Sulfur Trioxide-Amine Adducts

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Introduction

Many developments have taken place throughout the history of sulfonating reagents from the synthesis of stabilized liquid SO$_3$ to the production of chlorosulfonic acid (ClSO$_3$H) and the development of SO$_3$-amine adducts. Reactions involving SO$_3$ and ClSO$_3$H prove to be quite violent and are accompanied with heat liberation. With such violent reactivity characteristics a milder reagent was sought. It has been long known that chlorosulfonic acid added dropwise and at low temperatures reacts with aliphatic and aromatic tertiary amines to produce the corresponding sulfur trioxide amine complexes (O$_3$S$^-$–$N^+$R$_3$) for example, O$_3$S$^-$–N(CH$_3$)$_3$, O$_3$S$^-$–N(CH$_2$CH$_3$)$_3$ and O$_3$S$^-$–N(CH$_2$CH$_2$CH$_3$)$_3$.

Sulfur trioxide amine adducts are valuable and versatile sulfonating agents. They may be utilised in the sulfation of alcohols (formation of S–O bonds), sulfonation of acid sensitive heterocycles, aliphatic and aromatic hydrocarbons (formation of S–C bonds) and the sulfamation of amines (formation of S–N bonds). The resulting products have quite diverse properties, for example, they may act as dyes, detergents, sweeteners, pharmaceutical intermediates, herbicides etc.

Precautions

All reactions must be carried out in a fumehood. Care must be taken when utilizing concentrated acids, which are highly corrosive and fume as they react with the moisture in the air e.g. chlorosulfonic acid, sulfuric acid, liquid sulfur trioxide. Anhydrous conditions are essential and solvents must be refluxed over appropriate agents and distilled and stored over molecular sieves. Solid reagents are recrystallized from appropriate solvents while liquids are distilled. It is necessary that sulfur trioxide complexes are stored in airtight containers to minimize hydrolysis of the reagent by water vapour.

Abstracts

(A) A series of sulfur trioxide complexes have been employed in the sulfation of alcohols. There are many examples extensively documented in the literature; tert-butyl alcohol and 3-methyl-3-pentanol were successfully converted to their sulfate ester salts by reaction with SO$_3$-pyridine or SO$_3$-Et$_3$N.

(B) Sulfonation proceeds via oxidation of hydrocarbons resulting in carbocation formation. Saturated aliphatic hydrocarbons possessing a carbonyl component are sulfonated upon the carbon adjacent to this entity. Unsaturated hydrocarbons are sulfonated quite easily forming a $\beta$-sultone or a solvated carbenium ion. However, (2SO$_3$)-pyridine sulfonates butyric acid quantitatively.
(C) Sulfonation is also achieved via insertion of SO$_3$ into a carbon-metal bond. Organometallic compounds treated with SO$_3$–Me$_3$N, a mild reagent, react to produce the corresponding sulfonic acids – a process suitable for both aliphatic and aromatic hydrocarbons.$^{6}$

(D) 2% Cross-linked $p$-bromopolystyrene, may also be converted to its corresponding aromatic polystyrenesulfonic acid via sulfonation of its lithium derivative.$^7$

(E) Initially amines were sulfamated by treatment with chlorosulfonic acid.$^{8,9}$ Both mono- and disubstituted sulfamates may be prepared with ease. SO$_3$–Pyridine is used extensively in the sulfamation of primary amines.$^{10}$ The amine salt formed is treated with hydroxide and the alkali sulfamate extracted and re-crystallised finally from 95% ethanol.$^{11}$

(F) Electron donating groups, for example methyl, ethoxy etc., promote sulfamation however electron-withdrawing groups such as nitro substituents hinder the reaction. Utilizing $a$-picoline (2-methylpyridine) however overcomes this problem and the resulting sulfamate may be extracted in good yield.$^{11}$

(F) SO$_3$–amine adducts, such as SO$_3$–pyridine, may be affectively utilized in the reduction of aromatic (diphenyl sulfoxide) and aliphatic sulfoxides at room temperatures resulting in quantitative yields of the corresponding sulfide.$^{12}$

References