

Product Information

92718 1-(Trimethylsilyl)imidazole – Pyridine mixture for GC derivatization, LiChropur®

Storage temperature: 2-8°C

1-(Trimethylsilyl)imidazole (TMSI) is the strongest reagent for hydroxyls. It reacts quickly and smoothly with hindered and unhindered hydroxyl and carboxyl groups. TMSI+pyridine 1:4 (v/v) is useful for derivatizing wet sugar samples, hindered hydroxyl groups in steroids and, in conjunction with fluorinated acylation reagents, amino acids. For moderately hindered or slowly reacting compounds, pyridine is an especially useful solvent for TMSI because of its ability to act as an HCl acceptor in silylation reactions involving organochlorosilanes.

Features/Benefits

- Useful for derivatizing alcohols, hormones, fatty and other organic acids, phenols, prostaglandins, steroids, sulfonic acids, and thiols.
- Derivatizes sugars in the presence of small amounts of water.
- Useful in multiderivatization schemes containing hydroxyl and amine groups.
- TMSI does not react with amines or amides.
- TMS derivatives are thermally stable but more susceptible to hydrolysis than their parent compounds.

Typical Procedure

This procedure is intended to be a guideline and may be adapted as necessary to meet the needs of a specific application. Always take proper safety precautions when using a silylating reagent. TMSI+pyridine is extremely sensitive to moisture and should be handled under dry conditions.

Prepare a reagent blank (all components, solvents *except sample*), following the same procedure as used for the sample.

1. Weigh 1-10 mg of sample into a 5 mL reaction vessel. If appropriate, dissolve sample in solvent. If sample is in aqueous solution, evaporate to dryness, then use neat or add solvent.

2. Add silylating reagent. TMSI+pyridine can be used at full strength or with a solvent.* In most applications it is advisable to use an excess of the silylating reagent – at least a 2:1 molar ratio of TMSI to active hydrogen.
3. Allow the mixture to stand until silylation is complete. To determine when derivatization is complete, analyze aliquots of the sample at selected time intervals until no further increase in product peak(s) is observed.

Derivatization times vary widely, depending upon the specific compound(s) being derivatized. Many compounds are completely derivatized as soon as they dissolve in the reagent. Compounds with poor solubility may require warming. A few compounds will require heating at 70°C for 20-30 min. Under extreme conditions compounds may require heating for up to 16 h to drive the reaction to completion.

If derivatization is not complete, evaluate the addition of a catalyst, use of an appropriate solvent, higher temperature, longer time and/or higher reagent concentration.

Use a glass injection port liner or direct on-column injection when working with silylating reagents. Erratic and irreproducible results are more common when stainless steel injection ports are used.

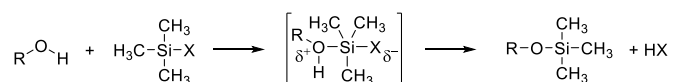
*Nonpolar organic solvents such as hexane, ether, and toluene are excellent solvents for the reagent and the reaction products; they do not accelerate the rate of reaction. Polar solvents such as pyridine, DMF, dimethylsulfoxide (DMSO), tetrahydrofuran (THF), and acetonitrile are more often used because they can facilitate the reaction. Pyridine is an especially useful solvent because it can act as an HCl acceptor in silylation reactions involving organochlorosilanes.



TMS derivatives and silylating reagents react with active hydrogen atoms. Do not analyze TMSI+pyridine derivatives on stationary phases with these functional groups (e.g. polyethylene glycol phases). Silicones are the most useful phases for TMS derivatives combining inertness and stability with excellent separating characteristics. Nonpolar silicone phases include SPB™-1 and SPB-5. Normal hydrocarbons (carbon-hydrogen analytes with single bonds) are separated by these phases. More polar phases, SPB-1701 and SP™-2250, separate carbon-hydrogen analytes that also contain Br, Cl, F, N, O, P, or S atoms or groups. A highly polar cyanopropylphenylsiloxane phase, SP-2330, is useful for separating fatty acid methyl esters or aromatics.

Mechanism¹⁻²

Silylation is the most widely used derivatization procedure for GC analysis. In silylation, an active hydrogen is replaced by an alkylsilyl group. Compared to their parent compounds, silyl derivatives generally are more volatile, less polar, and more thermally stable. Silyl derivatives are formed by the displacement of the active proton in -OH, -COOH, =NH, -NH₂ and -SH groups.



The general reaction for the formation of trialkylsilyl derivatives is shown above.

The reaction is viewed as a nucleophilic attack upon the Si atom of the silyl donor, producing a bimolecular transition state. The leaving group X (for TMSI, X = NCH=NCH=CH, for pyridine, X = C₅H₅N) must possess low basicity, the ability to stabilize a negative charge in the transition state, and little or no tendency for n (p-d) back bonding between itself and the silicon atom.

The ideal silyl leaving group X must be such that it is readily lost from the transition state during reaction, but possesses sufficient chemical stability in combination with the alkyl silyl group to allow long term storage of the derivatizing agent for use as required. As the formation of the transition state is reversible, the derivatization will only proceed to completion if the basicity of the leaving group X exceeds that of the group it replaces. The ease of derivatization of various functional groups for a given silyating agent follows this order: alcohol > phenol > carboxylic acid > amine > amide.

Within this sequence reactivity towards a particular silylating reagent will also be influenced by steric hindrance, hence the ease of reactivity for alcohols follows the order: prim. > sec. > tert., and for amines: prim. > sec.

Storage/Stability

Recommended storage conditions for the unopened product are stated on the label. Store in an amber bottle or ampule at 2-8°C in a dry, well ventilated area. Use only in a well ventilated area. Keep away from ignition sources. Properly stored, this mixture is stable indefinitely. Moisture will decompose both TMS reagents and derivatives. To exclude moisture, this reagent is packaged under inert gas. If you store an opened container or transfer the contents to another container for later reuse, add desiccant. **Before reuse, validate that your storage conditions adequately protected the reagent.**

References

1. K. Blau and J. Halket, *Handbook of Derivatives for Chromatography* (2nd ed.), John Wiley & Sons, New York, 1993.
2. D.R. Knapp, *Handbook of Analytical Derivatization Reactions*, John Wiley & Sons, New York, 1979.

Precautions and Disclaimer

This product is for R&D use only, not for drug, household, or other uses. Please consult the Safety Data Sheet for information regarding hazards and safe handling practices.

