



CERTIFICATE OF ANALYSIS

Laboratory Salinen Austria AG

PHARMASAL API

Sodium Chloride for pharmaceutical use
according to Ph. Eur, BP, USP, JP

Lot number: CRS060824
Retest date: 06.08.2027
Production date: 06.08.2024 - 10.08.2024

| | | Specification | Unit | Result | Unit |
|-------------------------------------|-------------|--------------------|--------|------------------|--------|
| Identification | Na+ | positive | | conforms | - |
| Identification | Cl- | positive | | conforms | - |
| Assay | NaCl | 99,5 - 100,5 | % | 99,97 | % |
| Bromides | Br- | <= 100 | ppm | <= 100 | ppm |
| Iodides | I- | <= 10 | ppm | <= 10 | ppm |
| | | | | conforms Ph. Eur | |
| Sulfate | SO4 2- | <= 200 | ppm | <= 200 | ppm |
| Phosphate | PO4 3- | <= 25 | ppm | <= 25 | ppm |
| Nitrite | NO2- | <= 0,01 | abs. | <= 0,01 | abs. |
| Heavy metals | as Pb | <= 3 | ppm | <= 3 | ppm |
| Iron | Fe | <= 2 | ppm | <= 2 | ppm |
| Aluminium | Al | <= 0,2 | ppm | <= 0,2 | ppm |
| Arsenic | As | <= 1 | ppm | <= 1 | ppm |
| Potassium | K | <= 500 | ppm | <= 500 | ppm |
| Barium | Ba | <= 10 | ppm | <= 10 | ppm |
| | | | | conforms Ph. Eur | |
| Magnesium & alkaline-earth metals | calc. as Ca | <= 100 | ppm | <= 100 | ppm |
| Ferrocyanides | [Fe(CN)6]4- | conforms | - | conforms | - |
| Insoluble matters | | <= 50 | ppm | <= 50 | ppm |
| Loss on drying | | <= 0,5 | % | <= 0,5 | % |
| Appearance of solution | | clear, colourless | | conforms | - |
| Acidity or Alkalinity | | conforms | | conforms | - |
| according to the regulations | | | | | |
| Residual Solvents | | Impossible due to | | conforms | - |
| according ICH-guideline | | production process | | | |
| Bacterial Endotoxins (Pyrogen free) | | < 5 | I.U./g | < 5 | I.U./g |
| TAMC | | <= 10 | CFU/g | <= 10 | CFU/g |
| TYMC | | <= 10 | CFU/g | <= 10 | CFU/g |

Appearance: white or almost white, crystalline powder or colorless crystals or white or almost white pearls
Solubility: freely soluble in water, practically insoluble in anhydrous ethanol

This lot conforms with the current Ph. Eur, USP, BP and JP monographs. In compliance with the guidelines on good manufacturing practice for active pharmaceutical ingredients (ICH Q7).
Store in a clean and dry place, nmt. 70% rel. Humidity.

It is suitable for the use in the manufacture of injectable dosage forms, peritoneal dialysis solutions, hemodialysis solutions and hemofiltration solutions.

Qualified Person: Birgit Spreitz
Date: 20.08.2024

Birgit Spreitz
20.08.2024



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