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REACTIONS OF β-NITROSTYRENES WITH GRIGNARD REAGENTS

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Abstract: α-Phenyl-β-nitrostyrene 1a and β-nitrostyrene 1b react with Grignard reagents to generate hydroximoyl halides 3 or nitrile oxides 4 after workup with ice cold concentrated aqueous HX acid solution. Carboxylic acids 5 are the only products isolated from 1b and products 3 or 4 are still obtained from 1a when concentrated sulfuric acid solution is used.

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Conjugated nitroalkenes are exceptional Michael acceptors to organometallic reagents. When nitroalkenes are added to Grignard reagents, high yields of nitroalkanes, *aci* nitro compounds or carbonyl compounds can be obtained.¹ Nef found that primary or secondary nitro compounds can be hydrolyzed, respectively, to aldehydes or ketones, by treatment of their conjugate bases with sulfuric acid. Primary nitro compounds are converted into carboxylic acids, through the Meyer reaction² when treated with concentrated sulfuric acid. Thus, either the Nef or the Meyer reaction occurs when the conjugate base is added to the strong acid (Scheme I).

α-Phenyl-β-nitrostyrene 1a and 2-aryl-1-nitroethenes 1b-1f react with Grignard reagents under a nitrogen atmosphere to give, via 1,4-addition, aci-anions 2. When aci-anions 2 are added to dilute aqueous hydrochloric acid (5%) solution, high yields (60-95%) of nitroalkanes are obtained. On the other hand, the hydroximoyl halides 3 or nitrile oxides 4 are generated when aci-anions 2 are added to³ ice cold concentrated aqueous HX acid solution (HCl 37%, HBr 48%, HI 57%). However, carboxylic acids 5 are the only products isolated from 2b-2f when 85% H₂SO₄ is used, although compounds 3 and 4 are stable enough to be separated when nitronate 2a is hydrolyzed in the same condition (Table 1). The generation of different products must be ascribed to steric effect in the aci-anion 2 and the presence of good trapping reagents such as halide. In the case of 2a, the two phenyl groups are presumed to block the attack of the water molecule so that the hydroximoyl halides 3 or nitrile oxides 4 can be isolated. On the other hand, the intermediates 3 and 4 are hydrolyzed to the carboxylic acids 5

Table 1 The addition of aci-anion 2 to concentrated $HX_{(aq)}$ or 85% $H_2SO_{4(aq)}$

	nitrostyrene	RMgX		(%	(% yield)	
Entry			HX _(aq)	3	4 5	
1	1a	(CH ₃) ₃ CMgCl	HCl		50	
2	1 a	(CH ₃) ₂ CHMgCl	HCl	77		
3	1 a	CH ₃ CH ₂ CH ₂ MgCl	HCl	96		
4	1a	CH ₃ CH ₂ CH ₂ MgBr	HBr	93		
5	la	CH ₃ MgCl	HC1	82		
6	1a	CH ₃ MgBr	НВг	85		
7	1a	CH ₃ MgI	ні	95		
8	la	C ₆ H ₅ CH ₂ MgCl	HCl	95		
9	1a	C ₆ H ₅ CH ₂ MgCl	HC1		95*	
10	1a	C ₆ H ₅ CH ₂ MgBr	HBr	94		
11	1a	c-C ₆ H ₁₁ MgBr	HBr		70	
12	1b	(CH ₃) ₂ CHMgCl	HCI	77		
13	1c	(CH ₃) ₃ CMgCl	HCl	56		
14	1d	(CH ₃) ₃ CMgCl	HC1	56		
15	1d	(CH ₃) ₂ CHMgCl	HC1	48		
16	1d	CH ₃ CH ₂ CH ₂ MgCl	HCl	75		
17	1 d	C ₆ H ₅ CH ₂ MgCl	HCl	93		
18	1 e	(CH ₃) ₃ CMgCl	HCl	30		
19	1e	(CH ₃) ₂ CHMgCl	HCl	92		
20	1f	(CH ₃) ₂ CHMgCl	HCl	50		
21	1a	(CH ₃) ₃ CMgCl	H ₂ SO ₄		62	
22	1a	CH ₃ CH ₂ CH ₂ MgBr	H ₂ SO ₄	89		
23	la	C ₆ H ₅ CH ₂ MgCl	H ₂ SO ₄	95		
24	1b	(CH ₃) ₃ CMgCl	H ₂ SO ₄		49	
25	1b	(CH ₃) ₂ CHMgBr	H ₂ SO ₄		55	
26	1 b	CH ₃ CH ₂ CH ₂ CHMgBr	H ₂ SO ₄		81	
27	1c	CH ₃ MgI	H ₂ SO ₄		44	
28	1f	C ₆ H ₅ CH ₂ MgCl	H ₂ SO ₄		53	

^{*} The $\mbox{CH}_2\mbox{Cl}_2$ extraction is nuetralized by NaHCO $_{3(aq)}$ solution.

when 2b-2f are treated with ice cold 85% sulfuric acid (Scheme II). Hydroximoyl halides 3 are believed from α -nitroso halides C because the transient intense blue color always is observed when nitronates 2 are added to concentrated acids. α -Nitroso halides C arise if halide (from RMgX, MgX₂ and/or HX_(aq)) trap the protonated nitronates A, analogous to water in the Nef reaction, or the nitroso cation B. A similar reaction has been reported in which α -chloronitroso compounds are formed from nitronate salts using dry hydrogen chloride. To the best of our knowledge, this is the first example of isolating hydroximoyl halides 3 and nitrile oxides 4 from the reactions of nitroalkenes with Grignard reagents.

It is known that nitrlle oxides 4 can undergo 1,3-dipolar cycloaddition with olefins or acetylenes to generate 2-isoxazoline or isoxazole respectively. 5 The reaction of pure nitrile oxides 4 or hydroximoyl halides 3 in the presence of triethylamine can react with diethyl fumarate, diethyl maleate or phenylacetylene to generate high yields (92-98%) of additional products. It is also easy to synthesize some intramolecular 1,3-dipolar cycloaddition products when β -nitrostyrenes 6 react with proper Grignard reagents (Scheme III).

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Ph
$$CNO$$
 + EtO_2C THF $reflux 3 hr.$ EtO_2C CO_2Et CH_2Ph EtO_2C CH_2Ph CH_2P

References and Notes

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- 3. The general procedure is to add nitroalkene (2 mmol) in 10 mL THF to 10 mmol of the Grignard reagent in 30 mL of THF at -20 °C. Within 10 minutes, the solution is added to ice cold saturated aqueous HX (HCl 37%, HBr 48%, HI 57%) solution and stirred 30 minutes. The solution is extracted with CH₂Cl₂, dried over MgSO₄, filtered and concentrated to give the products 3 or 4. The CH₂Cl₂ extraction can be neutralized by NaHCO_{3(aq)} or Et₃N to generate nitrile oxides 4 (entry 9) or furoxan (entry 15) after flash column separation in the absence of dipolarophile. The yields are based on NMR by using toluene or diiodomethane as internal standard.
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- 6. Compound **8**: MP 87 °C; 1 H-NMR(200MHz, CDCl₃) δ 7.40-7.12(m, 10H), 4.69-4.53(m, 1H), 4.00-3.78(m, 2H), 3.11-2.88(m, 2H), 2.18-2.01(m, 1H), 1.85-1.67(m, 1H); 13 C-NMR(200MHz, CDCl₃) δ 174.63, 144.23, 143.66, 128.86, 128.27, 127.61, 126.26, 127.03, 126.64, 75.45, 53.89, 53.67,43.95, 25.27. Calculated for C₁₈H₁₇NO: C, 82.10; H, 6.51; N, 5.32. Found: C, 81.99; H, 6.56; N, 5.11.