

Certificate of Quality



VWR®GEL LOADING PIPET TIPS

As per the manufacturer, the below product meets the following criteria:		
VWR Catalog Number	37001-152	
European Article Number	732-0509	
Lot Number	70052-640SQ-6192	
Description	VWR TIPS GEL 30MM 200UL PK576	
Date of Manufacture	10/7/2016	
Date of Sterilization	10/13/2016	
Date of Expiration	2019-10	
Country of Origin	Made in USA	

Quality System Compliance

The product conforms to written material specifications and was manufactured under the manufacturer's registered and audited ISO 9001 quality system, and underwent lot testing as outlined in the manufacturer's laboratory procedures. These products come with the highest standard of quality assurance.

QC Testing

Each lot of item was produced in a tightly controlled environment and subjected to the manufacturer's rigorous testing and performance procedures. The products meet all stated standards for precision, clarity, warp, centrifugation and freedom from contamination.

Product Specifications

Material:	Pipet Tip Material: Polypropylene	
	Certified free of Bisphenol A (BPA), phthalates and cytotoxic effects. Plastic resins used in product manufacturing have been tested for heavy metals using the prescribed USP method and confirmed to have levels lower than 1 PPM. Resins are USP Class-VI certified, RoHS, FDA regulation CFR 21, and are free of Substances of Very High Concern (SVHC), REACH compliant, and are FDA approved for food contact.	
Non-Pyrogenic Statement:	Product samples are exposed to endotoxin-free water and the resulting extraction fluid is tested for contamination using the kinetic turbidimetric Limulus Amoebocyte Lysate (LAL) assay protocol and USP guidelines. All products tested must display less than 0.05 EU/ml to be certified free of endotoxin.	
ATP Assay:	Product sample surfaces are tested for the presence of Adenosine triphosphate (ATP) using a controlled bioluminescence reaction to detect contamination. Luminescence data is compared to results generated by ATP-free surfaces and surfaces with known amounts of ATP as a positive control. The relative light units' result must indicate less than 2X10 ⁻¹² mg/ul of ATP for the product to be certified as ATP free.	
DNase & RNase Free:	Product samples are exposed to nuclease-free water and the resulting extraction fluid is tested for nuclease activity on commercially available 7.5 kb Poly(A) tailed RNA (1ug) and HindIII- digested DNA (1ug) with a one hour 37°C incubation in appropriate buffers. Results are visualized on an agarose gel with appropriate positive and negative controls. Extraction	

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	fluid samples must show no degradati occurred for the product to be certified as	ion of the nucleic acids by the extraction fluid has s RNase-free and DNAse-free.	
BSE/TSE Statement:	(Bovine Spongiform Encephalopathy/Transmissible Spongiform Encephalopathy)		
	The product contains resins that are processed under one of the following sets of conditions: Hydrogenation of Tallow at 200°C Hydrolysis at 260°C and 48 bar for 1.5 to 2 hours Vacuum distillation a 232°C		
	or		
	minutes. These conditions exceed the Commission Directive EC 98/16/EC of 1 76/768/EEC and further Amendment 419	and 300 PSIG for 2.5 hours Distilled at 232°C for 5 e European Union standard as listed in the 22nd March 5th, 1998 as annexed to council Directive EC 9 Annex II of 12 June 2001, and as agreed to by the ecommendations concerning Tallow derivatives for	
	The manufacturer does not store any products of animal origin, including animal proteins, in any manufacturing areas of their facility or warehouses.		
Sterilization Process:	Products from the specified lot number was processed by e-beam radiation at a dose range 15.2-32.0 kiloGray. This dosage is sufficient to guarantee a sterility assurance level of Sterility assurance levels are based on the probability of a positive, nonsterile part occur after irradiation. An SAL of 10 ⁻⁶ is the highest level of sterility able to be guaranteed according the ISO 11137 standard and represents a 1 in 1,000,000 chance of a nonsterile part occur. The Manufacturer's sterilization program is an ISO 11137 validated process, with biobut studies and radiation validation audits performed quarterly.		
	Specified Dosage Range:	15.20 - 32.00 kGy	
	Minimum Dosage Delivered:	16.1 kGy	
	Maximum Dosage Delivered:	30.2 kGy	
Latex Statement:	No Latex was used in the manufacturing and shipping materials used in the produce	g of the product. This also includes all the packaging ction of all items.	
		may come in contact with Latex would be from the are required to wear for the purpose of clean handling	

Disclaimer: VWR states that this declaration will not discharge the user from his obligation to ensure the product is suitable for the intended use.

and to avoid biological contamination of the products.

This Certificate was automatically generated and is valid without a Signature.

Date: 11/7/2016