



## Buffered Sodium Chloride Peptone Solution pH 7.0

LQ123

For the preparation of test suspension in accordance with harmonized methods of USP, EP, BP & JP.

### Composition\*\*

Ingredients	Gms / Litre
Peptone	1.000
Potassium dihydrogen phosphate	3.60
Disodium hydrogen phosphate	7.230
Sodium chloride	4.300

\*\*Formula adjusted, standardized to suit performance parameters

### Directions

Label the ready to use LQ123 bottle. Inoculate the sample and Incubate at specified temperature and time.

### Principle And Interpretation

The composition of this medium is as per USP (1) and is in accordance with the harmonized methodology of USP/EP/BP/JP (1,2,3,4). This medium is recommended for preparation of stable test strain suspension employed for validating the microbiological testing procedures of non-sterile products. The standardized stable suspensions are used so that the suitability of this test to detect microorganism in presence of product can be established. Non-fatty products insoluble in water and water-soluble products are diluted/dissolved using this solution. Peptone (meat or casein) serves as nutrient source and maintains the cell viability. Phosphates in the medium act as good buffering agents. Sodium chloride maintains the osmotic balance and cell integrity. Polysorbates reduce surface tension and also inactivate phenolic compound, if present in the test sample. Edel and Kampelmacher (6) noted that sub lethal injury to Salmonellae might occur in many food preservation processes. Pre-enrichment in Buffered Sodium chloride-Peptone solution pH 7.0 (LQ123L) at 35°C for 18-24 hours results in repair of injured cells (5). This medium supports the repair of injured cells that have sensitivity to low pH. It is also recommended for pre-enrichment and repair of injured cells (5).

### Quality Control

#### Appearance

Sterile clear Buffered Sodium Chloride Peptone Solution in bottle.

#### Colour

Colourless solution.

#### Quantity of medium

100ml of medium in bottle

#### pH

7.00- 7.00

#### Growth Promotion Test

In accordance with the harmonized method of USP/EP/BP/JP.

#### Sterility test

Passes release criteria

#### Cultural response

Cultural characteristics observed after an incubation at 30-35°C for 18-24 hours.

Organism	Growth	Inoculum (CFU)
<i>Candida albicans</i> ATCC 2091	luxuriant	50 -100
<i>Pseudomonas aeruginosa</i> ATCC 9027	luxuriant	50 -100
<i>Candida albicans</i> ATCC10231	luxuriant	50 -100
<i>Escherichia coli</i> ATCC 25922	luxuriant	50 -100
<i>Escherichia coli</i> ATCC 8739	luxuriant	50 -100
<i>Escherichia coli</i> NCTC 9002	luxuriant	50 -100
<i>Salmonella</i> Abony NCTC 6017	luxuriant	50 -100
<i>Salmonella</i> Typhimurium ATCC 14028	luxuriant	50 -100
<i>Bacillus subtilis</i> ATCC 6633	luxuriant	50 -100
<i>Staphylococcus aureus</i> ATCC25923	luxuriant	50 -100
<i>Staphylococcus aureus</i> ATCC 6538	luxuriant	50 -100
<i>Pseudomonas aeruginosa</i> ATCC 27853	luxuriant	50 -100

## Storage and Shelf Life

Store between 2-8°C. Use before expiry date on the label.

## Reference

1.The United States Pharmacopoeia, 2011, The United States Pharmacopoeial Convention. Rockville, MD. 2.British Pharmacopoeia, 2011, The Stationery office British Pharmacopoeia 3.European Pharmacopoeia, 2011, European Dept. for the quality of Medicines. 4.Japanese Pharmacopoeia, 2008. 5.Sadovski A.Y., 1977, J. Fd. Technol., 12:85. 6.Edel W. & Kampelmacher E.H.,1973, Bull, Wld. Hlth. Org., 48:167.



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