



pH is a scale of active hydrogen ion concentration in aqueous solution. In pure water, the concentration of hydrogen ions is 10^{-7} M. The pH scale converts this to a convenient positive number:

$$pH = -\log([a_{H^+}])$$

pH indicates acidity (pH<7), alkalinity (pH >7) or neutrality (pH = 7). Pure water undergoes autoprotolysis to yield equal numbers of hydrogen and hydroxide ions:

$$H_2O \longleftrightarrow H^+ + OH$$

The equilibrium constant at 25°C is:

$$K_W = [H^+] \times [OH^-] = 10^{-14}$$

pOH is the negative logarithm of the active hydroxide concentration. The equilibrium expression can be rewritten as:

$$pK_W = pH + pOH = 14$$

This equation applies for all aqueous systems. Very strong acids and bases can have pH's <0 and >14 respectively. Although pH is defined for aqueous solutions, measurements can be made in non-aqueous solutions provided the solvent is sufficiently conductive, or a conductive solvent is added to a non-conductive sample.

Construction

A standard pH electrode consists of two electrochemical half-cells usually co-axially combined into a single housing called a "combination electrode" to form an electrical circuit. The sensing half-cell comprises a pH sensitive glass membrane attached to a sealed insulating tube containing a solution of fixed pH in contact with a silver-silver chloride element. A voltage proportional to pH is measured

between the sensing half-cell and a reference half-cell which ideally maintains a stable potential independent of the sample. The following diagram represents the Intermediate Junction of Ionode's IJ Koala Series.



Figure 6

Part of the circuit is electronic (heavy lines) (see Figure 6). The remainder is ionic. Unlike other pH electrode measuring systems which rely on electron transfer such as antimony and quinhydrone, the pH glass electrode is insensitive to ORP.

pH glass is modified lithium silica which is a lithium ion conductor at room temperature. Hydrous glass ion exchange layers exist on both sides of the membrane (see Figure 7 - Figure 9). These allow reversible exchange between hydrogen and lithium ions.





Equal concentrations of H⁺ on either side of the membrane causes no potential to develop across the membrane (see Figure 7).



Figure 8

Unequal concentrations of H⁺on either side of the membrane cause the system to equalise the number of positive ions on either side of the membrane. This causes migration of a lithium ion through the membrane (see Figure 8).



Figure 9

As the negatively charged oxygen counter ions are fixed in the glass structure, a potential develops across the membrane (see Figure 9). This builds until the charge developed opposes further migration according to the Nernst equation.

The reference half-cell completes the circuit. The primary IJ reference is a silver/silver chloride element in a gelled saturated potassium chloride electrolyte. This makes solution contact with the secondary sleeve electrolyte through a low porosity wick. Usually, the sleeve electrolyte is concentrated potassium chloride. However, as the IJ reference has a double junction, it allows the use of other suitable sleeve electrolytes. Potassium chloride is the most commonly used reference electrolyte because of the similar size and mobility of the potassium and chloride ions. This minimizes charge separation when the sleeve electrolyte diffuses into the sample

Solution contact between the sleeve electrolyte and the sample is through the porous junction between the ground stem and the sleeve. This constriction allows uninhibited movement of ions between the sample and reference half cell to assure a repeatable reference junction potential. At the same time it does not grossly contaminate the sample with reference ions.



Figure 10

In common electrode designs, a porous restriction (typically a porous ceramic or fibre wick) is used to slow the flow (see Figure 10a).

If the restriction becomes clogged and movement of ions becomes inhibited, the electrode will appear to be stable in buffer solutions, but it may produce errors in non-ideal and low buffering capacity samples. The IJ KOALA reference system overcomes this by allowing free movement of ions past the restriction, allowing the junction to be cleaned and the sleeve electrolyte to be easily replaced. The result is a reference electrode system of assured reliability (see Figure 10b).

Output

The voltage output is related to the hydrogen ion activity by the Nernst equation:

$$E = E_0 + \frac{RT}{nF} \ln\left(\left[H^+ \right] \right)$$

This can be rewritten in linear form by substituting in the definition of pH and grouping all the constants to give:

$$E = E_0' - SLOPE(T) \times pH$$

 $E_0^{'}$ is referred to as the isopotential point. Theoretically, it is the pH which has no temperature dependence.

The IJ44 has been designed so that it is compatible with all modern pH meters.



Offset

Each pH electrode has a small offset which varies from electrode to electrode. It is due to an "asymmetry potential" across the membrane. It is usually within ±0.2pH for a new electrode, and requires correction by calibration in a buffer at or near pH 7. The offset increases as the electrode ages and is also an indication of the condition of the electrode, especially cleanliness. Most pH electrodes have pH0 value of close to pH 7.0.

Slope

The SLOPE is a function of temperature (T) and contains the conversion of the natural logarithm to the base ten logarithm. It is defined as the number of mV per unit of pH. It is manually or automatically (ATC) compensated in pH meters.

Table 1: Theoretical (Nernstian) Slope

Т	SLOPE(T)	Т	SLOPE(T)
(°C)	(mV)	(°C)	(mV)
0	54.197	30	60.149
5	55.189	35	61.141
10	56.181	38	61.737
15	57.173	40	62.133
20	58.165	45	63.126
25	59.157	50	64.118

A new electrode should have a slope between 95-102% of theoretical. Sub-Nernstian slope is corrected by calibration with a second buffer. As the pH membrane ages, the slope decreases. This affects the accuracy.

For best performance, the slope should be >95%. Some meters calculate and display % slope after calibration. It can also be accurately measured using the mV mode of a pH meter.

1) Measure the temperature of two buffers and select the pH values for the buffers at this temperature (see <u>Table 2</u>: pH Values of Standard Buffers at Various Temperatures)

2) Measure the mV readings for the two buffers. Allow several minutes for the electrode to attain thermal equilibrium with the buffers.

3) Divide the measured difference by the theoretical difference (see <u>Table 1</u>) and multiply by 100. Example:

Buffer 1 (pH 6.865 @ 25°C): +11 (mV) Buffer 2 (pH 4.008 @ 25°C): +174 (mV) SLOPE(T=25°C): 59.157 (mV/pH) % Slope = $\frac{(174-11)}{(6.865-4.008) \times 59.157} \times 100 = 96.4\%$

Note that ATC provided by pH meters only corrects the temperature term in the Nernst equation. It does not correct the temperature coefficients of buffers unless they are specified and the meter software calculates accordingly. With the exceptions of fully dissociated strong acids and bases, all samples have temperature coefficients as well. These vary depending on the nature of the sample and cannot be corrected by the pH meter. For best reporting, the temperature at which the measurement has been made should be stated.

Sodium Error

The selectivity of H^+ over Na^+ is extremely high, but small errors become apparent at high pH (i.e. low H^+ concentration) and high Na^+ concentration.

Corrections to the measurement can be made if the sodium concentration of the sample is known. The sodium error increases with aging.

A measured pH can be corrected by locating it on the x-axis and drawing a line straight up until it intersects the sodium concentration line of the sample (extrapolate between lines for sodium concentrations not drawn). Draw a line from the intersection directly over to the y-axis to determine the error in the measurement. The actual pH of the sample is determined by adding the error to the measured pH. For example, if a sample with 1M Na⁺ concentration measures pH 12.6, then the actual pH is 12.77.

The sodium error is larger in the "A" glass pH electrodes. As a consequence, "A" glass pH electrodes are best suited to pH measurements below pH12. The "A" glass would be preferred for measurements in strong acids, titrations and for measurement in lower conductivity samples.

A graph showing sodium error response for Ionodes' C Glass formulation is reproduced below.

Glass C Sodium Error



Response Time

This varies according to the pH glass composition, membrane wall thickness, pH meter filtering, buffering capacity of the sample, measurement temperature (longest at <10C), lack of thermal equilibrium between the electrode and buffers/samples, cleanliness of the junction, and the age, chemical exposure history and cleanliness of the membrane. Most increases in response time for new electrodes are due to the latter.

Impedance

pH membranes typically have resistances of 100 – 1000 MOhm. This requires the pH meter to have an input impedance of at least 10¹²Ohms in order to avoid signal loss and polarisation of the electrode. The latter occurs when a significant current flows through the electrode. This alters the Ag/AgCl/sat KCl half cell potentials and permanently damages the membrane. Polarisation can be caused by damp or shorted connectors, poor connector insulation, a low impedance meter or meter circuit malfunction.

THEORY

ORP Theory (/en/theory/orp-theory)

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Ion Selective Theory (/en/theory/ion-selective-theory)
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Conductivity Theory (/en/theory/conductivity-theory)

pH Theory (/en/theory/ph-theory)



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IJ Series Intermediate Junction Electrodes

Setting the benchmark in super-long life, versatility and value for money since 1972, the flagship of the lonode Lab sensor range is the renowned IJ Series.

Built to exacting standards and available in pH, ISE, ORP and Reference, the IJ Series benefits are many ... A removable sleeve and renewable junction for longer life and easy cleaning, an isolated secondary reference to effectively defeat contamination, an annular ground-glass junction to provide

fast response times and resistance to clogging, selectable electrolyte and an optional toughened glass spear membrane for direct penetration. An exceptional choice in virtually any sample, the IJ Series is unequalled in food, wine, dairy, contaminated waste samples, industrial, mining, petro-chemical, electro plating and countless more. The IJ Series also features models with solvent resistant bodies and high temperature housings for the most demanding of environments.

Replaceable electrolyte

Removeable sleeve

Avaliable for pH, ISE & ORP

Polypropylene body

Way back In 1972,

History of the Intermediate Junction

Back in 1972, Ionode's R&D Chemists and Engineers set out to develop a sensor that would solve the the common problems of contamination and clogging etc. but still be tough, fast and super reliable. Working together, they decided it was possible and they set to work on what would become the IJ Series ... a sensor that would defeat contamination, work in any sample and have a lifetime far in excess of other single or double junction sensors. The benchmark in sensor versatility, reliability and performance was then born in the IJ Series. Featuring an renewable intermediate junction for easy cleaning, Ionode's proprietary glass formulae and a tough yet flexible body, the IJ Series quickly earned it's reputation for having the longest life in the industry and being the best value sensor available.

More Junctions, Less Contamination

IJ have unique junction mechanics which protect the primary reference from the contamination associated with traditional gel electrodes whilst providing the low maintenance and long life of a refillable sensor.

Removeable Sleeve

IJ electrodes have a removeable sleeve to make cleaning quick and easy.



IJ's Explained Why you need an intermediate junction probe

IJ's have unique junction mechanics which protect the primary reference from the contamination associated with traditional gel electrodes whilst providing the low maintenance and long life of a refillable sensor.





The IJ Range The full range of IJ electrodes

Model	Parameter	Measuring Range	Temp. Range	Sensor Type	Temperatur ← Swipe to see more information Compensat
IJ14	Ref		0 - 60	Reference	n/a
IJ44- Spear	рН	рН 0 - 14	0 - 60	A glass spear	Optional
IJ44- HT- Spear	рН	pH 0 - 14	0 - 100	A glass spear	Optional
IJ40	рН	рН 0 - 14	0 - 60	A glass bullet	Optional

IJ40- ALK	рН	рН 0 - 14	0 - 60	C Glass bullet	Optional
IJ40- HT- ALK	рН	рН 0 - 14	0 - 100	C Glass bullet	Optional
IJ64	ORP	±2000mV	0 - 60	Platinum Wire	n/a
IJ64D	ORP	±2000mV	0 - 60	Platinum Disc	n/a
IJ64- HT	ORP	±2000mV	0 - 100	Platinum Wire	n/a
IJAg	Ag⁺, Cl⁻	±2000mV	0 - 60	Silver Billet	n/a
IJAu	ORP	±2000mV	0 -60	Gold Wire	n/a

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Model	Parameter	Measuring Range	Temp. Range	Main Interferences	Slope
IJ-F	F ⁻	0.2 - 19,000 ppm	0 - 60	OH-	57mV/decade ± 5mV
IJ-Ag2S	S ²⁻	0.003 - 32,000 ppm	0 - 60	Hg ^{+/2+}	27mV/decade ± 3mV
	Ag+	0.01 - 108,000 ppm			57mV/decade ± 5mV
IJ-Cl	CI⁻	1.8 - 35,500 ppm	0 - 60	Br⁻, I⁻, S²⁻, CN⁻, S ₂ O ₃ ²⁻	57mV/decade ± 5mV
IJ-Br	Br⁻	0.2 - 79,900 ppm	0 - 60	I⁻, S²⁻, CN⁻	57mV/decade ± 5mV
IJ-I	l-	0.005 - 127,000 ppm	0 - 60	S ²⁻ , CN⁻	57mV/decade ± 5mV
IJ-CN	CN	0.2 - 260,000 ppm	0 - 60	S ²⁻ , I⁻	57mV/decade ± 5mV