



## BATCH ANALYSIS CERTIFICATE - BP/Ph Eur GRADE

**Dosage Forms:** Suitable for the use in the manufacture of injectable dosage forms, peritoneal dialysis solutions, hemodialysis solutions and hemofiltration solutions.

**PRODUCT TYPE:** Pharmaceutical Sodium Chloride  
**CAS No.:** 7647-14-5  
**Molecular formula:** NaCl  
**Molecular Weight:** 58.44  
**BATCH/MFD<sup>1</sup>:** 13052025

**REPORT NO:** 32139  
**DATE PACKED:** 13-21/05/25  
**RETEST<sup>2</sup> DATE:** 13/05/2030

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### RESULTS

Tests Performed (Methods as per Pharmacopoeia)	RESULT			BP 2025/Ph Eur 11.7 (0193)
				SPECIFICATION
Appearance	Conforms			White or almost white, crystalline powder
Chlorides Identification	Conforms			Conforms
Sodium Identification	Conforms			Conforms
Appearance of Solution	Conforms			Clear and Colourless
Acidity	0.00	ml	(max)	NMT 0.5ml 0.01M NaOH
Alkalinity	0.20	ml	(max)	NMT 0.5ml 0.01M HCl
Bromides as Br	Conforms			100 ppm (max)
Ferrocyanides	Conforms			Conforms
Iodides	Conforms			Conforms
Nitrites	Conforms			Conforms
Phosphate as PO <sub>4</sub>	Conforms			25 ppm (max)
Sulphates as SO <sub>4</sub>	Conforms			200 ppm (max)
Aluminium as Al	Conforms			0.2 ppm (max)
Arsenic as As	Conforms			1 ppm (max)
Barium	Conforms			Conforms
Iron as Fe	Conforms			2 ppm (max)
Magnesium & Alkaline-earth metals	0.1	ml	(max)	NMT 2.5ml 0.01M EDTA (100 ppm max calculated as Ca)
Potassium as K	18	ppm	(max)	500 ppm (max)
Loss on Drying (at packing)	0.02	%w/w	(max)	0.5% w/w (max)
Bacterial endotoxins	<1.00	EU/g		<5 EU/g
Assay as NaCl	99.58			99.0-100.5% w/w
Residual Solvents	No class 1, 2 or 3 solvents are used in the manufacture of Pharmaceutical Sodium Chloride			
<b>ADDITIONAL ANALYSIS</b>	<b>ACTUAL RANGE</b>			< = Less Than
Filtration Rate (Minimum after 2 litres of 20.15% w/v brine solution)	Minimum	429	ml/min	NMT = Not more than
	Maximum	462	ml/min	u = micron

- Note:**
1. Batch/MFD in ddmmyyy format is the date manufacture commenced. This date is also the traceable batch code.
  2. Due to the stable nature of Sodium Chloride, Dominion Salt as permitted under PIC/s GMP guide PE-009 for API's, states a retest date in lieu of an expiry date.
  3. This product has been produced at Dominion Salt Mount Maunganui site
  4. All results reported are on a wet matter basis with the exception of the Assay test which is reported on a dried basis.
  5. The samples have been collected and analysed in accordance with our documented sample plan.
  6. All methods have been performed in Dominion Salt's on site Laboratory. Methods (unless stated) are documented in the Laboratory's Test Method Manual and will be made available upon request in writing to the Quality Manager
  7. Supply of this salt is subject to the terms of trade of Dominion Salt Limited upon request. The laws of New Zealand shall govern all disputes.
  8. Storage Conditions: Store unopened in clean, dry conditions

Conforms to Specification: TD

Release Date: 30.05.25

Approved by: \_\_\_\_\_

Eliska Veckova  
QHSE Officer

For Distribution use only:

"Life's most essential mineral, in the world's safest hands."

**BATCH ANALYSIS CERTIFICATE - INJ GRADE PYROGEN FREE U.S.P.**

*Dosage Forms:* Suitable for the use in the manufacture of injectable dosage forms, peritoneal dialysis solutions, hemodialysis solutions and hemofiltration solutions.

**PRODUCT TYPE:** Pharmaceutical Sodium Chloride  
**CAS No.:** 7647-14-5  
**Molecular formula:** NaCl  
**Molecular Weight:** 58.44  
**BATCH/MFD<sup>1</sup>:** 13052025

**REPORT NO:** 32140  
**DATE PACKED:** 13-21/05/25  
**RETEST<sup>2</sup> DATE:** 13/05/2030

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**RESULTS**


Tests Performed <i>(Methods as per Pharmacopoeia)</i>	RESULT			USP 2024 (47)-NF 42, Issue 1
				SPECIFICATION
Appearance	Conforms			White or almost white, crystalline powder
Chlorides Identification	Conforms			Conforms
Sodium Identification	Conforms			Conforms
Appearance of Solution	Conforms			Clear and Colourless
Acidity	0.00	ml	(max)	NMT 0.5ml 0.01M NaOH
Alkalinity	0.20	ml	(max)	NMT 0.5ml 0.01M HCl
Bromides as Br	Conforms			100 ppm (max)
Ferrocyanides	Conforms			Conforms
Iodides	Conforms			Conforms
Nitrites	Conforms			Conforms
Phosphate as PO <sub>4</sub>	Conforms			25 ppm (max)
Sulphates as SO <sub>4</sub>	Conforms			200 ppm (max)
Aluminium as Al	Conforms			0.2 ppm (max)
Arsenic as As	Conforms			1 ppm (max)
Barium	Conforms			Conforms
Iron as Fe	Conforms			2 ppm (max)
Magnesium & Alkaline-earth metals	0.1	ml	(max)	NMT 2.5ml 0.01M EDTA (100 ppm max calculated as Ca)
Potassium as K	18	ppm	(max)	500 ppm (max)
Loss on Drying (at packing)	0.02	%w/w	(max)	0.5% w/w (max)
Assay as NaCl	99.58			99.0-100.5% w/w
Residual Solvents	No class 1, 2 or 3 solvents are used in the manufacture of Pharmaceutical Sodium Chloride			

ADDITIONAL ANALYSIS	ACTUAL RANGE		
	Filtration Rate (Minimum after 2 litres of 20.15% w/v brine solution)	Minimum	429
	Maximum	462	ml/min

< = Less Than  
 NMT = Not more than  
 u = micron

- Note:**
1. Batch/MFD in ddmmyyyy format is the date manufacture commenced. This date is also the traceable batch code.
  2. Due to the stable nature of Sodium Chloride, Dominion Salt as permitted under PIC/s GMP guide PE-009 for API's, states a retest date in lieu of an expiry date.
  3. This product has been produced at Dominion Salt Mount Maunganui site
  4. All results reported are on a wet matter basis with the exception of the Assay test which is reported on a dried basis.
  5. The samples have been collected and analysed in accordance with our documented sample plan.
  6. All methods have been performed in Dominion Salt's on site Laboratory. Methods (unless stated) are documented in the Laboratory's Test Method Manual and will be made available upon request in writing to the Quality Manager
  7. Supply of this salt is subject to the terms of trade of Dominion Salt Limited upon request. The laws of New Zealand shall govern all disputes.
  8. Storage Conditions: Store unopened in clean, dry conditions

Conforms to Specification: TD

**Release Date:** 30.05.25  
 Approved by:   
 Eliska Veckova  
 QHSE Officer

For Distribution use only:

## BATCH ANALYSIS CERTIFICATE - JP GRADE

**Dosage Forms:** Suitable for the use in the manufacture of injectable dosage forms, peritoneal dialysis solutions, hemodialysis solutions and hemofiltration solutions.

**PRODUCT TYPE:** Pharmaceutical Sodium Chloride  
**CAS No.:** 7647-14-5  
**Molecular formula:** NaCl  
**Molecular Weight:** 58.44  
**BATCH/MFD<sup>1</sup>:** 13052025

**REPORT NO:** 32141  
**DATE PACKED:** 13-21/05/25  
**RETEST<sup>2</sup> DATE:** 13/05/2030

**PAGE:** 1 of 1

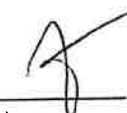
### RESULTS

Tests Performed <i>(Methods as per Pharmacopoeia)</i>	RESULT			JP XVIII supp.II (18th Edition)
				SPECIFICATION
Appearance	Conforms			White or almost white, crystalline powder
Appearance of Solution	Conforms			Clear and Colourless
Sodium Identification	Conforms			Conforms
Chlorides Identification	Conforms			Conforms
Acidity	0.00	ml	(max)	NMT 0.5ml 0.01M NaOH
Alkalinity	0.20	ml	(max)	NMT 0.5ml 0.01M HCl
Sulphates as SO <sub>4</sub>	Conforms			200 ppm (max)
Phosphate as PO <sub>4</sub>	Conforms			25 ppm (max)
Bromides as Br	Conforms			100 ppm (max)
Iodides	Conforms			Conforms
Ferrocyanides	Conforms			Conforms
Iron as Fe	Conforms			2 ppm (max)
Barium	Conforms			Conforms
Magnesium & Alkaline-earth metals	0.1	ml	(max)	NMT 2.5ml 0.01M EDTA (100 ppm max calculated as Ca)
Arsenic as As <sub>2</sub> O <sub>3</sub>	Conforms			2 ppm (max)
Loss on Drying (at packing)	0.02	%w/w	(max)	0.5% <sup>w/w</sup> (max)
Assay as NaCl	99.58			99.0-100.5% <sup>w/w</sup>
<b>ADDITIONAL ANALYSIS</b>	<b>ACTUAL RANGE</b>			< = Less Than
Filtration Rate (Minimum after 2 litres of 20.15% w/v brine solution)	Minimum	429	ml/min	NMT = Not more than
	Maximum	462	ml/min	u = micron

- Note:**
1. Batch/MFD in ddmmyyyy format is the date manufacture commenced. This date is also the traceable batch code.
  2. Due to the stable nature of Sodium Chloride, Dominion Salt as permitted under PIC/s GMP guide PE-009 for API's, states a retest date in lieu of an expiry date.
  3. This product has been produced at Dominion Salt Mount Maunganui site
  4. All results reported are on a wet matter basis with the exception of the Assay test which is reported on a dried basis.
  5. The samples have been collected and analysed in accordance with our documented sample plan.
  6. All methods have been performed in Dominion Salt's on site Laboratory. Methods are documented in the Laboratory's Test Method Manual and will be made available upon request in writing to the Quality Manager
  7. Supply of this salt is subject to the terms of trade of Dominion Salt Limited upon request. The laws of New Zealand shall govern all disputes.
  8. Storage Conditions: Store unopened in clean, dry conditions

Conforms to Specification: TD

**Release Date:** 30.05.25

Approved by:   
 Eliska Veckova  
 QHSE Officer

For Distribution use only:



## MICROBIOLOGICAL BATCH ANALYSIS CERTIFICATE

**PRODUCT TYPE:** Pharmaceutical Sodium Chloride  
**CAS No.:** 7647-14-5

**REPORT NO:** 32142  
**DATE PACKED:** 13-21/05/25  
**RETEST<sup>2</sup> DATE:** 13/05/2030  
**PAGE:** 1 of 1

**BATCH/MFD<sup>1</sup>:** 13052025

### RESULTS


TESTS PERFORMED (Methods as per Pharmacopoeia)	RESULT	SPECIFICATION
Bacterial Endotoxins (Pyrogen)	<1.00 EU/g	<5 EU/g
Microbial limit test:		
Total aerobic count	<10 CFU/g	not more than 100 CFU/g
Yeasts and Moulds	<1 CFU/g	not more than 100 CFU/g
Heat Resistant Microbes(@ 80°C for 10 mins)	ND Not detected/g	Not detected/g
Pathogenic Organisms:		
Escherichia Coli	ND Not detected/g	Not detected/g
Staphylococcus aureus	ND Not detected/g	Not detected/g
Pseudomonas aeruginosa	ND Not detected/g	Not detected/g
Salmonella spp.	ND Not detected/100g	Not detected/100g

ND = Not Detected or Absent

- Note: 1. Batch/MFD in ddmnyyyy format is the date manufacture commenced. This date is also the traceable batch code.  
 2. Due to the stable nature of Sodium Chloride, Dominion Salt as permitted under PIC/S GMP guide PE-009 for APIs, states a retest date in lieu of an expiry date .  
 3. All the above tests have been performed by laboratories contracted to Dominion Salt Mount Maunganui site (Original reports are available upon written request to the Quality Manager).  
 4. The samples have been collected and analysed in accordance with our documented sample plan.  
 5. < = less than > = greater than  
 6. Supply of this salt is subject to the terms of trade of Dominion Salt Limited upon request. The laws of New Zealand shall govern all disputes.

**Release Date:** 30.05.25

Conforms to Specification: TD

Approved by:   
 Eliska Veckova  
 QHSE Officer

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