S0040-4039(96)00468-6

Stereoselective Conjugate Addition of Carbon Nucleophiles to Chiral (E)-Nitroalkenes Bearing a γ -Stereocenter. Origins of the Observed anti Selectivity.

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Abstract. The reaction between different organometallic carbon nucleophiles and chiral (E)-nitroalkenes has been studied. Anti selectivities have been observed in all cases, SCF-MO calculations suggest that the observed stereocontrol relies on the anti disposition of a donor group (methyl or hydrogen) with respect to the C-C bond in formation. Copyright © 1996 Elsevier Science Ltd

Conjugate addition of organometallic carbon nucleophiles to electron deficient olefins has proven to be a very useful method for the synthesis of a wide range of chemically significant structures¹. In particular, the asymmetric version of this reaction has received great attention². Nitroalkenes are among the most convenient substrates for this reaction³, since the electron withdrawing nitro group induces a high electrophilic character to the olefinic moiety, and the nitro group is in turn very versatile⁴. However, the number of reports dealing with the interaction of chiral nitroalkenes with achiral nucleophiles is scarce⁵. In this paper we report our experimental and computational results on the 1,2-asymmetric induction observed in the reaction of nitro-

Scheme 1

alkenes⁶ 1a,b with several organometallic nucleophiles (Scheme 1). We have collected in Table 1 the most significant data among the different metals and reaction conditions tested. From these results it can be concluded that nitroalkene 1a is quite efficient as Michael acceptor both in terms of chemical yield and

stereoselectivity. For example, reaction of 1a with phenyl lithium takes place with a diastereomeric excess of 60% (Table 1, entry c). This stereocontrol was slightly improved by using an excess of lithium chloride (entry d). However, the best result for 1a was obtained using lithium diphenylcuprate as the nucleophile (entry e). The stereoselectivity of methyl nucleophiles was comparatively lower (entries g-1). Nitroalkene 1b induces in general a lower stereocontrol, and the best result was again obtained using lithium diphenyl cuprate (entry n) and lithium dimethyl cuprate (entry q). Therefore, according to our results, the best reagents for this kind of transformation appear to be lithium organocuprates of type LiR₂Cu. The structure of compound anti-2a (relative configuration only) was stablished by X-Ray diffraction analysis⁷ (Figure 1), and the stereochemistry

Table 1. Reaction of (E)-nitroalkene **1a,b** with several nucleophiles.

Entry	Product	Reagent ^a	Time	Yield	anti:syn ^c
			(h)	(%)b	
a	2a	PhMgBr (1.2)	1.5	70	77:23
b	2a	PhMgBr/CeCl ₃ (2.0)	4	75	83:17
С	2 a	PhLi (1.5)	3	90	80:20
ď	2a	PhLi/LiCl (1.5:5)	3	95	85:15
e	2a	PhLi/CuI (3.2:1.6)	2.5	86	91:9
f	2a	PhLi/CuCN (5.1:2.6)	6	90	77:23
g h	2 b	MeMgBr (2.0)	3	90	59:41
h	2 b	MeMgBr/CeCl ₃ (2.0)	20	71	57:43
i	2 b	MeLi (1.5)	5	76	76:24
j	2 b	MeLi/LiBr (1.5)	7	81	70:30
k	2 b	MeLi/CuI (3.2:1.6)	7	60	65:35
l	2 b	MeLi/Me ₂ .CuBr (1.0:1.0)	7	13	55:45
m	3a	PhLi (1.5)	2	90	50:50
n	3a	PhLi/CuI (3.2:1.6)	2.5	95	91:9
0	3 b	MeLi (1.5)	6	38d	60:40
p	3 b	MeLi.LiBr (1.5)	7.5	30d	50:50
_ <u>p</u>	3 b	MeLi/CuI (3.2:1.6)	7.5	90	72:28

^aNumbers in parentheses are the relative quantities with respect to 1 eq. of substrate. All runs were conducted at -78°C under an argon atmosphere. ^bYields of products 2 and 3 after isolation by flash chromatography (Silicagel 230-400 mesh, AcOEt/Hx 1:20). ^cIsomeric ratio determined by 300 MHz ¹H-NMR on the crude reaction mixture. ^dFormation of addition products over the nitro group was observed.

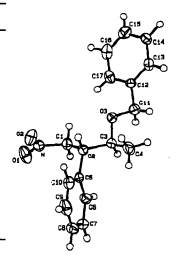


Figure 1. X-Ray structure of compound anti-2a

of the remaining products⁸ was elucidated by ¹H-NMR analysis of the crude reaction mixtures. The coupling constants between the C_{β} -H and C_{γ} -H protons were higher for the *anti*-isomers, as was expected from the more populated conformations of these stereoisomers. The chiral integrity of product *anti*-3a and therefore that of nitroalkene 1b was found to be \geq 98% on the basis of ¹⁹F-NMR experiments with the corresponding Mosher derivatives⁹.

The origins of the stereochemical outcome in the reaction between disubstituted (E)-Michael acceptors and nucleophiles are not clear at present, and several models have been developed 10,11 . Most of these models rely on qualitative arguments based on the preferred conformations of the acceptors and on sterical models. Dorigo and Morokuma 11 , however, have proposed a very interesting model for the conjugate addition of monomeric methyl-copper to (E)-2-pentenal. According to their proposal, an electron-donating group in the anti position of the corresponding TS (see Figure 2) is favored because it stabilizes the enal fragment. This effect is motivated by the strong interaction between the metal and the carbonyl group.

TS	anti	in	out	L	Product	ΔH_{f}	ΔG°
TS1	OH	Ĥ	Me	none	anti	7.0	-29.6
TS2	OH	Me	H	none	syn	6.1	-29.5
TS3	Me	H	OH	none	syn	6.3	-30.4
TS4	Me	OH	H	none	anti	5.0	-34.0
TS5	H	OH	Me	none	syn	4.8	-31.2
TS6	H	Me	OH	none	anti	3.8	-31.8
TS7	OH	Н	Me	H_2O	anti	-124.7	-170.5
TS8	OH	Me	Н	H_2O	syn	-126.0	-171:5
TS9	Me	H	OH	H_2O	syn	-125.9	-171.9
TS10	Me	OH	H	$H_2^{-}O$	anti	-127.6	-173.0
TS11	H	OH	Me	$\overline{\text{H}_2\text{O}}$	syn	-129.2	-174.8
TS12	H	Me	OH	H_2O	anti	-129.4	-175.5

Table 2. Energies of the transition structures TS1-12 computed at the RHF/PM3 level^{a,b} (see Figure 2)

Figure 2

Given that the reaction of phenyl lithium with 1a takes place with remarkable stereoselectivity and that the preference for the anti isomer is the maintained in the experiments reported in Table 1, we have studied the reaction between monomeric phenyl lithium and (S)-(E)-3-hydroxy-1-nitro-1-butene using the semiempirical Hamiltonian PM3¹¹ at RHF level of theory. The twelve possible six-membered ring¹² TSs depicted in Figure 2 have been located and characterized by harmonic analysis. The heats of formation and the standard free energies for TS1-12 have been collected in Table 2. We have found that the minimum energy TSs are those in which donating groups (hydrogen or methyl) are anti with respect to the incoming phenyl group. For non-solvated lithium atoms the most stable TSs are TS6 (hydrogen anti) if we consider the heat of formation, and TS4 (methyl anti) when the standard free energies are considered. Both of these TSs correspond to formation of the anti isomers, in agreement with our experimental results. Tetracoordination of lithium (TS7-12, see Table 2) results in a lower Lewis acid coordination between the metal and the nitro group and a higher $C_{Nu}...C_{\beta}$ bond distance (see Figure 3). In this case, TS12 (hydrogen anti) is the saddle point of lowest energy, particularly in terms of standard free energy (see Table 2). This saddle point also leads to the formation

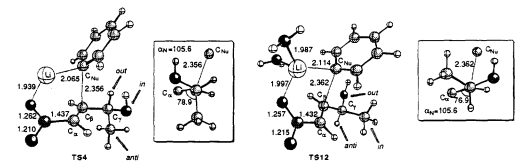


Figure 3. RHF/PM3 fully optimized structures of saddle points TS4 and TS12. Distances and angles are given in Å and deg., respectively. The insets correspond to views above the C_{α} - C_{β} bond. α_N is the C_{Nu} - C_{β} - C_{γ} bond angle.

^aAll energies are given in kcal/mol. ^bAll the saddle points were fully optimized using the eigenvector-following algorithm and were characterized by harmonic analysis, showing one imaginary frequency corresponding to formation of the C_{Nu}····C_B bond.

of the *anti* stereoisomer. Therefore, our calculations suggest that the observed preference for the *anti* stereoisomers is not related to allylic strain or Felkin-Ahn geometries, but to transition structures in which donating groups (methyl and, preferably, hydrogen¹⁴) are *anti* with respect to the carbon-carbon bond being formed (see Figure 3).

Acknowledgement. The present work has been supported by the Universidad del País Vasco-Euskal Herriko Unibertsitatea (Project UPV 170.215-EA156/94) and by the Gobierno Vasco-Eusko Jaurlaritza (Project GV 170.215-0119/94).

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- 6. For the preparation of (E)-nitroalkenes 1a,b see Ayerbe, M.; Cossío, F.P. Tetrahedron Lett. 1995, 36, 4447.
- 7. Full details of the crystal structure investigation of *anti-2a* have been deposited at the Cambridge Crystallographic Data Centre, Lensfield Road, UK.
- 8 anti-2a: m.p.: 98-100°C (from diethyl ether in hexanes); $\left[\alpha\right]_{25}^{D} = +118.1$ (c=0.2, CH₂Cl₂); IR (KBr) 1547, 1577 cm⁻¹; ¹H-NMR (CDCl₃) δ 7.41-7.13 (m, 10H); 5.04-4.95 (dd, J=12.8 Hz, J= 5.3 Hz, 1H); 4.40 (d, J= 11.5 Hz, 1H). 3.81-3.68 (m, 1H), 3.60-3.49 (m, 1H), 1.10 (d, J= 6.0 Hz, 3H); ¹³C-NMR (CDCl₃) δ 128.3, 127.9, 127.5, 127.2, 75.9, 75.7, 70.2, 50.4, 16.9.

anti-3a: m.p.: 68-70°C (from diethyl ether in hexanes); $[\alpha]_{25}^D = -43.1$ (c=0.5, CH₂Cl₂); IR (KBr) 1548, 1378 cm⁻¹; ¹H-NMR (CDCl₃) δ 7.37-7.21 (m, 5H), 4.96 (dd, J= 12.8 Hz, J'= 4.9 Hz, 1H), 4.68 (dd, J= 5.8 Hz, J'= 10.0 Hz, 1H), 4.26 (dt, J= 9.8 Hz, J'= 5.8 Hz, 1H), 3.80 (dd, J= 8.8 Hz, J'= 6.0 Hz, 1H), 3.62 (dd, J= 8.8 Hz, J'= 5.7 Hz, 1H), 3.56 (td, J= 9.8 Hz, J'= 4.9 Hz, 1H), 1.48 (s, 3H), 1.35 (s, 3H). ¹³C-NMR (CDCl₃) δ 136.3, 129.2, 128.4, 127.9, 110.4, 78.3, 77.0, 68.1, 48.9, 26.8, 25.4.

9. These experiments are summarized in the following scheme:

1:1 (blank experiment)

Reagents and conditions: i: LiAlH4 (4.0 eq), Et₂O, gentle reflux, 5 h. ii: (\pm)-MTPACl (1.2 eq.), NEt₃ (2.4 eq.), CH₂Cl₂, r.t., 16 h. iii: (S)-MTPACl (1.2 eq.) NEt₃ (2.4 eq.), CH₂Cl₂, r.t., 16 h). (MTPA: α -Methoxy- α -(trifluoromethyl)-phenylacetyl group).

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- 14. It has been suggested that C-H bonds are better donors than C-Me bonds: see Cieplack, A.S.; Tait, B.D.; Johnson, C.R. J.Am.Chem.Soc. 1989, 111, 8447.