Microwave-Assisted Sulfamide Synthesis

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Introduction

In the area of combinatorial library synthesis for medicinal agents there is a constant need for new methodologies [1]. The sulfamide compounds are noted for their broad and potent antibacterial activity [2-3]. The unsymmetric sulfamides appear to be more potent as protease inhibitors than the symmetric analogues due to the flipped conformation that occurs during binding [4]. Unfortunately, most syntheses focus on symmetric sulfamides. The few methods available for unsymmetrical compounds rely on low-yielding synthetic steps that are neither general nor selective [1,5]. A novel transition-metal-catalyzed process for making unsymmetric sulfamides that was recently reported has several limitations, especially with ortho-isomers [1]. Even though other available methods report high yields, they either require reagents that are not readily accessible or they focus on specific structures rather than a general procedure [6]. Winum and co-workers reported a novel sulfamoylating reagent used in the synth esis of sulfamides [8]. However, our study showed that using the sulfamoylating reagent added additional steps and resulted in slow, low-yielding reactions. In an effort to find a fast and general method for sulfamide synthesis we found that microwave heating facilitates the synthesis of sulfamides. This was accomplished in onepot reaction by a stepwise addition of CSI to tertbutanol at 0 °C to form the N-(tert butoxycarbonyl) sulfamoyl chloride intermediate 2 (scheme 1). Anilines or amines were added the reaction mixture was heated using microwave heating at 80 °C for five minutes. The resulting products were isolated using normal-phase flash chromatography with a good yield (table 1). Microwave synthesis provided great improvements in increasing product yield and decreasing reaction time [7].



The microwave assisted Mitsunobu reaction was used for alkylation of Bocsulfamides with different alcohols (Scheme 2) [9]. The reaction time depended on the structure of alcohols. For example, microwave irradiation of benzyl alcohol mixture with Boc - sulfamides, triphenyl -phosphine and diethyl azodicarboxylate (DEAD) in THF provided N-alkylated products in four minutes at 80 °C. In the case of 2- pyridinepropanol (compounds 10,12,14), one minute of microwave heating was enough to complete reaction.

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Table 1: One-pot microwave-assisted sulfamides synthesis

#	Amines	Products*	% Yield *	% Purity *	(M*+ Na)
з	<u>``</u>	°×k⁺	80	96	309.93
4	C ^{an,}	÷,0	55	97	295.94
5	$\mathbf{O}_{\mathbf{P}}$	+ ² ×P	62	93	308.93
6	8	ϕ_{χ^+}	67	96	334.97
7	00	o ^{ort} t	86	97	363.93

* Reactions were performed in the Biotage EMRYS[™] Liberator microwave system in 2-5 mL reaction vials at 80 ° C. b Yield of isolated product: All products were isolated on the Biotage Sp automated flash chromatography system (Flash 25+ M, 25 x 150, 40-63 mm, 60 Å), using ethyl acetate and hexane gradient . C Purity is calculated by HPLC (Waters, C8 4.6 x 50 mm, S-3 120 Å). . ^d Mass spectroscopy was carried out on a Micromass[®] ZQ (Waters). e 1H NMR data in CDCI3 was collected on a 500 MHz Bruker spectrometer.

The tert-butoxycarbonyl group removal is generally carried out with trifluoroacetic acid either neat or in combination with CH2Cl2 [14]. Since CF3COOH is volatile, harsh and corrosive, a search for an alternative method of deblocking is ongoing.

Recently was reported that Amberlyst 15, a strong acidic resin, can remove the Bocprotecting group and form salts with the deprotected amines [15]. This method has been used to facilitate the generation and purification of amines. However, this technique requires a long reaction time (12-24 hours). We decided to explore the scope and limitations of deblocking the BOC-group from sulfamides using silica-bonded phenylsulfonic acid, and the effects of microwave heating in altering the reaction time (scheme 3). Bocsulfamides were treated with Si-TsOH and heated by microwaves at 100 °C. In all the examples, the Boc- protecting group was completely removed within five minutes [16]. Here we report that microwave heating with Si-TsOH significantly shortens the Boc-removal time.

The formation of salts between the sulfamide and silica bonded acid depends on subsituents on the sulfamide nitrogen (pKa of sulfamides 7-11). The desired products were released from Si-TsOH surface using NH3/MeOH (scheme 3).





DIE 3:	ble 3: Microwave-assisted BOC- deblocking using Si-TsOH						
#	Products*	% Yield ⁵	M+ + Na ^c				
15		90	209.11				
16		75	195.03				
17		75	209.05				
18		61	235.63				
19		91	264.02				
20		71	325.38				
21		92	353.99				
22	Ċ ţ	89	354.42				

Table 3: Microwave-assisted BOC- deblocking using Si-TsOH

^a Reactions were performed in the Biotage EMRYSTM Liberator microwave system in 2-5 mL reaction vials. ^b Yield of isolated product. ^C Mass spectroscopy was carried out on a Micromass [®] ZQ (Waters). ^{e 1}H NMR data was collected on a 500 MHz Bruker spectrometer.



Conclusion

A general microwave assisted reaction in preparation of unsymmetric Boc -sulfamides is demonstrated. Also, an alternative method of Boc -removal from sulfamide was i ntroduced using Si -TsOH in conjunction with microwave heating. Boc- de blocked sulfamides were captured by Si-TsOH, depending on their pKa,. The captured sulfamides were released from surface of Si-TsOH by using NH3 in MeOH, followed by quick flash purification. This new method of microwave-assisted, Boc -cleavage group from sulfamides facilitates the preparation and purification of unsymmetric sulfamides.

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8. General procedure for Boc-sulfamides using microwave heating: In a typical experiment, chlorosulfonyl isocynate (0.24 ml, 2.7 mmol) was added dropwise to a solution of tert -butyl alcohol (0.26 ml, 2.7 mmol) in anhydrous dichloromethane (3 ml) in a sealed Pyrex tube under inert gas at 0 ° C. Amine (5.5 mmol) was then added and t he reaction was heated in a microwave cavity for 5 minutes at 80 °C. The reaction mixture was added to a Samplet TM cartridgeand purified by flash chromatography
8. General procedure for microwave assisted Boc-sulfamides cleavage with Si -TsOH: Method A: Silicabound p-toluenesulfonic acid (1.26g, 0.96 mmol) was added to the Boc - protected sulfamide (0.32 mmol) in 1:1 acetonitrile: DCM (4ml). The reaction was heated to 100 °C in a microwave cavity for 5 minutes. The reaction mixture was then loaded onto a silica column. Using the following conditions on flash chromatography yielded the desired compound



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