

GENERAL INFORMATION

Category: Sterile sampling bags

Family: Twirl'em

Lifespan: 5 years

TECHNICAL DESCRIPTION

Twirl'em bags have a practical and easy-to-use closing system. They are made of a flexible, strong and transparent plastic.



SPECIFIC INFORMATION

ITEM

Bag
Polyethylene blend
Transparent
89 x 178 mm / 3.5 X 7 po
3 mil.in / 0.0762 mm / 76.2 micron
230 ml / 8 oz
100 ml / 3 oz
Clear
Perforated line
Attachment with 1 round wire and 1 flat wire
Yes
Recyclable

PACKAGING INFORMATION

Outer box dimension : (W x D x H)	13.13 po x 8.06 po x 6.63 po 33 cm x 20 cm x 17 cm
Box weight:	7.00 LB / 3.18 KG
Conditioning:	1000 (2 x 500)
Storage condition:	Store in a dry place at room temperature

OTHER



EFL-3070

AVAILABLE DOCUMENTS

Data Sheet	Certificate of Compliance
Certificate of Analysis	Safety Data Sheet (SDS)
Certificate of Sterility	Pyrogen Declaration
DNase/RNase	

Reach out to us for additional resources, if applicable to this product.

DECLARATION	
CFIA	LABPLAS sampling bags are a solution that may be used in the CFIA Preventive Control Plan (PCP) for the seven principles of the HACCP system. The PCP is a Canadian federal initiative, under the Safe Food for Canadians Regulations (SFCR).
EU	The materials used to manufacture LABPLAS sampling bags meet, where applicable, the Eu No10/2011 standards for food contact with respect to particle migration.
DNase-free	This product is DNase-free. Sensitivity of 10-7 Kunitz units/µL
RNase-free	This product is RNase-free. Sensitivity of 10-9 Kunitz units/µL.
FDA	The plastic film used in the manufacture of the LABPLAS sampling bag meets the requirements of 21 CFR 177.1520 of the Food and Drug Administration.
Pyrogens	This product is non-pyrogenic at the endotoxin limit of 2.15 EU/device. Non- pyrogenicity is supported by endotoxin testing of randomly selected samples using the Limulus amebocyte lysate (LAL) gel assay according USP-NF <85> and <161> guidelines.
Sterile	Sterility is provided by dry heat during extrusion of the plastic at temperatures exceeding 428 F. The approach ensures a sterility assurance level (SAL) of 10-3. Continued process effectiveness is demonstrated through periodic sterility testing. Sterility testing follows the USP-NF <71> guideline.

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