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PRODUCT SPECIFICATIONS SHEET WORLD/GMP GRADE ETHYL ALCOHOL 96% (192 PROOF) Meets ACS/USP/EP/BP/JP Grade Monographs

With USP<232>, EMA and ICH Q3D Test Results Grain Derived Ethanol Catalog Number: 111WORLD192-Size Code*

*Individual package sizes have unique size codes

Manufactured in compliance with cGMP

	MONO-	F	TYPICAL	
TEST	GRAPH	SPECIFICATION	RESULT	
Assay (by GC, corrected for water)	ACS	NLT 95.0%	96.02%	
Assay (by relative density @20°C)	EP/BP ¹	95.1% - 96.9% (by volume)	96.02%	
Assay (by specific gravity@15°C)	JP	95.1% - 96.9% (by volume)	90.02%	
Proof	27CFR 30.23	Lot Analysis	192.0	
Characters Description	EP / BP JP	Ethanol is a clear, colorless volatile, flammable liquid. It is miscible with water and methylene chloride. It burns with a blue, smokeless flame. BP: about 78°C	Pass	
Identification A - Relative Density	EP/BP ¹	0.805 – 0.812 @ 20°C	0.808	
Specific Gravity	JP^1	d 15/15 0.80872 – 0.81601	0.81271	
Identification Test B	USP/EP/BP	Conforms to IR Spectra	Pass	
Identification 1	JP	Conforms to IR Spectra	Pass	
Identification Test C	EP/BP	An intense blue color appears on the paper and becomes paler after 10-15 minutes	Pass	
Identification Test D	EP/BP	A yellow precipitate is formed within 30minutes	Pass	
Solubility in Water	ACS	To Pass Test	Pass	
Color of Solution	USP	The Sample solution has the appearance of water or is not more intensely colored than the Standard solution	Pass	
Clarity of Solution	USP	Sample Solutions show the same clarity as that of water, or their opalescence is not more pronounced than that of	Pass	
Purity 1 – Clarity and Color of Solution	JP	the Standard solution. The mixture remains clear	Pass	
Appearance	EP/BP	Clear and Colorless, the dilution remains clear when compared with water	Pass	
Acidity or Alkalinity	USP/EP/BP	The solution is pink (30ppm, as acetic acid)	Pass	
Purity 2 – Acidity or alkalinity	JP	A light red color develops	Pass	
Titrable Acid	ACS	0.0005 meq/g max.	<0.0003 meq/g	
Titrable Base	ACS	0.0002 meq/g	<0.0001 meq/g	
Acetone/Isopropyl Alcohol	ACS	To Pass Test	Pass	
Methanol	ACS	0.1% max	<0.1%	
Substances Darkened by Sulfuric Acid	ACS	To Pass Test	Pass	
Substances Reducing Permanganate	ACS	To Pass Test	Pass	



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TEST	MONO- GRAPH	SPECIFICATION		TYPICAL RESULT
Limit of Nonvolatile Residue Residue after Evaporation Residue on Evaporation Purity 5 - Residue on Evaporation	USP ACS EP/BP JP	NMT 2.5 mg 0.001%, max 25 ppm, max NMT 2.5 mg		0.5mg 0.0006% <10 ppm 0.5mg
UV Absorbance Purity 4 - Other Impurities (absorbance)	USP/EP/BP JP	Examine between 235nm – 340nm. 240nm 250nm-260nm 270nm-340nm The spectrum shows a steadily described observable peaks or shoulders	0.40 max. 0.30 max. 0.10 max.	0.34 0.15 0.05 Pass
Organic Impurities Volatile Impurities Purity 3 – Volatile Impurities	USP EP/BP JP	Methanol Acetaldehyde and Acetal Benzene Sum of all other impurities	200 ppm max. 10ppm max 2ppm max. 300ppm max.	<10 ppm <1 ppm None Detected <50ppm

¹No USP specification for this assay



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Permitted Concentrations of Elemental Impurities Following Option 1 Guideline in drug products, drug substances and excipients¹

Reported in µg/g (ppm)

Element	Class	Oral Concentration µg/g	Parenteral Concentration µg/g	Inhalation Concentration μg/g	TYPICAL RESULT (in µg/g) (ppm)
Cd (Cadmium)	1	0.5	0.2	0.2	0.00
Pb (Lead)	1	0.5	0.5	0.5	0.00
As (Arsenic)	1	1.5	1.5	0.2	0.00
Hg (Mercury)	1	3	0.3	0.1	0.00
Co (Cobalt)	2A	5	0.5	0.3	0.00
V (Vanadium)	2A	10	1	0.1	0.00
Ni (Nickel)	2A	20	2	0.5	0.00
Tl (Thallium)	2B	0.8	0.8	0.8	0.00
Au (Gold)	2B	10	10	0.1	0.00
Pd (Palladium)	2B	10	1	0.1	0.00
Ir (Iridium)	2B	10	1	0.1	0.00
Os (Osmium)	2B	10	1	0.1	0.00
Rh (Rhodium)	2B	10	1	0.1	0.00
Ru (Ruthenium)	2B	10	1	0.1	0.00
Se (Selenium)	2B	15	8	13	0.00
Ag (Silver)	2B	15	1	0.7	0.00
Pt (Platinum)	2B	10	1	0.1	0.00
Li (Lithium)	3	55	25	2.5	0.00
Sb (Antimony)	3	120	9	2	0.00
Ba (Barium)	3	140	70	30	0.00
Mo (Molybdenum)	3	300	150	1	0.00
Cu (Copper)	3	300	30	3	0.00
Sn (Tin)	3	600	60	6	0.00
Cr (Chromium)	3	1100	110	0.3	0.00

¹Includes all requirements for ICH Q3D-Step 4 version, EMA (EP) 5.2 and USP <232> and <233> General Chapters. Form: Ethanol, Pure, 192, ACS/USP/EP/JP Rev. 2.5, 01/18, PJM

This product is for further commercial manufacturing, laboratory or research use, and may be used as an excipient or a process solvent for pharmaceutical purposes. It is not intended for use as an active ingredient in drug manufacturing nor as a medical device or disinfectant. Appropriate/legal use of this product is the responsibility of the user.