PREPARATION AND NUCLEOPHILIC SUBSTITUTION OF (E)-1-BROND-2-PHENYLSULFONYL-2-ALKENES AND 3-ACETOXY-2-PHENYLSULFONYL-1-ALKENES

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Summary - Vinylphenylsulfone reacts with aldehydes to yield sulfonylated secondary allylic alcohols which are converted to either primary allylic bromides, or secondary allylic acetates. Both react highly regionelectively with lithium cyanocuprates, or enolates.

We have recently reported the multicoupling ability of 3-bromo 2-ter-butylsulfonyl propene, which may be used either as an a^2/a^2 synthon², or as a d^2/a^2 synthon³⁻⁴ (in the presence of Zinc) (Scheme 1)

Scheme 1

$$+ so_{2}$$

$$+ so_{3}$$

It was of great interest to test the generality of this synthetic approach by extending this study to the case of higher homologs of substrate 1, namely compounds of type 2 and 3

where regionalectivity of attack has to be accounted for, since nucleophilic attack of $\frac{2}{2}$ or $\frac{3}{2}$ may follow a $S_{N}2$ or $S_{N}2$, pathway.

In this paper $\frac{1}{2}$, we report the preparation of substrates of type $\frac{1}{2}$ and $\frac{1}{2}$, and their regioselective use as multicoupling reagents.

Preparation of 2-Phonylsulfonyl-1-alken-3-ols

The starting alcohols can be prepared easily by the basic treatment of a vinyl sulfone in the presence of an aldehyde. He used the commercially available phenyl vinyl sulfone, instead of tert-Butyl vinyl sulfone.

This scheme has been disclosed in the case of $\alpha.8$ -ethylenic ketones, esters, nitriles, using 10% of 1,4-diezebicyclo [2.2.2] octane (DABCO) and we have checked that it may apply in our case.

At 25°, the reaction is slow, the more so if the aldehyde is q-substituted (pivalaldehyde reacts to the extent of 10% after 150 days) but in a number of cases the reaction can be used preparatively (see table 1). The yield depends mainly on the rate competition between the hydroxy alkylation, and the degradation of the aldehyde (aldols, polymers). With propionaldehyde, the hydroxy sulfone could not be entirely separated from such oligomers, by column chromatography. Among the various catalysts that we tried, DABCO proved to be the best, 1,8-diazabicyclo-[5,4,0] undec-7-ene(DBU) is too basic and leads to polymerisation. Heating (130°, sealed tube) speeds up the reaction but yields are lower by 10-20%. Extension of this reaction to vinyl pnenyl sulfoxide led to no reaction.

Table 1 - Synthesis of allylic alcohols from 4, RCHD, and 10% DABCO at 25°C

RCHU	Product	Reaction time	Yield
		(weeks)	
MeCH0	<u>5a</u>	2	84
EtCHO	<u>5b</u>	2	•
PrCHO	<u>5c</u>	4	75
BuCHO	<u>5d</u>	10	79
iBuCHO	<u>5e</u>	11	81
Ph-CHO	<u>5f</u>	3	57

Crotonaldehyde gave no isolable—product, and furfural led to 20% of product after 3 weeks. The allylic alcohols are readily converted to the corresponding acetates by acetic anhydride in the presence of a catalytic amount of $BF_3-Et_20^7$ (0°, 10 min), (Scheme 2) Scheme 2

$$R = \frac{\text{Ac}_20}{\text{BF}_3.0\text{Et}_2}$$
 $R = \frac{\text{Rehe}, 2a : 94\%}{\text{Ren.Pr}, 2c : 95\%}$

and to the corresponding bromides (only the E isomer) by N.Bromosuccinimide-dimethyl sulfide in dichloromethane $^{\rm B}$ (12 h at 25°C) in high yield (see Scheme 3 and Table 2) Scheme 3

S0₂Ph

R

NBS, Me₂S

Br

$$\frac{3}{2}$$
 $\frac{3}{2}$

R

Table 2 - Synthesis of 2-phonylsulfonyl 1-bromo-2-alkenes 3 from alcohols 5 a-c-d-e (Scheme 3)

Alcohol	R	Product	Yield %
<u>5a</u>	Me	<u>3a</u>	85
<u>5a</u> <u>5c</u>	Pr	<u>3c</u>	87
5 <u>d</u> 5 <u>e</u>	n.Bu	<u>3d</u>	87
<u>5e</u>	i.Bu	<u>3e</u>	82

Nucleophilic attack of the allylic acetates (2) and browides (3)

These allylic bromides and acetates have been reacted with ketone enolates and cyanocuprates 9 , with the assumption that in both cases, an addition-elimination pathway should direct the incoming nucleophile so that an overall " $^5\text{N}^2$ " reaction should take place. This proved to be the case. Moreover, the bulky sulfonyl moiety promotes not only a good regio— but also stereoselectivity, so that compounds 6 are exclusively of E configuration (see scheme 4 and table 3).

Scheme 4

Table 3 - Mucleophilic substitution of allylic acetates 2m and bromides 3m

Entry	Substrate	Nucleophile	Product ^a		Yield %
1	24	BuCuCNLi	Σ Bu	<u>6a</u>	82
2	<u>3a</u>	BuCuCNLi	Bu £	<u>7a</u>	89
3	<u>2a</u>	(2)-C ₆ H ₁₃ -CH=CHCuCNLi	Hex	<u>66</u>	78
4	<u>3a</u>	(Z)-C ₆ H ₁₃ -CH - CH-CuCNLi	Hex	<u>76</u>	77 ^b
5	<u>2a</u>	PhCuCNLi	Ph	<u>6c</u>	84
6	<u>3a</u>	PhCuCNLi	Ph \	<u>7c</u>	96
7	<u>2a</u>	Cyclo-C ₆ H ₁₁ -CuCNMgCl	Σ o.Hex	<u>6d</u>	95
8	<u>3a</u>	Cyclo-C ₆ H ₁₁ -CuC NMg Cl	C.Hex F	<u>7d</u>	94
9	<u>2a</u>	ter-BuCuCNLi	£ t.8u	<u>6e</u>	98
10	<u>3a</u>	ter-BuCuCNLi	t.Bu ∑	<u>7•</u>	35 ^c
11	<u>20</u>	Ph-C=CH ₂	COPP	<u>6f</u>	80
12	<u>3a</u>	OLi Ph-C=CH ₂	PhCO	<u>7f</u>	56
13	<u>2a</u>	OLi odi	X	<u>69</u>	89
14	<u>3a</u>		X-it-	<u>79</u>	78

e/ $_{\rm I}$ stands for ${\rm SO_2Ph}$; b/ with 10% of ${\rm \underline{6b}}$ in the presence of BF3-OEt2-see text; c/ with 64% of ${\rm \underline{6e}}$

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Thus starting from either the allylic acetate or bromide, one can get the corresponding vinyl-sulfone, either linear or branched. Such high regio- and stereoselectivity has been described in the case of Phenyl sulfonyl cyclo pentenol derivatives 10 . However in the case of a bulky nucleophile (table 3 entry 10) such as a ter-Butyl cyanocuprate, the primary bromide $\frac{3a}{2}$ leads to 65% of $\frac{6e}{2}$ via $S_{\rm H}^2$, together with $\frac{7e}{2}$ (35%). With (Z)-1-octenyl cyanocuprate (table 3 entry 4) $\frac{3a}{2}$ gives a 50/50 mixture of $\frac{6b}{2}$ and $\frac{7b}{2}$, but in the presence of $\frac{8F}{3}$, $\frac{0Et_2^{11}}{2}$, the $\frac{5}{2}$ ratio raises to $\frac{88}{12}$.

Amines can also be used as nucleophiles: in this case, our results are in good accordance with those of Doomes et al¹² who studied enalogous nucleophilic substitutions of 1-bromo (and 1-amino)-2-(methylsulfonyl)-3-phenyl-2-propenes. We observed that anilin in excess, or its chloromagnesium amide (made from butylmagnesium chloride), both give a mixture of S_N^2 and S_N^2 products (8 and 9) which slowly equilibrate to give the S_N^2 derivative 8 (scheme 5) in yield of 78%.

Scheme 5

Br PhNH₂ excess or 1/ PhNHMgBr
$$\frac{3e}{2/H_20}$$
 8

A similar trend is observed in the case of thiolates: lithium phenyl thiolate gives exclusively the primary thiol ether $\underline{10}$ corresponding to an overall $S_{\rm N}2$ reaction.

When allowed to react with one equivalent of sodium methyl thiolate (a more nucleophilic reagent than phenyl thiolate) $\underline{10}$ leads to a 20:80 mixture of $\underline{10}$ and $\underline{10}$, thus either an S_N^2 isoperative, or two S_N^2 substitutions eventually lead to the thermodynamically more stable sulfide $\underline{10}$, the latter pathway bring more probable (scheme 6).

Scheme 6

A further proof of the latter hypothesis is brought by the reaction of <u>3a</u> with lithium pyridine-2-thiolate, which should react first by the more nucleophilic thiolate moiety, and then could undergo an intramolecular nucleophilic attack by nitrogen: from the four possible educts, only two are formed, namley one colorless (<u>12</u>) showing a methylene unit in NMR, and the yellow <u>11</u>, showing an allylic methyl group.

Each of them, isolated by preparative t. l.c. dissolved in ether, and submitted after a short time, to analytical t.l.c. shows again two spots corresponding to 11 and 12.

These results also point to an exclusive attack by an S_N2' pathway.

Finally, as an illustration of the preceding stereo- and regionalective synthese, we have prepared a skipped (Z-Z) diene according to Scheme 8 :

Scheme 8

$$S0_2Ph$$
 BF_3-Et_20
 OAc
 O

sulfone $\underline{6b}$ being reductively desulfonylated, according to Julia's procedure 13, to (Z,Z)-2,5dodecadiene 10.

In conclusion, due to the presence of the phenylsulfonyl moiety, the easily available acetates of type $\underline{2}$, or browides of type $\underline{3}$ can be substituted regionalectively, leading to (Z)-1-2 disubstituted alkenes, or to 3-substituted terminal alkenes, once the sulfonyl moiety is discarded.

EXPERIMENTAL PART -

THF and other were distilled from sodium/benzophenone. Infrared spectra were recorded on a Perkin Elmer 4576 spectrometer. Proton NMR spectra were obtained at 100NHz with a Jeol MH100 and at 250MHz with a Bruker AM250. 13 C-NMR spectra were obtained with a Jeol FX90. Chemical shifts in COCl, solution are reported in ppm relative to tetramethylsilane as an internal standard. Gas chromatography was carried out with a Carlo Erba 2150 model equiped with an OVID1 (20 m) column. Merck 60 (70-230 mesh) silica gel was used for the fash chromatography

General procedure for the preparation of 2-ter-butylsulfonyl-1-alken-3 ols $\underline{5}$ In a dry erlemmeyer, flushed with Argon are placed 3 g (17.8 mmol) of vinylphenyl sulfone in 7 ml of the freshly distilled aldehyde. 0.2 g (1.78 mmol) of dry DABCO are then added (dissolution). The erlemmeyer is stoppered and left at room temperature. Reaction is followed by T.L.C. After disappearance of the phenylvinyl sulfone, the mixture is taken up by 100 ml CH_Cl_ and successively washed with a 1MHCl solution (30 ml). The organic layer is dried over $Mg50_{\perp}$. Solvent and excess aldehyde are evaporated under vacuum, and the resulting oil is chromatographed on silica. In the case of $\underline{5a}$ we operated on a 1 mole scale, and in this case, a careful evaporation of solvent and remaining acetaldehyde under high vacuum led to a product readily used for further transformations. Spectroscopic data are collected in table 4.

Z-Phenylsulfonyl-1-butan-3-ol 5a Yield (from 17.8 mmol of vinyl sulfone) : 3.17 g (84%). Chromatography with ether : CH_2Cl_2 : hexane/ 8:70:30 gives an oil. Found: C, 56.30; H, 5.65%. Calcd. for $C_{10}H_{12}SO_3$: C, 56.58; H, 5.70%.

2-Phenylsulfonyl-1-hexen-3-ol 5c

See general procedure. From n- butanal. Obtained 3.20 g (75%) of 5c as an oil. Eluent ether : CH₂Cl₂: hexame/ 8:70:30. Found: C, 60.50; H, 6.52%. Calcd. for C₁₂H₁₆SO₃: C, 59.97; H, 6.71%.

2-Phenylsulfonyl-1-hepten-3-ol 5d

See general procedure. From n-pentanal 3.57 g (79%) of $\underline{5d}$ are obtained as an oil. Same eluent as for $\underline{5c}$.

2-Phenylsulfonyl-5-Methyl-1-hexen-3-ol 5e

See general procedure. From isovaleraldehyde. 3.66 g of <u>Se</u> are obtained as an oil (same eluent as for 5c).

2-Phenylsulfonyl-3 phenyl-1-propen-3-ol $\underline{5f}$ See general procedure. From benzaldehyde. 2.79 g of $\underline{5c}$ are obtained as an oil (same eluent as for $\underline{5c}$). M.p. 78°C . Found: C, 65.58; H, 5.10%. Calc. for C15H14SO2: C, 65.67; H, 5.14%.

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Preparation of the secondary allylic acetates 2

5 mmol of the preceding alcohols are dissolved in 4 ml acetic anhydride, and to the solution maintained at 0°C, are added 0.1 ml BF $_3$ -0Et $_2$ 0 (0.8 mmol). The mixture turns organge progressively, and the reaction is over after 10-15min. The solution is then poured in CH $_2$ Cl $_2$ Cl $_3$ Cl $_3$ Cl $_4$

Acetate of 2-phenylsulfonyl-1-butan-3-ol $\underline{2a}$ From $\underline{5a}$. 1.19 g (94%) of $\underline{2a}$ are obtained as an oil. Eluent : ether : $\mathrm{CH_2Cl_2}$: hexane/3:70:30. Found : C, 56.28 ; H, 5.60%. Calcd. for $\mathrm{C_{12}H_{14}S0_4}$: C, 56.68 ; H, 5.55%.

Acetate of 2-phenylsulfonyl-1-hexen-3-ol $\frac{2c}{5c}$ From $\frac{5c}{5c}$. 1.34 g (95%) of $\frac{2c}{5c}$ are obtained as an oil. Same eluent as for $\frac{2a}{5c}$.

Preparation of 2-phonylsulfonyl-3-bross-1-alkenes 3

from alcohols 5 a, c, d, e. A solution of T.8 ml dimethylsulfide in 14 ml CH_Cl_ is added dropwise to a stirred suspension of 3.58 g (0.02 mol) of N-bromosuccinimide in 45 ml CH_Cl_ maintained at -20°C. The temperature is allowed to raise up to 0°C and the pale yellow suspension is stirred for 15 min. A solution of 19 mmol of the alcohol 5 in 18 ml CH_Cl_ is then added. The mixture is stirred at 25° for 12 h whereby a yellow solution is obtained, which is successively washed with a saturated solution of sodium hydrogenocarbonate (2 x 20 ml) then brine (20 ml) and dried over magnesium sulfate. Solvents are evaporated under vacuum and the residue is either recrystallized, or flash chromatographed on silica. Spectroscopic data are collected in table 4.

(E)-1-Browo-2-phenylsulfonyl-2-butene $\frac{3a}{5}$ From $\frac{5a}{5}$ are obtained 4.44 g (85%) $\frac{3a}{5}$ first chromatographed eluent ether : CH_2Cl_2 : hexane/5:70:30, then recrystallized From ether. m.p. : 73°C. Found : C, 43.55 ; H, 3.95%. Calcd. for $\text{C}_{10}\text{H}_{11}\text{SO}_2\text{Br}$: C, 43.65 ; H, 4.03%.

(E)-1-Browo-2-phenylsulfonyl-2-hexana $\frac{3c}{5c}$ From $\frac{5c}{5c}$. 5.00 g of $\frac{3c}{3c}$ are obtained as an oil. Same eluent as for $\frac{3a}{5c}$.

(E)-1-Bromo-2-phenylsulfonyl-2-heptene 3dFrom 5d. 5.24 g (87%) of 3d are obtained as an oil. Same eluent as for 3a.

(E)-1-Brown-2-phenylsulfonyl-5-methyl-2-hexene 3eFrom 5e, 4.94 g (82%) of 3e are obtained as an \overline{oil} . Same eluent as for 3a.

General procedure for the reaction of cyanocuprates with allylic-acetates 2 or bromide 3

To a stirred suspension of 358 mg (4 mmol) of copper cyanide in 15 ml THF at -80°, is added slowly a solution of 4 mmol of the lithium— or magnesium organometallics 1N in other or THF. The mixture is then stirred for 30 min at temperatures in the -30,-10°C range, according to each case. A solution of 2 mmol of the acetate 2 or bromide 3 in 5 ml THF is then added slowly at -80°C. In the case of 3a and Z octenylcyanocuprate, a solution of 4 mmol BF $_2$ -OEt $_2$ (0.5 ml) in 2 ml other is added at -80°C before introducing reagent 3. The mixture is allowed to warm slowly and is followed by T.L.C. When no more starting 2 or $\frac{3}{2}$ is detected, the mixture is hydrolyzed with a saturated solution of NH $_2$ Cl (20 ml) and 1 ml concentrated ammonia. The aqueous phase is extracted with other (2 x 20 ml). The organic phase, washed with a sat. solution of NH $_2$ Cl (20 ml) is dried over MgSO $_2$. Solvents are evaporated under vacuum, and the residue is chromatographed on silica. Spectroscopic data are collected in table 5.

(E)-3-Phenylsulfonyl-2-octene <u>6a</u> From butyl lithium and <u>7a</u>. Reaction time 1 hr at $-60^{\circ} \rightarrow -15^{\circ}$ C. 0.41 g (82%) of sulfone <u>6a</u> are obtained as an oil. Eluent: CH₂Cl₂: cyclohexane/70:30. Found: C, 66.70; H, 7.90%. Calcd. for C₁₄H₁₀SO₂: C, 66.63; H, 7.99%.

2-Phenylsulfonyl-3-methyl-1-hexene 7aFrom butyllithium and 3a. Reaction time 0.5 hr at -60°C. 0.45 g (89%) of 7a are obtained as an oil. Same eluent as for 6a.

3-Phenylsulfonyl-2(E),5(Z) dodecadiene $\underline{6b}$ From 2a,and (Z)1-octenyl-lithium cyapocuprate. The latter reagent, prepared from (Z)-1- iodo-1-octene by lithium/iodine exchange and then following the general procedure, with a reaction time of 1.5 hr at -15°C, led to 0.48 g (78%) of sulfone $\underline{6b}$, separated from isomer $\underline{7b}$ during chromatography (eluent ether: $\underline{CH_2Cl_2}$: cyclohexane/1:70:30. (2)-2-Phenylsulfonyl-3-methyl-1,4 undecadiene 7b From 3a and (2):1 octenyl cuanocuprate (prepared as above) in the presence of 2 equivalents of BF_3-Et_30 added prior to the bromide 3a. Reaction time 2 hr at $-60^{\circ} \rightarrow -10^{\circ}C$. 0.47 g of 7b are obtained (77%) as an oil; Same eluent as for 6b.

(E)-2-Phenylsulfonyl-1-phenyl-2-butene $\underline{6c}$ From 2a and lithium phenyl cyanocuprate. Reaction time: 1 hr at -60° \rightarrow -45°C. 0.46 g (84%) of $\underline{6c}$ are obtained as an oil. Same eluent as for $\underline{6b}$.

2-Phenylsulfonyl-3-phenyl-1-butene $\underline{7c}$ From $\underline{3a}$ and lithium phenyl cyanocuprate. Reaction time 0.25 hr at -60°C. 0.52 g (96%) of $\underline{7c}$ are obtained (oil). M.p. : 48°C. Found : C, 70.49 ; H, 5.88%. Calcd. for $C_{16}H_{16}SO_2$: C, 70756 ; H, 5.92%.

(E)-2-Phenylsulfonyl-1-cyclohexyl-2-butane $\underline{6d}$ From $\underline{2a}$ and lithium cyclohexyl cuanocuprate. Reaction time 2 hr at -80° \rightarrow -30°C. 0.53 g (95%) of $\underline{6d}$ are obtained as an oil. Eluent CH₂Cl₂: hexane/70:30.

From 3a and lithium cyclohexyl cyanocuprate. Reaction time 1 hr at -60°C. 0.52 g (94%) of 7d are obtained as an oil. Same eluent as for 6d.

(E)-4-Phenylsulfonyl-2,2-dimethyl-4-hexene 6e

From 2a and lithium ter-butyl cyanocuprate. Reaction time 2.2 hr at -60° \rightarrow -45°C. 0.49 g (98%) of $\frac{6e}{}$ are obtained as an oil. Eluent: CH₂Cl₂: cyclohexane/70:30. Found: C, 67.00; H, 8.05. Calcd. for $C_{14}^{}H_{20}^{}SO_{2}$: C, 66.63; H, 7.99%.

2-Phenylsulfonyl-3,4,4 trimethyl-1-pentene 7e From 3a and lithium ter-butyl cyanocuprate. Reaction time 0.25 hr at -60°C. 0.17 g of $\frac{7e}{2}$ (35%) are obtained as an oil, separated from 0.32 g $\frac{6e}{2}$. Eluent CH₂Cl₂: cyclohexane/70:30.

General procedure for the reaction of lithium enclates with acetate 2s and bromide 3s

2-Phonylsulfonyl-3-cyclohexyl-1-butene 7d

To a stirred solution of 361 mg (3.6 mmol) of diisopropylamine in 7 ml THF at $-80\,^{\circ}$ C is added 3.2 ml of a 1N solution of n-Butyl lithium and the temperature is raised up to $-40\,^{\circ}$ C. After 15 min, the solution is cooled to $-80\,^{\circ}$ C and 3 mmol of the ketone in 2 ml THF are added. The solution is then stirred at $-60\,^{\circ}$ C for 0.5 hr and a solution of 2 mmol of 2a or 3a in 3 ml THF is added at $-80\,^{\circ}$ C. The reaction is followed bu t.l.c. while temperature is allowed to raise. The mixture is then hydrolyzed with a saturated NH Cl solution (20 ml). After extraction of the aqueous layer by ether 20 ml, the organic phase is washed with sat. NH Cl (20 ml), dried over MgSO and solvents are evaporated under vacuum. The residue is chromatographed by flash chromatography. See spectroscopic data in table 5.

(E)-3-Phenylsulfonyl-6-phenyl-2-hexen-6-one $\frac{6f}{hr}$ From $\frac{2a}{2}$ and acetephenone. Reaction time 2.5 $\frac{6f}{hr}$ at -60°- $\frac{1}{2}$ -10°C, 0.5 g of $\frac{6f}{2}$ (80%) are obtained. Eluent CH₂Cl₂: cyclohexene/80:20, m.p.: 86°C (CH₂Cl₂/pentane).

4-Phenylsulfonyl-3-methyl-1-phenyl-4-penten-1-one $\frac{7f}{From 3e}$ and acetophenone. Reaction time 2.5 hr at $\frac{7}{-60^\circ}$ \rightarrow -10°C. Obtained : 0.35 g (56%) of $\frac{7f}{From 3e}$ cyclohexane/80:20. m.p. : 133°C (CH₂Cl₂/pentene). Found : C, 68.50 ; H, 5.70%. Calcd. for $C_{18}H_{18}SO_3$: C, 68.76 ; H, 5.77%.

(E,E)-6-Phenylsulfonyl-1-(2,6,6-trimethyl-1-cyclohexenyl)-1,6-octadien-3-one $\underline{6g}$ From -ionone and $\underline{2a}$. Reaction time 3.5 hr at -60° \longrightarrow -25°C. 0.69 g (89%) of $\underline{6g}$ are obtained as an oil. Eluent ether: CH₂Cl₂: cyclohexene/2:70:30. Found: C, 71.38; H, 7.75%. Calcd. for $C_{23}H_{30}SO_3$: E, 71.46; H, 7.82%.

(Z)-6-Phenylsulfonyl-1-(Z,6,6-trimethyl-1-cyclohexenyl)-5-methyl-1,6-heptadien-6-one 7g From -ionone and 3a. Reaction time 1.5 hr at -60° \rightarrow -20°C. 0.60 g of 7g are obtained as an oil. Same eluent as for 6g.

2-Phenylsulfonyl-3-(N-phenylamino)-1-butene 8 and (E)-2-Phenylsulfonyl-1-(N-phenylamino)-2-butene 9 A solution of 0.40 g (4.3 mmol) of anilin in 5 ml THF is added to a stirred solution of 0.55 g of sulfone 3a (2 mmol) in 5 ml THF at -78°C. The mixture is warmed up to -50°C, and the reaction, followed by t.l.c., is over after 30 min. 10 ml of saturated NH_Cl solution and 25 ml CH_Cl, are added. The organic phase is washed with brine (20 ml), bried over NgSO, and evaporated under vacuum. Chromatography of the residue (ether : CH_Cl): hexane/2.70:30) gives 0.48 g (85%) of a 57/43 mixture of 8 (oil) and 9 (solid, m.p.: 88°C).

2-Phenylsulfonyl-1-phenylthio-prop-2-ene 10

5ame general procedure as for the reaction of enolates, from 2a and lithium phenyl thiolate (prepared from thiophenol and n-butyllithium at - 80° in THF) reaction time 1 h at -40°. 0.596 mg of 10 are obtained (98%).

m.p. : 52°C.

 3 7.40 to 8.11(m,5H(phenyl)) 27.85(d), 14.29 and 13.93 (a and e)

3(2-pyridylthio)-2-phenylsulfonylbut-1-ene 12 (and thione 11)
Same procedure as for 10. Preparative T.L.C. with ether : CH2Cl2 : cyclohexane/10:70:30 allows the isolation of $\frac{11}{11}$ (yellow) and $\frac{12}{12}$ (colorless) (oils). As they equilibrate rapidly, the NMR spectra show the caracteristic following signals: $\frac{12}{12}: \begin{array}{ll} \text{H RMN} & \text{(CDCl}_3, \text{§}): 1.62(\text{d},3\text{H}, J=7.5\text{Hz}(\text{CH}_3-\text{CH}), 4.89(\text{q},1\text{H},J=7.5\text{Hz}, (\text{CH}-\text{CH}_3)), 6.3(\text{s},1\text{H}, \text{H-C}=\text{trans to 50}_2), 6.5 to 8.1(\text{m},10\text{H}) \\ & \text{[he 13C NMR spectrum shows no signal above 160 ppm.} \\ \hline 11: \begin{array}{ll} \text{H RMN} & \text{§}: 2.04(\text{d},3\text{H},J=7.5\text{Hz},\text{CH}_3-\text{CH}=), 5.43(\text{s},2\text{H},(-\text{CH}_3-\text{CH}=)), 6.51 to 8.10 \text{ m}. \\ \hline \end{array}$

13_{C RMN} (CDC1₃,\$): 180.8 (C=5)

mixture 11 + 12 : found : C, 59.06 ; H, 5.00%. Calcd. for $C_{15}H_{15}S_{2}O_{2}N$: C, 58.99 ; H, 4.95%

(Z,Z)-2,5-dodecadiene 13 (2,2)-2,5-dodecadiene $\frac{13}{13}$ 13 According to Julia's procedure , 1 mmol of $\underline{6b}$ (0.306 g) gave 0.121 g (73%) of $\underline{10}$ (oil).

IR: (neat) 3010,2960,2920,2860,1650,1585,1465,1455,1440,1400,1380,1365,1305,1270,1225,1150,1090, 1025,965,900,740,700

Table 4 - Spectroscopic data of alcohols 5, acetates 2, bromides 3

Compound	1H NMR CDC13- Sfrom TMS	13 C RMN(CDC1 $_3$, δ from TMS	1.R.ª
a so	1.22(d,3H,J=6.4Hz(d)), 3.30,s,1H,(0H),4.45(m, 1H,(c)),6.05(s,1H,(Ha trans/50 ₂)),6.29(s,1H, (Ha cis/50 ₂)),7.4 to 8.35,m,5h(Ph)	154.44(b),139.26(e), 133.40(h),129.08(f), 127.89(g),123.78(a), 64.47(c),22.62(d)	3480,2975,2930, 1580,1445,1375, 1300,1175,1135, 1100,1075,1035, 955,915,835,750, 685,620
JON 5c	0.78(t,3H,J=6.8Hz,(f)) 1.11 to 1.68(m,4H,(d,e)) 3.39(d,1H,OH),4.44(m,1H,(c)),6.18(s,1H(Ha trans/SO ₂)),6.45(s,1H,(Ha cis/SO ₂),7.5-8.4(m,5H,phenyl)	153.68(a),139.26(g), 133.69(j),129.28(i), 128.00(h),124.81(b) 68.05(c),38.46(d),18.35 (e),15.55(f)	3500,2960,2925, 2870,1580,1445, 1380,1300,1170, 1140,1115,1080, 1025,970,915,775, 755,695
h sos f sd	0.78(t,3H,J=7Hz,(g)), 1.0 to 1.6(m,6H,(d,e,f)) 3.68(d,1H,J=4.8Hz,0H), 4.49(m,1H(c)),6.12(s,1H, Ha trans/S0 ₂)),6.50(s, 1H,Ha cis/S0 ₂)),7.45- 7.85(m,5H,ph@nyl)	153.68(b),139.29(h), 133.66(k),129.25(j), 128.00(1),124.84(æ), 68.32(c),36.08(d),27.23 22.17	3500,2950,2925, 2865,1580,1445, 1375,1300,1165, 1135,1075,955, 750,685
h Son	0.78(larged,6H,(f,g)), 1.08-1.92(m,3H,(d,e)), 3.42(d,1H,J=5Hz,0H), 4.47(m,1H,(c)),6.12(s, 1H(Ha trans/50 ₂)),6.33 (s,1H,Ha cis/50 ₂)),7.35- 7.95(m,5H,pheny1)	154.16(b),139.11(h), 133.63(k),129.19(i), 127.97(j),124.45(a), 66.53(c),47.73(e),24.37 (d),23.12 and 21.36(f,g)	3490,2960,2875, 1585,1465,1435, 1385,1370,1305, 1170,1145,1080, 960,920,770,750,

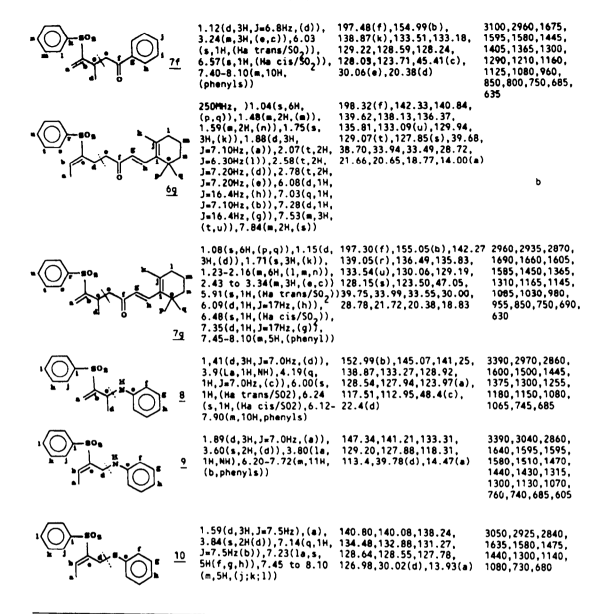
a/ as film (neat) for liquids, or KBr plates (solids)

Table 5 - Spectroscopic data of products 6 and 7, from nucleophilic addition of cuprates and enolates to 2a and 3a

Compound	¹ H NMR spectra	13C NMR spectra	1.R.ª
80	0.75-2.5(m,11H,(d-h)), 6a 1.85(d,3H,J=7.5Hz,(a)), 7.05(q,1H,J=7.5Hz(b)), 7.5-8.1(m,5H,phenyl)	142.50,140.32(c and i), 136.99,133.05(b and 1), 129.09,128.08(k and j), 36.69,28.32,26.30,22.18, 14.11, 13.90	3060,2950,2920, 2860,1640,1580, 1445,1300,1155, 1130,1080,755, 720,690
a so	0.75(t,3H,J=7.0Hz,(h)), 1.02(d,3H,J=7.5Hz,(d)), 7a 1.35(m,6H,(e,f,g),2.49 (m,1H,(c)),5.88(s,1H(Hatrans/50 ₂)),6.51(s,1H, (Hacis/50 ₂)),7.5-8.15 (m,5H,phen91)	156.27(a),139.05(i), 133.39(1),129,10(k), 128.29(j),122.19(b), 36.47,33.46,28.87, 22.28,21.75(d),13.85(h)	2960,2920,2850, 1445,1380,1300, 1175,1148,1125, 1080,950,840, 750,690,630

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\$ 60 m	0.9(t,3H,J=7.0Hz,(1)), 1.32(m,8H(h,i,j,k)), 1.89(d,3H,J=7.5Hz,(a)), 3.09(d,2H,J=7.0Hz,(d)), 4.89-5.58(m,2H,(e,f)), 7.14(q,1H,J=7.5Hz,(b)), 7.45-8.1(m,5H,(phenyl))	141.08,140.01,137.71 133.06(p),131.66,129.01, (o),128.09(n),124.48, 31.73(d),29,29,28.99, 27.29,24.46,22.61,14.06	2960,2925,2855, 1640,1585,1450, 1380,1315,1305, 1155,1135,1085, 1000,970,900,785, 725,690
70	0.87(t,3H,J=7.0Hz,(1)), 1.2(m,11H,d,h,i,j,k), 1,70(m,2H,(g)),3.45(m, 1H,(c)),5.15(m,2H(e,f)), 5.86(s,1H(Ha trans/50 ₂), 6.40(s,1H,Ha cis/50 ₂), 7.40~8.0(m,5H,pheny1)	155.56(b),139.62(m), 133.30(p),131.34,130.89, 129.01(o),128.24(n), 123.14(a),32.06(c),31.67, 29.23,28.90,27.05,22.58, 22.14(d),14.06(1)	2920,2850,1580, 1440,1310,1300, 1165,1140,1080, 745,685
65	1.77(d,3H,J=7.5Hz,(a)), 3.72(s,2H,(d)),7.11(m, 5H,(f,g,h)),7.30(q,1H, J=7.5Hz,(b)),7.5-8.1(m, 5H,(j,k,1))	140.90,139.91,139.20, 136.55,132.83,128.80, 128.18,127.97,127.82, 126.18,31.61(d),14.51(a)	3060,3030,2920, 1640,1595,1580, 1490,1480,1445, 1300,1285,1145, 1125,1080,735, 700,685,630
1 00 a 7c	1.34(d,3H,J=7.13Hz,(d)), 3.83(q,1H,J=7.13Hz,(c)), 5.82(s,1H,(Ha trans/ 50 ₂)),6.5(s,1H,Ha cis/ 50 ₂)),6.85-7.10(m,5H, (f,g,h)),7.2 à 7.7(m,5H, (j,k,1))	155.14(b),141.94,139.35, 133.04(1),120.80,128.27, 127.94,127.05,126.54, 124.51,39.48(c),22.05(d)	3068,3015,,2970, 2930,1600,1580, 1490,1445,1300, 1160,1130,960, 905,740,690,645
90 g 6d	1.0 to 1.80(la,11H,(e,f,g,h)),1.86(d,3H,J=7.5Hz,(a)),2.19(d,2H,J=7.0Hz,(d)),7.2(q,1H,J=7.5Hz,(b)),7.5-8.1(#,5H,phenyl)	140.93,140.42,138.28, 132.94(1),128.95(k), 127.88(j),36.97(e),33.64 33.10,26.21,26.13(h), 14.66(a)	3030,2920,2850, 2260,1640,1585, 1480,1305,1215, 1180,1160,1140, 1120,1085,1000, 910,760,735,690, 620
1 To To To	0.93(d,3H,J=7.5Hz,(d)), 0.90 à 1.98(la,m,11H, (e,f,g,h)),2.34(m,1H, (c)),5.82(s,1H,(Ha trans/50 ₂)),6.48(s,1H, (Ha cis/50 ₂)),7.5-8.1(m 5H,(phenyl7)	155.53(b),139.14(i), 133.36(1),129.04(k), 128.36(j),122.67(a), 41.98(c),39.15(e),31.10, 28.96,26.13,18.44	2915,2825,1450, 1390,1315,1305, 1175,1150,1120, 1080,1025,1000, 950,895,865,840, 765,750,690,650
1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	0.99(s,9H,(f)),1.86(d, 3H,J=7.5Hz,(a)),2.31(s, 2H,(d)),7.14(q,1H, J=7.5Hz,(b)),7.5-8.1(m, 5H,(phenyl))	142.21,141.22,141.14, 132.76d(j),128.94(i), 127.70(h),38.52(e),33.44 (d),30.54(f),16.12(e)	3060,2950,1640, 1580,1475,1445, 1300,1195,1080, 760,750,735,685
3 - 80 e	0.87(s,9H,(f)),0.90(d, 3H,J=7.5Hz,(e)),2.55(q, 1H,J=7.5Hz,(c)),6.0(s,1H (Ha trans/SO ₂),7.5-8.10 (m,5H,(phenyI))	155.53(b),139.35(g), 133.30(j),129.04, ,128.47(h,i),123.98(a), 42.25(c),33.96(e), 27.71(f),18.20(d)	3060,2950,1640, 1580,1475,1445, 1300,1145,1120, 1080,1065,750, 685,620
5 <u>6f</u>	1.87(d,3H,J=7.5Hz,(e)), 2.73(m,2H,(d)),3.21(m, 2H,(e)),7.14(q,1H, J=7.5Hz(b)),7.30-8.05 (m,5H,(phenyl))	198.12(f),140.61,139.45 (k),138.38,136.17,133.04 (n),129.05(m),128.40, 127.77,37.13(d),20.47(e), 13.92(a)	3060,2960,2930, 2910,2850,1970, 1895,1815.1725. 1700,1585,1575, 1480,1445,1405, 1375,1360,1340, 1300,1290,1275, 1215,1200,1185, 1150,1130,1085, 1015,1005,970, 940,925,860,850, 780,765,755,730,690,650



- (a) as film (neat) for liquids, or KBr plates for solids
- (b) not recorded

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