



OTIC FILE_COPY

OFFICE OF NAVAL RESEARCH

Contract N00014-86-K-0284

R&T Code 413c024---01

Technical Report No. 17

Liquid Crystalline Copoly(vinylether)s Containing 4(4')-Methoxy-

4'(4)-Hydroxy- α -Methylstilbene Constitutional Isomers as Side Groups

By

Virgil Percec and Dimitris Tomazos Department of Macromolecular Science Case Western Reserve University Cleveland, OH 44106-2699



This document has been approved for public release and sale; its distribution is unlimited.





87 8 4 0 5 2

EC.	JAITY CLAS	SIFICATION OF	THIS PAGE	REPORT DOCU	MENTATION 6	PAGE		
1a	REPORT SE	CURITY CLASSI			16 RESTRICTIVE	MARKINGS	<u></u>	
	Unclassified							
28 .					Available for publication			
2ъ	DECLASSIFICATION / DOWNGRADING SCHEDULE ERFORMING ORGANIZATION REPORT NUMBER(S)			ULE	Distribution unlimited			
4 P				5. MONITORING ORGANIZATION REPORT NUMBER(S)				
	763 	hnical Pep	port No. 17					
6a	NAME OF PERFORMING ORGANIZATION see Western Deserve University			6b. OFFICE SYMBOL (If applicable) 48566	78. NAME OF MONITORING ORGANIZATION ONR			
6c.	ADDRESS (City, State, and	I ZIP Code)		76. ADDRESS (Cit	y, State, and Z	IP Code)	
	.10)4 C10	0 Adelbert veland, Ob	5 Road A 44106		Office of Arlington	f Naval Rea n, VA - 22:	search 217	
8a.	NAME OF ORGANIZA	FUNDING / SPOI	NSORING	8b. OFFICE SYMBOL (If applicable)	9. PROCUREMENT INSTRUMENT IDENTIFICATION NUMBE		NUMBER	
97	ADDRESS (City, State, and ZIP Code)				10 SOURCE OF F	UNDING NUM	BERS	
ος.		Office of Naval Besearch FOR H - Opingr			PROGRAM	PROJECT	TASK	WORK UN
σ.	011 202	he of Nav SH. Quiney	va: Besearen V		ELEMENT NO.	NO.	NO	ACCESSIO
11	Off Puc Arl TITLE (Incl Li a - PERSONAL	H. Quine H. Quine ington, V. ude Security C and Crysta Methylsti AUTHOR(S) REPORT	val Research y A 22217 Issufication) alline Copoly Ibone Constit V. Perece and	y(vinylether)s Co outional (somers 1 D. Tomazos	ELÉMENT NO. NOOU14-86 ontaining 4(4 as Side GRoup	NO K-0284 ')-Methoxy ps	NO 413c024 -4'(4)-Hydro	ACCESSION
11 12 134	Off FOC Arl TITLE (Incl LI C PERSONAL PERSONAL SUPPLEME	H. Quine ington, V. ude Security C and Crysts Methylsti AUTHOR(S) REPORT	val Research y A 22217 Tassification) alline Copoly lbone Constit V. Perece and 13b. Time FROM 10N	/(vinylether)s Co cutional Isomers 1 D. Tomazos COVERED TO	ELÉMENT NO. NOOU14-86 Ontaining 4(4 as Side GRoup 14. DATE OF REPO	NO. K-0284 ')-Methoxy ps ORT (Year, Mon	NO 413c0??4 -4'(4)-Hydro th, Day) 15. PAG	ACCESSION
11 12 134	Off FUC Arl TITLE (Incl Lin a - PERSONAL TYPE OF SUPPLEME	Nor of Nar H. Quine ington, V. ude Security C and Crysta Methylsti AUTHOR(S) REPORT	Val Research y <u>A 22217</u> <u>Jassification</u>) alline Copoly <u>Ibone Constit</u> <u>V. Percee and</u> <u>13b. TIME</u> <u>FROM</u> 'ION	/(vinylether)s Co outional Isomers 1 D. Tomazos COVERED TO	ELÉMENT NO. NOOU14-86 Ontaining 4(4 as Side GRoup	NO. K-0284 ')-Methoxy ps ORT (Year, Mon	NO 413c0?4 -4'(4)-Hydro th, Day) 15. PAG	ACCESSION
11 12 134 16	Off FOC Arl TITLE (Incl LI a - PERSONAL TYPE OF SUPPLEME	Not of National Action of National Action of National Action of the second of the seco	val Research y A 22217 Issufication) alline Copoly Ibone Constit V. Perece and 13b. TIME FROM_ NON	/(vinylether)s Co cutional isomers 1 D. Tomazos COVERED TO 18. SUBJECT TERMS	ELÉMENT NO. NOOU14-86 Ontaining 4(4 as Side GRoup 14. DATE OF REPO	NO. K-0284 ')-Methoxy ps ORT (Year, Mon	NO 413c0?4 -4'(4)-Hydro th, Day) 15. PAG	ACCESSION
11 12 13a 16	Off FUC Arl TITLE (Incl Li a - PERSONAL TYPE OF SUPPLEME FIELD	Nor of Nav H. Quine Lington, V. Ude Security C and Crysts Methylsti AUTHOR(S) REPORT NTARY NOTAT	<pre>val Hesearch y A 22217 lassification) alline Copoly lbone Constit V. Perene and 13b. TIME FROM TON CODES SUB-GROUP</pre>	<pre>/(vinylether)s Co cutional isomers d D. Tomazos COVERED TO</pre>	ELÉMENT NO. NOOU14-86 Ontaining 4(4 as Side GRoup 14. DATE OF REPO (Continue on revers thermotropic	NO. K-0284 ')-Methoxy ps ORT (Year, Mon Liquid er,	NO 413c024 -4'(4)-Hydro th, Day) 15. PAG and identify by b	ACCESSION DXy- GE COUNT Nock number) D1ymers
11 12 13a 16	Off FUC Arl TITLE (Incl Li a - PERSONAL TYPE OF SUPPLEME FIELD	No. of Nav H. Quine ington, V. ude Security C and Crysts Methylsti AUTHOR(S) REPORT NTARY NOTAT	val Research y A 22217 Issufication) alline Copoly Ibone Constit Y. Perece and 13b. TIME FROM TION CODES SUB-GROUP	/(vinylother)s Co cutional Isomers d D. Tomazos COVERED TO 18. SUBJECT TERMS Side-chain	ELÉMENT NO. NOOU14-86 ontaining 4(4 as Side GRoup 14. DATE OF REPO (Continue on revers thermotropic	NO. K-0284 ')-Methoxy ps ORT (Year, Mon Se if necessary liquid er	NO 413c024 -4'(4)-Hydro th, Day) 15. PAG and identify by t ystalline po	ACCESSION
11 12 13a 16	Off Fuc Arl TITLE (Incl Li Q - PERSONAL TYPE OF SUPPLEME FIELD	H. Quine H. Quine ington, V. ude Security C and Crysta Methylati AUTHOR(S) REPORT NTARY NOTAT COSATI GROUP (Continue on	<pre>val Hesearen y A 22217 lessification) alline Copoly lbone Constit V. Perene and 13b. TIME FROM TION CODES SUB-GROUP reverse if necessar</pre>	<pre>/(vinylother)s Co cutional isomers d D. Tomazos COVERED TO</pre>	ELÉMENT NO. NOOU14-86 Intaining 4(4 as Side GRoup 14. DATE OF REPO (Continue on revers thermotropic number)	NO. K-0284 ')-Methoxy ps ORT (Year, Mon e if necessary liquid er	NO 413c0/24 -4'(4)-Hydro th, Day) 15. PAG and identify by b ystalline po	ACCESSION DXY - GE COUNT Nock number) D1ymers
11 12 13a 16 17	Off Fuc Arl TITLE (Incl Li a - PERSONAL TYPE OF SUPPLEME FIELD	H. Quine H. Quine Lington, V. Ude Security C and Crysta Methylati AUTHOR(S) REPORT NTARY NOTAT COSATI GROUP (Continue on	<pre>val Hesearen y A 22217 lassification) alline Copoly lbone Constit V. Perece and 13b. TIME FROM TION CODES SUB-GROUP reverse if necessar</pre>	<pre>/(vinylether)s Co outional isomers d D. Tomazos COVERED TO 18. SUBJECT TERMS Side-chain y and identify by block</pre>	ELÉMENT NO. NOOU14-86 Intaining 4(4 as Side GRoup 14. DATE OF REPO (Continue on revers thermotropic number)	NO. K-0284 ')-Methoxy ps ORT (Year, Mon liquid er,	NO 413c0/94 -4'(4)-Hydro th, Day) 15. PAG and identify by b ystalline po	ACCESSION DXY
11 12 13 16 17 19	off For Arl TITLE (Incl Lif a - PERSONAL TYPE OF SUPPLEME FIELD	H. Quine H. Quine ington, V. ude Security C and Crysta Methylati AUTHOR(S) REPORT NTARY NOTAT COSATI GROUP (Continue on Artiched	<pre>val Nesearch y A 22217 lassification) alline Copoly lbone Constit V. Perece and 13b. TIME FROM TION CODES SUB-GROUP reverse if necessar .</pre>	(vinylether)s Co sutional isomers 1 D. Tomazos COVERED TO 18. SUBJECT TERMS Side-chain y and identify by block	ELÉMENT NO. NOOU14-86 Intaining 4(4 as Side GRoup 14. DATE OF REPO (Continue on revers thermotropic number)	NO. K-0284 ')-Methoxy ps ORT (Year, Mon liquid er,	NO 413c0/24 -4'(4)-Hydro th, Day) 15. PAG and identify by t ystalline po	ACCESSION DXY
11 12 134 16	Off FUC Arl TITLE (Incl Li a - PERSONAL TYPE OF SUPPLEME FIELD	H. Quine H. Quine ington, V. ude Security C and Crysta Methylati AUTHOR(S) REPORT NTARY NOTAT COSATI GROUP (Continue on Continue on	val Nesearch y <u>A 22217</u> lassification) alline Copoly lbone Constit V. Perece and 13b. TIME FROM TION CODES SUB-GROUP reverse if necessar	(vinylether)s Co sutional isomers 1 D. Tomazos COVERED TO 18. SUBJECT TERMS Side-chain by and identify by block	ELÉMENT NO. NOOU14-86 Intaining 4(4 as Side GRoup 14. DATE OF REPO (Continue on revers thermotropic number)	NO. K-0284 ')-Methoxy ps ORT (Year, Mon liquid er	NO 413c024 -4'(4)-Hydro th, Day) 15. PAG and identify by t ystalline po	ACCESSION DXY- GE COUNT Nock number) D1ymers
11 12 134 16	οrf For Arl TiTLE (Incl Li α - PERSONAL TYPE OF SUPPLEME FIELD	Mar of Nav M. Quine Lington, V. Ude Security C and Crysta Methylati AUTHOR(S) REPORT NTARY NOTAL COSATI GROUP (Continue on	<pre>val Hosearch y A 22217 lassification) alline Copoly lbone Constit V. Perece and 13b.TIME FROM NON CODES SUB-GROUP reverse if necessar .</pre>	(vinylether)s Co sutional isomers 1 D. Tomazos COVERED TO 18. SUBJECT TERMS Side-chain by and identify by block	ELÉMENT NO. NOOU14-86 Intaining 4(4 as Side GRoup 14. DATE OF REPO (Continue on revers thermotropic number)	NO. K-0284 ')-Methoxy ps ORT (Year, Mon liquid er,	NO 413c024 -4'(4)-Hydro th, Day) 15. PAG and identify by t ystalline po	ACCESSION DXY- GE COUNT Nock number) D1ymers
11 12 13 16 17 19	οrf Fuc arl TiTLE (Incl Li α - PERSONAL TYPE OF SUPPLEME FIELD	H. Quine H. Quine Lington, V. Ude Security C and Crysta Methylsti AUTHOR(S) REPORT NTARY NOTAL COSATI GROUP (Continue on	<pre>val Hosearch y A 22217 lassification) alline Copoly lbone Constit V. Perece and 13b.TIME FROM NON CODES SUB-GROUP</pre>	(vinylether)s Co sutional isomers 1 D. Tomazos COVERED TO 18. SUBJECT TERMS Side-chain ry and identify by block	ELÉMENT NO. NOOU14-86 Intaining 4(4 as Side GRoup 14. DATE OF REPO (Continue on revers thermotropic number)	NO. K-0284 ')-Methoxy ps ORT (Year, Mon ie if necessary liquid er	NO 413c024 -4'(4)-Hydro th, Day) 15. PAG and identify by t ystalline po	ACCESSION DXY- GE COUNT Nock number) D1ymers
11 12 13 16 17 19	οrf For Arl TiTLE (Incl Li α - PERSONAL TYPE OF SUPPLEME FIELD	Market Ma	val Nesearen y <u>A 22217</u> Jassification) alline Copoly <u>Ibene Constit</u> <u>Y. Perece and</u> <u>13b. TIME FROM</u> <u>ION</u> CODES <u>SUB-GROUP</u>	(vinylether)s Co cutional isomers 1 D. Tomazos COVERED TO 18. SUBJECT TERMS Side-chain y and identify by block	ELÉMENT NO. NOOU14-86 Ontaining 4(4 as Side GRoup 14. DATE OF REPO (Continue on revers thermotropic number)	NO. K-0284 ')-Methoxy ps ORT (Year, Mon ie if necessary liquid er	NO 413c024 -4'(4)-Hydro th, Day) 15. PAG and identify by b ystalline po	ACCESSION DXY- GE COUNT Nock number) D1ymers
11 12 134 16	οrf Fuc arl TiTLE (incl Li α - PERSONAL TYPE OF SUPPLEME FIELD	Market Ma	val Nesearen y <u>A 22217</u> lassification) alline Copoly lbene Constit Y. Perece and 13b. TIME FROM NON CODES sub-GROUP 	(vinylether)s Co sutional isomers 1 D. Tomazos COVERED TO 18. SUBJECT TERMS Side-chain by and identify by block	ELÉMENT NO. NOOU14-86 Intaining 4(4 as Side GRoup 14. DATE OF REPO (Continue on revers thermotropic number)	NO. K-0284 ')-Methoxy ps ORT (Year, Mon ie <i>if necessary</i> liquid er	NO 413c024 -4'(4)-Hydro th, Day) 15. PAG and identify by t ystalline po	ACCESSION
11 12 134 16 17 19	O DISTRIBU	Tion of Nav H. Quine Lington, V. Ude Security C and Crysta Methylati AUTHOR(S) REPORT NTARY NOTAL COSATI GROUP (Continue on Artschod	A 22217 Jassification) alline Copoly bone Constit Y. Perece and 13b. TIME FROM NON CODES SUB-GROUP reverse if necessar	(vinylether)s Co sutional isomers 1 D. Tomazos COVERED TO 18. SUBJECT TERMS Side-chain by and identify by block	ELÉMENT NO. NOOU14-86 Intaining 4(4 as Side GRoup 14. DATE OF REPO (Continue on revers thermotropic number)	NO. K-0284 ')-Methoxy ps ORT (Year, Mon liquid er, liquid er,	NO 413c024 -4 * (4) - Hy dro th, Day) 15. PAG and identify by t ystalline po	ACCESSION DXY- GE COUNT Nock number) D1ymers

ABSTRACT

Ethoxy vinyl ethers containing 4-methoxy-4'-hydroxy- α -methylstilbene and 4-hydroxy-4'-methoxy- α -methylstilbene constitutional isomers as side groups were synthesized by phase transfer catalyzed etherification of a mixture containing the above mentioned isomers with 2-chloroethyl vinyl ether. Cationic copolymerization of various ratios between the two constitutional isomeric monomers led to a mixture of two copolymers which were separated by fractional precipitation. One copolymer exhibits a nematic mesophase, the other exhibits two smectic mesophases.

Accesion For M NTIS CRA&L DTIC TAB [] Upannounced Justification By Distribution/ Availability Codes Avail and for Dist Special DTIC FOPY NSPECTED

prg. 1

Liquid	Crystalline	Copoly(vinylether)s	Containing
4(4')-Met	hoxy-4'(4)-Hydro	oxy-α-Methylstilbene	Constitutional
Isomers a	s Side Groups		

The second s

Virgil Percec and Dimitris Tomazos Department of Macromolecular Science Case Western Reserve University Cleveland, OH 44106, USA

SUMMARY

Bethoxy vinyl ethers containing 4-methoxy-4'-hydroxy- α methylstilbene and 4-hydroxy-4'-methoxy-a-methylstilbene constitutional isomers as side groups were synthesized by phase transfer catalyzed etherification of a mixture containing the above mentioned isomers with 2-chloroethyl vinyl ether. Cationic copolymerization of various ratios between the two constitutional isomeric monomers led to a mixture of two copolymers which were separated by fractional precipitation. One copolymer exhibits a nematic mesophase, the other exhibits two smectic mesophases.

INTRODUCTION

of liquid crystalline polymers The synthesis (LCP) containing mesogenic units which exhibit either constitutional (1. 2) or conformational (3, 4) isomerism is of current laboratory. interest to our The copolymerization of monomer-pairs containing **mesogenic** units which exhibit constitutional isomerism is of interest both because it can be used to tailor phase transitions (1) and because it can provide qualitative information on the degree of decoupling of the side-groups from the main chain (2).

This paper presents our first results on the synthesis and cationic copolymerization of ethoxy vinyl ethers containing 4-methoxy-4'-hydroxy-a-methylstilbene and 4-hydroxy-4'-methoxy -a-methylstilbene constitutional isomers as side-groups.

EXPERIMENTAL

Materials

All the reagents were commercially available and were used as received unless otherwise specified. Methylene chloride used in the polymerization reaction was purified by conventional techniques used to achieve the purity required for carbocationic polymerization. 4,4'-Dihydroxy- α -methylstilbene (HMS) was synthesized and purified as previously reported (5). Its purity was higher



Figure 1. 200 MHz⁻¹H-NMR spectra of two representative sample: of MHMS ((CD₃)₂CO, TMS)

than 99.5% (HPLC). 4(4')-Methoxy-4'(4)-hydroxy- α -methylstilbene (MHMS) was synthesized as previously reported (1). Its composition was determined by 200 MHz ¹H-NMR spectroscopy (Fig. 1).

Synthesis of Monomers and Polymers

Scheme 1 describes the synthesis of monomers and polymers. Synthesis of Vinyl Ethers (VE)

Ethoxy vinyl ethers were prepared by a general procedure previously used for the synthesis of various vinyl ethers (6). An example follows. To a stirred mixture containing MHMS (2.0 g, $8.3x10^{-3}$ mole), 2-chloroethyl vinyl ether (2.1 ml, $2.1x10^{-1}$ mole), NaOH (0.42 g, $1.1x10^{-3}$ mole), toluene (15 ml) and DMSO (1 ml) and heated at 80° C, was added tetrabutylammonium hydrogen sulfate (TBAH, 0.1 g, $2.9x10^{-4}$ mole). The temperature was increased to 95° C and the reaction mixture was stirred overnight. After cooling to room temperature, the reaction



TABLE 1: CATIONIC COPOLYMERIZATION OF VINYL ETHERS BY BF .ET 0 SOLVENT, CH CL ; POLYMERIZATION TIME, 1 HR.

MONOMER AND COMPOSITION	[M] (MOLE/L)	3 (11.10 (MOLE7L)	POLYMERIZATION TEMP.(°C)	Conv.	-3 Mn.10	Mw/Mn
VE-1(x/y=1/2)	0.147	7.4	-10	88.0	3.7	1.97
VE-2(x/y=3/1)	0.147	7.4	-20	88.0	4.2	1.95
VE-3(x/y=1.3/1)	0.141	7.4	-10	83.3	3.2	2.16

ALC: NO.

mixture was washed with NaOH 10% aqueous solution, water, dried over MgSO₄ and the solvent was evaporated in a rotavapor. The obtained solid was recrystallized once from methanol and once from heptane to yield 1.65 g (64%) of white crystals. Purity was higher than 99.0% (HPLC). DSC traces of two different monomer compositions are presented in Fig. 2. 200 MHz ¹H-NMR (CDCl₃, δ , TMS, ppm): 2.24 (s, -CH₃), 3.84 (s, CH₃O-), 4.07 (t, -CH₂O-), 4.23 (t, -CH₂OPh), 4.10 and 4.30 (m, CH₂=), 6.50 - 6.60 (d of d, -CH=), 6.71 (s, Ph-CH=), 6.89 -7.47 (m, 8 aromatic protons).

Polymerizations

の一部である

The copolymerization of the monomers was performed in methylene chloride by using BF₃.Kt₂O as initiator. Experimental details are as previously reported (6). Table I presents the polymerization results. Techniques

The experimental techniques used in the characterization of monomers and polymers were previously described (3, 5).

RESULTS AND DISCUSSION

Figure 1 presents the NMR spectra of two different compositions of MHMS. MHMS-1 (less soluble than MHMS-2) has a 1/2 mole ratio between the x and y isomers (i.e., x/y=1/2), while MHMS-2 has x/y=3/1. The DSC thermograms of the corresponding vinyl ethers are shown in Figure 2. Both mixtures exhibit two meltings due to the two individual isomers present in the mixture, and a monotropic nematic mesophase.

The DSC thermograms of three different copolymers (compositions are available on the figure) show in each case a

glass transition temperature and three liquid crystalline transitions. With the exception of the cooling scan of PVE-1 and heating scan of PVE-2, it is difficult to separate the enthalpy changes associated with each transition in part. Nevertheless, an inspection of the heating and cooling thermograms, shows that the second LC transition from the heating scan is less suppercooled on the cooling scan than the last transition. The enthalpies associated with these two transitions (Fig. 3 cooling scans) would suggest two nematic mesophases.

However, the GPC traces of all three polymers exhibit a bimodal molecular weight distribution. The two fractions of these polymers can be separated by precipitation in acetone (giving PVE-i with $\overline{M}n$ of about 7,000) and by evaporating the acetone layer (giving PVB-s with $\overline{M}n$ of about 2,000 - 3,000). Figure 4 presents typical GPC traces for the unfractionated and fractionated PVE-2. A reinvestigation by DSC of the PVE-i



Figure 3. Heating (left side) and cooling (right side) DSC therm grams of unfractionated PVE. Enthalpy changes are in kcal/mru or mole in all figures.



1.0.0.0.0

į



Figure 5. DSC thermograms of the fr-Figure 7. DSC thermograms of the actionated samples of PVE-2.

PVE-2s as a function of sample thermal history.

and PVE-s shows clearly that all three copolymers consist of a mixture of two copolymers (Fig. 5). The low molecular weight fraction (PVE-s) is always nematic. The high molecular weight fraction (PVE-i) exhibits two smectic mesophases. Although the enthalpy change associated with the isotropization of the snectic mesophase (Fig. 5, PVE-2i) is lower than the expected values. the optical polarization microscopy reveals a texture which is characteristic for smectic mesophases i.e., batonnets (Fig. 6). This result is not unexpected since we have previously reported that smectic mesophases obtained from copolymers containing constitutional isomeric mesogens present low **isotropization** enthalpies (2). Optical polarized micrographs of the low molecular weight fractions exhibit textures characteristic for nematic mesophases (Fig. 6).

An additional interesting behavior of the low molecular weight fractions is presented in Fig. 7. As obtained by solvent evaporation, the polymer presents two meltings (Fig. 7A). The subsequent heating and cooling scans present only the nematic mesophase (curve B). This is due to the kinetically controlled crystallization process. On annealing at 70°C, the transitions аге polymer crystallizes, and the melting overlapping the isotropization temperature (curves C, D). Nevertheless, when the polymer is stirred in acetone and the acetone insoluble fraction does not undergo filtered. crystallization anymore. The acetone soluble fraction contains oligomers which do not exhibit liquid crystallinity even after quenching, but it crystallizes very fast. Its Mn is always about 1,000.



Figure (. Typical optical polarization micrographs (magnification, 300x) of the nematic texture exhibited by PVE-2s after extraction with acetone: A) after 1 second at 78°C; B) after 1 min at 78°C; C) after 3 min at 78°C; and D) the smectic texture exhibited by PVE-2i after 2 hr at 112°C on cooling scans.

Although at the first site this copolymerization system looks unusual, in our oppinion it does not represent an unexpected result. It is well documented that carbocationic polymerization induced by undefined initiators like $BF_3.Et_2O$ can give rise to multiple initiating systems which initiate and propagate with different rate constants (i.e., at least ion-pairs and free ions) (7). Two sets of rate constants are certainly going to give rise to two different sets of reactivity ratios for the same monomer pair, and therefore, a

single copolymerization experiment can provide two copolymers with different compositions. In this particular case, one composition gives a nematic copolymer while the other composition gives a smectic copolymer. Since smectics are not miscible with nematics they can be easily separated by additional fractionation. This supposition requires experiments, eventually with two different individual constitutional isomers, since 'H-NMR spectroscopy can not discriminate between the two isomers of the disubstituted HMS.

Last but not least, it is interesting to mention that the copolyacrylates and the copolymethacrylates containing the same mesogenic units and two methylenic units in the flexible spacer give only nematic mesophases, independent on their composition (2). The formation of smectic polymers by replacing the ester interconnecting group with an ether one, supports our previous statement that the degree of decoupling of the side-groups from the main chain is not dependent on the spacer length only, but also on the nature of the backbone-spacer interconnecting group, and on the nature of the polymer backbone (6, 8).

ACKNOWLEDGEMENTS

Financial support from the Office of Naval Research is gratefully acknowledged.

REFERENCES

- 1. V. Percec, J. M. Rodriguez-Parada, C. Bricsson and H. Nava, Polym. Bull., <u>17</u>, 353(1987)
- 2. V. Percec, C. S. Hsu and D. Tomazos, J. Polym. Sci: Part A: Polym. Chem., submitted
- 3. C. S. Hsu, J. M. Rodriguez-Parada and V. Percec, Makromol. Chem., <u>188</u>, 1017(1987)
- 4. B. Hahn and V. Percec, Macromolecules, in press
- 5. V. Percec, H. Nava and H. Jonsson, J. Polym. Sci: Part A: Polym. Chem., <u>25</u>, 1943(1987)
- 6. J. M. Rodriguez-Parada and V. Percec, J. Polym. Sci: Part A: Polym. Chem., <u>24</u>, 1363(1986)
- 7. J. P. Kennedy and R. Marechal, "Carbocationic Polymerization", J. Wiley, New York, 1982
- 8. C. S. Hsu and V. Percec, Makromol. Chem., Rapid Commun., in press

