

Technical Data Sheet

Rev. 14252

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2mL CryoClear[™] Sterile, Barcoded Cryogenic Vials

Item# 3002 - Internal thread, round bottom, self-standing









Item# 3002 (shown actual size)



All CryoClear™ cryogenic vials are sold in tamper evident, resealable plastic bags. 50 vials per bag, 10 bags per case.

Description

Globe Scientific's CryoClear™ vials are designed for cryogenic storage and transport of biological specimens at temperatures as low as -196°C. This innovative line features outstanding leak resistance and purity due to the special thermoplastic elastomer layer that is co-molded with the screw cap. The caps feature a star shaped top cavity that is engineered to work with automated capping/decapping equipment. Each vial has a unique printed barcode for automated data collection, accurate sample inventory, and to conceal the sample's identity. The vials have printed graduation marks for exact measurements and a large white writing area for manual specimen identification.

Features and Benefits

- Screw caps are molded with a thermoplastic elastomer (TPE) layer to provide a 100% leak-proof seal
- Star shaped top cavity is engineered for use with automated capping/decapping equipment
- Unique barcode printed on each vial for automated data collection
- White writing surface for specimen identification
- Printed graduations for accurate measurements
- Vials and caps are autoclavable
- Produced from medical grade raw materials that will not discolor after re-sterilizing
- Chemical resistant polypropylene vial
- Tamper evident packaging
- Self-standing vials interlock in Globe workstation

Products

Item#	Description	Packaging
3002	2mL, sterile cryogenic vial,	50 per bag,
	internal thread, round bottom,	500 per case
	self-standing	

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Technical Specifications

Material: Vial: Medical/FDA Polypropylene (PP)

Cap: Medical/FDA Polyethylene (PE) co-molded with a

thermoplastic elastomer (TPE) layer

Sterilization: Beta radiation

SAL level: 10-6

Autoclavable: Vials and caps
Temperature range: -196° to +121°C

Mold release agents: None used in manufacturing

Maximum psi: 14.5 psi

Maximum centrifugation speed: 17530 xg (To obtain RPM, verify the diameter of the

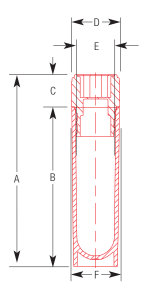
centrifuge and convert)

Printing on the vial is resistant to the following common laboratory chemicals:

Isopropyl alcohol: 5-100% Butanol: 100% DMSO: 10-20% Bleach: 5%

Acetic acid: 10% Hydrochloric acids: 10%

NaOH: 10%



2mL - Internal Thread				
Α	Overall Height	48.4mm		
В	Vial Height	40.2mm		
С	Cap Height	8.3mm		
D	Cap O.D.	12.3mm		
Ε	Vial I.D.	10.0mm		
F	Vial O D	12.5mm		

Product Certifications

Manufacturing environment: Produced, assembled and packaged in a ISO 7 cleanroom (Class 10,000) in accordance with quality guidelines ISO 13485:2004,

ISO 14644, ISO 14698 (Federal Standard 209), ISO 14000 and ISO 18000

Material: Virgin raw materials are tested according to the "United States Pharmacopia" (USP) and with the "drug master file" (DMF) at the FDA Certified

nder USP Class VI

CONEG: This product meets CONEG requirements and therefore does not contain heavy metals in the color concentrate

Sterilization: Sterilzed in accordance with EN ISO 11137

DNase, RNase and pyrogen free: Every lot is certified* by an independent laboratory to be free of DNase, RNase and pyrogens

 $\label{thm:equality:equality:equal} \mbox{Human DNA and ATP free:} \quad \mbox{Every lot is certified* by an independent laboratory to be free of Human DNA and ATP}$

Non-cytotoxic: This product is certified* to be free of cytotoxins

Non-hemolytic: This product is certified* to be free of articles causing hemolysis

In vitro Diagnostic Medical Device: Complies with Directive 98/79/EC of the European Parliament and the Council of 27 October 1998 on in vitro diagnostic medical devices

Leak proof: Vials are certified leak proof at a pressure of 95 kPa (0.95 bar, 14.5 psi) by applying a closure force of not less than 8 cNm

IATA (International Air Can be used as a primary receptacle for the transport of diagnostic specimens as outlined by the IATA Dangerous Goods Regulations,

Transportation Association): Part 6.3,5

*Certificate of Analysis available upon request.



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