



Instruction Manual

HI 4007

HI 4107

Chloride Ion

Selective Electrode

Half-cell

Combination

HI 4007 Chloride Half-cell

HI 4107 Chloride Combination Electrode

I. Introduction:

The Hanna HI 4007 and HI 4107 are ion selective electrodes designed for the measurement of chloride ions in aqueous solutions. The HI 4007 is a solid state half-cell sensor that requires a separate reference. The HI 4107 is a combination ion selective electrode.

II. Specifications

Type: Solid State electrode with a Silver Chloride pellet.

Ion(s) measured: Chloride (Cl^-)

Measurement range: 1.0 M to 5×10^{-5} M
35,450 to 1.8 ppm

Interfering ions: Cyanide, and Mercury ions must be absent. Ratio of interfering ion to Cl^- must be less than the ratio indicated below:

1.0	for I^-	iodide
3.5	for Br^-	bromide
	for CO_3^{2-}	carbonate
1.0	for OH^-	hydroxide
0.01	for $\text{S}_2\text{O}_3^{2-}$	thiosulfate

Operating Temperature: 0-80°C

Operating pH: 2-11 pH

Dimensions: 12 mm (OD) X 120 mm
nominal insertion
(0.47" X 4.72")

Connection: BNC

III. Theory of Operation:

The HI 4007 or HI 4107 chloride electrodes are potentiometric devices used for the rapid determination of free chloride ions in ground water, boiler water and drinking water. The electrode functions as a sensor or ionic conductor. The HI 4007 requires a separate reference electrode to complete its electrolytic circuit. The HI 4107 has a reference electrode incorporated in its design. The silver chloride pellet is practically insoluble in the test solutions being measured and produces a potential change due to changes in the sample's ion activity. When the ionic strength of the sample is fixed by the addition of ISA, the voltage is proportional to the concentration of chloride ions in solution and the electrode follows the Nernst equation.

$$E = E_0 + 2.3 RT/nF \log A_{\text{ion}}$$

E = observed potential

E_0 = Reference and fixed internal voltages

R = gas constant (8.314 J/K Mol)

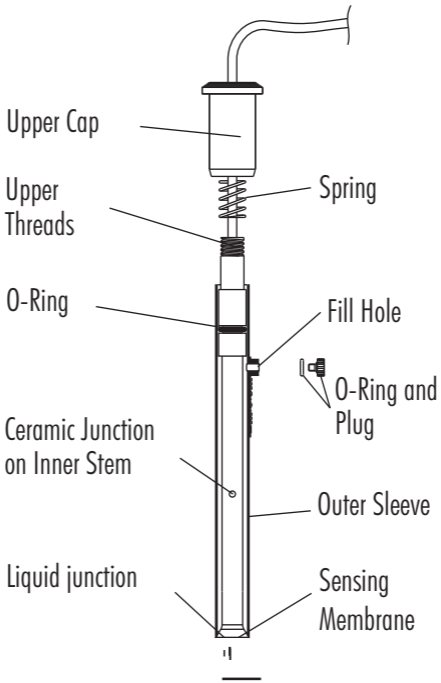
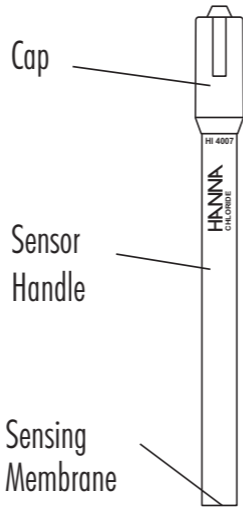
n = Charge on ion (-1)

A_i = ion activity in sample

T = absolute temperature in K

F = Faraday constant (9.648×10^4 C/equivalent)

IV. Design elements of the HI 4007 and HI 4107 electrodes



V. Equipment required:

- Hanna HI 5315 Double Junction Reference Electrode with HI 7072 Fill Solution for HI 4007.
- Hanna HI 4222 pH/ISE/mV meter or other suitable ion or pH/mV meter. (Note: log/linear graph paper is useful if an ISE (ion) meter is not available).
- Hanna HI 180 Magnetic Stirrer or equivalent with Teflon coated stirring bars (HI 731320). (Note: isolate beakers from stirrer motor heat by placing insulating material such as foam or cork between them).
- Hanna HI 76404 Electrode Holder or equivalent.
- Plastic beakers (HI 740036P) or other suitable measurement vessel.

VI. Solutions Required for Chloride Measurements

Select appropriate Hanna Instruments standard and ISA from the list below:

0.1 M Sodium Chloride Standard, 500mL HI 4007-01

100 ppm Chloride standard, 500 mL HI 4007-02

1000 ppm Chloride standard, 500 mL HI 4007-03

ISA, 500 mL HI 4000-00

Using volumetric pipettes and glassware make dilutions to bracket the concentration of the samples. Standards with concentrations $< 10^{-3}$ M (35.5 ppm) should be prepared daily.

Two mL of Hanna ISA for Halide electrodes (HI 4000-00) should be added to 100 mL of sample or standard.

A user prepared acetate pH buffer may also be used to adjust samples to approximately 4 pH. Add 10 mL per 100 mL sample or standard. Prepare 250 mL by dissolving 19.5 g ammonium acetate and 15.0 mL glacial acetic acid in 200 mL deionized water and diluting to volume.

A user supplied oxidizing reagent may be used to suppress interferences such as hydroxide, sulfide, bromide, ammonia or cyanide. Add 10 mL to 100 mL standard or sample. Permit sample to mix 10 minutes before measurements are taken. Discard samples and standards promptly as the chloride in these will oxidize upon standing making them unreliable. Do not permit electrodes to remain in this solution for long periods of time. Work in a hood and protect eyes and skin. Prepare oxidizing reagent by diluting 6.25 mL analytical grade concentrated nitric acid into approximately 800 mL deionized water. Mix carefully. Dilute to 1 liter.

VII. General Guidelines

- Calibration standards and sample solutions should have the same ionic strength. ISA, acetate buffer or oxidizing reagent should be added to both samples and standards in the same ratio.
- Calibration standards and sample solutions should be at same temperature.
- Thermally insulate beaker with standard or sample from magnetic stirrer.
- Calibration standards and sample solutions should be stirred at the same rate using identical sized TFE coated stir bars.
- Rinse electrode pair with distilled or deionized water between samples and gently dry off using soft disposable absorbent toweling. Do not rub sensor surface.
- Presoaking chloride sensor in a dilute standard will optimize response. Do not use concentration above $10^{-3}M$.
- A scratched, pitted, or tarnished pellet surface can cause drift, a loss of low level response, or poor repeatability. Optimum response can be restored by removing the damaged surface with the microabrasive strip HI 4000-70.

- Avoid large changes in temperature (thermal shock) as it may damage the sensor.

HI 4007

- Remove protective cover from sensor tip.
- Prepare HI 5315 reference electrode by filling electrolyte reservoir with HI 7072 fill solution.
- Place sensor and reference electrodes into electrode holder and connect cable connectors to meter.

HI 4107

- Remove the protective plastic wrap that covers the ceramic junction before assembling sensor for the first time.
- HI 7072 reference fill solution should be added daily to electrolyte reservoir before electrode use.
- During measurement always operate electrode with the fill hole open.
- During normal use, fill solution will slowly drain out of the tapered cone junction at the lower portion of the electrode. Excessive loss (> 4 cm drop within 24 hours) is not normal. If this occurs verify cap is tightened and the interface between the internal cone and outer body is free of debris.
- Add fill solution daily to maintain a good head pressure. For optimum response, this level should be maintained and not be allowed to drop more than 2-3 cm (1-inch) below fill hole. It must cover the ceramic found on the inner stem.
- If an erratic measurement occurs, check to see if foreign matter is seen trapped near the internal cone. Drain and refill with fresh fill solution.

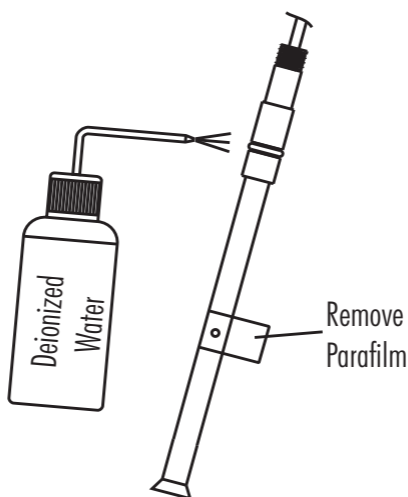
VIII. Electrode Preparation

HI 4007

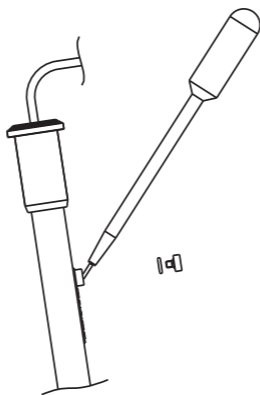
1. Remove protective cover from sensor tip.
2. Prepare reference electrode by filling outer electrolyte reservoir with HI 7072.
3. Place sensor and reference electrodes into electrode holder and connect cable connectors to meter.

HI 4107

1. Unwrap plastic film seal found over ceramic junction on inner stem and discard. This is only used for shipping and long term storage.
2. Rinse inner stem with deionized water making certain to wet the o-ring found on the inner stem.



3. Reassemble electrode by gently pushing the inner assembly into the outer body, sliding spring down cable, and screwing cap into place.
4. Remove fill hole cover and o-ring on fill hole spout.
5. Using the dropper pipette provided, add a few drops HI 7072 fill solution to the electrode, wetting the o-ring and rinsing out the fill solution chamber.



6. Holding the body of the electrode gently press upper cap with your thumb. This permits the fill solution to drain out of the body. Release cap and verify electrode returns to its original position. (You may need to gently assist for this to occur).



7. Tighten the electrode cap onto the body and fill electrode body until fill solution volume is just below fill hole.
8. Position electrode in a Hanna HI 76404 electrode holder (or equivalent) and connect plug to meter.

IX. Quick Check of Electrode Slope

- Connect sensors to pH/mV/ISE meter
- Place meter in mV mode.
- Place 100 mL of DIW into a beaker with stir bar.
- Place reference and measuring half-cell into prepared sample.
- Add 1 mL of a standard (0.1 M or 1000 ppm standard) to beaker. Record the mV value when stable.
- Add an additional 10-mL of standard to the solution. Record the mV when reading has stabilized. This value should be less than the previous noted (more negative).
- Determine the difference between the two-mV values. An acceptable value for this slope is -56 ± 4 mV.

X. Corrective action

- Verify protective cap has been removed.(HI 4007)
- Verify plastic film has been removed from inner stem.(HI 4107)
- Verify electrodes are connected properly to meter and meter is powered.
- Verify dilute standards are freshly made and stored. Remake solutions if appropriate.
- If the sensor slope just misses the suggested slope window, soaking the sensor in a dilute standard may solve the problem. (Choose 10^{-3} M chloride or 100 ppm standard).
- A scratched sensing surface can be polished with HI 4000-70 polishing strip. Cut off approximately 1 inch of the micro-abrasive strip. Wet the frosted side with deionized water and place against damaged membrane of the electrode. Place your thumb against the shiny backing and slowly rotate back and forth while applying gentle pressure. Continue polishing until you are satisfied with the surface.

If dark deposits appears on polishing strip move the paper slightly and continue polishing.

- If the sensor slope just misses the suggested slope window, soaking the sensor in a standard may solve the problem.
- If the membrane is damaged, the response becomes extremely sluggish, or the slope of the electrode has decreased significantly, and procedures above have not helped, the sensor should be replaced.

XI. Direct Calibration and Measurement

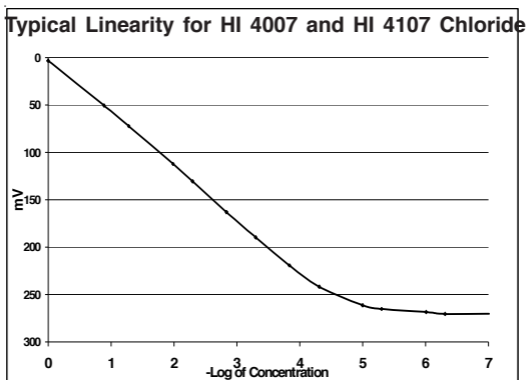
This method is a simple procedure for measuring many samples. A direct reading ISE meter (HI 4222 or equivalent) determines concentration of the unknown by a direct reading after calibrating the meter with the standards. The meter is calibrated with two or more freshly made standards that are in the linear measurement range of the unknowns. More calibration standards are required in non-linear regions. Unknowns are read directly. At very low levels of chloride, special precautions must be employed for reproducible measurements. Water used for standards must be chloride free and sensors and glassware must be rinsed repeatedly with this water to prevent carry over. In the region where the electrode response appears curved, more calibration points are needed, and calibration will need to be repeated more frequently.

A pH/mV meter in mV mode with semi log graph paper may also be used. Two or more freshly prepared standards that are in the measurement range of the unknowns are measured in mV mode on the meter.

These values are plotted on the semi-log paper and the points are connected to form a straight-line curve. When samples are measured, their mV values are converted to concentration by following the mV to the concentration axis on the semi-log plot.

Procedure

- 1) Follow sections VIII and IX to prepare sensors for measurement.
- 2) Follow section VI to prepare standards/ solution. Standards should bracket and fall within the range of interest.
Two mL HI 4000-00 ISA, or 10 mL acetate pH buffer or 10 mL of oxidizing reagent is added to 100 mL of both samples and standards. Add stir bar and mix before taking measurements. Measurements using the oxidizing reagent must be made 10 minutes after mixing and not permitted react longer.
- 3) Follow section VII; General Guidelines to optimize test set-up.
- 4) During calibration it is best to start with lower concentration samples first. Wait for a stable measurement before recording values. Slightly longer equilibrations are required at lower concentrations .
- 5) To prevent carry over and contamination of samples, rinse sensors with DIW and blot dry between samples.



XII. Other Measurement Techniques

Known Addition (for Cl⁻)

An unknown concentration can be determined by adding a known amount (volume and concentration) of measured ion to a known volume of the sample. This technique is called Known Addition. The method can use an ideal sensor slope, but actual determined slopes at the temperature of measurement should be used if known. This method is preprogrammed in the Hanna HI 4222 pH/ISE/mV meter, which simplifies the method greatly.

Example: Chloride ion determination with known addition.

1. A 50 mL sample of unknown (V_{sample}) is placed in a clean plastic beaker with a chloride sensor. Add 2 mL of acetate pH buffer or HI 4000-00 ISA (V_{ISA}) OR if chloride I electrode interferences are known to exist, add 50 mL oxidizing reagent (V_{ISA}). Mix well. and record the stable mV value. (mV 1)
2. 5 mL (V_{std}) of 10^{-3}M (C_{std}) standard is added to the beaker and the mV value decreases. The unknown chloride concentration in the original sample (C_{sample}) can then be determined by the following equation.

$$C_{\text{sample}} = \frac{C_{\text{standard}} V_{\text{standard}}}{(V_T)10^{\Delta E/S} - (V_{S'})} \left(\frac{V_{S'}}{V_{\text{sample}}} \right)$$

$$(V_{\text{sample}} + V_{\text{standard}} + V_{\text{ISA}}) = V_T$$

$$(V_{\text{sample}} + V_{\text{ISA}}) = V_{S'}$$

3. The procedure can be repeated with a second standard addition to verify slope and operation of the method.

Titration

A chloride electrode may be used as an indicator to follow the progress of a chloride titration with silver nitrate. The electrode can be used in colored samples where other indicators suffer from interferences. During the titration the sensor follows the decrease in chloride concentration while small additions of silver nitrate titrant are added. The silver reacts with the chloride ions forming a precipitate of silver chloride. At the stoichiometric end point, a large change in mV occurs. Measurements may be automated by use of the Hanna Titrator HI 901 or titrated manually.

XIII. pH

The HI 4107 and HI 4007 electrodes may be used in solutions with pH values between 2 and 11. Samples that fall beyond this range should be adjusted with acetate pH buffer. See Section VI.

XIV. Storage and Care of the HI 4007 and HI 4107 sensors

The HI 4007 sensor can be stored in very dilute standards ($< 10^{-4}$ M) for short periods of time and should be stored dry with the protective cap on when not in use.

The model HI 4107 combination electrode can be left in dilute standards ($< 10^{-4}$ M) for short time periods.

For long term storage, the electrode should be drained and washed of salts with distilled or deionized water. Unscrew the upper cap and move outer sleeve up cable. Wrap the ceramic junction on the inner stem with Parafilm® or other sealing wrap. Place the protective cap provided over the sensor membrane. Store dry disassembled electrode in storage box provided with electrode.

XV. Conversion tables

For Cl⁻

Multiply by

Moles/L (M) to ppm (mg/L)

3.545×10^4

ppm (mg/L) to M (moles/L)

2.821×10^{-5}



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WARRANTY

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