INTRODUCTION TO CO2 MEASUREMENTS IN SEA WATER

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METHODS FOR MEASURING THE CO2 PARAMETERS IN SEA WATER

USUAL PARAMETERS MEASURED

- Total dissolved inorganic carbon $C_T = [CO_2] + [HCO_3^-] + [CO_3^{2-}]$
- Total hydrogen ion concentration (pH)

 $pH = -lg[H^+]$

- Partial pressure of $CO₂$ (in air in equilibrium with sea water) $p(CO_2) = x(CO_2) p = [CO_2] / K_0$
- Total alkalinity

 $A_T = [HCO_3^-] + 2[CO_3^{2-}] + [B(OH)_4^-] + [OH^-] - [H^+]$

TOTAL DISSOLVED INORGANIC CARBON

• Carbon dioxide mass balance equation $C_T = [CO_2] + [HCO_3^-] + [CO_3^{2-}]$] *T, p independent*

UNITS: moles per kilogram of solution (usually µmol kg⁻¹)

TOTAL DISSOLVED INORGANIC CARBON

• Carbon dioxide mass balance equation $C_T = [CO_2] + [HCO_3^-] + [CO_3^{2-}]$

] *T, p independent*

- 1. Acidify a known amount of sample
- 2. Extract the $CO₂$
- 3. Measure the amount of $CO₂$
	- Coulometry
	- Infra-red analyzer
	- Other

TOTAL HYDROGEN ION CONCENTRATION (PH)

• Species concentration, [H+]

 $pH = -lg[H^+]$

] *T, p dependent*

UNITS: pH is dimensionless

But, total hydrogen ion concentration is in moles per kilogram of solution

TOTAL HYDROGEN ION CONCENTRATION (PH)

• Species concentration, [H+]

 $pH = -lg[H^+]$

] *T, p dependent*

- 1. Electrometric measurement of pH
- 2. Spectrophotometric measurement of pH

ELECTROMETRIC PH MEASUREMENT

A direct electrometric technique using an electrode sensitive to hydrogen ion in a "pH cell";

The Nernst equation underlies this approach to measuring pH:

$$
E = E^*(S, T, p) - \left(\frac{RT\ln 10}{F}\right) \lg[H^+]
$$

$$
= E^*(S, T, p) + \left(\frac{RT\ln 10}{F}\right) pH
$$

$$
pH(X) = pH(S) - \frac{E_X - E_S}{RT\ln 10/F}
$$

]

For this operational approach to give an accurate estimate of [H+], it is essential that the calibration standard be matched closely to the sample in composition. **IT RARELY IS !**

To use autonomously (not constant *S, T, p*), it is essential that the functional form of *E**(*S, T, p*) be known well. **IT OFTEN ISN'T!**

SPECTROPHOTOMETRIC PH MEASUREMENT

A colorimetric technique in which an $\int \frac{\mathcal{E}_1(I^{2-})}{\sqrt{I^{1.1}}}$ $A_1 / A_2 - \varepsilon_1 (H I^{-}) / \varepsilon_2 (H I^{-})$ indicator dye is added to the solution $\sqrt{ }$ \setminus $\sqrt{ }$ \setminus $pH = -lg K(HI^{-})$ $\frac{1}{2}$ + lg $\overline{}$ \lfloor $\varepsilon_{2}(\mathrm{HI}^{-})$ $1-(A_1/A)\varepsilon_2(I^{2-})/\varepsilon_1(I^{2-})$ $\overline{\mathcal{K}}$ $\overline{}$ $\overline{\mathcal{K}}$ $\overline{}$ and the pH inferred from the resulting absorbance spectrum 2.5 45000
40000 **Spectrum of** *m***-cresol** $H₂I$ **purple in Tris buffer in** 2.0 **seawater** 35000 **Scaled Absorbance** Scaled Absorbance Extinction coefficient 30000 1.5 25000 20000 $H₁$ 1.0 15000 HI^- = $H^+ + I^{2-}$ 10000 0.5 $A_{\lambda} = \varepsilon_{\lambda}(\mathrm{H}\mathrm{I}^{-}) [\mathrm{H}\mathrm{I}^{-}] + \varepsilon_{\lambda}(\mathrm{I}^{2-}) [\mathrm{I}^{2-}]$] 5000 0.0 Ω 400 450 500 550 600 650 700 750 500 550 600 700 750 450 650 400 wavelength / nm wavelength / nm

POTENTIAL IMPLICATIONS OF IMPURE DYE

ARTICLE

pubsacs.org/est

Purification and Characterization of meta-Cresol Purple for Spectrophotometric Seawater pH Measurements

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pid and precise measurements acid-base indicators used in couracy. This work describes eta-cresol purple (mCP), and nodynamic equilibrium conperature (T) and salinity (S) . on concentration scale (pH_F) nce ratios $(R = \lambda A/\lambda A)$ as

Thus far the uncertainty introduced by dye impurities has only been characterized at a single temperature (25 °C) and salinity (35) .

COMPARISON OF TECHNIQUES

 $*$ The Honeywell DuraFET® seems to have a significantly more stable calibration than a conventional pH cell.

PARTIAL PRESSURE OF CO₂ (IN AIR THAT IS IN EQUILIBRIUM WITH THE WATER SAMPLE)

• Species concentration, $[CO₂]$

 $CO₂(g) = CO₂(aq)$

 $[CO₂]$ $x(\mathrm{CO}_2)\cdot p$

mole fraction of CO2

equilibration pressure

 $p(CO_2) = x(CO_2) p = [CO_2] / K_0$

T, p dependent

UNITS: pressure units (usually µatm)

PARTIAL PRESSURE OF CO₂ (IN AIR THAT IS IN EQUILIBRIUM WITH THE WATER SAMPLE)

• Species concentration, [CO₂]

 $p(CO_2) = x(CO_2) p = [CO_2] / K_0$ **T, p dependent**

- A. Gas phase equilibration
	- Measure $x(CO_2)$ in the gas phase using NDIR
	- Measure *x*(CO₂) in the gas phase using GC
- B. Membrane equilibration (with gas or external solution)
	- Measure $x(CO_2)$ in the gas phase using NDIR
	- Measure pH in external solution (*see pH section*)

* These values are for commercial systems when working well.

TOTAL ALKALINITY

 $A_{\text{T}} = [\text{HCO}_3^-] + 2[\text{CO}_3^{2-}] + [\text{B(OH)}_4^-] + [\text{OH}^-] - [\text{H}^+]$

T, p independent

UNITS: moles per kilogram of solution (usually µmol kg⁻¹)

TOTAL ALKALINITY

• Hydrogen ion mass-balance equation $A_T = [HCO_3^-] + 2[CO_3^{2-}] + [B(OH)_4^-] + [OH^-] - [H^+]$] *T, p independent*

Acidimetric titration

- Closed-cell
- Open-cell
- a. Can also add acid in one aliquot (single point method)
- b. Can use either a pH cell or spectrophotometry to estimate [H+] during titration.
- c. Ideally use NLLS to locate equivalence point; can use Gran technique

COMPARISON OF TECHNIQUES

* These values are for commercial systems when working well.

STATUS OF CURRENT MEASUREMENT PRACTICE

SAMPLING & ANALYSIS STRATEGIES

Discrete samples (collected from desired locations, preserved, then analyzed later in a laboratory)

Automated on-line analysis (*e.g.* a stream of sea water pumped into the instrument)

Instruments operated in a profiling mode, *e.g.* as part of a CTD package

Autonomous instruments able to carry out their analyses remotely in the ocean

DEVELOPMENT STATE OF $CO₂$ MEASUREMENT SYSTEMS

Level 0 – No working system available

- Level 1 Prototype system available in single lab
- Level 2 2nd generation prototype in use
- Level 3 Home-built systems in a number of labs
- Level 4 Can be purchased commercially
- Level 5 Commercially available, reliable, and fully supported

ESTIMATED UNCERTAINTIES[†] FOR DISCRETE MEASUREMENTS

[†] Based on measuring surface oceanic CO₂ levels

Contents lists available at ScienceDirect

Marine Chemistry

journal homepage: www.elsevier.com/locate/marchem

An inter-laboratory comparison assessing the quality of seawater carbon dioxide measurements Will make available in the Dropbox

MARINE CHEMISTRY

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article info abstract

Article history: Received 25 April 2014 Received in revised form 7 January 2015 Accepted 4 February 2015 Available online 11 February 2015

Keywords: Carbonate chemistry Intercomparison Carbon dioxide Alkalinity Dissolved inorganic carbon

pH Ocean acidification

Seawater $CO₂$ measurements are being made with increasing frequency as interest grows in the ocean's response to changing atmospheric $CO₂$ levels and to climate change. The ultimate usefulness of these measurements depends on the data quality and consistency. An inter-laboratory comparison was undertaken to help evaluate and understand the current reliability of seawater $CO₂$ measurements. Two seawater test samples of different $CO₂$ content were prepared according to the usual method for the creation of seawater reference materials in the Dickson Laboratory at Scripps Institution of Oceanography. These two test samples were distributed in duplicate to more than 60 laboratories around the world. The laboratories returned their measurement results for one or more of the following parameters: total alkalinity (A_T) , total dissolved inorganic carbon (C_T) , and pH, together with information about the methods used and the expected uncertainty of the measurements. The majority of laboratories reported A_T and C_T values for all their measurements that were within 10 µmol kg⁻¹ of the assigned values (i.e. within \pm 0.5%), however few achieved results within 2 µmol kg⁻¹ (i.e. within \pm 0.1%), especially for C_T . Results for the analysis of pH were quite scattered, with little suggestion of a consensus value. The high- $CO₂$ test sample produced results for both C_T and pH that suggested in many cases that $CO₂$ was lost during analysis of these parameters. This study thus documents the current quality of seawater $CO₂$ measurements in the various participating laboratories, and helps provide a better understanding of the likely magnitude of uncertainties in these measurements within the marine science community at the present time. Further improvements will necessarily hinge on adoption of an improved level of training in both measurement technique and of suitable quality control procedures for these measurements.

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RECENT INTER-LABORATORY PROFICIENCY STUDY

Assigned values for total alkalinity, total dissolved inorganic carbon, and pH (25 °C; total scale) for the two test samples. Values are expressed as *mean* ± *standard deviation* (*number of analyses*).

Bockmon & Dickson, 2015

Bockmon & Dickson, 2015

RECENT INTER-LABORATORY PROFICIENCY STUDY

Bockmon & Dickson, 2015

FASTER, BETTER, CHEAPER

In 1992, NASA Administrator Daniel Goldin began the agency's "Faster, Better, Cheaper" initiative.

The popular consensus on "Faster, Better, Cheaper" is often expressed in the supposedly self-evident saying: "faster, better, cheaper — pick two."

Is this necessarily true for seawater carbonate system measurements?

We clearly understand the meanings of "faster" and "cheaper", but how should we define "better"?

DESIRE FOR HIGH-QUALITY MEASUREMENTS

WHAT IS QUALITY?

Quality is fitness for purpose.

Fitness for purpose: the property of data produced by a measurement process that enables a user of the data to make technically correct decisions for a stated purpose.

Fitness for purpose therefore refers to the magnitude of the uncertainty associated with a measurement in relation to the needs of the application area.

Has implications for the level of resources needed!

WHEN ASSESSING "BETTER", UNCERTAINTY IS NOT THE ONLY MEASURE!

YOU SHOULD CONSIDER:

- 1. Overall uncertainty required
- 2. Cost of measurement(s)
	- a. Cost of equipment (purchase / maintenance)
	- b. Cost of training
	- c. Cost of analysis (time & materials)
- 3. Convenience of measurement
	- a. Availability of equipment (purchase / maintenance)
	- b. Sample size required
	- c. Time until results are available
- 4. Cost of making a wrong measurement.

http://cdiac.ornl.gov/oceans/Handbook_2007.html

FOR OPEN-OCEAN MEASUREMENTS

YOU SHOULD CONSIDER:

- 1. Overall uncertainty required AS GOOD AS POSSIBLE!
- 2. Cost of measurement(s) MONEY NO OBJECT?
	- a. Cost of equipment (purchase / maintenance)
	- b. Cost of training
	- c. Cost of analysis (time & materials)
- 3. Convenience of measurement WOULD BE NICE!
	- a. Availability of equipment (purchase / maintenance)
	- b. Sample size required
	- c. Time until results are available
- 4. Cost of making a wrong measurement. VERY HIGH!

FOR OCEAN ACIDIFICATION MEASUREMENTS

YOU SHOULD CONSIDER:

- 1. Overall uncertainty required STILL NEEDS THOUGHT
- 2. Cost of measurement(s) BE NICE IT IT WERE CHEAP!
	- a. Cost of equipment (purchase / maintenance)
	- b. Cost of training
	- c. Cost of analysis (time & materials)
- 3. Convenience of measurement PLEASE!!
	- a. Availability of equipment (purchase / maintenance)
	- b. Sample size required
	- c. Time until results are available
- 4. Cost of making a wrong measurement. NEEDS THOUGHT

KEY QUESTIONS TO ASK ABOUT ANY ANALYTICAL METHOD

KEY QUESTIONS

- 1. How does the method work?
- 2. How is it calibrated?
- 3. What is the overall uncertainty of the measured value?
- 4. Does it provide the quality I need?
- 5. What does the method cost to use?

WE SHALL DISCUSS THIS MORE DURING THE WEEK

