

## SPECIFICATION

### Propylene Glycol Ph. Eur.\* / USP\* Propylene Glycol USP/EP

Product code: 029

Testing specifications: Ph. Eur.\* / USP\* / LSM 029\*

The material meets all requirements of Ph. Eur.\*, USP\*, JP\*, FCC\* and (EU) No 231/2012 (E1520)

Parameter	Ph. Eur.*		USP*	
	Specification	Test method	Specification	Test method
Assay	—	—	≥ 99.5%**	GC, LSM 029**
Characters	clear, colourless	Ph. Eur.*	—	—
Identification	A / B / C / FT-IR	Ph. Eur.*	A/B/C**	USP*
Boiling point	184 – 189°C	Ph. Eur.*	—	—
Appearance	clear, colourless	Ph. Eur.*	—	—
Acidity	≤ 0.05ml 0.1 N NaOH	Ph. Eur.*	≤ 0.20ml 0.10 N NaOH	USP*
Oxidising substances	≤ 0.2ml 0.05 N Sodium thiosulphate	Ph. Eur.*	—	—
Reducing substances	conforms Ph. Eur.*	Ph. Eur.*	—	—
Chloride	—	—	≤ 70 ppm	USP*
Sulfate	—	—	≤ 60 ppm	USP*
Relative density / Specific gravity	$d_{20}^{20}$ : 1.035 – 1.040	Ph. Eur.*	$d_{25}^{25}$ : 1.035 – 1.037	USP*
Refractive index $n_D^{20}$	1.431 – 1.433	Ph. Eur.*	—	—
Residue on ignition	—	—	≤ 3.5 mg / 50 g	USP*
Sulphated ash	≤ 0.01% m/m	Ph. Eur.*	—	—
Water	≤ 0.2%	Ph. Eur.*	≤ 0.2%	USP*
Limit of diethylene glycol and ethylene glycol: - DEG - EG	—	—	≤ 0.10% ≤ 0.10%	GC, LSM 029**
Additional Parameters				
Parameter	Specification	Test method		
Assay	≥ 99.80% m/m	GC, LSM 029**		
Monoethylene Glycol	≤ 0.01% m/m	GC, LSM 029**		
1,2-Butanediol	≤ 0.01% m/m	GC, LSM 029**		
1,3-Propanediol	≤ 0.01% m/m	GC, LSM 029**		
Diethylene Glycol	≤ 0.01% m/m	GC, LSM 029**		
Di- and Tripropylene Glycol	≤ 0.1% m/m	GC, LSM 029**		

\*\* The Requirements of USP Identity B/C and Assay are covered by internal validated GC method LSM 029.

\*current version

Compiled by: 16.12.2019  Dr. Katja Teufel QC-Manager	Approved by: 17.12.2019  Dr. Philipp Hoch QA-Manager	Released by: 20.12.2019  Dr. Frank Milek Qualified Person (GMP)	Effective:  01.01.2020	Supersedes:  01.01.2017
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<b>Storage:</b>	It is recommended to store the material at ambient temperatures in closed containers and protected from sunlight and other sources of UV light.
<b>Shelf life:</b>	24 months
<b>Manufacturer:</b>	Dow Deutschland
<b>Manufacturing Site:</b>	Stade (Germany)

#### **Regulatory Compliance:**

##### **BSE/TSE / GMO / Kosher status / Aflatoxins:**

The material has no BSE/TSE risk, is not derived from GMO, is kosher and complies with the German "Aflatoxin Verbotverordnung".

##### **Residual solvents (Ph. Eur.\* 5.4 / USP\* <467> / ICH Q3C\*):**

Neither solvents of class 1 - 3 are used during manufacturing of this product, nor can they occur within the manufacturing process.

##### **Elemental Impurities (Ph. Eur.\* 5.20 / ICH Q3D\*):**

At least three independent batches have been analysed for elemental impurities according to guideline ICH Q3D. All determined values of elemental impurities were below the level of 30% of the permitted concentrations for parenteral application according to table A.2.2 of the guideline ICH Q3D. More detailed information is available upon request.

##### **Batch certification:**

Every batch is analysed according to all parameters of this specification (except characters). The Certificate of Analysis (CoA) provides all results like above including release date, date of manufacture, residual solvents statement and elemental impurities statement. All CoAs are signed by a Qualified Person according to EU-GMP or a responsible QA/QC-Manager.

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