



Material Safety Data Sheet for Mononucleosis Test Device

1. Identification of the substance/preparation and of the company/undertaking

Product: Henry Schein One Step+ Mononucleosis Test Device

DISTRIBUTOR:

Henry Schein, Inc.
135 Duryea Road
Melville, NY 11747

Further information available from: (800) 472-4346

Information in case of emergency: CHEMTREC, (800) 424-9300

2. Composition/information on ingredients

Test Device contains dry reagents (bovine erythrocyte extracted antigen, buffers, desiccant). Stored in a sealed foil pouch.

3. Hazards identification

Physico-Chemical Hazard: None

Health Hazard: Each Device contains approximate sodium azide concentration = <0.02% (7.5 ng/test), buffer with sodium azide 0.09% (1 mg/mL), positive control with human plasma and sodium azide 0.09% (1 mg/mL), and negative control with human plasma and sodium azide 0.09% (1 mg/mL). They are all below accepted limit.

Environmental Hazard: None

4. First-aid measures

The pouched product is a plastic device with only dry reagents being supplied; therefore accidental contact via spillages, etc. can only occur with the sample tested on the Device. If the user contacts the sample, normal personal hygiene measures should be taken.

5. Fire-fighting measures

Not Applicable

6. Accidental release measures

Not Applicable.

7. Handling and storage

Store at 2-30°C.

8. Exposure controls/personal protection

Use good laboratory practice, including laboratory gloves and safety spectacles.

9. Physical and chemical properties

The product comprises insert membrane and pad materials in which the chemical reagents are supplied in dry form and which, along with desiccant, are housed in a plastic casing and foil pouched. The product is odorless.

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10. Stability and reactivity

Stable under conditions of use until the expiry date indicated on the corresponding label.

11. Toxicological information

Not Applicable

12. Ecological information

Not Applicable.

13. Disposal considerations

The used Device should be disposed of via an auto clave or by incineration as for other waste containing biological material.

14. Transport information

Not Applicable.

15. Regulatory information

Not classified as hazardous according to EEC criteria.

16. Other information

None.