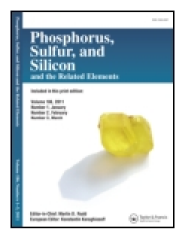
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Chiral Bis(oxazoline)-copper Catalyzed Enantioselective Imidation of Sulfides

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Chiral Bis(oxazoline)-copper Catalyzed Enantioselective Imidation of Sulfides

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Prochiral sulfides reacted with PhI=NTs in the presence of a catalytic amount of Cu(I) salt together with a chiral 4,4'-disubstituted bis(oxazoline) ligand to afford the corresponding chiral sulfimides.

KEYWORDS: sulfide, sulfimide, sufenamide, bis(oxazoline) ligand

INTRODUCTION

Quite recently, chiral sulfimides were demonstrated to be useful in organic synthesis as methylidene transfer reagents to prochiral carbonyl groups, leading to optically active epoxides.¹ Despite the fact that other optically active organosulfur compounds such as sulfonium ylides and sulfoxides have many asymmetric synthetic applications, the chemistry of their nitrogen analogues, sulfimides, has been much less investigated.^{2,3} We now report that the asymmetric reaction of various sulfides 1 with PhI=NTs in the presence of Cu(I) catalyst derived from CuOTf and the chiral 4,4'-disubstituted bis(oxazoline) ligand 3 produces the corresponding chiral sulfimides 2.⁴

RESULTS AND DISCUSSION

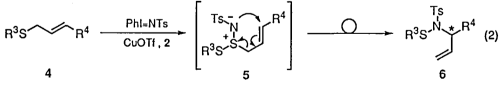
Our initial attempts were to explore and exploit a new imidation method of sulfides 1 with PhI=NTs using Cu(I) salt as catalyst, eqn. (1). The synthesis of sulfimides was carried out by treatment of suitable sulfides with 1 equiv PhI=NTs in MeCN as solvent in the presence of 5 mol% of Cu(I) triflate (CuOTf). Representative results are summarized in Table I. Treatment of methyl *p*-tolyl sulfide with PhI=NTs at 25 °C for 26 h afforded the corresponding sulfimide in 83 % isolated yield (entry 1). Similarly, a variety of sulfides could be converted smoothly to the corresponding sulfimides.

In addition, using a chiral bis(oxazoline) ligand we applied the direct imidation of various sulfides into optically active sulfimides. Thus, in the imidation of benzyl 1-

	_				Cat		ABLE I hesis of sulfu	nides.
3^{1} S B^{2} $-$	S ₂ Phl≖NTs [−] NTs +S			(1)		1		2
R ^{est} CuC	DTf (5 m	nol%) R ^{1.}	$^{3}R^{2}$	(1)	R1		R ² y	ield/%
1			2		p-Tol		Me	83
					Ph		Ph	79
	\circ	\sim_{0}			Ph		Bn	82
	く !!				PhCH ₂	CH ₂	Bn	50
•.	YN	7			4-McO	C ₆ H ₄	Bn	70
	Ph	Ph			1-Nap		Bn	54
		3			4.NO-C	C ₆ H ₄ CH ₂	Ph	72
					4-11020	-6114 C112	•••	12
Synthesi	TABLE s of chira	E 11 1 sulfimides.				TABLE		
Synthesi 1						TABLE	E III sulfenamides	
Synthesi 1 R ¹		l sulfimides.	cc / %		Synthesi	TABLE	E III sulfenamides	6
1	s of chira	l sulfimides. 2		 	Synthesi 4 R ³	TABLE is of chiral	E III sulfenamides	6
1	s of chira	l sulfimides. 2 yield / %	cc / %	· Pł	Synthesi 4 R ³	TABLE is of chiral R ⁴	E III sulfenamides (yield / %	6 ee %
1 R ¹ 1-Nap	s of chira R ² Bn	l sulfimides. 2 yield / % 75	cc / % 71	Pł 2-	Synthesi 4 R ³	TABLE is of chiral R ⁴ Ph	E III sulfenamides yield / % 40	6 ee/% 27
1 R ¹ 1-Nap Ph	s of chira R ² Bn Bn	l sulfimides. 2 yield / % 75 78	cc / % 71 64	Pł 2-	Synthesi 4 R ³ NO ₂ C ₆ H ₄ Nap	TABLE is of chiral R ⁴ Ph Bn	E III sulfenamides yield / % 40 37	5 6 27 43

naphthyl sulfide, the best enantiomeric excess (71 % ee) was obtained when the reaction was performed in toluene in the presence of 6 mol% chiral ligand 3 and 5 mol% CuOTf at 25 °C for 48 h to afford the corresponding chiral sulfimide in 75 % isolated yield (Table II). Other results are also shown in the table.

When the reaction was applied to allylic sulfides 4, the expected aziridination to the double bond did not occur at all and, instead, the chiral allylic sulfenamides 6 were obtained selectively in good yields, Table III. This fact shows that the nitrogen attack occurred only at the sulfur atom to give the chiral allylic sulfimide intermediates 5 and the [2,3] sigmatropic rearrangement with chirality transfer followed, eqn. (2) (Table III).



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