00	77.117

Table 10. Recommended heat capacity data for amorphous polytetrafluoroethylene

traffuoroethylene	
Temp	Heat capacity
(K)	[J/(K mol)]
100.00	22.060
180.00	32.960
190.00	35.280
$200.00 \ (T_g)$	37.590
210.00	39.910
220.00	42.220
230.00	44.980
240.00	46.855
250.00	47.610
260.00	48.368
270.00	49.128
273.15	49.368
280.00	49.891
290.00	50.656
298.15	51.282
300.00	51.424
310.00	52.194
320.00	52.967
330.00	53.743
340.00	54.521
350.00	55.302
360.00	56.086
370.00	56.872
380.00	57.660
390.00	58.451
400.00	59.245
410.00	60.042
420.00	60.840
430.00	61.642
440.00	62.446
450.00	63.253
460.00	64.062
470.00	64.874
480.00	65.386
490.00	65.718
500.00	66.050
510.00	66.382
520.00	66.714
530.00	67.046
540.00	67.378
550.00	67.710
560.00	68.042
570.00	68.374
580.00	68.706
590.00	69.038
600.00	69.370
610.00	69.702
620.00	70.034
	70.366
630.00 640.00	70.300 70.698
640.00	
700.00	72.690

3.2. Recommended Heat Capacities of Heteroatom Backbone Polymers

3.2.1. Introduction

Recommended heat capacities for polyesters derived by substitution of the COO- group in the backbone chain of polyethylene were available in the ATHAS data bank 1980 for polyglycolide, polycaprolactone and poly (ethylene terephthalate). Since then Lebedev et al. 35 have reported heat capacities on a series of polylactones covering polyglycolide and polycaprolactone, polyprioplactone, polybutyrolactone, polyvalerolactone, polyundecanolactone, polytridecanolactone and polypentadecanolactone. In addition, heat capacities are also available for poly (ethylene oxalate), 36 poly (butylene adipate), 37 poly (trimethylene adipate), 38 poly (ethylene sebacate), 39 poly (hexamethylene sebacate) 38 and poly (trimethylene sebacate). 38 All the investigations were reevaluated and found to meet our standards of acceptability. The various investigations are discussed individually and the data listed as recommended heat capacities.

3.2.2. Polylactones

The reported data of Lebedev et al.35 on polyglycolide were very high and did not fit into any addition scheme with other polymers. The author had suggested (CH2-COO-)x as the repeating unit. Since PGL is synthesized, however, from the bimolecular lactide instead of the unstable 3-atom lactone, the heat capacities given authors must have referred (CH2-COO-CH2-COO-) and not (CH2-COO-). Hence, the reported data were halved at all temperatures and the recommended data given in Tables 12 and 13 fit now the addition scheme of all the other polylactones. The details of the measurements on the various polylactones are given in Table 2.

Data were retrieved at the standard data bank intervals from the tabulated, smoothed data presented by the author. For this, the heat capacities from 10 to 200 K (in case of PGL) and 150 K for all other polylactones was interpolated using the spline-function technique. Above these temperatures up to just below $T_{\rm g}$, the already smoothed data were curve-fitted into linear equations which are listed in Table 11 along with the equations for the melt. The recommended experimental data are given in Tables 12 and 13 and are plotted in Figs. 2 and 3. Heat capacity increases from PGL to PPDL, while the glass transition temperatures (Table 14) decrease up to PVL and then increase up to PPDL, reaching a value close to that of polyethylene.

3.2.3. poly (Ethylene Oxalate)

One investigation has been reported that deals with the heat capacity of *poly* (ethylene oxalate).³⁶ Details of this investigation which meets our data standards of acceptable data (discussed in Ref. 1), are given in Table 2. Heat capacity of an amorphous sample has been measured over the temperature range of 8 to 360 K. The data on these samples are given in Table A19.

To obtain the recommended data, the author's tabulated, smoothed data from 10 to 240 K were interpolated using the Spline function technique. From 250 K up to 300 K, C_n was curve-fitted into a linear equation:

Table 11. Results of curve fitting literature data on heat capacities of polylactones to derive the recommended data

Polymera	State	Equation ^b	Temperature range (K) ^c
PGL	Solid	$C_p = 13.89503 + 0.171812 T$	210 - 318
	Molten	$C_p = 100.8 + 0.0380966 T$	318 – 550
PPL	Solid	$C_p = 15.70001 + 0.2480 T$	160 - 249
	Molten	$C_p = 91.4315 + 0.146407 T$	249 - 400
PBL	Solid	$C_p = 18.93358 + 0.32953567$	160 – 214
	Molten	$C_p = 124.817 + 0.103469 T$	214 - 350
PVL	Solid	$C_p = 24.80002 + 0.3741713 T$	160 – 207
	Molten	$C_p = 131.310 + 0.17399 T$	207 - 350
PCL	Solid	$C_p = 25.98002 + 0.454714 T$	160 – 209
	Molten	$C_p = 119.595 + 0.291496 T$	209 - 350
PUDL	Solid	$C_p = 36.8026 + 0.8141549 T$	160 – 227
	Molten	$C_p = 241.226 + 0.400712 T$	227 – 400
PTDL	Solid	$C_p = 40.08001 + 0.97025 T$	160 - 229
	Molten	$C_p = 323.459 + 0.370269 T$	229 - 370
PPDL	Solid	$C_p = 50.80290 + 1.067743 T$	160 - 251
	Molten	$C_p = 389.612 + 0.387217 T$	251 - 370

^aThe structure and abbreviations of the polymers are given in the appendix.

$$C_p = 37.04763 + 0.3088571 T J/(K mol)$$
 (15)

The RMS deviation was $\pm 0.04\%$. This equation was used to extend the data up to the glass transition temperature (306 K). To obtain melt data, the results above the glass transition were curve-fitted into the equation:

$$C_p = 133.003 + 0.179037 T J/(K mol)$$
 (16)

The RMS deviation was $\pm 0.2\%$. Melt data from 306 (T_g) up to 360 K were obtained from this equation. The recommended heat capacities above and below T_g are given in Table 15.

3.2.4. poly (Butylene Adipate)

One investigation has been reported in the literature, which deals with the heat capacity of poly (butylene adipate).³⁷ The reported heat capacities for both the solid and liquid appear somewhat higher than what is expected if one doubles the C_p data for poly (δ -valerolactone) considering that PBA is actually double the repeat unit of this polymer. However, since these are the only available data, they are incorporated in the data bank as preliminary data. The details of the measurement are given in Table 2. The data on the semicrystalline sample are given in Table A20 and 21. Heat capacity of a

semicrystalline sample has been reported. Below the glass transition temperature (T_g 199 K), heat capacity of amorphous and crystalline polymers can be assumed to be the same. To obtain the preliminary recommended data from 80 to 199 K (T_g), the author's tabulated data were refitted to the equation:

$$C_p = \exp[-9.1043 \times 10^{-3} (\ln T)^3 + 0.216717 (\ln T)^2 - 0.674235 (\ln T) + 4.24806] \text{ J/(K mol)}$$
 (17)

In the molten state (199 to 450 K), heat capacity data were obtained by curve fitting the data into the linear equation:

$$C_p = 296.844 + 0.325835 T \text{ J/(K mol)}$$
 (18)

The preliminary recommended data below and above the glass transition temperature were obtained using Eqs. (17) and (18), respectively, and are given in Tables 16 and 17

3.2.5. poly (Trimethylene Adipate), poly (Hexamethylene Sebacate) and poly (Trimethylene Succinate)

Only one investigation³⁸ is reported in the literature that deals with the heat capacities of the melts of poly(trimethylene adipate), poly(hexamethylene sebacate)

 $^{{}^{\}mathrm{b}}C_{\mathrm{p}}$ is in J/(K mol).

Temperature range over which the equation is valid.

TABLE 12. Recommended heat capacity data for semicrystalline polylactones below the glass transition temperature in [J/(K mol)]

Temp (K)	PGL	PPL	PBL	PVL	PCL	PUDL	PTDL	PPDL
	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
0.00	0.000	1.390	1.470	1.510	2.420	3.380	3.300	3.470
10.00	2.184	3.240	4.316	4.313	6.130	8.349	9.408	9.829
15.00	3.980	5.650	7.870	8.000	10.720	14.811	17.360	18.480
20.00		8.374	11.612	12.103	15.667	22.157	26.169	28.415
25.00	5.960		15.300	16.330	20.660	29.932	35.260	38.890
30.00	8.020	11.200				29.932 45.563	53.210	59.660
40.00	12.120	16.560	22.220	24.570	30.170			79.330
50.00	16.040	21.220	28.410	32.160	37.950	60.180	70.450	
60.00	19.570	25.540	33.990	39.080	45.740	73.610	86.320	97.580
70.00	22.720	29.600	39.050	45.370	53.020	86.010	100.700	114.400
80.00	25.550	33.400	43.650	51.100	59.610	97.450	113.900	129.900
90.00	28.110	36.910	47.870	56.320	65.620	108.000	126.000	144.100
100.00	30.470	40.130	51.760	61.110	71.140	117.800	137.200	157.200
110.00	32.670	43.050	55.400	65.520	76.250	126.900	147.700	169.200
120.00	34.740	45.710	58.850	69.630	81.020	135.400	157.700	180.400
130.00	36.690	48.190	62.140	73.520	85.550	143.500	167.200	190.900
140.00	38.530	50.550	65.340	77.250	89.910	151.300	176.500	200.900
150.00	40.280	52.840	68.480	80.910	94.190	158.900	185.600	210.800
160.00	41.960	55.370	71.580	84.668	98.734	167.080	195.330	221.640
170.00	43.580	57.850	74.900	88.409	103.281	175.220	205.030	232.320
180.00	45.150	60.330	78.220	92.151	107.829	183.360	214.730	243.000
190.00	46.690	62.820	81.540	95.893	112.376	191.500	224.430	253.670
200.00	48.220	65.300	84.860	99.635	116.923	199.630	234.130	264.350
210.00	49.976	67.780	88.170	NA	NA	207.770	243.830	275.030
220.00	51.694	70.260	NA	NA	NA	215.917	253.535	285.710
230.00	53.412	72.740	NA	NA	NA	NA	NA	296.380
240.00	55.130	75.220	NA	NA	NA	NA	NA	307.060
250.00	56.848	NA	NA	NA	NA	NA	NA	317.739
260.00	58.566	NA	NA	NA	NA	NA	NA	NA
270.00	60.284	NA	NA	NA	NA	NA	NA	NA
273.15	60.825	NA	NA	NA	NA	NA	NA	NA
280.00	62.002	NA	NA	NA	NA	NA	NA	NA
290.00	63.721	NA	NA	NA	NA	NA	NA	NA
298.15	65.121	NA	NA	NA	NA	NA	NA	NA
300.00	65.439	NA	NA	NA	NA	NA	NA	NA
310.00	67.157	NA	NA	NA	NA	NA	NA	NA

The abbreviations used for the polymer names have been described in the appendix.

and poly (trimethylene succinate). The details of the measurements are given in Table 2. The heat capacity data is given in Table $\Delta 22$. The only available heat capacities are given in Table 18 as preliminary, recommended data.

3.2.6. poly (Ethylene sebacate)

In the ATHAS data bank $(1980)^8$ heat capacities of semicrystalline poly (ethylene sebacate) were available only above the glass transition temperature ($T_g = 245 \text{ K}$) and a recommendation could, therefore, not been made. Since then, Wunderlich *et al.*³⁹ have measured heat capacities of five differently crystallized samples, extending the range of measurement to 120 K. The details of the measurements are given in Table 2. The new data are given in Table A23. To obtain the recommended heat capacities below T_g (245 K), the equation given by the authors for the data, curve-fitted between 120 and 220 K, was used. Heat capacities of the melt were obtained from

the equation of the curve-fitted melt data between 350–410 K. The recommended heat capacities below and above $T_{\rm g}$ are given in Table 19.

3.2.7. Aliphatic Polyamides

In the ATHAS data bank (1980)⁸ heat capacities of nylon 6, nylon 6.6 and nylon 6.12 were listed. Since then Wunderlich et al. 40 have reported heat capacities of nylon 6, nylon 6.6, nylon 11, nylon 12, nylon 6.9, nylon 6.10, and nylon 6.12. Heat capacities of nylon 6 and nylon 6.6 already recommended in the 1980 data bank remain unaltered, as the new measurements showed a deviation of only about 2% from the previous data which is considered within the experimental error. The details of the new measurements on semicrystalline samples are given in Table 2. All new heat capacities are based on only one investigation⁴⁰ and are given in Table A24 and A25. Table 20 lists the recommended heat capacities for the solid

TABLE 13. Recommended heat capacity data for molten polylactones in [J/(K mol)]

Temp (K)	PGL	PPL	PBL	PVL	PCL	PUDL	PTDL	PPDL
210.00	NA	NA	NA	167.848	180.809	NA	NA NA	NA
220.00	NA	NA	147.580	169.588	183.724	NA	NA	NA
230.00	NA	NA	148.615	171.328	186.639	333.390	408.620	NA
240.00	NA	NA	149.650	173.068	189.554	337.397	412.323	NA
250.00	NA	128.033	150.684	174.808	192.469	341.404	416.026	NA
260.00	NA	129.497	151.719	176.547	195.384	345.411	419.728	490.288
270.00	NA	130.961	152.754	178.287	198.299	349.418	423.431	494.161
273.15	NA	131.423	153.080	178.835	199.217	350.680	424.597	495.380
280.00	NA	132.425	153.788	180.027	201.214	353.425	427.134	498.033
290.00	NA	133.890	154.823	181.767	204.129	357.432	430.836	501.905
298.15	NA	135.083	155.666	183.185	206.505	360.698	433.854	505.061
300.00	NA	135.354	155.858	183.507	207.044	361.440	434.539	505.777
310.00	NA	136.818	156.892	185.247	209.959	365.447	438.242	509.649
320.00	112.991	138.282	157.927	186.987	212.874	369.454	441.944	513.521
330.00	113.372	139.746	158.962	188.727	215.789	373.461	445.647	517.394
340.00	113.753	141.210	159.996	190.467	218.704	377.468	449.350	521.266
350.00	114.134	142.674	161.031	192.207	221.619	381.475	453.052	525.138
360.00	114.515	144.138	NA	NA	NA	385.482	456.755	529.010
370.00	114.896	145.602	NA	NA	NA	389.489	460.458	532.882
380.00	115.277	147.066	NA	NA	NA	393.497	NA	NA
390.00	115.658	148.530	NA	NA	NA	397.504	NA	NA
400.00	116.039	149.994	NA	NA	NA	401.511	NA	NA
410.00	116.420	NA						
420.00	116.801	NA						
430.00	117.182	NA						
440.00	117.563	NA						
450.00	117.943	NA						
460.00	118.324	NA						
470.00	118.705	NA						
480.00	119.086	NA						
490.00	119.467	NA						
500.00	119.848	NA						
510.00	120.229	NA						
520.00	120.610	NA						
530.00	120.991	NA						
540.00	121.372	NA						
550.00	121.753	NA						

The abbreviations used for the polymer names have been described in the appendix.

and for the melt. Solid heat capacity data are the smoothed experimental data. Recommended melt data were obtained from the equation:

$$C_p = N_{\rm C}(7.4506 + 0.0745 T) + N_{\rm N}(86.8483 - 0.0226 T) (\pm 1.22\%) \text{ J/(K mol)}$$
 (19)

where $N_{\rm N}$ is the number of amide groups and $N_{\rm C}$ is the number of methylene groups developed from an addition scheme on melt heat capacities of all aliphatic homopolyamides and copolymers. The equation describes the experimental results with an accuracy of $\sim 1\%$. Using this equation heat capacities were extrapolated up to the glass transition temperature as between $T_{\rm g}$ and $T_{\rm m}$ heat capacity varies with thermal history of the sample.

The glass transition temperatures show no definite pattern of change on going from nylon 6 to nylon 6.12. It is 313 K for nylon 6; nylon 11 and nylon 12 have a value of 316 K and 314 K, respectively; while nylon 6.6, nylon 6.9, nylon 6.10 and nylon 6.12 have their glass transition tem-

peratures at 323 K, 331 K, 323 K and 319 K, respectively. The recommended value of ΔC_p for 100% amorphous nylons obtained by subtracting solid C_p data (calculated from an approximate vibrational spectrum fitted to experimental heat capacities)⁴⁰ from the recommended melt data was 68, 74, 110, 118, 141 for nylon 11, nylon 12, nylon 6.9, nylon 6.10 and nylon 6.12 respectively.

TABLE 14. Heat capacity change at the glass transition temperature of the various polylactones

Polymer	$T_{g}(K)$	ΔC_p [J/(Kmol)]
Polyglycolide	318	44.4
Polypropiolactone	249	50.4
Polybutyrolactone	214	57.4
Polyvalerolactone	207	65.1
Polycaprolactone	209	59.5
Polyundecanolactone	227	110.6
Polytridecanolactone	229	146.0
Polypentadecanolactone	251	168.0

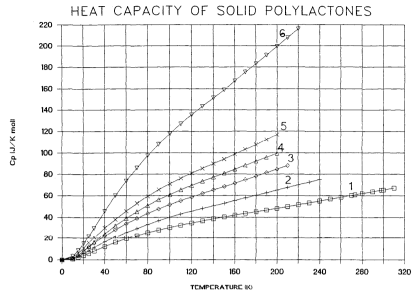


Fig. 2. Recommended heat capacities below the glass transition temperature for the following polylactones: 1. polyglycolide, 2. $poly(\beta$ -propiolactone), 3. $poly(\gamma$ -butyrolactone), 4. $poly(\delta$ -valerolactone), 5. $poly(\epsilon$ -caprolactone), and 6. polyundecanolatone.

HEAT CAPACITY OF MOLTEN POLYLACTONES

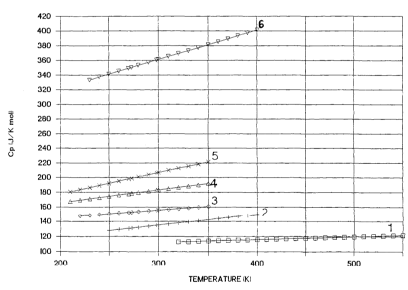


Fig. 3. Recommended heat capacities for the following molten polylactones: 1. polyglycolide, 2. poly (β propiolactone), 3. poly (γ butyrolactone), 4. poly (δ-valerolactone), 5. poly (ς-caprolactone), and 6. polyundecanolactone.

TABLE 15a. Recommended heat capacity data for amorphous poly (ethylene oxalate) below the glass transition

TABLE 16. Recommended heat capacity data for semicrystalline poly-(butylene adipate) below the glass transition

0.00	0.000
10.00	1.830
15.00	5.223
20.00	9.180
25.00	13.175
30.00	17.070
40.00	24.630
50.00	31.850
60.00	37.950
70.00	43.320
80.00	48.350
90.00	53.210
100.00	57.930
110.00	62.480
120.00	66.850
130.00	71.050
140.00	75.060
150.00	78.900
160.00	82.570
170.00	86.110
180.00	89.580
190.00	93.030
200.00	96.530
210.00	100.100
220.00	103.700
230.00	107.300
240.00	110.800
250.00	114.262
260.00	117.350
270.00	120.439
273.15	121.412
280.00	123.528
290.00	126.616
298.15	129.133
300.00	129.705
$306.00 \ (T_g)$	131.558

Temp	Heat capacity	
(K)	[J/(K mol)]	
80.00	108.734	
90.00	118.245	
100.00	127.737	
110.00	137.217	
120.00	146.690	
130.00	156.161	
140.00	165.632	
150.00	175.107	
160.00	184.587	
170.00	194.074	
180.00	203.570	
190.00	213.075	
199.00 (T_g)	221.639	

TABLE 17. Recommended heat capacity data for molten poly (butylene adipate)

Temp (K)	Heat capacity [J/(K mol)]	
199.00 (Tg)	361.685	
200.00	362.011	
210.00	365.269	
220.00	368.528	
230.00	371.786	
240.00	375.044	
250.00	378.303	
260.00	381.561	
270.00	384.819	
273.15	385.846	
280.00	388.078	
290.00	391.336	
298.15	393.992	
300.00	394.595	
310.00	397.853	
320.00	401.111	
330.00	404.370	
340.00	407.628	
350.00	410.886	
360.00	414.145	
370.00	417.403	
380.00	420.661	
390.00	423,920	
400.00	427.178	
410.00	430.436	
420.00	433.695	
430.00	436.953	
440.00	440.211	
450.00	443.470	

TABLE 15b. Recommended heat capacity data for molten *poly* (ethylene oxalate)

Temp (K)	Heat capacity [J/K mol)]	
306.00 (Tg)	187.788	
310.00	188.504	
320.00	190.295	
330.00	192.085	
340.00	193.876	
350.00	195.666	
360.00	197.456	

3.2.8. poly (Diethyl Siloxane)

In the ATHAS data bank (1980)9 no recommendation of experimental heat capacities could be made as the data reported by Karasz et al.9 on sample 1 and Turdakin et al.9 on sample 2 differed by 2-9%. Since then additional heat capacity measurements have been reported and the details of these are given in Table 2. The crystallinity reported by the authors⁴¹ on sample 3 was found to be in error since an incorrect value of ΔC_p for the sample reported was used. Hence, before proceeding with the comparison of the C_p data, it was considered necessary to reevaluate the crystallinity. The value of ΔC_p for sample 3 was recalculated to be 7.374 J/(K mol) and this together with the ΔC_p for 100% amorphous PDES (obtained from the author's melt data and data for the solid calculated from an approximate vibrational spectrum fitted to author's data) results in a crystallinity for the semicrystalline PDES of 78.6% as listed in Table 2.

The heat capacity data for samples 3 to 9 over the different temperature ranges are given in Tables A26-A27. Table A26 lists the heat capacity data below the glass transition temperature. For samples 1 and 2 data have already been deposited with the Physics Auxiliary Publication Service of the American Institute of Physics as described in Ref. 9. Table A27 shows the data for the condis crystal, liquid crystal and the melt.

Recommended experimental heat capacities below 40 K were obtained by curve-fitting the tabulated data points reported by Lebedev *et al.*⁴¹ From 10 to 20 K C_p was obtained from the equation of the smoothed data.

$$C_p = \exp[-18.90025 + 18.16324 \ln T - 5.681586 \ln T^2 + 0.6641430 \ln T^3] (\pm 0.6\%) J/(K mol) (20)$$

From 25 to 40 K curve-fitted data were obtained from the equation:

$$C_p = -8.0921 + 0.9564671 T + 4.187165 \times 10^{-4} T^2$$

(± 2.2%), J/(K mol) (21)

Heat capacities reported by Karasz⁹ were higher than both Lebedev⁴¹ and Turdakin's⁹ data in the temperature range 70 to 100 K. At 70 K, for example, it was $\sim 9\%$ higher while at 90 K it was $\sim 3\%$ higher. Therefore, to determine the recommended experimental data from 50 K-135 K (T_g), an average of only Turdakin and Lebedev's data over the temperature of overlap (60–100 K), together with the heat capacity data of Lebedev at 50 K and 110 K was fit into the equation:

$$C_p = -7.778002 + 1.064166 T - 0.00230083 T^2$$

(± 0.2%) J/(K mol) (22)

The data of Turdakin at 110 K and 120 K appeared to be associated with the glass transition, as it was 1–2% higher than that of Lebedev. Hence, only Lebedev's data at 120 and 130 K were fit into a linear equation:

$$C_p = 6.418823 + 0.6749153 T (\pm 0.4\%) J/(K mol)$$
 (23)

and this equation was used to obtain the heat capacity from 120 to 135 K (T_e).

The recommended data for molten *poly* (diethyl siloxane) were obtained by averaging the melt data reported by Lebedev from 300 to 330 K and the work carried out in our laboratory⁴² from 290–360 K. Averaged data were fit into a linear equation:

$$C_p = 96.87783 + 0.2284644 T (\pm 0.2\%) \text{ J/(K mol)}$$
 (24)

This equation was used to extend the data to the glass transition temperature. Recommended heat capacities below $T_{\rm g}$ (from 10 to 135 K) were obtained using Eqs. 20 to 23 (Table 21). The heat capacities above $T_{\rm g}$, up to 360 K can be represented by Eq. (24) (Table 22). The use of Eq. (24) to approximate the heat capacities of the condis and liquid crystalline state is a reasonable approximation.⁴²

Table 18. Recommended heat capacity data for molten poly(trimethylene adipate), poly(hexamethylene sebacate) and poly(trimethylene succinate)

Temp (K)	Heat capacity [J/(K mol)] PTMA PHMS		
310.00	348.401	NA	291.014
320.00	352.371	NA	293.544
330.00	356.341	NA	296.074
340.00	360.312	565.884	298.604
350.00	364.282	573.434	301.134
360.00	368.252	580.985	303.664
370.00	NA	588.536	NA
380.00	NA	596.087	NA
390.00	NA	603.638	NA
400.00	NA	611.188	NA

Table 19a. Recommended heat capacity data for semicrystalline poly(ethylene sebacate) below the glass transition

Temp (K)	Heat capacity [J/(K mol)]	
120.00	151.338	
130.00	159.607	
140.00	167.876	
150.00	176.145	
160.00	184.414	
170.00	192.683	
180.00	200.952	
190.00	209.221	
200.00	217.490	
210.00	225,759	
220.00	234.028	
230.00	242.297	
240.00	250.566	
$245.00 (T_g)$	254.701	

TABLE 19b. (cont'd.): Recommended heat capacity data for molten poly(ethylene sebacate)

Temp (K)	Heat capacity (J/K.mol)
245.00 (T _g)	408.760
250.00	411.540
260.00	417.100
270.00	422.660
273.15	424.411
280.00	428.220
290.00	433.780
298.15	438.311
300.00	439.340
310.00	444.900
320.00	450.460
330.00	456.020
340.00	461.580
350.00	467.140
360.00	472.700
370.00	478.260
380.00	483.820
390.00	489.380
400.00	494.940
410.00	500.500

3.3. Recommended Heat Capacities of Carbon Backbone Polymers with Pendant Aromatic/Aliphatic Groups

3.3.1. Introduction

The series of polymers covered under this section include p-substituted polystyrenes and series of polyitaconates having aliphatic as well as cyclic pendant side chains. In the ATHAS data bank (1980) heat capacity of polystyrene has already been critically evaluated.⁵ An update on this was described in Sec. 2. New heat capacity data are reported for poly(p-fluorostyrene), poly(p-chlorostyrene), poly(p-bromostyrene), poly(p-iodostyrene) and poly(p-methyl styrene) and will be discussed in the next section. The change in heat capacity at T_g (Table 23) for poly(p-chlorostyrene) and poly(p-bromostyrene) is close to that of polystyrene [$\Delta C_p = 30.8 \text{ J/(K mol)}]$. poly(p-iodostyrene) and poly(p-methyl styrene) have a slightly higher T_g as expected due to the bulky substituted group.⁴³

Heat capacity determinations have also been reported on a series of poly (di-n-alkyl itaconates) with side chain lengths varying from 1 to 10 carbon atoms and for poly (dicyclo-alkyl itaconates). Polyitaconates having linear side chains exhibit two jumps in the heat capacity. The low-temperature inflection in the heat capacity curves of poly (di-n-heptyl itaconate) to poly (di-n-decyl itaconate) has been referred to as $T_g^{\ L}$ and attributed to the independent motions of the side chains. The polymer softens and is about 60–70 K above the $T_g^{\ L}$ (Table 24).

The heat capacity data will be discussed in the subsequent sections.

3.3.2. Substituted Polystyrenes

Wunderlich et al.⁴³ have carried out a detailed analysis of heat capacities of the various substituted polystyrenes and unsubstituted polystyrenes. The details of this investigation which meet our standards of acceptable data are listed in Table 2. The equations of the curve-fitted data are given in Table 25, and heat capacity data are given in Tables A28 and A29. The recommended experimental heat capacities based on only one investigation are given in Tables 26 and 27, and plotted in Figs. 4 and 5.

3.3.3. Polyitaconates

Only two investigations are reported in the literature 44,45 which deal with the heat capacities of poly (dinalkyl itaconates). The details of the measurements which meet our standards of acceptable data are given in Table 2. Heat capacities retrieved from the graphs are given in Tables A30-A32. Table 28 gives the various curve-fitted equations. The preliminary recommended data for poly (dimethyl itaconate) and poly (di-n-propyl itaconate) are given in Tables 29 and 30 and are plotted in Fig. 6. Heat capacities of all other available solid poly (di-n-alkyl itaconates) are given in Table 31 and are plotted in Fig. 7.

Heat capacity measurements have also been reported by Cowie and coworkers^{46,47} for *poly* (dicycloalkyl itaconates) with cycloalkyl groups such as cyclooctyl, cyclodecyl and cyclododecyl. The details of the measurements which meet our standards of acceptable data are given in Table 2. Tables A33 to A34 give the heat capacities for the three itaconates. The curve-fitted equations of the smoothed data are given in Table 32. The preliminary recommended heat capacities for *poly* (dicylodecyl itaconate) and *poly* (dicyclo dodecyl itaconate) are given in Table 31 and in Fig. 7 while the heat capacity data for *poly* (dicyclooctyl itaconate) are given in Table 33.

3.3.4. poly (p-Methacryloyloxybenzoic acid)

Only one investigation⁴⁸ is reported for the heat capacity of poly(p-methacryloyloxy benzoic acid). The details of the measurements which meets our standards of acceptable data are given in Table 2. Heat capacities for an amorphous sample have been reported. The data below the glass transition temperature (T_g 316 K) are given in Table A35. The experimental data are directly used from 10 to 80 K, while the heat capacities from 80 to 310 K were fit into the equation:

$$C_p = \exp[0.0840357 (\ln T)^3 - 1.17815 (\ln T)^2 + 6.33259 (\ln T) - 7.81629] (\pm 0.4\%) J/(K mol)$$
 (25)

The preliminary, recommended heat capacities from 10 to 310 K are given in Table 34.

TABLE 20. Recommended heat capacity data for nylons

Temp (K)	Nylon 11	Nylon 12	Nylon 6.9	Nylon 6.10	Nylon 6.12
		Below	, T _g		
230.00	238.210	254.020	321.530	340.870	381.780
240.00	249.260	264.100	332.440	354.280	395.570
250.00	259.990	273.910	343.660	368.180	410.520
260.00	271.300	283.950	355.330	381.970	425.770
270.00	282.660	294.770	368.400	395.520	439.750
280.00	294.130	305.980	381.850	409.720	458.630
290.00	306.970	318.560	395.270	426.590	475.420
300.00	320.910	334.490	408.080	441.820	494.480
310.00	342.100	358.690	425.330	463.600	520.210
320.00	NA	NA	454.510	497.000	NA
330.00	NA	NA	469.639	NA	NA
		Mel	lt		
320.00	392.522	423.813	NA	NA	659.882
330.00	399.746	431.782	NA	607.279	671.350
340.00	406.970	439.751	584.476	617.257	682.818
350.00	414.194	447.720	593.709	627.235	694.286
360.00	421.418	455.689	602.942	637.213	705.754
370.00	428.642	463.658	612.175	647.191	717.222
380.00	435.866	471.627	621.408	657.169	728.690
390.00	443.090	479.596	630.641	667.147	740.158
400.00	450.314	487.565	639.874	677.125	751.626
410.00	457.538	495.534	649.107	687.103	763.094
420.00	464.762	503.503	658.340	697.081	774.562
430.00	471.986	511.472	667.573	707.059	786.030
440.00	479.210	519.441	676.806	717.037	797.498
450.00	486.434	527.410	686.039	727.015	808.966
460.00	493.658	535.379	695.272	736.993	820.434
470.00	500.882	543.348	704.505	746.971	831.902
480.00	508.106	551.317	713.738	756.949	843.370
490.00	515.330	559.286	722.971	766.927	854.838
500.00	522.554	567.255	732.204	776.905	866.306
510.00	529.778	575.224	741.437	786.883	877.774
520.00	537.002	583.193	750.670	796.861	889.242
530.00	544.226	591.162	759.903	806.839	900.710
540.00	551.450	599.131	769.136	816.817	912.178
550.00	558.674	607.100	778.369	826.795	923.646
560.00	565.898	615.069	787.602	836.773	935.114
570.00	573.122	623.038	796.835	846.751	946.582
580.00	580.346	631.007	806.068	856.729	958.050
590.00	587.570	638.976	815.301	866.707	969.518
600.00	594.794	646.945	824.534	876.685	980.986

3.4. Recommended Data on Heat Capacity of Polymers with Aromatic Groups in the Backbone

3.4.1. poly (p-Phenylene)

Only one investigation⁴⁹ is reported in the literature which deals with the heat capacity of poly (p-phenylene). The heat capacity of a semicrystalline sample has been measured from 80 to 300 K. The details of the measurements are given in Table 2 and the heat capacity data are given in Table A36. The glass transition temperature of poly (p-phenylene) is unknown, while the melting temperature is believed to be above 1000 K. The preliminary recommended heat capacities were obtained by curve-fitting

the data from 80 to 300 K into Eq. (26), and are given in Table 35:

$$C_p = \exp[8.080832 - 3.371739 (\ln T) + 0.6658902 (\ln T)^2 - 0.03270898 (\ln T)^3] J/(K mol)$$
 (26)

Figure 8 shows the plot of the heat capacity over this temperature range.

3.4.2. poly (Thio-1,4-Phenylene)

Only one investigation is reported in the literature which deals with heat capacities of *poly* (thio-1,4-phenylene).⁵⁰ The details of the measurements are given

in Table 2. Heat capacity of PTP over the range of measurement is given in Tables A37 and A38. The recommended data below the glass transition temperature (363 K) were obtained by extrapolating the author's curve-fitted data from 220 up to T_g . Both semicrystalline and amorphous heat capacities were reported to be in close agreement. Above $T_{\rm g}$, a rigid amorphous fraction dependence of heat capacity was reported for the semicrystalline polymers obtained by isothermal non-isothermal crystallization. Hence, the melt data from 540 to 600 K were extrapolated down to T_g and represent the recommended data above the glass transition temperature. The recommended data for the solid and melt are given in Tables 36 and 37. The solid heat capacities are plotted in Figure 8.

3.4.3. poly (p-Xylylene)

Only one investigation is reported in the literature which deals with heat capacities of poly(p-xylylene).⁵¹ The details of the measurements are given in Table 2. The data are given in Table A39. The preliminary recommendated data for a sample of high crystallinity $\sim 90\%$, below and above the glass transition temperature (286 K) is given in Table 38. Above T_g , the semicrystalline data is included as it is the only available data and it is for a sample of high crystallinity. Figure 8 shows the heat capacities when compared with poly(p-phenylene) and poly(thio-1,4-phenylene).

TABLE 21. Recommended heat capacity data for semicrystalline poly-(diethyl siloxane) below the glass transition

Temp (K)	Heat capacity [J/(K mol)]	
0.00	0.000	
10.00	2.477	
15.00	6.112	
20.00	10.789	
25.00	16.081	
30.00	20,979	
40.00	30.837	
50.00	39.678	
60.00	47.789	
70.00	55,440	
80.00	62.630	
90.00	69.360	
100.00	75.630	
110.00	81.440	
120.00	87.409	
130.00	94.158	
$135.00 \ (T_{\rm g})$	97.532	

Table 22. Recommended heat capacity data for molten poly(diethyl siloxane)

Temp	Heat capacity	
(K) [*]	[J/(K mol)]	
135.00 (T _g)	127.721	
140.00	128.863	
150.00	131.147	
160.00	133.432	
170.00	135.717	
180.00	138.001	
190.00	140.286	
200.00	142.571	
210.00	144.855	
220.00	147.140	
230.00	149.425	
240.00	151.709	
250.00	153.994	
260.00	156.279	
270.00	158.563	
273.15	159.283	
280.00	160.848	
290.00	163.133	
298.15	164.994	
300.00	165.417	
310.00	167.702	
320.00	169.986	
330.00	172.271	
340.00	174.556	
350.00	176.840	
360.00	179.125	

TABLE 23. Heat capacity change at the glass transition temperature of the various substituted polystyrenes

Polymer ^a	$T_{\mathbf{g}}$ (K)	ΔC_P [J/(K mol)]
PS	273	30.86
PFS	384	33.3°
PCS	406	31.1 ^d
PBS	410	31.9 ^d
PIS	424	37.9 ^d
PMS	380	34.6 ^d

^aFor the structure and abbreviations used for the various polymers see the table given in appendix.

TABLE 24. Transitions in poly (di-n-alkyl itaconates)

Polymera	T _g ^L (K)	C _p ^L [J/(K mol)]	Tg ^u (K)	$\frac{\Delta C_p^{\mathbf{u}}}{[J/(K \text{ mol})]}$
PDMI	_	_	377	54.23
PDPI		~	304	57.84
PDHI	172	45.64	248	13.04
PDOI	178	99.12	240	14.16
PDNI	187	183.36	245	11.46

^aFor the structure and abbreviations used for the various polymers see the table given in appendix.

^bData from reference 5.

Data obtained with the DuPont DSC at a heating rate of 20 K/min.

^dData obtained with Perkin-Elmer DSC-2 at a heating rate of 10 K/min.

TABLE 25. Equations used to derive the recommended heat capacities of various substituted polystyrenes

Polymer ^a	State	Equation ^b (RMS dev.)	Temperature range (K) ^c	Number of measurements	
PFS	Solid	$C_p = 44.84 + 5.21 \times 10^{-3} T + 7.476 \times 10^{-4} T^2 (0.9\%)$	130 - 384	3	
PCS	Solid	$C_p = -4.20 + 0.4866 T (0.3\%)$	300 - 406	3	
	Molten	$C_p = 112.57 + 0.2775 T(0.3\%)$	406 - 550	3	
PBS	Solid	$C_p = 6.33 + 0.4649 T (1.5\%)$	300 - 410	. 4	
	Molten	$C_p = 113.94 + 0.2801 T (1.2\%)$	410 - 550	4	
PIS	Solid	$C_p = 7.18 + 0.4945 T (0.5\%)$	300 - 424	4	
	Molten	$C_p = 139.02 + 0.2730 T (0.5\%)$	424 - 550	4	
PMS	Solid	$C_p = -3.54 + 0.5138 T (0.2\%)$	300 - 380	2	
	Molten	$C_p = 90.85 + 0.3564 T (0.3\%)$	380 - 500	2	

^aThe structure and abbreviations of the polymers are given in the appendix.

TABLE 26. Recommended heat capacity data for amorphous substituted polystyrenes below the glass transition temperature in [J/(K mol)]

Temp	PS	PFS	PCS	PBS	PIS	PMS
(K)						
130.00	57.887	58.152	NA	NA	NA	NA
140.00	61.426	60.222	NA	NA	NA	NA
150.00	65.009	62.443	NA	NA	NA	NA
160.00	68.650	64.812	NA	NA	NA	NA
170.00	72.358	67.331	NA	NA	NA	NA
180.00	76.142	70.000	NA	NA	NA	NA
190.00	80.011	72.818	NA	NA	NA	NA
200.00	83.971	75.786	NA	NA	NA	NA
210.00	88.028	78.903	NA	NA	NA	NA
220.00	92.187	82.170	NA	NA	NA	NA
230.00	95.992	85.586	NA	NA	NA	NA
240.00	100.140	89.152	NA	NA	NA	NA
250.00	104.428	92.868	NA	NA	NA	NA
260.00	108.836	96.732	NA	NA	NA	NA
270.00	113.346	100.747	NA	NA	NA	NA
273.15	114.785	102.042	NA	NA	NA	NA
280.00	117.944	104.911	NA	NA	NA	NA
290.00	122.617	109.224	NA	NA	NA	NA
298.15	126.475	112.850	NA	NA	NA	NA
300.00	127.357	113.687	141.780	145.800	155.530	150.600
310.00	132.154	118.299	146.646	150.449	160.475	155.738
320.00	137.001	123.061	151.512	155.098	165.420	160.876
330.00	141.893	127.973	156.378	159.747	170.365	166.014
340.00	146.825	133.034	161.244	164.396	175.310	171.152
350.00	151.791	138.245	166.110	169.045	180.255	176.290
360.00	156.788	143.605	170.976	173.694	185.200	181.428
370.00	161.813	149.114	175.842	178.343	190.145	186.566
380.00	NA	154.773	180.708	182.992	195.090	NA
390.00	NA	NA	185.574	187.641	200.035	NA
400.00	NA	NA	190.440	192.290	204.980	NA
410.00	NA	NA	NA	NA	209.925	NA
420.00	NA	NA	NA	NA	214.870	NA

The abbreviations used for the polymer names have been described in the appendix.

 $^{{}^{\}mathrm{b}}C_{p}$ is in J/(K mol). Temperature range over which the equation is valid.

3.4.4. poly (Oxy-3-bromo-2,6-dimethyl-1,4-phenylene)

Only one investigation²⁹ is reported in the literature which deals with the measurement of heat capacities of poly (oxy-3-bromo-2,6-dimethyl-1,4-phenylene). The details of the measurement are given in Table 2 and the only available heat capacity data are given in Table A40. The preliminary recommended heat capacities are given in Table 39 and plotted in Fig. 9. To extend the heat capacities up to T_g (559 K) the data from 480 to 520 K were curve-fitted into the equation:

$$C_p = 52.15326 + 0.396001 T (J/K mol)$$
 (27)

and, using this equation, extended to 559 K (T_g). The ΔC_p at T_g has been reported by the authors²⁰ as 18 J/(K mol) and is much lower than poly (oxy-2,6-dimethyl-1,4-phenylene) whose ΔC_p is 28.3 J/(K mol) at 482 K.

3.4.5. poly[Oxy-2,6-bis (1-methylethyl)-1,4-phenylene]

Only one investigation is reported in the literature which deals with heat capacities of poly [oxy-2,6-bis(1-methylethyl)-1,4-phenylene].⁵² The details of the measurement are given in Table 2. Heat capacity data on samples whose state has not been identified by the authors can thus be used only as a preliminary set of data. Table A41 gives the heat capacities below and above the glass transition temperature. Recommended heat capacities below the glass transition temperature (425.5 K) were obtained by extrapolating the author's curve-fitted equation from 270 to 425.5 K (Table 40). Figure 9 shows the heat capacities when compared with that of poly (oxy-2,6-dimethyl-1,4-phenylene) and poly (oxy-3-bromo-2,6-dimethyl-1,4-phenylene).

3.4.6. poly (Ethylene Terephathate)

In the ATHAS data bank (1980),⁸ the recommended heat capacity was evaluated for only amorphous poly (ethylene terephthate) as the available data was limited to samples of either low crystallinity or completely amorphous samples. Since then Collocot et al.⁵³ have reported heat capacities below 15 K. The heat capacity data retrieved from the graph are given in Table A42. Based on the author's extrapolated data for 100% crystalline and 100% amorphous PET it is now possible to obtain recommended heat capacities for the two limiting states in the low temperature range where amorphous heat capacity is usually higher than the crystalline data.

The new low temperature amorphous data were found to be in close agreement with Choy's data (sample 15, Ref., 8) while Assfalg's (sample 8, Ref., 8) showed larger deviations. Therefore, to arrive at a better recommended set of heat capacities in the low temperature region (1.2–15 K) for amorphous PET, the data of Collocot and Choy were averaged in the region of overlap and curve-fitted into the equation:

$$C_{pa} \exp \left[-4.879994 + 3.464046 \left(\ln T\right) - 0.1205552 \left(\ln T\right)^2 - 0.06363761 \left(\ln T\right)^3\right]$$
(± 0.94%) J/(K mol) (28)

This equation represents the new C_p data from 1.2 to 15 K. Above 15 K up to 590 K heat capacities of 100% amorphous PET were recommended earlier.⁸

For crystalline PET, the smoothed heat capacity data retrieved from the graph were used. From 1.2 to 15 K the smoothed data can be represented by the equation:

$$C_{pc} = \exp[-4.790307 + 3.27316 (\ln T) + 0.03140159 (\ln T)^2 - -0.100217 (\ln T)^3]$$

$$(\pm 2\%) \text{ J/(Kmol)}$$
(29)

An attempt was also made to arrive at a recommended set of heat capacities for crystalline PET above 110 K. From 20 to 110 K the heat capacities can be assumed to be close to that for amorphous PET, as only amorphous PET data is available in the literature. Between 110 K and 230 K heat capacities for only one semicrystalline sample (no 24, Ref., 8, crystallinity 0.39) were available. These data seem to be in error as they are much lower than the heat capacity for other samples of comparable crystallinities at temperatures of overlap above 240 K. Hence, it was not possible to arrive at a recommended of heat capacity for crystalline PET between 110 and 240 K. From 240 K to 520 K heat capacities were available for semicrystalline PET where crystallinity ranged between 0.01 to 0.52. Above 240 K the heat capacity of crystalline PET obtained from the two-phase model $C_p = w^c C_p^c +$ $(1 - w^{c}) C_{p}^{a}$, showed erratic variations. This indicates that in order to determine heat capacity for crystalline PET at temperatures above 110 K, data for samples of higher crystallinities are needed. The new recommended heat capacities from 1.2 to 15 K for 100% crystalline and 100% amorphous PET are given in Table 41.

3.4.7. poly (Butylene Terephthalate)

Four investigations have been reported in the literature on the heat capacity measurements of five semicrystalline and one quenched sample of *poly* (butylene terephthalate) (PBT).⁵⁴⁻⁵⁷ Details of the three investigations⁵⁴⁻⁵⁶ which meet our standards of acceptability are given in Table 2. Data on these samples below and above the glass transition and for the melt are given in Tables A43 to A46. Heat capacity data on semicrystalline PBT reported by Bair *et al.*⁵⁷ has not been included, as a very high heating rate of 40 °C/min has been used and the reported data are much higher than those reported by other authors.

A detailed study on PBT carried out in our laboratory⁵⁴ has revealed the existence of rigid amorphous fractions of the semicrystalline samples crystallized from glass and from melt between 275 and 490 K. The glass transition temperatures of amorphous PBT samples quenched in a

TABLE 27. Recommended heat capacity data for molten substituted polystyrenes in [J/(K mol)]

Temp (K)	PS	PFS	PCS	PBS	PIS	PMS
380.00	195.934	NA	NA	NA	NA	NA
390.00	198.587	NA	NA	NA	NA	229.846
400.00	201.240	NA	NA	NA	NA	233.410
410.00	203.893	NA	226.345	NA	NA	236.974
420.00	206.546	NA	229.120	231.582	NA	240.538
430.00	209.199	NA	231.895	234.383	256.410	244.102
440.00	211.852	NA	234.670	237.184	259.140	247.666
450.00	214.505	NA	237.445	239.985	261.870	251.230
460.00	217.158	NA	240.220	242.786	264.600	254.794
470.00	219.811	NA	242.995	245.587	267.330	258.358
480.00	222.464	NA	245.770	248.388	270.060	261.922
490.00	225.117	NA	248.545	251.189	272.790	265.486
500.00	227.770	NA	251.320	253.990	275.520	269.050
510.00	230.423	NA	254.095	256.791	278.250	NA
520.00	233.076	NA	256.870	259.592	280.980	NA
530.00	235.729	NA	259.645	262.393	283.710	NA
540.00	238.382	NA	262.420	265.194	286.440	NA
550.00	241.035	NA	265.195	267.995	289.170	NA
560.00	243.688	NA	NA	NA	NA	NA
570.00	246.341	NA	NA	NA	NA	NA
580.00	248.994	NA	NA	NA	NA	NA
590.00	251.647	NA	NA	NA	NA	NA
600.00	254.300	NA	NA	NA	NA	NA

The abbreviations used for the polymer names have been described in the appendix.

HEAT CAPACITY OF SOLID POLYSTYRENES

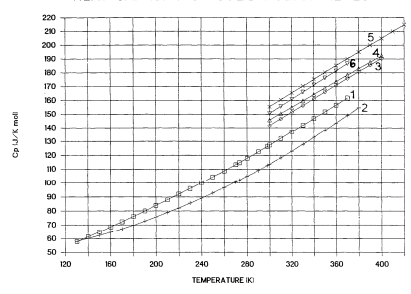


Fig. 4. Recommended heat capacities for the following solid polymers: 1. polystyrene, 2. poly (p-fluorostyrene), 3. poly (p-chlorostyrene), 4. poly (p-bromostyrene), 5. poly (p-iodostyrene), and 6. poly (p-methyl styrene).

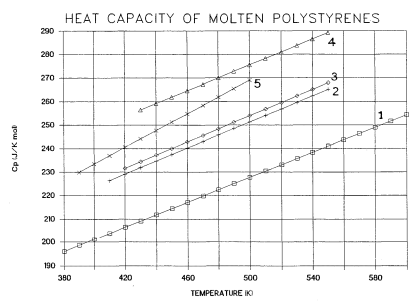


Fig. 5. Recommended heat capacities for the following molten polymers: 1. polystyrene, 2. poly (p-fluorostyrene), 3. poly (p-chlorostyrene), 4. poly (p-bromostyrene), 5. poly (p-iodostyrene), and 6. poly (p-methyl styrene).

cooling bath of glass beads was found to be 248 K, which is much lower than that reported for semicrystalline samples, where $T_{\rm g}$ varies between 310–325 K. Amorphous samples showed a second $T_{\rm g}$ at about 310 K with a ΔC_p of 20 J/(K mol). Since the lower limit of the DSC used by us was 220 K and amorphous samples showed a $T_{\rm g}$ at 248 K, C_p data below the first $T_{\rm g}$ could not be retrieved. Due to the presence of a crystallization exotherm before the second $T_{\rm g}$, the heat capacities above $T_{\rm g}$ are no longer due to the amorphous melt.

As a preliminary set of heat capacities of semicrystalline PBT (below the glass transition at 320 K) the data for samples 1 and 3 from 150–240 K were averaged over the region of overlap and fit into the equation:

$$C_p = -6.938805 + 0.9502766 T - 4.876461 \times 10^{-5} T^2 (\pm 0.11\%) \text{ J/(K mol)}$$
 (30)

Between 240 to 310 K the averaged data for samples (1,4–6) were smoothed by fitting into the equation:

$$C_p = 13.13337 + 0.8544043 T (\pm 0.15\%) J/(K mol) (31)$$

To obtain heat capacities of the melt, the data of sample 1 (which is an average of 7 measurements) were fit into the equation:

$$C_p = 219.6479 + 0.452209 T (\pm 0.1\%) J/(K mol)$$
 (32)

and this equation was used to extrapolate the data up to $T_{\rm g}$. The data on samples 2-6 have not been included, due to deviations above 1% introduced during the retrieval of

data from the graph. Heat capacity data on amorphous PBT (sample 2) is not recommended, since it is associated with crystallization and shows the dependence of heat capacity on the rigid amorphous fraction.

Preliminary recommended data below $T_{\rm g}$ (248 K) obtained from Eq. (30), between 1st $T_{\rm g}$ and 2nd $T_{\rm g}$ at 320 K obtained from Eq. (31) and heat capacities of the melt obtained from Eq. (32) are given from 248 K up to 570 K in Tables 42 and 43 respectively.

3.4.8. poly (4-Hydroxybenzoic Acid)

Only one investigation is reported in the literature⁵⁸ which deals with heat capacities of commercial, semicrystalline poly (4-hydroxybenzoic acid). The details of the measurement are given in Table 2. Heat capacities are given for a semicrystalline sample below the glass transition region and in the glass transition region in Tables A47 and A48. The recommended heat capacities (Table 44) from 170 to T_g (434 K) are included in the ATHAS data bank as a preliminary set of data as from 290 K up to 420 K the reported heat capacities are associated already with the beginning of a broad glass transition. The data from 390 to 420 K were curve-fitted into the equation:

$$C_p = 19.28989 + 0.362003 T (\pm 0.04\%) J/(K mol)$$
 (33)

This was used to obtain C_p data from 390 to 434 K (T_g). From 500 to 650 K the author has reported C_p data of the melt obtained from an empirical addition scheme in

HEAT CAPACITY OF LINEAR MACROMOLECULES

TABLE 28. Results of curve fitting literature data on heat capacity of poly (di-n-alkyl itaconates) to derive the recommended data

Polymer ^a	State		Equation ^b (RMS dev.)	Temperature range (K) ^c
PDMI	Solid	(i)	$C_p = -6.68519 + 1.038724 T - 0.0010872 T^2 (0.8\%)$	110–290
		(ii)	$C_p = 64.5583 + 0.47983 T (0.3\%)$	300–377
	Molten		$C_p = 213.0095 + 0.22989 T (0.2\%)$	377- 450
PDPI	Solid	(i)	$C_p = 29.5446 + 1.284867 T - 0.00099196 T^2 (0.6\%)$	110- 210
		(ii)	$C_p = 75.4368 + 0.84752 T (0.4\%)$	220- 304
	Molten		$C_p = 283.0105 + 0.35491 T (0.1\%)$	304 410
PDHI	Solid		$C_p = 215.2651 - 0.8023134 T + 0.00795689 T^2 (0.6\%)$	110– 170
PDOI	Solid		$C_p = 74.58574 + 1.67328 T (0.7\%)$	110– 170
PDNI	Solid		$C_p = 19.2201 + 2.033779 T (O.2\%)$	110– 180
PDDI	Solid		$C_p = 100.2314 + 1.6497 T (1.0\%)$	110- 170

^aThe structure and abbreviations of the polymers are given in the appendix.

TABLE 29a. Recommended heat capacity data for amorphous poly (dimethyl itaconate) below the glass transition

TABLE 29b. Recommended heat capacity data for molten *poly* (dimethyl itaconate)

Temp (K)	Heat capacity [J/(K mol)]	Temp (K)	Heat capacity [J/(K mol)]	
110.00	94.419	377.00 (T _g)	299,680	
120.00	102.306	380.00	300.370	
130.00	109.975	390.00	302.669	
140.00	117.427	400.00	304.968	
150.00	124.661	410.00	307.267	
160.00	131.678	420.00	309.566	
170.00	138.477	430.00	311.865	
180.00	145.059	440.00	314.164	
190.00	151.424	450.00	316.463	
200.00	157.571			
210.00	163.500			
220.00	169.212			
230.00	174.707			
240.00	179.984			
250.00	185.044			
260.00	189.887			
270.00	194.512			
273.15	195.923			
280.00	198.919			
290.00	203.109			
298.15	207.619			
300.00	208.507			
310.00	213.305			
320.00	218.104			
330.00	222.902			
340.00	227.700			
350.00	232.499	·		
360.00	237.297			
370.00	242.095			
$377.00 \ (T_g)$	245.454			

 $^{{}^{\}mathrm{b}}C_{p}$ is in J/(Kmol).

Temperature range over which the equation is valid.

which C_p data of several copolymers of POB and PON are evaluated.

$$C_p = n \operatorname{ON}(173.65 + 0.285 T) + n \operatorname{OB}(132.31 + 0.179 T)$$
 (34)

Since this is not truly experimental, the data are not recommended. The recommended heat capacities are plotted in Figure 10.

3.4.9. poly (2,6-Hydroxynaphthoic Acid)

Only one investigation⁵⁸ is reported in the literature which deals with heat capacities of commercial, semicrystalline poly (2,6-hydroxynaphthoic acid). The details of the measurements are given in Table 2. Heat capacity data are given in Tables A49 (below $T_{\rm g}$), A50 (in the glass transition region) and A51 (melt). To obtain recommended heat capacities up to glass transition (399 K), the data between 170 and 280 K were extrapolated as above 290 K the reported data are associated with the beginning of the glass transition. Heat capacity of the melt from 399 K ($T_{\rm g}$) up to 650 K was obtained by extrapolating author's curve-fitted data between 500 and 640 K. The recommended heat capacities are given in Table 45 and 46 and are plotted in Fig. 10.

3.4.10. poly (Ethylene-2,6-naphthalene Dicarboxylate)

Only one investigation⁵⁹ is reported in the literature which deals with the heat capacity measurement of poly (ethylene-2,6-naphthalene dicarboxylate) (PEN). Heat capacity of semicrystalline samples crystallized from the melt and from the glassy state under widely ranging crystallization conditions, and of amorphous samples prepared with different cooling rates have been reported. The details of this investigation are given in Table 2. Heat capacity data below T_g and of the melt are given in Tables A52 and A53. The author's curve-fitted equation for the heat capacity data between 220-300 K was used to obtain the heat capacity of semicrystalline PEN from 220 up to 390 K (glass transition temperature). The glass transition temperature on cooling changes logarithmically $(T_g = 382.3 + 2.03 \ln q, q \text{ in K/min})$. Amorphous PEN shows a higher heat capacity above 300 K up to the glass transition. The authors have reported the difference to be $\sim 3\%$ [9–10 J/(K/mol)] above 340 K. Between $T_{\rm g}$ and $T_{\rm m}$ a rigid amorphous fraction (up to 0.2) dependent on the crystallinity was reported. This fraction starts to gain mobility at 430 K and decreases in amount, to reach zero after melting of the low melting crystals. Since amorphous samples were associated with crystallization which occurred between 460-510 K, recommended heat capacity of the melt were obtained from 390 K (T_e) up to 600 K by extrapolating the author's curve-fitted equation (540-600 K) for the melt. The change in heat capacity at the glass transition was found to be 81.57 J/(K mol), close to the value of 80.1 J/(K mol) estimated by the authors. The recommended heat capacity below and above $T_{\rm g}$ are given in Tables 47 and 48.

Table 30a. Recommended heat capacity data for amorphous poly(di-n-propyl itaconate) below the glass transition

Temp	Heat capacity
(K)	[J/(K mol)]
110.00	158.877
120.00	169.444
130.00	179.813
140.00	189.983
150.00	199.955
160.00	209.729
170.00	219.304
180.00	228.681
190.00	237.859
200.00	246.839
210.00	255.621
220.00	261.892
230.00	270.367
240.00	278.842
250.00	287.318
260.00	295.793
270.00	304.268
273.15	306.938
280.00	312.743
290.00	321.219
298.15	328.126
300.00	329.694
$304.00 \ (T_g)$	333.084

TABLE 30b. Recommended heat capacity data for molten poly(di-n-propyl itaconate)

Temp (K)	Heat capacity [J/(K mol)]	
304.00 (Tg)	390.902	
310.00	393.032	
320.00	396.581	
330.00	400.130	
340.00	403.679	
350.00	407.228	
360.00	410.777	
370.00	414.326	
380.00	417.875	
390.00	421.424	
400.00	424.973	
410.00	428.522	

Temp (K)	PDHI	PDOI	PDNI	PDDI	PDCYDI	PDCYDDI
110.00	223.289	258.647	242.936	281.698	203.786	239.776
120.00	233.567	275.380	263.274	298.195	225.792	256.298
130.00	245.436	292.113	283.611	314.692	247.952	273.505
140.00	258.896	308.846	303.949	331.189	270.265	291.398
150.00	273.948	325.578	324.287	347.686	292.730	309.978
160.00	290.591	342.311	344.625	364.183	315.348	329.242
170.00	308.826	359.044	364.963	380.680	338.119	349.193
180.00	NA	NA	385.300	397.177	361.043	369.830
190.00	NA	NA	NA	413.674	384.120	391.152
200.00	NA	NA	NA	430.171	407.350	413.160
210.00	NA	NA	NA	NA ·	430.733	435.854
220.00	NA	NA	NA	NA	454.268	459.234
230.00	NA	NA	NA	NA	475.543	483.299
240.00	NA	NA	NA	NA	496.822	508.050
250.00	NA	NA	NA	NA	518.101	533.488
260.00	NA	NA	NA	NA	539.380	559.610
270.00	NA	NA	NA	NA	560.659	586.419
273.15	NA	NA	NA	NA	567.362	595.006
280.00	NA	NA	NA	NA	581.938	613.914

TABLE 31. Recommended heat capacity data for amorphous polyitaconates below the glass transition temperature in (J/K.mol)

The abbreviations used for the polymer names have been described in the appendix.

3.4.11. poly (Oxy-1,4-phenylene-oxy-1,4-phenylene-carbonyl-1,4-phenylene)

Four investigations⁶⁰⁻⁶³ have been reported in the literature which deal with heat capacities of poly (oxy-1,4-phenylene-oxy-1,4-phenylene-carbonyl-1,4-phenylene) (PEEK). Three⁶¹⁻⁶³ meet our standards of acceptable data. The details of the measurements are given in Table 2. Table A54 lists the heat capacity data below $T_{\rm g}$ (419 K) while A55 gives the melt data. Heat capacities of semicrystalline samples crystallized at a wide range of conditions, and quenched with liquid N2 were measured by Cheng and Wunderlich.⁶² Below 240 K the heat capacities of semicrystalline and amorphous samples were identical, and from 340 to 410 K the heat capacity showed dependence on crystallinity. Heat capacity of amorphous PEEK reported on sample 4 by Kemmish et al. 61 agree below $T_{\rm g}$ to $\sim 1\%$ with Cheng and Wunderlich. 62 Above $T_{\rm g}$ (419 K) and below $T_{\rm m}$ the heat capacity showed dependence on the rigid-amorphous fraction (fraction of amorphous material that does not contribute to the increase in C_p at T_g). To obtain the recommended heat capacity data below $T_{\rm g}$ the author's⁶² curve-fitted data from 130 to 240 K were used.

From 340 to 410 K the data for 100% crystalline and 100% amorphous were retrieved from the graph of heat capacity vs crystallinity. Table 49 shows the heat capacity as a function of crystallinity from 340 to 410 K and this was smoothed further by curve-fitting into the following equations:

$$C_p^c = 52.56964 + 0.8790476 T [\pm 0.2\%, J/(K mol)]$$
 (35)

$$C_p^a = 28.74449 + 0.9758333 T [\pm 0.13\%, J/(K mol)]$$
 (36)

Equation (35) was used to obtain the heat capacity of crystalline PEEK from 250 K to 410 K and Eq. (36) was used to obtain heat capacity of amorphous PEEK from 250 to 419 K $(T_{\rm g})$.

Melt heat capacities reported by Wunderlich and Cheng (sample 4) and Kemmish (sample 1) agreed to $\sim 2\%$. Since the measurements on sample 4 were more extensive (average of 14 runs in the temperature range of 600–680 K), their data were used as recommended heat capacities from 419 K ($T_{\rm g}$) up to 680 K. The recommended set of data below and above $T_{\rm g}$ are listed in Table 50 and Table 51 and are plotted in Fig. 11.

3.4.12. poly[Oxy-1,4-phenylene-sulfonyl-1,4-phenylene-oxy-1,4-phenylene-(1-methylidene)-1,4-phenylene]

Recommended experimental heat capacity data of this bisphenol based amorphous polysulfone are based on two out of three investigations reported in the literature. 21,64,65 Details of the investigations included are given in Table 2. Heat capacities have also been measured by Privalko *et al.* 65 from 308 to 523 K, but only graphical data of polysulfone (chemical structure not revealed) has been presented for the temperature dependence of a reduced heat capacity. Data on the two samples discussed here in the temperature regions below and above the glass transition are given in Tables A56 and A57. To obtain the recommended data below T_g (458.5 K), the tabulated data on sample 2 from 10 to 350 K were smoothed by curve fitting into the following equations:

From 10–30 K:
$$C_p = \exp[-5.947623 + 4.872284 (\ln T) - 0.7156859 (\ln T)^2 + 0.04178513 (\ln T)^3] (\pm 0.4\%)$$
 (37)

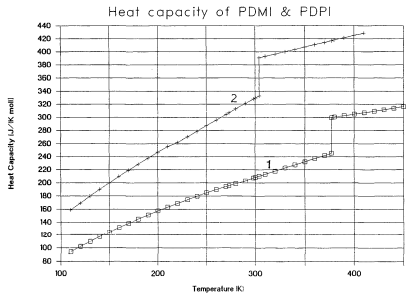


Fig. 6. Recommended heat capacies for the following two polymers:
1. poly(dimethyl itaconate) and 2. poly(di-n-propyl itaconate).



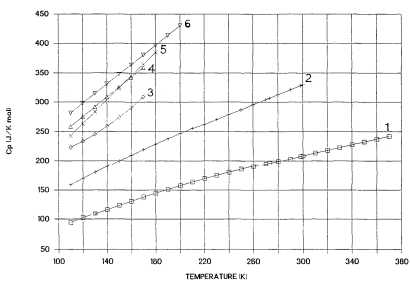


Fig. 7. Recommended heat capacities for several solid polyitaconates: 1. poly(dimethyl itaconate), 2. poly(di-n-propyl itaconate), 3. poly(di-n-hexyl itaconate, 4. poly(di-n-octyl itaconate), 5. poly(di-n-nonyl itaconate), and 6. poly(di-n-decyl itaconate).

TABLE 32. Results of curve fitting literature data on heat capacity of poly(dicycloalkyl itaconates) to derive the recommended data

Polymer ^a	State		Equation ^b (RMS dev.)	Temperature range (K) ^c
PDCyOI	Solid	(i)	$C_p = 9.00687 + 1.896353 T - 0.00011835 T^2 (0.6\%)$	110 -200
		(ii)	$C_p = 17.34212 + 1.99247 T - 0.00082074 T^2 (0.4\%)$	210 – 310
		(iii)	$C_p = 119.4146 + 1.41202 T (0.2\%)$	320 - 390
	Molten		$C_p = 423.197 + 0.80616 T (0.1\%)$	390 - 440
PDCyDI	Solid	(i)	$C_p = -28.2002 + 2.02489 T - 0.00076432 T^2 (O.8\%)$	110 – 230
		(ii)	$C_p = -13.876 + 2.1279 T (0.3\%)$	240 – 280

^aThe structure and abbreviations of the polymers are given in the appendix.

From 40–60 K: $C_p = -64.21593 + 5.459890 T - 0.0586861 T^2 + 0.000291628 T^3 (\pm 0.47\%)$ (38)

TABLE 33a. Recommended heat capacity data for amorphous poly (dicyclo octyl itaconate) below the glass transition

From 70–190 K: $C_p = 26.148 + 1.43144 T + 0.001090276 T^2 - 2.558776 \times 10^{-6} T^3 (\pm 0.8\%)$	(39)			
From 200–310 K: $C_p = -50.30969 + 2.162671 T - 0.001161756 T^2 (\pm 0.17\%)$	(40)			
From 320–350 K: $C_p = -239.516 + 3.3455461 T - 0.003009135 T^2 (\pm 0.4\%)$	(41)			
From 360–450 K: $C_p = 271.6364 + 0.8409091 T$ (± 1.6%)	(42)			
The heat capacities reported by Richardson ²¹ on sample 1 were higher than that of sample 2 in the region of overlap. For example at 380 K it was $\sim 5\%$ higher while at				

The heat capacities reported by Richardson²¹ on sample 1 were higher than that of sample 2 in the region of overlap. For example at 380 K it was ~ 5% higher while at 440 K it was higher by ~ 8%. Since the error limit of the measurements on sample 2 above 330 K was 5%, the data on sample 1 from 380–440 K and the data points from 340 to 370 K of sample 2 were further smoothed by curve-fitting into the equation:

$$C_p = 59.02983 + 1.460377 T (\pm 0.6\%)$$
 (43)

Recommended heat capacities below the glass transition were obtained as follows. From 10 to 30 K Eq. (37), 40–60 K Eq. (38), 70–190 K Eq. (39), 200–310 Eq. (40), 320–330 K Eq. (41) and from 340 to 458.5 K ($T_{\rm g}$) Eq. (42) was used. Heat capacities of molten polysulfone [Eq. (43)] are recommended. The data below and above $T_{\rm g}$ are listed in Tables 52 and 53, and plotted in Fig. 12.

Temp	Heat capacity
(K)	[J/(K mol)]
110.00	216.174
120.00	234.865
130.00	253.533
140.00	272.177
150.00	290.797
160.00	309.394
170.00	327.966
180.00	346.516
190.00	365.041
200.00	383.543
210.00	399.566
220.00	415.961
230.00	432.193
240.00	448.260
250.00	464.163
260.00	479.902
270.00	495.477
273.15	500.349
280.00	510.887
290.00	526.134
298.15	538.438
300.00	541.216
310.00	556.134
320.00	571.259
330.00	585.380
340.00	599.500
350.00	613.620
360.00	627.740
370.00	641.860
380.00	655.980
$390.00 \ (T_g)$	670.100

 $^{{}^{\}mathrm{b}}C_{p}$ is in J/(K mol).

Temperature range over which the equation is valid.

TABLE 33b. Recommended heat capacity data for molten *poly* (dicyclooctyl itaconate)

TABLE 35. Recommended heat capacity data for semicrystalline poly-(p-phenylene) below the glass transition

octyl itaconate)		(p-phenylene) below the glass transition		
Temp (K)	Heat capacity [J/(K mol)]	Temp (K)	Heat capacity [J/(K mol)]	
390.00 (Tg)	737.599	80.00	28.220	
400.00	745.660	90.00	30.312	
410.00	753.722	100.00	32.486	
420.00	761.784	110.00	34.729	
430.00	769.845	120.00	37.031	
440.00	777.907	130.00	39.387	
		140.00	41.792	
		150.00	44.241	
		160.00	46.733	
		170.00	49.263	
		180.00	51.831	
		190.00	54.434	
TABLE 34 Recommended heat capacit	y data for <i>poly</i> (p-methacryloyloxy	200.00	57.071	
benzoic acid) below the gla	ass transition	210.00	59.740	
		220.00	62.440	
Temp	Heat capacity	230.00	65.171	
(K)	[J/(K mol)]	240.00	67.930	
	[6,(42,333)]	250.00	70.717	
10.00	3.800	260.00	73.531	
15.00	9.112	270.00	76.371	
20.00	15.150	273.15	77.271	
25.00	21.383	280.00	79.236	
30.00	27.630	290.00	82.126	
40.00	40.050	298.15	84.499	
50.00	49.950	300.00	85.040	
60.00	60,870			
70.00	70.890			
80.00	79.950			
90.00	88.272			
100.00	96.346	TABLE 36. Recommended heat c	apacity data for semicrystalline poly-	
110.00	104.249	(thio-1,4-phenylene) b	elow the glass transition	
120.00	112.040		-	
130.00	119.765	Temp	Heat capacity	
140.00	127.461	(K)	[J/(K mol)]	
150.00	135.160	()	[-,(,)]	
160.00	142.887	220.00	80.263	

150.663

158.505

166.429

174.448

182.575

190.820

199.191

207.699

216.350

225.152

234.112

236.967

243.235

252.529

260.233

261.999

271.650

Temp (K)	Heat capacity [J/(K mol)]
220.00	80.263
230.00	84.053
240.00	87.848
250.00	91.651
260.00	95.466
270.00	99.297
273.15	100.507
280.00	103.147
290.00	107.019
298.15	110.193
300.00	110.916
310.00	114.840
320.00	118.794
330.00	122.781
340.00	126.802
350.00	130.860
360.00	134.957
$363.00 (T_s)$	136.193

170.00

180.00

190.00

200.00

210.00

220.00

230.00

240.00

250.00

260.00

270.00

273.15

280.00

290.00

298.15

300.00

310.00

Table 37. Recommended heat capacity data for molten poly (thio-1,4-phenylene)

Heat capacity Temp [J/(K mol)] (K) $363.00 (T_g)$ 165.374 166.254 370.00 167.511 380.00 168.769 390.00 400.00 170.026 410.00 171.283 172.541 420.00 173.798 430.00 440.00 175.056 450.00 176.313 460.00 177.570 470.00 178.828 180.085 480.00 490.00 181.343 182.600 500.00 510.00 183.857 185.115 520.00 530.00 186.372 540.00 187.630 550.00 188.887 560.00 190.144 191.402 570.00 580.00 192.659 193.917 590.00 600.00 195.174

Table 39. Recommended heat capacity data for amorphous poly (oxy-2,6-dimethyl,5-bromo-1,4-phenylene) below the glass transition

Temp (K)	Heat capacity [J/(K mol)]
310.00	173.811
320.00	177.913
330.00	182.004
340.00	186.086
350.00	190.158
360.00	194.220
370.00	198.272
380.00	202.315
390.00	206.347
400.00	210.370
410.00	214.383
420.00	218.386
430.00	222.379
440.00	226.363
450.00	230.337
460.00	234.300
470.00	238.254
480.00	242.234
490.00	246.194
500.00	250.154
510.00	254.114
520.00	258.074
530.00	262.034
540.00	265.994
550.00	269.954
$559.00 (T_g)$	273.518

Table 38 Recommended heat capacity data for semicrystalline (crystallinity = 90%) poly(p-xylylene)

Temp	Heat capacity
(K)	[J/(K mol)]
	73
220.00	95.241
230.00	100.179
240.00	105.148
250.00	110.149
260.00	115.181
270.00	120.243
273.15	121.000
280.00	125.337
$286.00 \ (T_g)$	128.409
290.00	130.460
298.15	134.664
300.00	135.622
310.00	140.817
320.00	146.045
330.00	151.308
340.00	156.604
350.00	161.933
360.00	167.297
370.00	172.694
380.00	178.125
390.00	183.589
400.00	189.087
410.00	194.619

Table 40. Recommended heat capacity data for poly(oxy-2,6-bis(1-methylethyl)-1,4-phenylene) below the glass transition

Temp (K)	Heat capacity [J/(K mol)]
270.00	227.707
273.15	229.833
280.00	234.456
290.00	241.206
298.15	246.707
300.00	247.955
310.00	254.705
320.00	261.455
330.00	268.204
340.00	274.954
350.00	281.703
360.00	288.453
370.00	295.203
380.00	301.952
390.00	308.702
400.00	315.451
410.00	322.201
420.00	328.951
$425.50 (T_g)$	332.663

TABLE 41. Recommended heat capacity data for poly (ethylene terephthalate)

TABLE 43. Recommended heat capacity data for molten poly (butylene terephthalate)

Temp (K)	Heat capacity [J/(K mol)]	
	Crystalline	Amorphous
1.2	0.014180	0.014224
1.4	0.021517	0.023980
1.6	0.030947	0.037436
1.8	0.042689	0.055108
2.0	0.056945	0.077458
3.0	0.172138	0.271379
4.0	0.373338	0.619381
5.0	0.672904	1.124794
6.0	1.078123	1.774736
7.0	1.592487	2.548069
8.0	2.216689	3.420507
9.0	2.949358	4.367650
10.0	3.787629	5.366644
15.0	9.404000	10.51606

Temp	Heat capacity
(K)	[J/(K mol)]
248.00 (T _e)	331.796
250.00	332.700
260.00	337.222
270.00	2/1 7//

TABLE 42. Heat capacity data for semicrystalline poly (butylene terephthalate) below the glass transition

Temp (K)	Heat capacity [J/(K mol)]
150.00	134.505
160.00	143.857
170.00	153.199
180.00	162.531
190.00	171.853
200.00	181.166
210.00	190.469
220.00	199.762
230.00	209.045
240.00	218.190
$248.00 (T_g)$	225.026
250.00	226.734
260.00	235.278
270.00	243.823
273.15	246.514
280.00	252.367
290.00	260.911
298.15	267.874
300.00	269.455
310.00	277.999
$320.00\ (T_{\rm g})$	286.543

(K)	[3/(12 11101)]
248.00 (T _g)	331.796
250.00	332,700
260.00	337.222
270.00	341.744
273.15	343.169
280.00	346.266
290.00	350.789
298.15	354.474
300.00	355.311
310.00	359.833
320.00	364.355
330.00	368.877
340.00	373.399
350.00	377.921
360.00	382.443
370.00	386.965
380.00	391.487
390.00	396.009
400.00	400.532
410.00	405.054
420.00	409.576
430.00	414.098
440.00	418.620
450.00	423.142
460.00	427.664
470.00	432.186
480.00	436.708
490.00	441.230
500.00	445.752
510.00	450.274
520.00	454.797
530.00	459.319
540.00	463.841
550.00	468.363
560.00	472.885
570.00	477.407

TABLE 44. Recommended heat capacity data for semicrystalline *poly* (4-hydroxybenzoic acid) below the glass transition

TABLE 46. Recommended heat capacity data for molten poly(2,6-hydroxynaphthoic acid)

Temp (K)	Heat capacity [J/(K mol)]
170.00	70.762
180.00	74.739
190.00	78.716
200.00	82.693
210.00	86.670
220.00	90.647
230.00	94.624
240.00	98.601
250.00	102.578
260.00	106.555
270.00	110.532
273.15	111.784
280.00	114.509
290.00	118.400
298.15	121.835
300.00	122.600
310.00	126.700
320.00	131.800
330.00	135.100
340.00	139.500
350.00	144.100
360.00	148.500
370.00	152.600
380.00	156.900
390.00	160.471
400.00	164.091
410.00	167.711
420.00	171.331
430.00	174.951
$434.00 \ (T_g)$	176.399

Temp (K)	Heat capacity [J/(K mol)]
399.00 (Tg)	274.140
400.00	274.480
410.00	277.884
420.00	281.288
430.00	284.691
440.00	288.095
450.00	291.498
460.00	294.902
470.00	298.306
480.00	301.709
490.00	305.113
500.00	308.517
510.00	311.920
520.00	315.324
530.00	318.727
540.00	322.131
550.00	325.535
560.00	328.938
570.00	332.342
580.00	335.745
590.00	339.149
600.00	342.553
610.00	345.956
620.00	349.360
630.00	352.763
640.00	356.167
650.00	359.571

TABLE 45. Recommended heat capacity data for semicrystalline poly (2,6-hydroxynaphthoic acid) below the glass transition

Table 47. Recommended heat capacity data for semicrystalline poly(ethylene-2,6-naphthalene dicarboxylate) below the glass transition

Temp	Heat capacity
(K)	[J/(K mol)]
170.00	104.318
180.00	110.269
190.00	116.220
200.00	122.171
210.00	128.122
220.00	134.073
230.00	140.024
240.00	145.975
250.00	151.926
260.00	157.877
270.00	163.828
273.15	165.703
280.00	169.779
290.00	175.730
298.15	180.580
300.00	181.681
310.00	187.632
320.00	193.583
330.00	199.534
340.00	205.485
350.00	211.436
360.00	217.387
370.00	223.338
380.00	229.289
390.00	235.240
399.00 (T _g)	240.596

Temp (K)	Heat capacity [J/(K mol)]
220.00	203.045
230.00	211.317
240:00	219.620
250.00	227.956
260.00	236.324
270.00	244.725
273.15	247.377
280.00	253.152
290.00	261.633
298.15	268.546
300.00	270.115
310.00	278.596
320.00	287.078
330.00	295.559
340.00	304.041
350.00	312.522
360.00	321.004
370.00	329.485
380.00	337.967
$390.00 (T_g)$	346.448

TABLE 48. Recommended heat capacity data for molten *poly* (ethylene-2,6-naphthalene dicarboxylate)

Temp (K)	Heat capacity [J/(K mol)]
390.00 (Tg)	428.020
400.00	432.114
410.00	436.208
420.00	440.302
430.00	444.396
440.00	448.490
450.00	452.585
460.00	456.679
470.00	460.773
480.00	464.867
490.00	468.961
500.00	473.055
510.00	477.149
520.00	481.243
530.00	485.337
540.00	489.431
550.00	493.526
560.00	497.620
570.00	501.714
580.00	505.808
590.00	509.902
600.00	513.996

Table 49. Results of curve fitting the heat capacity data for poly(oxy-1,4-phenylene-oxy-1,4-phenylenecarbonyl-1,4-phenylene) as a function of crystallinity into the equation $C_p = w^C C_p^C + (1-w^C) C_p^a$

Temp (K)	Heat capacity [J/(K mol)]	
	crystalline	amorphous
340.00	350.550	360.191
350.00	360.000	370.382
360.00	370.000	380.085
370.00	378.880	390.298
380.00	387.000	400.000
390.00	394.440	409.000
400.00	403.500	418.000
410.00	413.330	429.500

TABLE 50. Recommended heat capacity data for poly(oxy-1,4-phenylene-oxy-1,4-phenylenecarbonyl-1,4-phenylene) below the glass transition

Temp (K)	Heat capacity [J/(K mol)]	
	Crystalline	Amorphous
130.00	152.016	152.016
140.00	161.309	161.309
150.00	171.008	171.008
160.00	181.161	181.161
170.00	191.808	191.808
180.00	202.990	202.990
190.00	212.743	212.743
200.00	223.213	223.213
210.00	234.007	234.007
220.00	244.811	244.811
230.00	255.627	255.627
240.00	266.453	266.453
250.00	272.332	272.703
260.00	281.122	282.461
270.00	289.912	292.219
273.15	292.681	295.293
280.00	298.703	301.978
290.00	307.493	311.736
298.15	314.658	319.689
300.00	316.284	321.494
310.00	325.074	331.253
320.00	333.865	341.011
330.00	342.655	350.769
340.00	351.446	360.528
350.00	360.236	370.286
360.00	369.027	380.044
370.00	377.817	389.803
380.00	386.608	399.561
390.00	395.398	409.319
400.00	404.189	419.078
410.00	412.979	428.836
419.00 $(T_{\rm g})$	NA	437.619

TABLE 51. Recommended heat capacity data for molten poly (oxy-1,4phenylene-oxy-1,4-phenylene carbonyl-1,4-phenylene)

TABLE 52. Recommended heat capacity data for amorphous poly (oxy-1,4-phenylene-sulfonyl-1,4-phenylene-oxy-1,4-phenylene-(1-methylidene)-1,4-phenylene) the glass transition

Temp (K)	Heat capacity [J/(K mol)]
419.00 (Tg)	515.765
420.00	516.260
430.00	521.215
440.00	526.170
450.00	531.125
460.00	536.080
470.00	541.035
480.00	545.990
490.00	550.945
500.00	555.900
510.00	560.855
520.00	565.810
530.00	570.765
540.00	575.720
550.00	580.675
560.00	585.630
570.00	590.585
580.00	595.540
590.00	600.495
600.00	605.450
610.00	610.405
620.00	615.360
630.00	620.315
640.00	625.270
650.00	630.225
660.00	635.180
670.00	640.135
680.00	645.090

Temp (K)	Heat capacity [J/(K mol)]					
0.00	0.000					
10.00	7.290					
15.00	16.910					
20.00	28.470					
25.00	41.010					
30.00	54.640					
40.00	78.950					
50.00	98.520					
60.00	115.410					
70.00	130.813					
80.00	146.331					
90.00	161.943					
100.00	177.636					
110.00	193.393					
120.00	209.199					
130.00	225.039					
140.00	240.898					
150.00	256.759 272.609 288.430					
160.00						
170.00						
180.00	304.209					
190.00	319.930					
200.00	335.754					
210.00	352.618					
220.00	369.249					
230.00	385.648					
240.00	401.814					
250.00	417.748					
260.00	433.450					
270.00	448.919					
273.15	453.744					
280.00	464.157					
290.00	479.161					
298.15	491.218					
300.00	493.934					
310.00	508.474					
320.00	522.950					
330.00	536.819					
340.00	554.370					
350.00	572.460					
360.00	584.600					
370.00	601.010					
380.00	620.303					
390.00	633.032					
400.00	645.762					
410.00	658.492					
420.00	671.221					
430.00	683.951					
440.00	696.680					
450.00	709.410					
$458.50 \ (T_{\rm g})$	720.230					

Table 53: Recommended heat capacity data for molten poly (oxy-1,4-phenylene-sulfonyl-1,4-phenylene-oxy-1,4-phenylene-(1-methylidene)-1,4-phenylene)

Temp	Heat capacity
(K)	[J/(K mol)]
458.50 (T _g)	822.712
460.00	824.065
470.00	833.082
480.00	842.098
490.00	851.115
500.00	860.132
510.00	869.149
520.00	878.166
530.00	887.182
540.00	896.199

Heat capacity of phenylene polymers

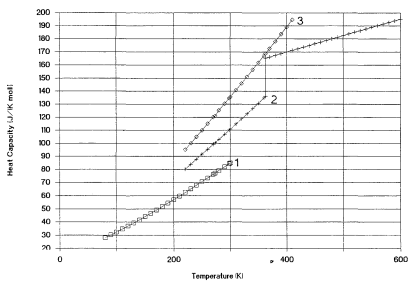


Fig. 8. Recommended heat capacities for several polymers: 1. poly(p-phenylene), 2. poly(thio-1,4-phenylene), and 3. poly(p-xylylene).

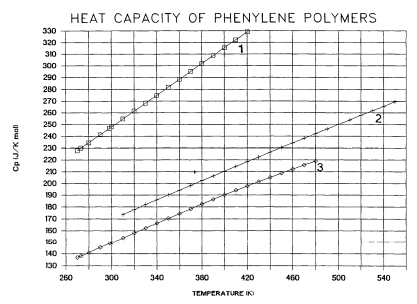


Fig. 9. Recommended heat capacities for several polymers:
1. poly[oxy-2,6-bis (1-methyl ethyl)-1,4-phenylene],
2. poly (oxy-3-bromo-2,6-dimethyl-1,4-phenylene), and 3. poly (oxy-2,6-dimethyl phenylene).

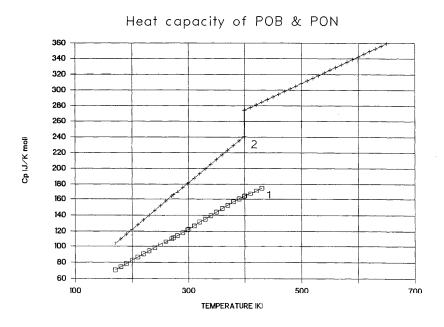


Fig. 10. Recommended heat capacities for two polymers below and above the glass transition: 1. poly (4-hydroxybenzoic acid), and 2. poly (2,6-hydroxynaphthoic acid).

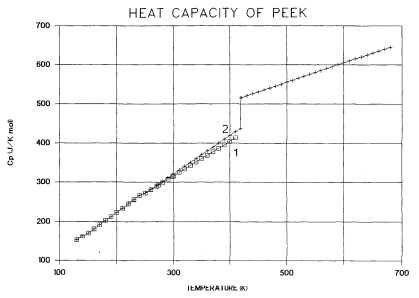


Fig. 11. Recommended heat capacities for 1. crystalline, and 2. amorphous *poly* (oxy-1,4-phenylene-oxy-1,4-phenylene).

HEAT CAPACITY OF POLYSULFONE

Fig. 12. Recommended heat capacities for amorphous poly[oxy-1,4-phenylene-sulfonyl-1,4-phenylene-oxy-1,4-phenylene-(1-methylidene)-1,4-phenylene].

TEMPERATURE (K)

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Appendix

ATHAS table of thermal properties of linear macromolecules^a

Polymer $T_g(K)$	$\Delta C_p^{\ b}$	T _m °(K)	$\Delta H_{ m f}^c$	SHG d	S _o e	Θ_1	θ ₃	N ^f	C_p^{g}
Polyethylene (c) – (a) 237 PE (CH ₂ -)	_ 10.5(1)	414.6	4.11	× ×	0 3.0	519 519	158 80	2 2	0.1–410 0.1–600
Polypropylene (c) – (a) 270 PP (CH ₂ – CHCH	- 19.2(2)*	460.7 -	8.70 -	×	0 5.2	714 633	91 78	7 7	10.0–460 10.0–600
poly (2-Methyl-1,3- (a) 278 PMP (CH ₂ – CCH	pentadiene) 34.3(?)	-	_	?	?	?	?	?	230–320
poly (1-Butenylene (c) (a) 171 PBUT cis (CH=C	27.2(3)*	284.7 - ·)	9.20 -	× ×	0 18	589 589	87 ?	8 8	30–171** 171–350
poly (1 Butenylene (c) – (a) 190 PBUT trans (CH=	28.0(3)*	437 - ₂ -)	3.73 ^h	×	0 16	599 599	95 ?	8	30–180** 360–500
poly (Vinyl fluorida (c) – (a) 314 PVF (CH ₂ – CHF	- 17.0(2)*	503.2	7.54 	× ×	0 9.4	440 440	105	4 4	80-314** 480-530i
poly (Vinylidene fl (c) – (a) 212 PVF2 (CH ₂ -CF ₂	_ 21.2(2)*	483.2 —	6.70 -	×	0 5.1	346 346	66 ?	4 4	5-212** 450-580i
Polytrifluoroethyle (c) – (a) 280 PTRIF (CHF – Cl	- 13.8(2)*	495.2	5.44 -	× ×	0 12.7	315 315	56 ?	4 4	25-280** 480-600i
Polytetrafluoroeth (c) – (a) 200 PTFE (CF ₂ -CF ₂	9.4(1)	605	4.10 ^j 	× ×	0 3.3	250 250	54 ?	2 2	0.3–280 180–700
Polyoxymethylene (c) – (a) 190 POM (O-CH ₂ -	- 28.2(2)*	457.2 -	9.79 -	× ×	0 3.4	232 232	117 ?	2 2	0.1-390 190-600
Polyoxyethylene (c) – (a) 206 POE (O – CH ₂ – C	38.2(3)* CH ₂ -)	342 _	8.66 -	× ×	0 8.1	353 353	114 ?	4 4	10–342 206–450
Polyglycolide (c) – (a) 318 PGL (CH ₂ – CO –	- 44.4(?)	506 -	11.1	× ×	0 7.3	521 521	98 ?	6 6	10–318** 318–550
poly (β-Propiolacto (c) – (a) 249 PPL [(CH ₂) ₂ – CC	- 50.4(?)	366	10.2	× ×	0 11	522 522	85 ?	8 8	10-249** 249-400

ATHAS table of thermal properties of linear macromolecules^a - Continued

Polymer $T_g(K)$	$\Delta C_p^{\ b}$	T _m °(K)	 ΔH _f °	SHG ^d	S _o e	θ ₁	Θ_3	N ^f	C_p^{8}
	<u>ыс</u> р	1 m (N)		5110		<u> </u>		14	С _р °
poly (γ -Butyrolactone) (c) – (a) 214 PBL [(CH ₂) ₃ -CO-O	- 57.5(4) -]	337.5	13.2	× ×	0 18	474 474	96 ?	10 10	10–214** 214–350
poly (δ-Valerolactone) (c) – (a) 207 PVL [(CH ₂) ₄ – CO – O	- 65.1(5) -]	331	18.2 -	× ×	0 30	502 502	101 ?	12 12	10–207** 207–350
poly (ε-Caprolactone) (c) – (a) PCL [(CH ₂) ₅ – CO – O	- -1	342.2	19.0	×	0	491	101	14	10–209**
Polyundecanolactone (c) – (a) 227 PUDL [(CII ₂) ₁₀ – CO –	- 110.6(11) O-]	365 -	38.2 _	×	0 61	528 528	105 ?	24 24	10-227** 227-400
Polytridecanolactone (c) – (a) 229 PTDL [(CH ₂) ₁₂ – CO –	_ 146.0(13) O-]	368 _	46.0 –	×	0 68	519 519	112 ?	28 28	10–229** 229–370
Polypentadecanolactone (c) – (a) 251 PPDL [(CH ₂) ₁₄ – CO –	- 168.0(15)	370.5	56.0 -	× ×	0 93	525 525	114 ?	32 32	10–251** 251–380
poly (Trimethylene adip (a) ? PTMA [O - (CH ₂) ₃ - O	?	_ 2)4-CO-]	-	?	?	?	?	?	310–360
poly (Butylene adipate) (c) – (a) 199 PBAD [O-(CH ₂) ₄ -O	– 140.0(?) – CO – (CH ₂	328.8 - 2) ₄ -CO-]	?	× ?	0 ?	514 514	(108) ?	24 24	80–199** 199–450
poly (Ethylene oxalate) (c) - (a) 306 PEOL [O-(CH ₂) ₂ -O	- 56.2(?) -CO-CO-	450 -]	23	? ×	0 ?	533 533	? 89	12 12	? 10–360
poly (Ethylene sebacate) (c) - (a) 245 PES [O-(CH ₂) ₂ -O-6	- 154.0(12)	356.2 	31.9 _	× ×	0 (22)	514 514	(125) (80)	28 28	120-245** 245-410
poly (Hexamethylene sel (a) ? PHMS [O-(CH ₂) ₆ -O	?	_ 2)8 – CO –]	-	?	?	?	?	?	340400
poly (Trimethylene succi (a) ? PTMS [O-(CH ₂) ₃ -O-	?	_) ₂ -CO-]	. –	?	?	?	?	?	310–360
Nylon 11 (c) – (a) 316 NYLON11 [CH ₂) ₁₀ – CO	- 68.4(?)* O-NH-]	493	44.7 	× ×	0 78	420 420	(67) ?	24 24	230–321** 470–550
Nylon 12 (c) – (a) 314 NYLON12 [(CH ₂) ₁₁ – C	_ 74.3(?)* CO – NH –]	500	48.4 -	× ×	0 82	455 455	(67) ?	26 26	230–310** 460–540

ATHAS table of thermal properties of linear macromolecules* - Continued

Polymer	T _g (K)	$\Delta C_{\rho}^{\ b}$	T _m °(K)	$\Delta H_{ m f}^c$	SHG ^d	S _o e	Θ_1	θ ₃	N ^f	$C_{\rho}^{\mathbf{g}}$
Nylon 6,9 (c) (a) NYLON6	331	_ 109.5(?)* ₂) ₆ – NH – Co	500 - O-(CH ₂) ₇ -C	69 - O-]	× ×	0 114	579 579	(84)	34 34	230–332** 500–590
` '	- 323 1186		506 - CO – (CH ₂) ₈ – C	71.7 _ CO-]	× ×	0 120	543 543	(84) ?	36 36	230–322** 510–590
Nylon 6,1 (c) (a) NYLON6	_ 319	_ 141.4(?)* I ₂₎₆ – NH – C	520 - CO - (CH ₂) ₁₀ -	80.1 - CO-]	× ×	0 124	533 533	(84) ?	40 40	230–319** 319–600
(c) (a)	hyl siloxane) - 135 - Si(CH ₂ CH ₃)	_ 29.2(2) ₂ -]	282.7	1.84 ^k 	×	0 8.4	480 480	87 ?	14 14	10-135** 135-360
	ne ¹ - 373 - CHC ₆ H ₅ -)	- 30.8(1+1)	516.2	10.0	× ×	0 4.4	284 284	110 48	6 6	? 0.1–600
(a)	uorostyrene 384 – CH(C ₆ H ₄ F)	33.3(1+1)	_	-	×	?	284	(48)	6	130–384
(a)	lorostyrene 406 2 – CH(C ₆ H ₄ Cl	31.1(1+1))-]	_	-	×	?	284	(48)	6	300–550
(a)	omostyrene 410 2 – CH(C ₆ H ₄ Br	31.9(1+1) ·)-]	_	-	×	?	284	(48)	6	300–550
` '	lostyrene 424 – CH(C ₆ H ₄ I) -	37.9(1+1) -]		-	×	?	284	(48)	6	300–550
(a)	ethylstyrene 380 ₂ – CH(C ₆ H ₄ C	34.6(1+1) [H ₃)-]	_	-	×	?.	284	(48)	6	300–550
(a)	ethacryloyloxyb 316 A [CH ₂ – CCl	60.0(?)		-	?	?	?	?	?	10–310
(a) `	ethyl itaconate 377 H ₂ – C(CH ₂ CC	54.2(?)	_ OCH3)-]		?	?	557	(67)	20	110-450
(a)	Propyl itacona 304 I ₂ – C(CH ₂ CO	57.8(?)	_ H₃)(COOCH₂C	_ CH ₂ CH ₃) –]	?	?	428	(67)	28	110-410
(a) 1	Heptyl itacona 172 ^m CH ₂ – C[CH ₂ C	43.96(?)	_ H₃][COO(CH₂]	_)6CH3] —	?	?	582	(67)	44	110–170
(a) 1	Octyl itaconate 78 ^m CH ₂ – C[CH ₂ C	95.76(?)	_ H₃][COO(CH₂)	_)7CH3]	?	?	518	(67)	48	110–170

ATHAS table of thermal properties of linear macromolecules^a - Continued

Polymer $T_g(K)$ ΔC_p^b	T _m °(K)	$\Delta H_{\mathrm{f}}^{c}$	SHG ^d	S,e	Θ_1	θ ₃	<i>N</i> ^f	C_p^{g}
poly (di-n-Nonyl itaconate) (a) 187 ^m 177.6(?) PDNI - CH ₂ - C[CH ₂ COO(CH ₂)8	_ CH ₃][COO(C	_ H ₂) ₈ CH ₃]−	?	?	589	(67)	52	110–180
poly (di-n-Decyl itaconate) (a) ? ? PDDI - CH ₂ -C[CH ₂ COO(CH ₂) ₉ 0	_ CH₃][COO(C	– H₂)9CH₃] <i>–</i>	?	?	585	(67)	56	110–200
poly (Dicyclo octyl itaconate) (a) 390 ^m 67.5(?) PDCYOI – CH ₂ – C[CH ₂ COO(CH	_ [COO(CF	_ H ₂) ₈] —	?	?	?	?	?	110–440
poly(Dicyclo decyl itaconate) (a) ? ? PDCYDI - CH ₂ -C[CH ₂ COO(CH	_ [COO(Cl	 H ₂) ₁₀]	?	?	?	?	?	110–280
poly (Dicyclo dodecyl itaconate) (a) ? ? PDCYDDI -CH ₂ -C[CH ₂ COO(C	_ CH2)20][COO(9	- CH2)20] -	?	?	?	?	?	100–280
$poly(p ext{-Phenylene})$ (c) : (a) ? ?(0+1) PPP (C ₆ H ₄ -)	>1000 _	?	× ?	0 ?	544 544	(54) (40)	3 3	80–300** ?
poly (Thio-1,4-phenylene) (c) (a) 363 29.2(0+1) PTP (S-C ₆ H ₄ -)	593) –	8.65 -	× ×	0 (4.2) ₅ 66	566 (40)	(54) 5	5	220–360** 360–600
poly(p-Xylylene) (c) (a) 286 37.6(1+1) PPX $(CH_2 - C_6H_4 - CH_2)$	700)* –	10.0 ⁿ	× ?	0 ?	562 562	(54) (40)	7 7	220–410** (286–410)
poly (Oxy-2,6-dimethyl-1,4-phenylen (c) (a) 482° 31.9(1+1) PPO [O-C ₆ H ₂ (CH ₃) ₂ -]	580	5.95 -	? ×	0 (8)	.564 564	(54) (40)	5 5	80–482** 482–570
poly (Oxy-3-bromo -2,6-dimethyl-1,4- (a) 559 18(?) – PPBO [O – C ₆ HBr(CH ₃) ₂ –]	phenylene) –	?	?	?	?	?		310-559
poly [oxy-2,6-bis (1-methylethyl)-1,4- (c) 426 17.6(?) PPPRO [O - $C_6H_2(CH_2CH_2CH_3)_2$ -	? ?	?	- -	?	?	?	?	270–426**
poly (Ethylene terephthalate) (c) (a) 342 77.8(4+1) PET $[O - (CH_2)_2 - O - CO - C_6H_4$		26.9 -	×	0 22	586 586	54 44	15 15	1.0–10 1.0–590
poly (Butylene terephthalate) (c) (a) 248 ^p 107(6+1) - PBT [O - (CH ₂) ₄ - O - CO - C ₆ H ₄ -	518.2 - -CO-]	32.0 ×	× (10)	0 542	542 (40)	(54) 19	19	150-310** 248-570
poly (4-Hydroxybenzoic acid) (c) (a) 434 34.0(1+1) PHBA (O-C ₆ H ₄ -CO-)	_ 	q -	x x	0 ?	823 823	(54) (25)	7 	170–434** –
poly (2,6-Hydroxynaphthoic acid) (c) – – (a) 399 33.5(1+1) PHNA (O – C ₁₂ H ₈ – CO –)	- ı -	r -	× ×	0 ?	640 640	(54) (27) 9	9	170–399** 399–650

ATHAS table of thermal properties of linear macromolecules^a - Continued

Polymer	$T_g(K)$	$\Delta C_p^{\ b}$	T _m °(K)	ΔH _f °	SHG⁴	S _o e	Θ_1	θ ₃	N ^t	C_{ρ}^{8}
poly (Ethy	dene-2,6-nar	ohthalene dicar	boxylate							
(c) `	- '		610	25.0	×	0	600	(54)	17	220-390**
(a)	390	81.6(4+1)	_	_	×	(10)	600	(30)	17	390-600
PÉN [O-	$-(CH_2)_2-C$	$-CO-C_{12}H_8$	-CO-]							
poly (4,4'-)	Isopropylide	ne diphenylene	ecarbonate)							
(c)	_	-	608.2	33.6	×	0	569	(54)	14	?
(a)	424	48.8(2+2)		_	×	(25)	569	40	14	0.4-750
PC [C ₆ H ₄	$-C(CH_3)_2$	-C ₆ H ₄ -O-C	0-0-]							
poly (Oxy-	1,4-phenyle	ne-oxy-1,4-pher	ylene-carbo	onyl-1,4-pheny	lene)					
(c)	_	_	668.2	37.4	×	0	560	(54)	15	130-419**
(a)	419 ^s	78.1(1+3)	-	_	×	(17)	560	(40)	15	419-680
PEEK [O	- C ₆ H ₄ - O	-C ₆ H ₄ -CO-	$C_6H_4-]$							
poly (Oxy-		ne-sulfonyl-1,4-	phenylene-c	oxy-1,4-pheny	lene-(1-methy	lidene)-				
(a)	459	102.5(?)	_	_	?	?	708	70	26	10-540
PBISP IC	-C4H4-S	$O_2 - C_6 H_4 - O_7$	- C6H4 - C(C	CHab - CaHa-	-1					

- a. This table includes all the new and updated data collected and measured by Advanced Thermal Analysis Laboratory data bank as of August 31, 1989. The detailed table for all the available polymers in the literature is available with us. Please correspond with us about improvements, new data, errors, etc. In the table, 'a' represents amorphous, and 'c' represents 100% crystalline and '**' represents heat capacity of semicrystalline polymers. The abbreviations used for the polymer names are computer file names. The structure of the repeating unit enclosed within the parenthesis shows one single bond on the right side instead of two half bonds cut by a parenthesis as making this is not possible on our computer.
- b. The change in the heat capacity in J/(K mol) at T_g is as available in the ATHAS recommended experimental data bank. ** represents that the change in the heat capacity is derived from the difference in the experimental liquid C_p and C_p calculated using ATHAS computation scheme. The first number within parenthesis refers to "small" beads. Their heat capacity increase at the glass transition temperature is about 11.3 J/(K mol). The second number refers to "large" beads. Their heat capacity increase at the glass transition temperature is double or triple that of a "small" bead. The average per "small" bead or equivalent of the listed polymers is 11.5 + 1.7 J/(K mol).
- c. The melting temperature is the "best available estimate" of the equilibrium melting temperature. Heat of fusion in the unit of kJ per mole of repeating units computed for 100% crystallinity.
- d. Thermodynamic functions, such as enthalpy, entropy, and Gibbs free energy etc. × represents that the data on these thermodynamic functions is available based on the ATHAS "recommended data."
- e. Residual entropy in amorphous state at absolute zero in units of J/(K mol).
- f. The number of skeletal vibrational modes used in the Tarasov equation with the theta temperatures of the previous two columns. The value of theta temperature in parentheses is derived from the data on polymers having similar backbone structure. Group vibrations are usually tabulated in the given references.
- g. Temperature range of recommended experimental heat capacity data. Computations of heat capacities of solids are usually carried out from 0.1 to 1000 K. For references see columns to the left.
- h. PBUT trans has an additional condis state at lower temperature. Crystal/condis crystal transition is at 356 K, and heat of transition is 7.8 kJ/mol, see Refs.(31).
- i. Between $T_{\rm g}$ and $T_{\rm m}$ melt $C_{\rm p}$ data is not extrapolated as liquid heat capacities of fluorinated polymers are non-linear. For details see Ref. (25).
- j. PTFE has additional room-temperature crystal/crystal/condis crystal transitions at 292 K and 303 K, combined heat of fusion is 850 J/mol, see Ref.(32)
- k. PDES has an additional condis state at a lower temperature. Crystal/condis crystal transition is at 206.7 K, and heat of transition is 2.72 kJ/mol. See Refs.(42).
- 1. For deuterated amorphous solid polystyrene and ring-only deuterated polystyrene heat capacities lead to Tarasov theta 3 and theta 1 temperatues of 55, 244 K and 49, 278 K, respectively. Thermodynamic functions S, H, and G are listed in Ref. (43).
- m. The glass transition temperature listed has been assigned to the relaxation process of the n-alkyl/cycloalkyl side groups. T_g^U has been assigned to the backbone. For details see Ref. (44-47).
- n. PPX has two lower first order transitions leading to condis crystals at 504 K and 560 K with heat of transition being 5.0 and 1.5 kJ/mol respectively, see Ref.(51).
- o. Semicrystalline PPO shows the existence of a rigid-amorphous phase which governs the thermal properties from T_g to Tm. Fusion, superheating and annealing are directly linked to this. For details see Ref. (30).
- p. The glass transition temperature of quenched PBT is 248 K and change in C_p at T_g is 107 J/(K mol). Semicrystalline PBT has a T_g between 310-325 K and change in C_p at 320 K is 77 J/(K mol), and shows the existence of rigid- amorphous fraction. For details see Ref. (38).
- q. PHBA shows a disordering transition at 616.5 K and heat of transition is 3.8 kJ/mol. For details see Ref. (58).
- r. PHNA shows a disordering transition at 614.5 K and heat of transition is 0.4 kJ/mol. For details see Ref. (58)
- s. Above T_g , poorly crystallized samples show a rigid-amorphous fraction that does not contribute to increase in heat capacity at T_g . For details see Ref. (62).