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NDP-066

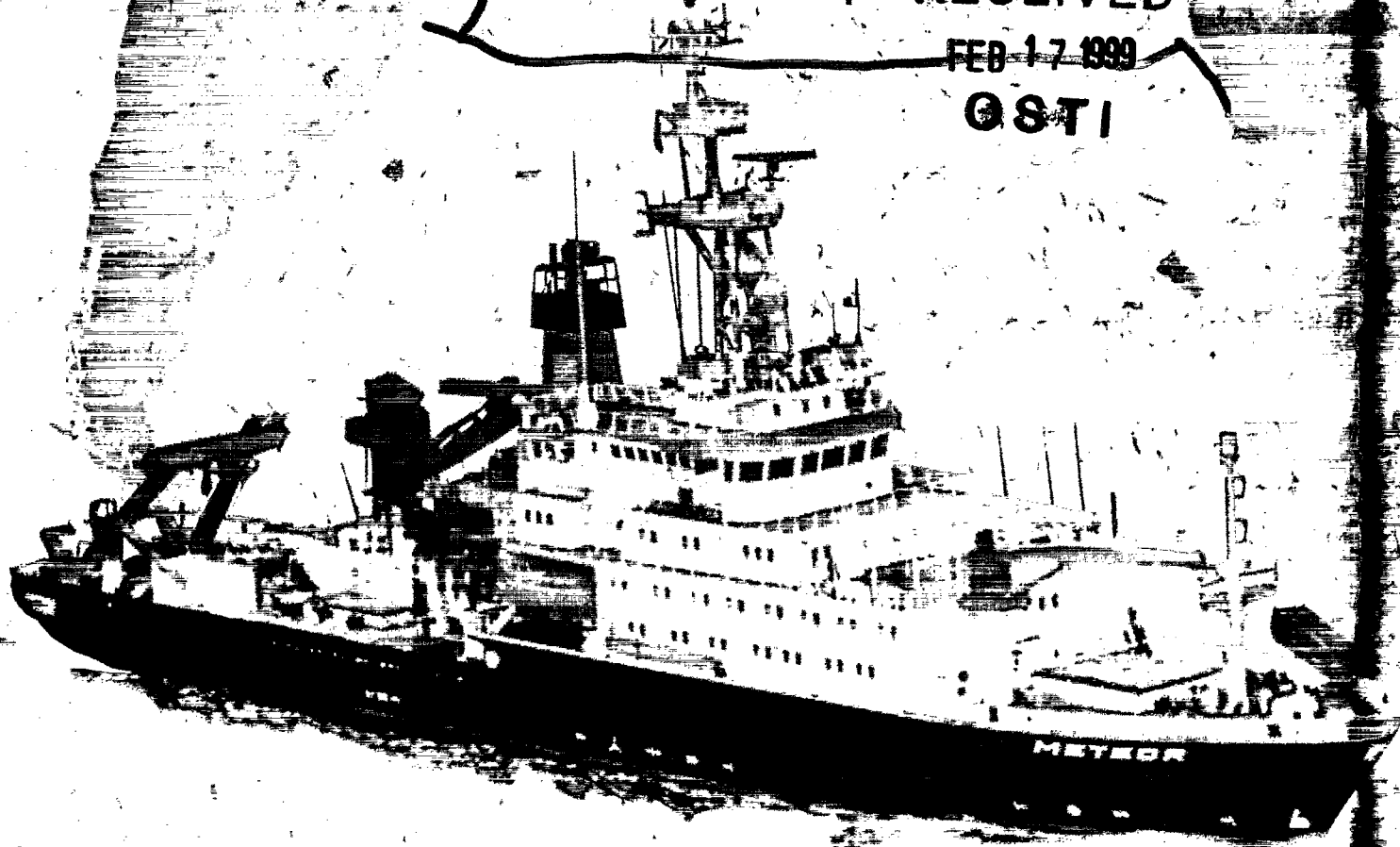
Carbon Dioxide, Hydrographic, and Chemical Data Obtained During the R/V Meteor Cruise 22/5 in the South Atlantic Ocean

(WOCE Section A10, December 1992 - January 1993)

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**CARBON DIOXIDE, HYDROGRAPHIC, AND CHEMICAL DATA OBTAINED
DURING THE R/V *METEOR* CRUISE 22/5 IN THE SOUTH ATLANTIC OCEAN
(WOCE SECTION A10, DECEMBER 1992-JANUARY 1993)**

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**Environmental Sciences Division
Publication No. 4814**

Date Published: December 1998

**Prepared for the
Environmental Sciences Division
Office of Biological and Environmental Research
U.S. Department of Energy
Budget Activity Numbers KP 12 04 01 0 and KP 12 02 03 0**

**Prepared by the
Carbon Dioxide Information Analysis Center
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managed by
LOCKHEED MARTIN ENERGY RESEARCH CORP.
for the
U.S. DEPARTMENT OF ENERGY
under contract DE-AC05-96OR22464**

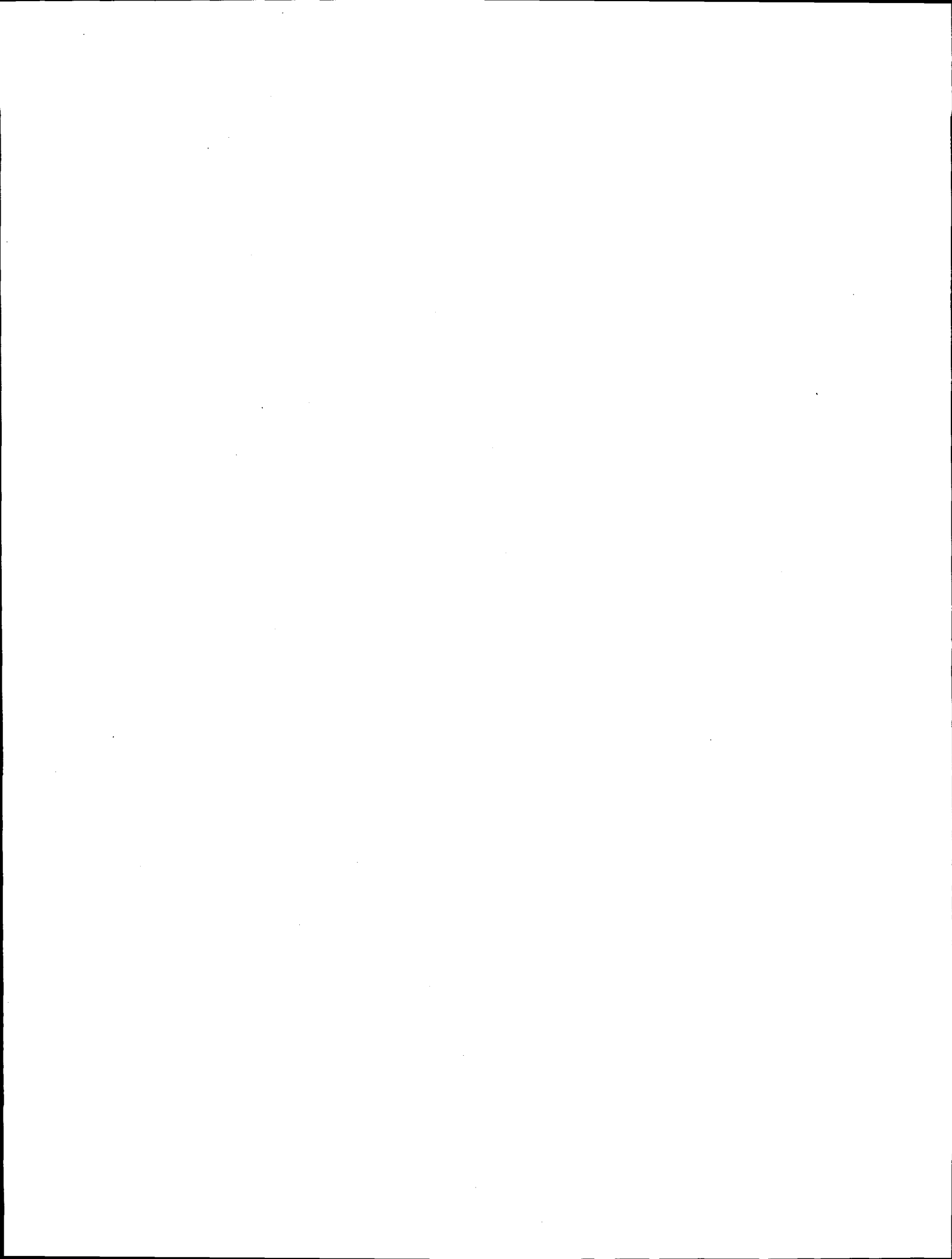
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ABSTRACT

Johnson, K. M., B. Schneider, L. Mintrop, D. W. R. Wallace, and A. Kozyr (ed.). 1998 Carbon Dioxide, Hydrographic, and Chemical Data Obtained During the R/V *Meteor* Cruise 22/5 in the South Atlantic Ocean (WOCE Section A10, December 1992–January 1993). ORNL/CDIAC-113, NDP-066. Carbon Dioxide Information Analysis Center, Oak Ridge National Laboratory, U.S. Department of Energy, Oak Ridge, Tennessee. 66 pp.

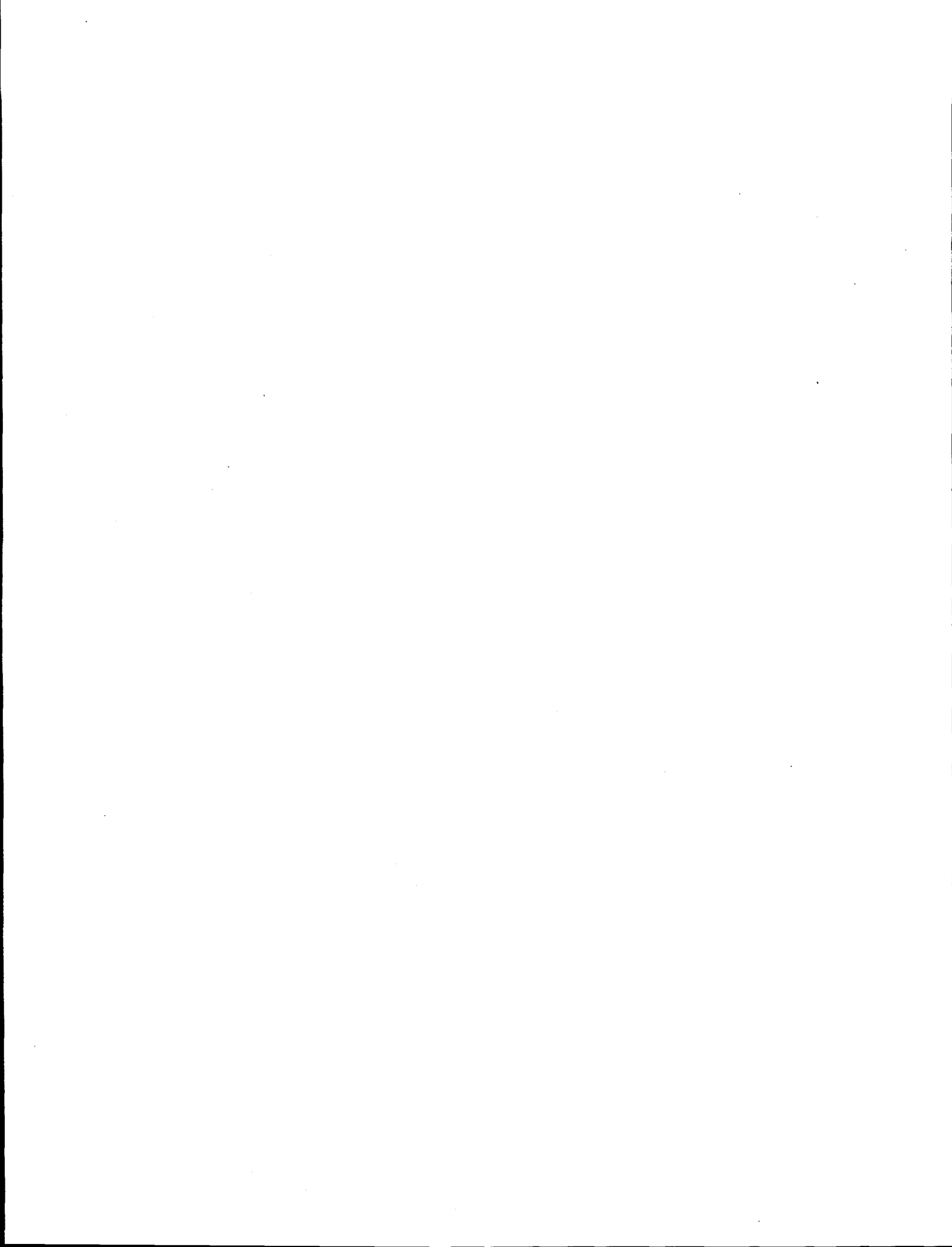
This data documentation discusses the procedures and methods used to measure total carbon dioxide (TCO_2) and total alkalinity (TALK) at hydrographic stations, as well as the underway partial pressure of CO_2 (pCO_2) during the R/V *Meteor* Cruise 22/5 in the South Atlantic Ocean (Section A10). Conducted as part of the World Ocean Circulation Experiment (WOCE), the cruise began in Rio de Janeiro on December 27, 1992, and ended after 36 days at sea in Capetown, South Africa, on January 31, 1993. Measurements made along WOCE Section A10 included pressure, temperature, and salinity [measured by conductivity, temperature, and depth (CTD) sensor], bottle salinity, bottle oxygen, phosphate, nitrate, nitrite, silicate, chlorofluorocarbons (CFC-11, CFC-12), TCO_2 , TALK, and underway pCO_2 .

The TCO_2 was measured by using two Single-Operator Multiparameter Metabolic Analyzers (SOMMAs) for extracting CO_2 from seawater samples that were coupled to a coulometer for detection of the extracted CO_2 . The overall precision and accuracy of the analyses was $\pm 1.9 \mu\text{mol/kg}$. Samples collected for TALK were measured by potentiometric titration; precision was $\pm 2.0 \mu\text{mol/kg}$. Underway pCO_2 was measured by infrared photometry with a precision of $\pm 2.0 \mu\text{atm}$. The work aboard the R/V *Meteor* was supported by the U.S. Department of Energy under contract DE-ACO2-76CH00016, and the Bundesministerium für Forschung und Technologie through grants 03F0545A and MFG 099/1.

The R/V *Meteor* Cruise 22/5 data set is available free of charge as a numeric data package (NDP) from the Carbon Dioxide Information Analysis Center. The NDP consists of three oceanographic data files, three FORTRAN 77 data-retrieval routines, a documentation file, and this printed documentation, which describes the contents and format of all files as well as the procedures and methods used to obtain the data. Instructions on how to access the data are provided.

Keywords: carbon dioxide; coulometry; World Ocean Circulation Experiment; South Atlantic Ocean; hydrographic measurements; alkalinity; partial pressure of carbon dioxide; carbon cycle

PART 1:
OVERVIEW



1. BACKGROUND INFORMATION

The World Ocean Circulation Experiment (WOCE) Hydrographic Program (WHP) is a major component of the World Climate Research Programme (WCRP). The overall goal of the WCRP is to better understand the role of the ocean in climate and climatic changes resulting from both natural and anthropogenic causes. The need for this experiment arose from serious concern over the rising atmospheric concentration of carbon dioxide (CO_2) and its effect on the heat balance of the global atmosphere. The increasing concentration of this gas may intensify the earth's natural greenhouse effect and alter the global climate in ways that are not well understood. The complex and poorly understood circulation patterns and biogeochemical cycles of the ocean yield a complex and uneven distribution of CO_2 . Although total CO_2 (TCO_2) is not an official WOCE measurement, a coordinated effort, supported in the United States by the U.S. Department of Energy (DOE), is being made on WOCE cruises through 1998 to measure the global spatial and temporal distributions of TCO_2 as well as other related parameters. Goals of the survey are to estimate the meridional transport of inorganic carbon in a manner analogous to the oceanic heat transport estimates (Bryden and Hall 1980; Brewer et al. 1989; Roemmich and Wunsch 1985) and to build a database suitable for carbon-cycle modeling and the subsequent estimation of anthropogenic CO_2 increase in the oceans. The CO_2 survey is taking advantage of the sampling opportunities provided by the WHP cruises during this period. The final data set is expected to cover ~23,000 stations.

This report discusses the results of the research vessel (R/V) *Meteor* Cruise 22/5 along the WOCE zonal section A10 (along ~30° S) (Fig. 1). The expedition started in Rio de Janeiro, Brazil, on December 27, 1992, and ended in Capetown, South Africa, on January 31, 1993. This section is one of four zonal sections (A8, A9, A10, and A11) completed in the South Atlantic Ocean during the WOCE survey. The large-scale three-dimensional distribution of temperature, salinity, and chemical constituents, including the carbonate system parameters, will be plotted using the data from these sections. Knowledge of these parameters and their initial conditions will enable researchers to determine heat and water transport as well as carbon transport. An understanding of this transport will contribute to the understanding of processes that are relevant to climate change. This section in the South Atlantic subtropical gyre is especially relevant to understanding CO_2 transport because it crosses both the Brazil and the Benguela boundary currents.

This data documentation is the result of the cooperative efforts of chemical oceanographers from Brookhaven National Laboratory (BNL), U.S.A; Baltic Sea Research Institute [Institut für Ostseeforschung (IOW)], Germany; and the Institute for Marine Sciences of University of Kiel [Institut für Meereskunde Kiel (IfMK)], Germany.

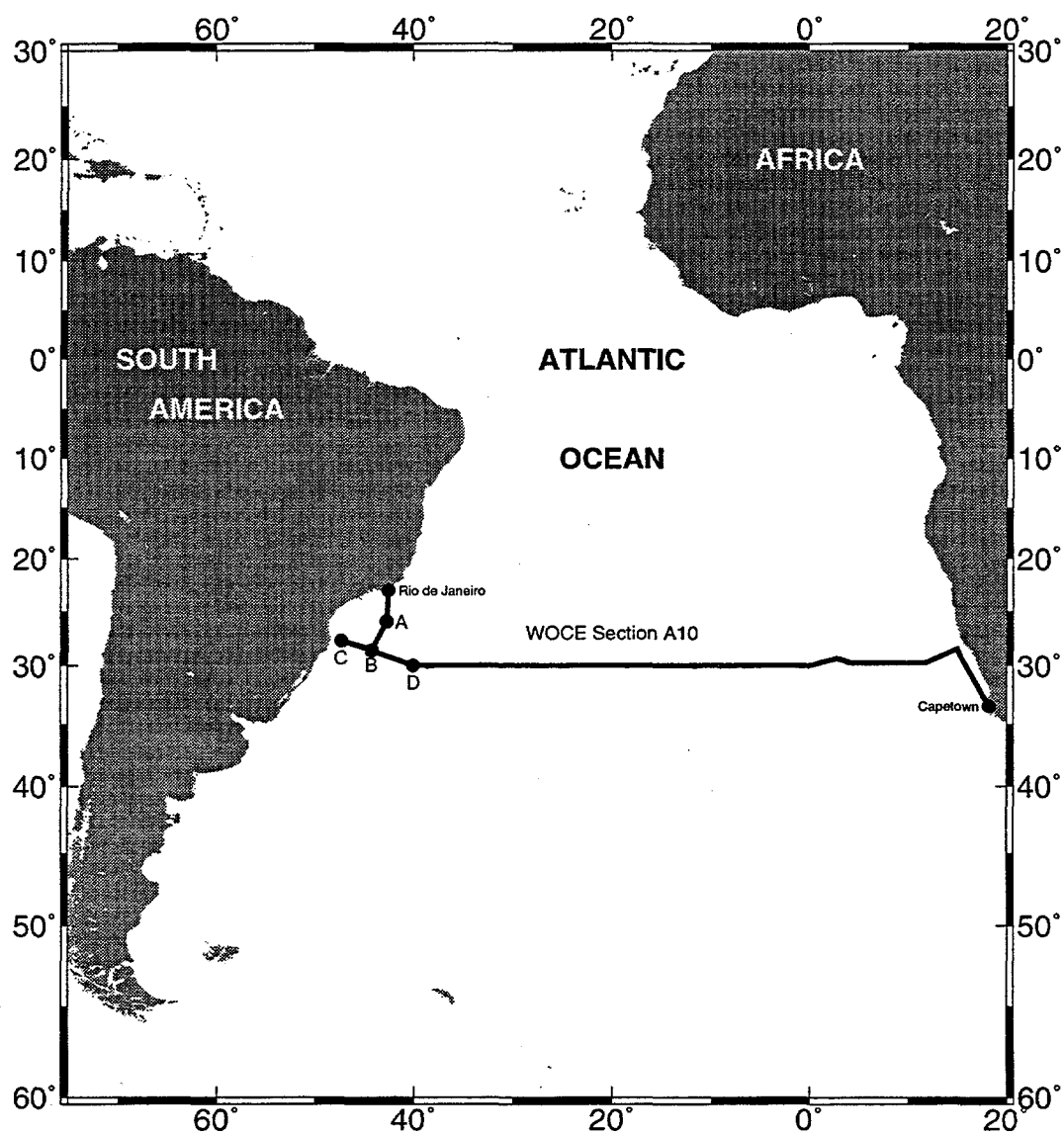


Figure 1. The cruise track during R/V *Meteor* Cruise 22/5 (WOCE Section A10). A, B, C, and D on the map designate waypoints, which are described in Section 2.3.

2. DESCRIPTION OF THE EXPEDITION

2.1 R/V *Meteor*, Technical Details and History

The R/V *Meteor* is owned by the Federal Republic of Germany through the Ministry of Research and Technology (BMFT), which financed its construction. It is operated by the German Research Foundation (DFG), which provides about 70% of its operating funds (the BMFT supplies the remainder). DFG also plans the scientific cruises and appoints the chief scientists. The Operations Control Office of the University of Hamburg is responsible for management, logistics, execution, and supervision of ship operations. These functions are exercised by direct cooperation with expedition coordinators and the managing owners, the Reedereigemeinschaft Forschungsschiffahrt GmbH, located in Bremen, Germany. The latter is responsible for hiring, provisioning, and coordinating ship maintenance. Used for ocean research, primarily in the Atlantic and Indian Oceans, the R/V *Meteor* routinely carries scientists from many different countries. The *Meteor* was completed in 1986 in Travemünde, Germany. The basic features of the vessel follow:

Port of registration	Hamburg
Call sign	DBBH
Classification	GL+100A4E2+MC Auto
Operator	University of Hamburg, Institute for Ocean Research
Built	1985–1986 at Schlichting Werft, Travemünde
Basic dimensions:	
<i>Gross registered tonnage</i>	3990 t
<i>Net registered tonnage</i>	1284 t
<i>Displacement</i>	4780 t
<i>Length overall</i>	97.50 m
<i>Beam</i>	16.50 m
<i>Draught maximum</i>	5.60 m
<i>Service speed</i>	12 kn
<i>Depth main deck</i>	7.70 m
Personnel	Crew: 32; scientists: 30
Main engine	4 × Mak6M 322 = 4 × 1000 kW at 750 rpm
Propulsion	Diesel-electrical, tandem-motor = 2 × 1150 kW
Fuel consumption	Approximately 12.0 t IFO-80 per day at service speed
Maximum cruise duration	60 days
Nautical equipment	Integrated navigation system with data transfer to position computer, echo sounder synchronization and supervision, and data-processing facility
Science quarters	20 laboratories on the main deck with ~400 m ² of working space for multidisciplinary research

Meteor (I) was constructed in 1925, the first research and survey vessel of that name. Owned by the German navy, it was based in Wilhelmshaven. One of its first expeditions was the German Atlantic Ocean Expedition of 1925–1927, which was organized by the Institute of Marine Research in Berlin. Thereafter, the vessel was used for German physical, chemical, and microbiological marine investigations and for navy surveying and fisheries protection duties.

Meteor (II) was planned after the 1950s; it was operated by the Deutsche Forschungsgemeinschaft (German Science Community) in Bad Godesberg and the Deutsches Hydrographisches Institut (German Hydrographic Institute) in Hamburg. Commissioned in 1964, *Meteor* (II) participated in the International Indian Ocean Expedition.

Multipurpose *Meteor* (III), used for the cruise described in this documentation, was completed in 1986, replacing *Meteor* (II). Based in Hamburg, it is used for German ocean research worldwide and for cooperative efforts with other nations in this field. The vessel serves scientists of all marine disciplines in all of the world's oceans.

2.2 R/V *Meteor* Cruise 22/5 Information

R/V *Meteor* Cruise 22/5 information follows:

Ship name	<i>Meteor</i>
Cruise/leg	22/5
WOCE Section	A10
Location	Rio de Janeiro, Brazil, to Capetown, South Africa
Dates	December 27, 1992, to January 31, 1993
Funding	German Science Community; Federal Ministry of Research and Technology, Bonn, Germany; and U.S. DOE
Chief Scientist	Dr. Reiner Onken, IfMK
Master	Martin Kull

Parameters measured	Institution	Principal investigators
CTD ¹ , salinity, XBT ² , and XCP ³	IfMK	R. Onken
Nutrients	IOW, IfMK	B. Schneider, and H. Johannsen
Oxygen	IOW, IfMK	B. Schneider, and H. Johannsen
Chlorofluorocarbons (CFCs)	UBT ⁴	W. Roether, and A. Putzka
Tritium, helium, and radiocarbon	UBT	W. Roether, and A. Putzka
TCO ₂	BNL, IOW	K. Johnson, and B. Schneider
TALK	IfMK	L. Mintrop
Underway pCO ₂	IOW	B. Schneider
ADCP ⁵	IfMK	R. Onken

¹Conductivity, temperature, and depth sensor.

²Expendable bathythermograph.

³Expendable current profiler.

⁴University of Bremen, Tracer Oceanography Laboratory.

⁵Acoustic Doppler current profiler.

2.3 Brief Cruise Summary

On December 26, 1992, K. Johnson of BNL arrived in Rio de Janeiro, where he joined Drs. B. Schneider and L. Mintrop and their CO₂ group consisting of A. Morak, U. Karbach, A. Korves, and J. Morlang who were already onboard the R/V *Meteor*. Setting up of the equipment began immediately by locating the coulometry systems in the Universal Laboratory, the underway pCO₂ system in the Geo-Laboratory, and the alkalinity system in one of the ship's chemistry laboratories. The R/V *Meteor* departed Rio de Janeiro at 6 p.m. on December 27, 1992. Work began almost immediately with ADCP and XBT measurements across the Brazil Current and the occupation of test station no. 620 (waypoint A). Then R/V *Meteor* headed to waypoint B and once again crossed the Brazil Current, where additional ADCP and XBT measurements were made (see Fig. 1). After R/V *Meteor* crossed the Brazil Current for the second time, the ship began the hydrographic stations at waypoint C. Between waypoints C and B the interval between stations was ~10 nautical miles (nm), and after waypoint B it rose to ~30 nm. The CO₂ sampling began at test station no. 620. The 30° S parallel was reached at waypoint D, and for the next few weeks the R/V *Meteor* sailed eastward along the 30° S parallel, passing over portions of the Vema Channel, the eastern part of the Rio Grande Rise, the Argentine Basin, the southern Brazil Basin, the Mid-Atlantic Ridge, the Angola Basin, the Walvis Ridge, and the Cape Basin. There was a small interruption in the work schedule to allow a New Year's Eve celebration. On January 8, nets laid by Spanish fishing boats on the western edge of the Rio Grande Rise caused the R/V *Meteor* to make a detour. (However, this meeting with the fishing boats also resulted in a trade between captains in which the R/V *Meteor* received fresh swordfish and tuna.) A small northward jog was made over the Walvis Ridge in order to sample around topographical features. The intervals between stations as the ship steamed eastward varied between 9 and 45 nm to limit the difference between the bottom depth at consecutive station depths to ≤1000 m. At 11° 30' E, the ship veered slightly to the east-northeast in order to avoid the South African 200-nm exclusion zone because permission to sample in these waters had not been obtained. After this turn, the station resolution was reduced to 20 nm until the last station on the African shelf at a depth of ~200 m. The measurement phase concluded on January 28, 1993, and the R/V *Meteor* steamed to Capetown, where it arrived on the afternoon of January 30. Aside from some light rain and intermittent cloudiness at the beginning of the cruise, the weather remained mostly sunny with summer temperatures and calm seas throughout. Closer to the coast of Africa, swells of ~5 m originating from subantarctic low-pressure areas were experienced but without any loss of work time.

Two Single-Operator Multiparameter Metabolic Analyzers (SOMMAs) were used for measuring TCO₂ concentration during this cruise. One was supplied by BNL and another by the IfMK. In addition, two potentiometric alkalinity titrators from IfMK were run in parallel, and an infrared (IR)-based system belonging to IfMK was deployed to measure underway pCO₂. The TCO₂ concentration was analyzed in 1425 samples from 57 of 100 CTD stations (57%) occupied during the cruise (Fig. 2). In addition, 116 coulometric measurements for the Certified Reference Material (CRM) and duplicate analyses were made during the cruise. The TALK concentration in 665 samples was determined by potentiometric titration during the cruise. Not all stations could be sampled for TCO₂ and TALK because of the time required for analysis; however, the goal of 50% station coverage for CO₂ samples was surpassed, and on average 1.5 stations were sampled per day by the CO₂ group. The standard WOCE parameters (oxygen, nutrients, and salinity) were analyzed on all samples, and the tracer samples included CFC's, helium, tritium, and radiocarbon. The underway pCO₂ system operated continuously.

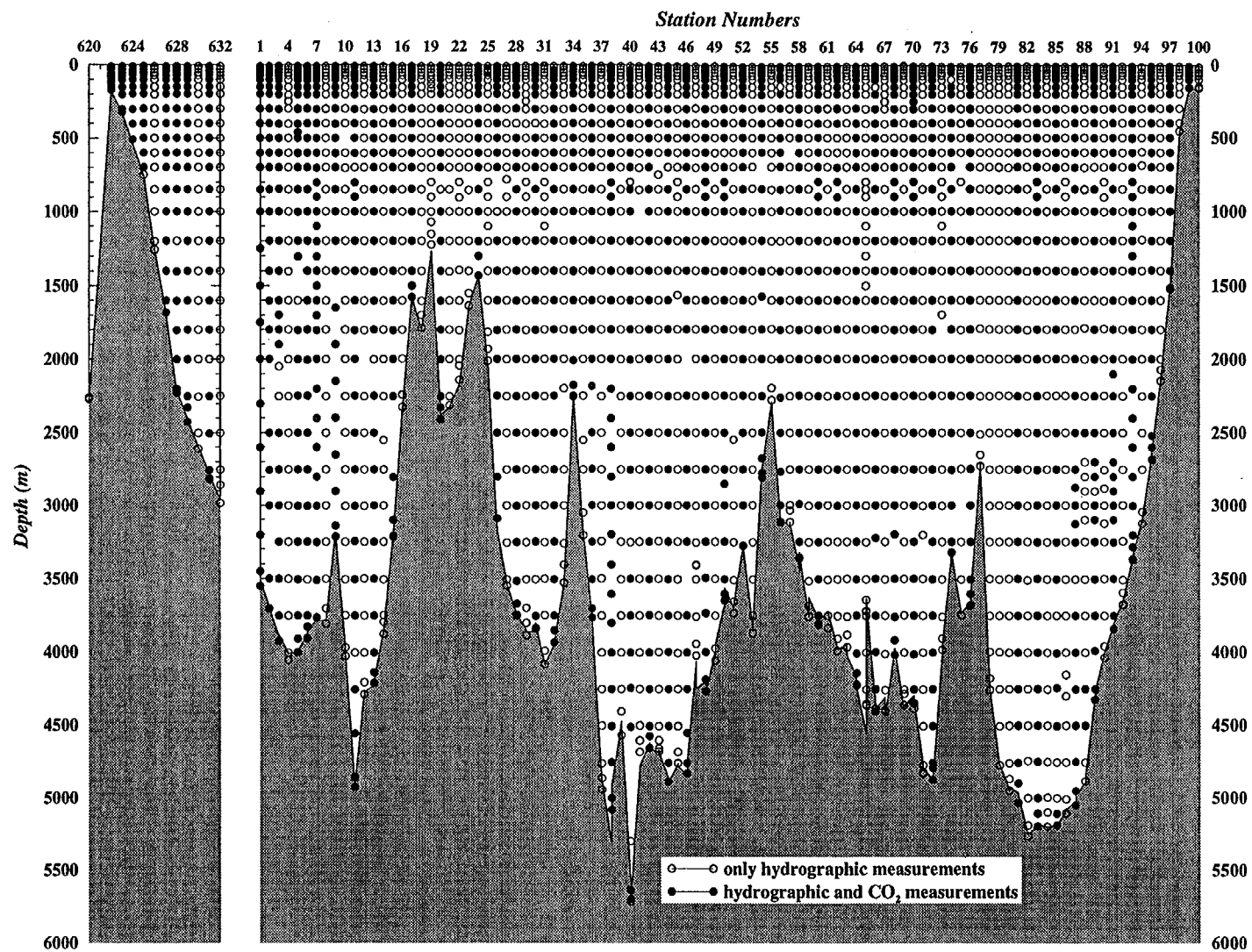


Figure 2. Sampling depths at all hydrographic stations occupied during R/V *Meteor* expedition along WOCE Section A10.

3. DESCRIPTION OF VARIABLES AND METHODS

3.1 Hydrographic Measurements

Water samples were collected with a General Oceanics rosette equipped with twenty-four 10-L Niskin bottles mounted on a Neil Brown Mark III CTD instrument provided by the IfMK. For stations deeper than 3500 m, two separate CTD/rosette casts were launched. For stations with a depth less than 3500 m, one CTD/rosette cast of up to 24 bottles was lowered. Surface currents down to 300 m, surface temperature, and surface salinity were measured continuously during the cruise with a hull-mounted ADCP and a thermosalinograph. In between CTD stations, XBTs were routinely launched. Over the boundary currents, XBTs were launched every half hour, and over the Benguela Current, the XBT launches were supplemented with free-falling XCPs.

No serious problems were experienced with the CTD/rosette systems during the cruise. Repeated checks on board and several careful verifications using the complete bottle data sets have been carried out, and the sampling pressures have been assigned to each sample. Reversing thermometers of both the electronic (SIS, Kiel) and the mechanical (Gohla Precision, Kiel) types were also read at the completion of each cast. The processing and quality control of CTD and bottle data followed the published guidelines in the *WOCE Operations Manual* (WHPO 91-1, 1991).

The CTD pressure, temperature, and conductivity data were processed and corrected according to laboratory calibrations. Pressure values are expected to be accurate to ± 3 dbar; temperature values to $\pm 0.002^\circ\text{C}$. Salinity for selected Niskin bottles (about one in every three) was also determined on a Guildline Autosol model 8400A that was standardized weekly with International Association for the Physical Sciences of the Ocean (IAPSO) water. These data were also used to process the CTD data, and the final salinity data are expected to be accurate to ± 0.002 . Bottle oxygen was determined by Winkler titration after the technique of Carpenter (1965) with the modifications of Culbertson et al. (1991), using standards and blanks run in seawater. The precision of the analyses determined from parallel analyses ($n = 10$) of samples at and well below saturation is $\pm 0.4\%$. The concentrations of nitrate, nitrite, phosphate, and silicate dissolved in seawater were determined for samples collected in high-density polyethylene screw-capped bottles by a continuous-flow method with an autoanalyzer. Precision was as follows: silicate, $\pm 1.3\%$; phosphate, $\pm 1.5\%$; and nitrite/nitrate, $\pm 1.1\%$. Preweighed standards were used to prepare the nutrient working standards aboard the ship.

3.2 TCO_2 Measurements

The TCO_2 was determined by using two automated dynamic headspace sample SOMMA processors with coulometric detection of the CO_2 extracted from acidified samples. A description of the SOMMA-coulometry system and its calibration can be found in Johnson et al. (1987), Johnson and Wallace (1992), and Johnson et al. (1993). Details concerning the coulometric titration procedure can be found in Huffman (1977) and Johnson et al. (1985). Samples were collected in 300-mL precombusted (450°C for 24 h) glass standard Biological Oxygen Demand (BOD) bottles and analyzed for TCO_2 during the cruise. During the cruise the samples were not poisoned with HgCl_2 as per normal operating procedure (DOE 1994), but they were analyzed within 24 h of collection. Before analysis, samples were kept in darkness in a cold room and subsequently thermally equilibrated for at least 3 h to the analytical temperature. Analyses of duplicate samples separated in time by up to 8 h showed no evidence of any significant biological

consumption or production of CO_2 during storage. The CRMs were supplied by Dr. A. Dickson of Scripps Institution of Oceanography (SIO) (DOE 1994) and were also routinely analyzed. CRMs from batches 7 and 11 were available for this work (batch 7: $S = 37.120$, $\text{TCO}_2 = 1926.41 \pm 0.82 \mu\text{mol/kg}$; batch 11: $S = 38.5$, $\text{TCO}_2 = 2188.77 \mu\text{mol/kg}$). The CRMs were made from filtered sterile salt solutions spiked with Na_2CO_3 , and their TCO_2 concentrations were determined by vacuum-extraction/manometry in the laboratory of Dr. C. D. Keeling at SIO.

Seawater introduced from an automated "to deliver" (TD) pipette into a stripping chamber was acidified, and the resultant CO_2 from continuous gas extraction was dried and coulometrically titrated on a model 5011 UIC coulometer. The coulometers were adjusted to give a maximum titration current of 50 mA, and they were run in the counts mode [the number of pulses or counts generated by the coulometer's voltage-to-frequency converter (VFC) during the titration was displayed]. In the coulometer cell, the hydroxyethylcarbamic acid formed from the reaction of CO_2 and ethanolamine was titrated coulometrically (electrolytic generation of OH^-) with photometric endpoint detection. The product of the time and the current passed through the cell during the titration (charge in Coulombs) is related by Faraday's constant to the number of moles of OH^- generated and thus to the moles of CO_2 that reacted with ethanolamine from the acid.

Each system was controlled with an IBM-compatible personal computer equipped with two RS232 serial ports, a 24-line digital input/output card, and an analog-to-digital card. The latter were manufactured by Real Time Devices (State College, Pa.). These were used to control the coulometer, barometer (BNL system only), solid-state control relays, and temperature sensors, respectively. The temperature sensors (model LM34CH, National Semiconductor, Santa Clara, Calif.), with a voltage output of 10 mV/ $^\circ\text{F}$ built into the SOMMA, were calibrated against thermistors certified to 0.01 $^\circ\text{C}$ (PN CSP60BT103M, Thermometrics, Edison, N.J.) using a certified mercury thermometer as a secondary standard. These sensors monitored the temperature of SOMMA components, including the pipette, gas sample loops, and the coulometer cell. The SOMMA software was written in GWBASIC Version 3.20 (Microsoft Corp., Redmond, Wash.), and the instruments were driven from an options menu appearing on the personal computer (PC) monitor. With the coulometers operated in the counts mode, conversions and calculations were made by using the SOMMA software rather than the programs and the constants hardwired into the coulometer circuitry.

The BNL SOMMA-coulometry system was calibrated with pure CO_2 by using hardware that consisted of an 8-port gas sampling valve (GSV) with two sample loops of known volume [determined gravimetrically by the method of Wilke et al. (1993)]. This GSV was connected to a source of pure CO_2 through an isolation valve with the vent side of the GSV plumbed to a barometer. When a gas loop was filled with CO_2 , the mass (moles) of CO_2 contained therein was calculated by dividing the loop volume (V) by the molar volume of CO_2 at the ambient temperature and pressure. The molar volume of CO_2 [$V(\text{CO}_2)$] was calculated iteratively from an expression using the instantaneous barometric pressure (P), loop temperature (T), gas constant (R), and the first virial coefficient $B(T)$ for pure CO_2 :

$$V(\text{CO}_2) = RT / P[1 + B(T) / V(\text{CO}_2)] . \quad (1)$$

The gas calibration factor (CALFAC)—the ratio of the calculated mass to that determined coulometrically—was used to correct the subsequent titrations for small departures from 100% recoveries (DOE 1994). Pressure was measured with a barometer, model 216B-101 Digiquartz Transducer (Paroscientific, Inc., Redmond, Wash.), which is factory-calibrated for pressures between 11.5 and 16.0 psia. The standard operating procedure was to make gas calibrations daily for each newly prepared titration cell (normally, one cell per day and three sequential calibrations per cell).

The "to deliver" volume (V_{cal}) of the BNL SOMMA sample pipette was determined (calibrated) gravimetrically during the cruise by periodically collecting aliquots of deionized water dispensed from the pipette into preweighed serum bottles. The serum bottles were crimp sealed and returned to shore, where they were reweighed on a model R300S (Sartorius, Göttingen, Germany) balance. The apparent weight (g) of water collected (W_{air}) was corrected to the mass in vacuo (M_{vac}) from

$$M_{vac} = W_{air} + W_{air} (0.0012/d - 0.0012/8.0), \quad (2)$$

where 0.0012 is the sea level density of air at 1 atm, d is the density of the calibration fluid at the pipette temperature and sample salinity, and 8.0 is the density of the stainless steel weights. V_{cal} was calculated by using the following equation:

$$V_{cal} = M_{vac}/d. \quad (3)$$

The V_{cal} of the pipette for the BNL system was 20.6114 ± 0.0024 mL ($n = 23$) at a mean temperature of 14.67°C (hereafter the calibration temperature t_{cal}). During the cruise, the mean pipette temperature was $14.95 \pm 0.97^\circ\text{C}$, and the vast majority of samples were analyzed at a measurement temperature (t) that was within 1°C of this calibration temperature. The sample volume (V_t) at the measured pipette temperature was calculated from the expression

$$V_t = V_{cal} [1 + a_v (t - t_{cal})], \quad (4)$$

where a_v is the coefficient of volumetric expansion for pyrex-type glass ($1 \times 10^{-5} ^\circ\text{C}^{-1}$) and t is the temperature of the pipette at the time of a measurement.

The BNL coulometer was periodically electronically calibrated as described in Johnson et al. (1993, 1996) and the DOE *Handbook of Methods* (1994). Briefly, at least two levels of current (usually 50 and 2 mA) were passed through an independent and very precisely known resistance (R) for a fixed time. The voltage (V) across the resistance was continuously measured and the instantaneous current (I) across the resistance was calculated from Ohm's law and integrated over the calibration time. Then, the number of pulses (counts) accumulated by the VFC during this time was compared to the theoretical number computed from the factory-calibration of the VFC [frequency = 10^5 pulses (counts) generated/sec at 200 mA] and the measured current. If the VFC was perfectly calibrated at the factory, the electronic calibration procedure would yield a straight line passing through the origin (intercept = 0) with a slope of 1. For the BNL coulometer, the mean electronic calibration slope during the R/V *Meteor* Cruise 22/5 was 0.999616 ± 0.000056 ($n = 12$, r.s.d. = 0.006%) with an intercept of -0.000533 $\mu\text{mol}/\text{min}$. From the factory calibration of the VFC and the value of the Faraday (96489 Coulomb/mol), a scaling factor of 4.82445×10^3 counts/ μmol was derived. The theoretical number of micromoles of carbon titrated (M) from samples or the gas loops was

$$M = [\text{Counts}/4824.45 - (\text{Blank} \times T_t) - (\text{INT}_{ec} \times T_t)]/\text{SLOPE}_{ec}, \quad (5)$$

where T_t is the length of the titration in minutes, *Blank* is the system blank in $\mu\text{mol}/\text{min}$, INT_{ec} is the intercept from electronic calibration in $\mu\text{mol}/\text{min}$, T_t is the time in minutes during the titration where current flow was continuous, and SLOPE_{ec} is the slope from electronic calibration. Note that the slope obtained from the electronic calibration procedure applied for the entire length of the titration, but the intercept correction applied only for the period of continuous current flow

(usually 3–4 min) because the electronic calibration can be carried out only for periods of continuous current flow. The TCO_2 concentration in $\mu\text{mol/kg}$ was calculated from

$$\text{TCO}_2 = M \times \text{CALFAC} \times [1/(V_i \times \rho)] \times d_{\text{Hg}} \times \text{CF}_{\text{crm}}, \quad (6)$$

where ρ is the density of sea water in g/mL at the measurement temperature and sample salinity calculated from the equation of state given by Millero and Poisson (1981), d_{Hg} is the correction for sample dilution with bichloride solution (for this cruise $d_{\text{Hg}} = 1.0$ for the BNL and Kiel analyses because HgCl_2 was not used), and CF_{crm} is a correction factor based on the daily liquid calibration by CRM analysis ($\text{CF}_{\text{crm}} = 1.0$ for all BNL analyses; no correction based on the CRM data).

The BNL SOMMA-coulometry system was equipped with a conductance cell (Model SBE-4, Sea-Bird Electronics, Inc., Bellevue, Wash.) for salinity measurement as described by Johnson et al. (1993). The conductance cell was factory calibrated, but SOMMA-measured salinities were continuously compared with the CTD salinities to ensure that the salinities of the analyzed samples matched the assigned salinities. Generally, agreement between CTD and SOMMA salinities was 0.02 or better.

A leak in the gas calibration hardware of the BNL system was discovered on January 12, 1993. It affected the gas calibrations by diluting the CO_2 calibration gas during the gas calibration procedure so that CALFACs determined between December 28, 1992, and January 12, 1993, were in error by approximately +0.1%. These CALFACs caused an error of +2 $\mu\text{mol/kg}$ in the CRM analyses. Repairs were made on January 12, and from this point through January 28, daily CALFACs were determined and used to calculate the values of CRM and TCO_2 . The mean CALFAC for the period January 12–28 was 1.004270 ± 0.000818 ($n = 12$). This CALFAC was used to recalculate the values of CRM and TCO_2 for the period from December 28 through January 11.

The IfMK system did not possess a gas calibration system, and gas calibration was not carried out during the cruise. This IfMK system was calibrated at the IfMK in Kiel prior to the cruise with liquid standards (Na_2CO_3 solutions) according to the method of Goyet and Hacker (1992). A mean CALFAC ($1.005 \pm 0.07\%$) was obtained in the laboratory from the ratio (true TCO_2 / measured TCO_2). This value was used in Eq. 6 to calculate the CRM and TCO_2 values throughout the cruise. During the calibration and at-sea work, the pipette volume, V_{cal} (also determined prior to the cruise), used for the IfMK system was 25.2347 mL at 20.02°C (see Eq. 4). This TD pipette volume was not redetermined gravimetrically during the cruise. Instead, an additional CF_{crm} based on the daily (cell-specific) CRM results was used to account for changes in pipette volume and/or system response by multiplying the TCO_2 sample results by the ratio (see Eq. 6):

$$\text{CF}_{\text{crm}} = \text{CRM (certified)} / \text{CRM (measured)} .$$

In summary, the IfMK system was calibrated as follows: a daily (cell-specific) CF_{crm} was applied to the water sample analysis results based on a laboratory-determined constant CALFAC (1.005) and a constant value of V_{cal} (25.2347 mL at 20.02°C). The IfMK coulometer was not electronically calibrated during the cruise, and the theoretical response (Slope = 1, Intercept = 0) was assumed in Eq. 5 for all calculations. Note, however, that the CRM analysis results from the IfMK system (also referred to as the Kiel system or Kiel SOMMA) (Fig. 3) were calculated with $\text{CF}_{\text{crm}} = 1$ in order that the variability of the CRM analyses and the magnitude of CF_{crm} could be assessed.

Scatter plot showing the difference from certified value ($\mu\text{mol kg}^{-1}$) versus day number (relative to Jan 1, 1993) for Kiel SOMMA and BNL SOMMA. The plot shows two data series: Kiel SOMMA (open squares) and BNL SOMMA (filled circles). A horizontal dashed line is drawn at y=0. The x-axis ranges from -5 to 35, and the y-axis ranges from -6 to 6. Kiel SOMMA data points are generally higher than BNL SOMMA data points, especially in the early days.

Figure 3. The distribution of differences between the measured and certified values of the CRM analyzed by the BNL SOMMA-coulometry system (closed circles) and the IFMK (Kiel) SOMMA-coulometry system (open squares) during R/V *Meteor* Cruise 22/5. The IFMK results have been calculated based on the pre-cruise calibration only.

Problems were encountered with the pinch-valve tension of the IfMK System. This valve controls the delivery of samples to the stripper. Although it always operated, it affected the analytical results by periodically allowing additional samples to be injected into the stripper. The weak valve tension prevented the complete sealing of the tubing connecting the pipette to the stripper. The resulting errors for the CRM analyses were on the order of 0.1 to 0.5%. The valve tension was adjusted during the cruise, and the effect of these errors on data quality was minimized because poor results for the CRM analyses caused by the malfunctioning pinch valve prompted repair of the system before any samples were run. However, the possibility remained that during the sample analyses periodic pinch valve failures may have occurred, and this prompted extensive quality assurance-quality control (QA-QC) of the data.

The first phase of the QA-QC procedure was an assessment of accuracy using the data from the CRM analyses. These data are shown in Fig. 3 and summarized in Table 1. For the BNL system, during the period from December 29, 1992, through January 11, 1993, a constant CALFAC (1.004270) was used to calculate CRM TCO₂, whereas between January 12 and 28, 1993, a daily (cell-specific) CALFAC was used to calculate CRM TCO₂. For the IfMK system, a constant CALFAC (1.005) was used for all calculations.

Table 1. Summary of CRM TCO₂ determinations made during R/V Meteor Cruise 22/5

System	No. (n)	Batch ^a	Mean $\mu\text{mol/kg}$	St. Dev. $\mu\text{mol/kg}$	Diff ^b	Dates	CALFAC	Outliers ^c
BNL	14	7	1926.7	0.65	+0.27	12/29–01/11	Constant	1
BNL	16	11	2188.7	0.89	–0.11	01/12–01/28	Daily	0
IfMK	18	7	1928.1	1.57	+1.68	12/30–01/12	Constant	1
IfMK	11	11	2191	1.88	+2.20	01/13–01/28	Constant	3

^aThe CRM were from Batch 7 and 11 with salinities of 37.12 and 38.50, and TCO₂ of $1926.41 \pm 0.82 \mu\text{mol/kg}$ ($n = 13$) and $2188.77 \pm 0.56 \mu\text{mol/kg}$ ($n = 5$), respectively.

^bThe mean difference between measured and certified TCO₂.

^cAn outlier is defined as a CRM analysis with an error $\geq 5.0 \mu\text{mol/kg}$.

Mean errors in the BNL system were significantly lower than the consistently positive errors observed in the IfMK system. For the BNL system, an outlier was obtained on January 4 (CRM bottle no. 2), but a second CRM (no. 275) run on the same cell gave a satisfactory result, and this cell was subsequently used to run samples. For the IfMK system, outliers were observed on January 7 and 12 (no. 353 and no. 8). In each case, a second CRM analysis (no. 318 and 244) gave satisfactory results, and the system was operated as normal. On January 20 two consecutive CRM analyses (no. 370 and 112) were classified as outliers, but a third CRM (no. 312) analysis gave a satisfactory result and the system was operated. Overall, 5 of 64 CRM analyses from Table 1 (7.8%) were classed as outliers, but 4 of these 5 outliers were obtained on the IfMK system which was further evidence for the slightly better performance of the BNL system. The greater number of outliers on the IfMK system could possibly be the result of malfunctioning of the IfMK pinch valve as described earlier. In general, the CRM results on the BNL system were identical to the manometric reference analyses at SIO. The BNL system response remained

constant over the duration of the cruise (see Fig. 3) whether an average CALFAC (12/28/1992–01/11/1993) or a cell-specific CALFAC (01/12/1993–01/28/1993) was used to calculate CRM TCO₂. These results confirm a similar finding obtained when mean CALFACs were used to calculate the TCO₂ data of the R/V *Meteor* Cruise 18/1 (WOCE Leg A1E) (Johnson et al. 1996).

The second phase of the QA-QC procedure was an assessment of sample precision on each system (instrument-specific precision). The system precision data are given in Table 2. For these data, “within-sample” precision is the average absolute difference between two replicates analyzed from the same sample bottle, “between-sample” precision is the average absolute difference between duplicate sample bottles taken from the same Niskin bottle, and “between-Niskin” precision is the average absolute difference of analyses of samples taken from two Niskin bottles that were closed at the same depth. The IfMK group assessed instrument-specific precision by periodically running two replicates from the same bottle (“within-sample”), whereas precision on the BNL system was assessed by running one replicate from each of two sample bottles filled from the same Niskin bottle (“between-sample”). The pooled standard deviation (S_p^2) is the square root of the pooled variance from the “between-sample” replicates ($n = 2$) according to Youden (1951):

$$S_p^2 = \sqrt{\frac{\sum_{i=1}^K \left(\sum_{j=1}^{n_i} (x_{ij} - \bar{x}_j)^2 - \left[\sum_{j=1}^{n_i} (x_{ij} - \bar{x}_j) \right]^2 / n_i \right)}{\sum_{i=1}^K n_i - K}}$$

where K is the number of samples analyzed, and $\sum_{i=1}^K n_i - K$ are the degrees of freedom for the calculation.

Table 2. Summary of sample precision for TCO₂ analyses made during R/V *Meteor* Cruise 22/5

System	Mean precision and St. Dev. (μmol/kg) ^a			S_p^2 ($K, n, \text{d.f.}$)
	within-sample (n)	between-sample (n)	between-Niskin (n)	
BNL	0	1.04 ± 1.11 (53)	1.26 ± 1.41 (12)	1.07 (53, 106, 53)
IfMK	1.16 ± 1.62 (46)	0.98 ± 0.36 (6)	1.53 ± 2.04 (5)	0.73 (6, 12, 6)
Combined		1.03 ± 1.06 (59)	1.34 ± 1.55 (17)	1.04 (59, 118, 59)

^aMean precision is $\left[\sum_{x=1}^n \text{abs}(x_1 - x_2) \right] / n$, where n is the number of comparisons between duplicate analyses, x_1 and x_2 .

Table 2 shows that there was no significant difference between the precision estimated using the three different methods; however, the standard deviation of the between-sample estimates was the lowest of the three methods. The same pattern was found for the TCO₂ data of the R/V *Meteor* Cruise 18/1 (WOCE Section A1E) when within-sample and between-sample precisions were

compared (see Johnson et al. 1996); these data were also consistent with results for other WOCE Sections (Johnson et al. 1995; Johnson et al. 1996). For the instrument-specific S_p^2 , K is the number of between-sample samples analyzed on the same instrument, n is the total number of replicates analyzed from K samples, and $n - K$ is the degrees of freedom (d.f.).

The third phase of the QA-QC procedure was to assess the performance of the systems by comparing results from aliquots of the same sample analyzed on each system. The precedent was set by the R/V *Meteor* Cruise 18/1 TCO_2 data set when two SOMMAs were also run in parallel to generate the data set (Johnson et al. 1996). For the R/V *Meteor* 18/1 data, a method-specific S_p^2 , assuming homogeneous variance, was calculated from aliquots of the same sample analyzed on each system. The same calculation was made for the applicable R/V *Meteor* Cruise 22/5 samples, and the method-specific precision (S_p^2) for this cruise calculated from 31 such samples ($K = 31$, $n = 2$, d.f. = 31) was $\pm 1.92 \mu\text{mol/kg}$. This is a more conservative estimate of overall cruise-wide precision than the instrument-specific precision shown in Table 2. For any measurement, irrespective of the instrument it was made on, the precision was $\pm 1.92 \mu\text{mol/kg}$. This includes all sources of error-random as well as any uncorrected systematic errors (bias).

Figure 4 is a histogram that shows the frequency distribution of the differences between aliquots of 31 samples that were measured on both systems. The mean and standard deviation of the mean difference was $0.81 \pm 2.46 \mu\text{mol/kg}$ (BNL - IfMK TCO_2 results) with most of the differences falling within the $\pm 1.0 \mu\text{mol/kg}$ range (Fig. 4). The IfMK calibration procedure therefore appears to have been successful in eliminating any overall system bias seen for the IfMK CRM analyses given in Table 1. For the CRM, the BNL system (gas calibrated) gave more accurate results than the IfMK system (not gas calibrated), and no corrections have been made to any of the sample data analyzed on the BNL system based on the CRM results. In summary, the mean difference between aliquots of the same sample analyzed on both systems was $< 1.0 \mu\text{mol/kg}$, and the method-specific pooled variance ($S_p^2 = \pm 1.92 \mu\text{mol/kg}$) calculated from Youden (1951) is a credible estimate of precision and accuracy for the R/V *Meteor* Cruise 22/5 data set generated by two systems run in parallel but calibrated differently.

The fourth step in the QA-QC procedure, the at-sea to onshore comparison, involved analyzing replicates of the same sample in real time at sea and later, after storage, on shore. This procedure was carried out on 14 samples collected at 7 stations. The onshore analyses were made by vacuum extraction/manometry in the laboratory of Dr. C. D. Keeling at SIO. The results of the comparison are given in Table 3 (Guenther et al., personal communication, 1998).

On the BNL system the initial comparisons (Jan. 13-15, $n = 4$, mean error $-1.93 \mu\text{mol/kg}$) were consistent with the precision and accuracy ($\pm 1.92 \mu\text{mol/kg}$) of the method, but larger differences were observed after January 15. The mean difference for the cruise was $-3.54 \mu\text{mol/kg}$ [for the R/V *Meteor* Cruise 18/1, the corresponding results were $-2.13 \mu\text{mol/kg}$ ($n = 7$) with a method-specific precision and accuracy of $\pm 1.65 \mu\text{mol/kg}$]. Overall, the ship-to-shore difference is clearly not depth dependent. The poorest results were the very negative differences for samples collected on January 17 at station 62 and run on the IfMK system. There were other reasons to suspect the accuracy of the shipboard analyses from station 62 made on the IfMK system, so these samples have been averaged separately in Table 3. Note that only 3 of the remaining 12 differences were within the analytical precision of the shipboard method and these occurred early on in the cruise; 6 of the 12 were essentially within 2 standard deviations ($\pm 3.84 \mu\text{mol/kg}$), but 3 differed by more than 2 standard deviations. All of the differences were negative. The CRM differences were not nearly as large as the ship-to-shore sample differences. The length of time the samples were stored prior to analysis onshore was also not correlated with the at-sea vs onshore differences. The reason for the difference between shipboard and shore-based analyses remains to be determined.

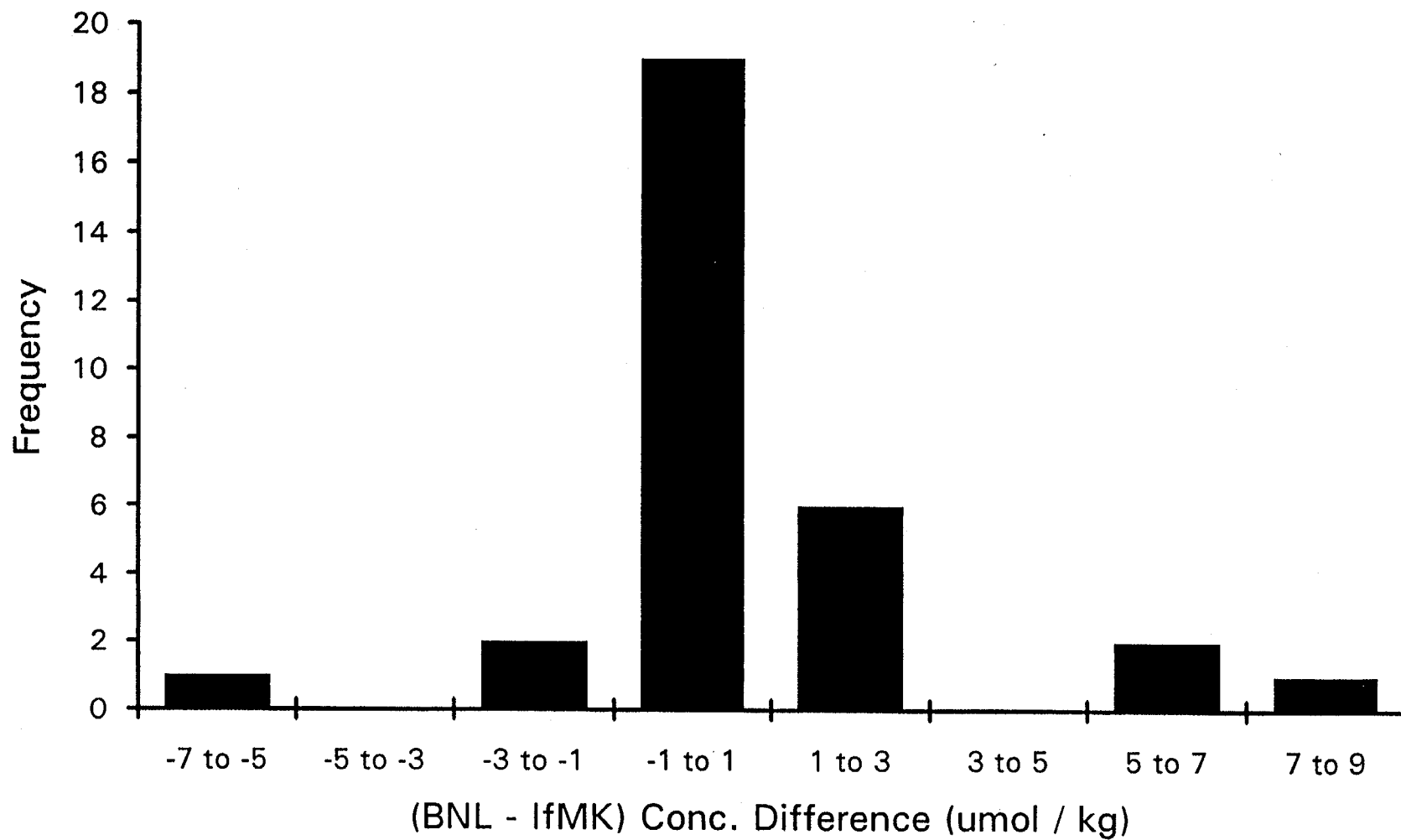


Figure 4. A histogram showing the frequency and distribution of the magnitude of the differences for 31 samples for which aliquots were analyzed on both measurement systems (BNL and IfMK) during R/V *Meteor* Cruise 22/5.

Table 3. Comparison of at-sea analyses of TCO₂ by coulometry and the onshore analyses of TCO₂ by manometry on aliquots of the same sample

Date (1993)	Station (no.)	Niskin (no.)	Depth (m)	At-sea ($\mu\text{mol/kg}$)	Onshore ($\mu\text{mol/kg}$)	Storage ^a (mo)	Difference ($\mu\text{mol/kg}$)	CRM diff ^b
BNL analyses								
1/13	48	318	10.2	2045.68	2047.49	11	-1.81	0.18
1/13	48	308	3002	2188.44	2189.98	10	-1.54	0.18
1/15	54	323	10.7	2044.96	2047.51	5	-2.55	-0.76
1/15	54	301	2808	2200.83	2202.68	5	-1.85	-0.76
1/19	68	323	10.4	2064.83	2068.02	10	-3.19	-0.14
1/19	68	307	3003	2200.1	2203.27	10	-3.17	-0.14
1/21	76	208	12.4	2057.86	2060.61	4	-2.75	1.12
1/21	76	306	3003	2203.31	2207.06	4	-3.75	1.12
1/24	85	213	11.8	2041.48	2047.61	3	-6.13	-0.68
1/24	85	312	3002	2200.73	2207.66	4	-6.93	-0.68
1/27	93	213	12	2029.99	2033.69	3	-3.70	-0.39
1/27	93	305	3004	2200.52	2205.61	3	-5.09	-0.39
Mean (n = 12)						6	-3.54	-0.11
St. Dev.						± 3	± 1.71	± 0.69
IfMK analyses								
1/17	62	208	12	2046.45	2054.22	4	-7.77	2.04
1/17	62	307	3004	2190.57	2202.12	5	-11.55	2.04
Mean (n=2)						4	-9.66	
St. Dev.						± 1	± 2.67	

^aStorage refers to the elapsed time (in months) between sample collection and onshore analysis by manometry.

^bThe SIO difference between the determined and certified CRM TCO₂ for the specific coulometer cell used to titrate the at-sea replicate sample.

The data given in Tables 2 and 3 suggested that further QA-QC analysis of the data was justified. As described above, the two SOMMA systems used during R/V *Meteor* Cruise 22/5 employed different calibration strategies, and the number of replicate samples analyzed on both instruments was insufficient to assess bias on a station-by-station basis. As an additional cross-check on the intercomparability of TCO₂ concentrations measured using the two analytical systems, the correlation of the TCO₂ was compared with other measured oceanographic parameters. Brewer et al. (1995) and Wallace (1995) have previously noted that strong multivariate relationships exist between TCO₂ and other hydrographic parameters (e.g., temperature, salinity, oxygen, and nutrients). These relationships are remarkably robust over basin-scales and have been used to examine the temporal buildup of CO₂ in the oceans (Wallace 1995; Wallace et al. 1996; Holfort et al. 1998) and to interpolate sparse data (Brewer et al. 1995).

Multiple linear regressions were initially performed for TCO₂ data collected from three geographical sections of the R/V *Meteor* Cruise 22/5. Earlier work had suggested that regression fits varied slightly from one ocean basin to another. The section was therefore broken down into three groups of stations: those occupied in zone 1 defined as being west of 13° W (west of the Mid-Atlantic Ridge; southern Brazil Basin); stations occupied in zone 2, defined as being between 13° W and 3° E (between the Mid-Atlantic and Walvis Ridges; Southern Angola Basin); and stations occupied in zone 3, defined as being east of 3° E (east of the Walvis Ridge; Northern Cape Basin). For each group of stations, all samples collected from below 200 m for which TCO₂ had been measured (on either system) were extracted, and a stepwise multiple linear regression was performed with TCO₂ as the dependent variable and the wide range of other measured hydrographic parameters as independent variables. The regression models determined that only potential temperature, salinity, apparent oxygen utilization (AOU), and silicate were significant predictors [this is the same group of parameters as found previously by Wallace (1995) for the WOCE Section A9 along 19° S]. In addition, a single regression was performed for all of the data collected below 200 m from the entire section. The regression parameters for these four different geographical groupings of stations (the defined zones 1, 2, and 3 and the entire section) are presented in Table 4. For each of these four geographical groupings, two sets of regression coefficients are presented: one was derived from a regression that employed the measured silicate concentration as an independent variable and one for a regression that did not use silicate as a predictor.

This initial exercise was not particularly satisfactory, as illustrated by the regression coefficients that varied significantly from one geographical zone to another and depended on whether silicate was employed as a predictor (Table 4). For example, the AOU coefficient varied from 0.33 to 0.63 when silicate was employed as an independent variable, and it reached as high as 0.86 when silicate was not used as a predictor. The salinity coefficient was even more variable, ranging from -2 to +24. In general, the potential temperature, AOU, and salinity coefficients were stable across the geographical groupings when silicate was not used in the regression: the inclusion of silicate caused the other coefficients to vary significantly. Use of silicate as a predictor could also shift the coefficients for the other parameters outside of their "oceanographically reasonable" ranges. For example, the AOU coefficient, if interpreted to reflect the respiratory quotient for organic material, should be 0.68 (Takahashi et al. 1985), 0.69 (Anderson and Sarmiento 1994), or 0.77 (Redfield et al. 1963). Clearly the AOU coefficient derived from regression no. 2A falls well outside this accepted range. Likewise, even the sign of the salinity coefficient is variable. "Oceanographic reasoning" suggests that there should be a positive partial correlation between TCO₂ and salinity because of the strong positive correlation between carbonate alkalinity and salinity in the ocean (the countervailing tendency of CO₂ gas solubility to decrease with increasing salinity is a relatively minor effect). The use of silicate did significantly reduce the overall standard error of the predictions (Table 4) and eliminated or markedly reduced certain systematic patterns in the distribution of the residuals with depth (results not shown).

Table 4. Summary of initial multiple regression results with (A) and without (B) silicate as an independent variable.

Regression no.	Longitude range	Intercept	Coefficients				Standard error
			Pot. temp.	Salinity	AOU	SiO ₄	
0A	All section	1987.32	-4.296	4.045	0.62	0.482	4.37
0B	All section	1370.4	-6.254	22.038	0.827		6.88
1A	Zone 1	1975.58	-4.259	4.37	0.633	0.462	4.77
1B	Zone 1	1395.65	-5.957	21.178	0.864	0.462	7.12
2A	Zone 2	2858.72	-2.531	21.163	0.327	1.126	4.44
2B	Zone 2	1316.58	-6.623	23.822	0.754	1.126	4.42
3A	Zone 3	2201	-3.949	-2.246	0.621	0.538	2.65
3B	Zone 3	1540.28	-7.049	17.431	0.764	0.538	7.76

In order to examine further the influence of silicate, the residuals evaluated from regressions based upon only potential temperature, salinity, and AOU against silicate were plotted. This plot (not shown) showed that for silicate concentrations between 0 and -40 $\mu\text{mol/kg}$, the residuals averaged zero and there was no discernible trend; however, for silicate concentrations greater than -40 $\mu\text{mol/kg}$, there was a very clear positive correlation of the residuals with silicate. On the basis of this it was decided to define a new parameter, the "silicate index" (I_{Si}), as

$$I_{\text{Si}} = ([\text{SiO}_4] > 40) \times ([\text{SiO}_4] - 40) .$$

This index is equal to zero for SiO₄ concentrations less than 40 $\mu\text{mol/kg}$, and it is equal to $[\text{SiO}_4] - 40$ when silicate is greater than or equal to 40 $\mu\text{mol/kg}$.

The results of regressions using the silicate index, potential temperature, salinity, and AOU as independent variables are presented in Table 5. It can be seen that use of the silicate index, rather than the silicate concentration, makes the regression coefficients for the other parameters much more consistent from one geographical zone to another (see Table 4). Given the overall consistency of fit, it is right to use a single regression equation to predict the TCO₂ over the entire section (Regression no. 0, Table 5).

The distribution of residuals arising from this single section-wide regression equation is presented separately for the three defined geographical zones in Figure 5. Separate symbols are employed for the residuals derived from measurements made on the BNL and IfMK SOMMA systems. In general, there was little or no systematic structure apparent in the residual distribution (except perhaps at the very low TCO₂ concentrations that are found close to the surface where seasonal effects may be significant), and the regression fits the data from all three zones reasonably well.

Table 5. Summary of multiple regression results when the Silicate Index (I_{Si}) was used as a predictor

Regression no.	Longitude range	Intercept	Coefficients				Standard error
			Pot. temp.	Salinity	AOU	I_{Si}	
0	All section	1706.33	-5.747	12.474	0.722	0.434	4.59
1	Zone 1	1704.61	-5.565	12.443	0.746	0.422	4.98
2	Zone 2	1863.67	-6.075	8.217	0.618	0.77	4.17
3	Zone 3	1964.15	-5.867	5.135	0.688	0.464	2.99

In order to assess the intercomparability of TCO_2 measurements made on the two SOMMA systems on a station-by-station basis, the mean residuals (calculated for each station) have been plotted (Fig. 6) on the basis of the section-wide fit. This plot permits an assessment of the overall consistency of the measured TCO_2 with the other measured hydrographic parameters over the entire cruise. The plot demonstrates the following:

1. There is some slight spatial structure to the station-mean residuals, with the mean residual in zone 2 ($12.9^\circ W < \text{longitude} < 3^\circ E$) being $\sim 1\text{--}2 \mu\text{mol/kg}$ higher than for the rest of the section. No corresponding trend in the CRM analyses on the BNL system was seen (see Fig. 3), and it was therefore hypothesized that this slight variation is "real" and is associated with different origins for water masses in this zone.
2. In general, there is no consistent difference between the station-mean residuals on the basis of measurements made with the BNL and IfMK SOMMA systems. The overall consistency of the two sets of measurements appears to be better than $\pm 2 \mu\text{mol/kg}$, which is consistent with the accuracy and precision bounds ($\pm 1.9 \mu\text{mol/kg}$) for the overall data set. This confirms that the cruise-wide calibration of the TCO_2 data analyzed with the two instruments was nearly identical.
3. Three stations (625, 46, and 62) appear to be outliers from the overall pattern. These stations have a mean residual that is significantly different from the overall mean of the station-mean residuals analyzed with the BNL SOMMA. All of these three stations were measured by means of the IfMK SOMMA. While the possibility could not be ruled out that these deviations arise from errors in the measurement of the independent variables used in the regressions (e.g., the oxygen or silicate analyses or from "real" oceanographic variability), it was hypothesized that they were the result of calibration error of the TCO_2 analysis at these stations.

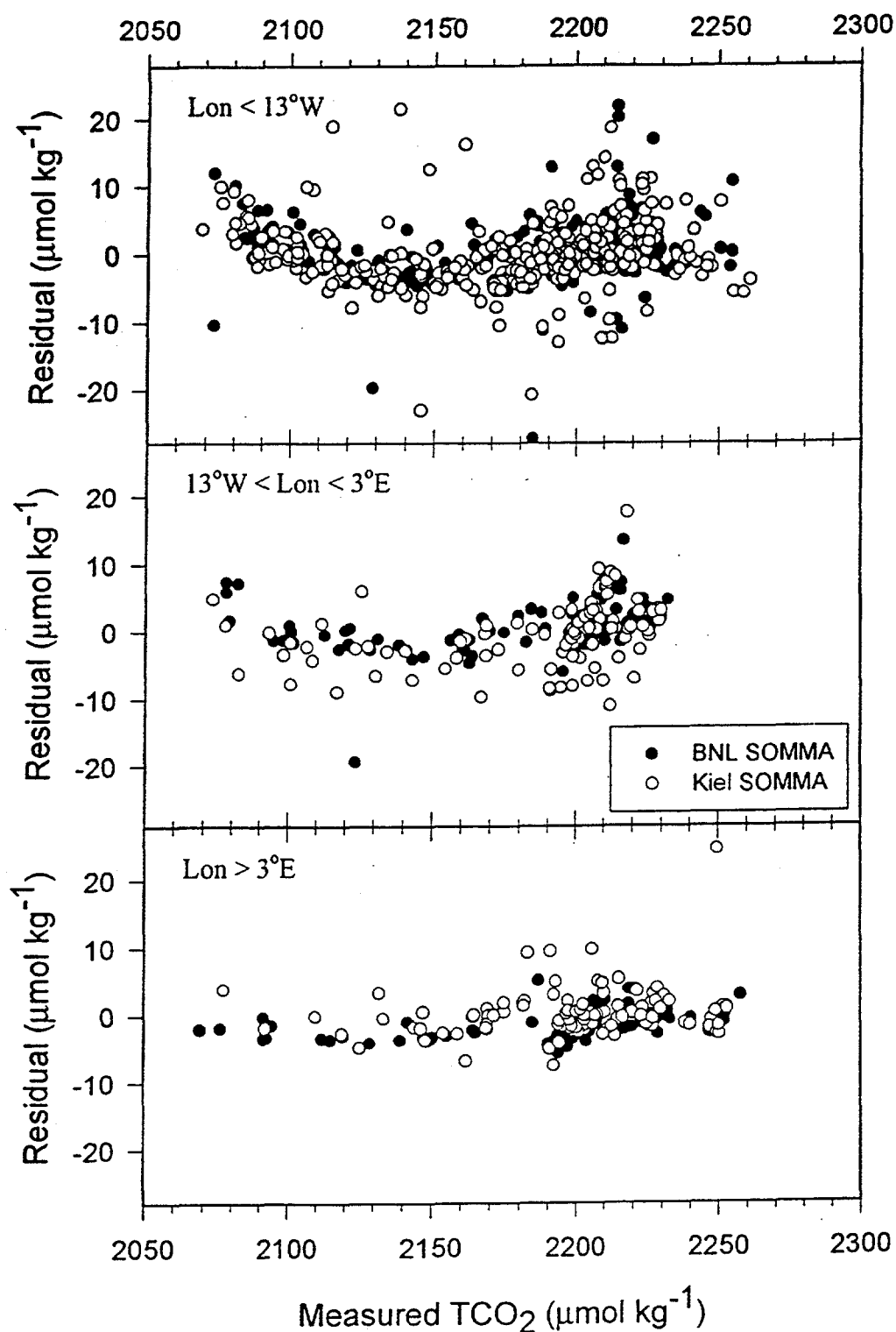


Figure 5. Residuals (observed – predicted) of TCO_2 versus the observed TCO_2 concentration for three separate geographical groupings of stations occupied during R/V *Meteor* Cruise 22/5. The residuals were evaluated against a multiple linear regression equation derived from all the TCO_2 data collected during the cruise (at depths > 200 m). The independent variables used in the regression were potential temperature, salinity, apparent oxygen utilization, and a silicate index. The regression coefficients used were those presented for regression no. 0 in Table 5.

Meteor 22; Cruise-Wide Regression

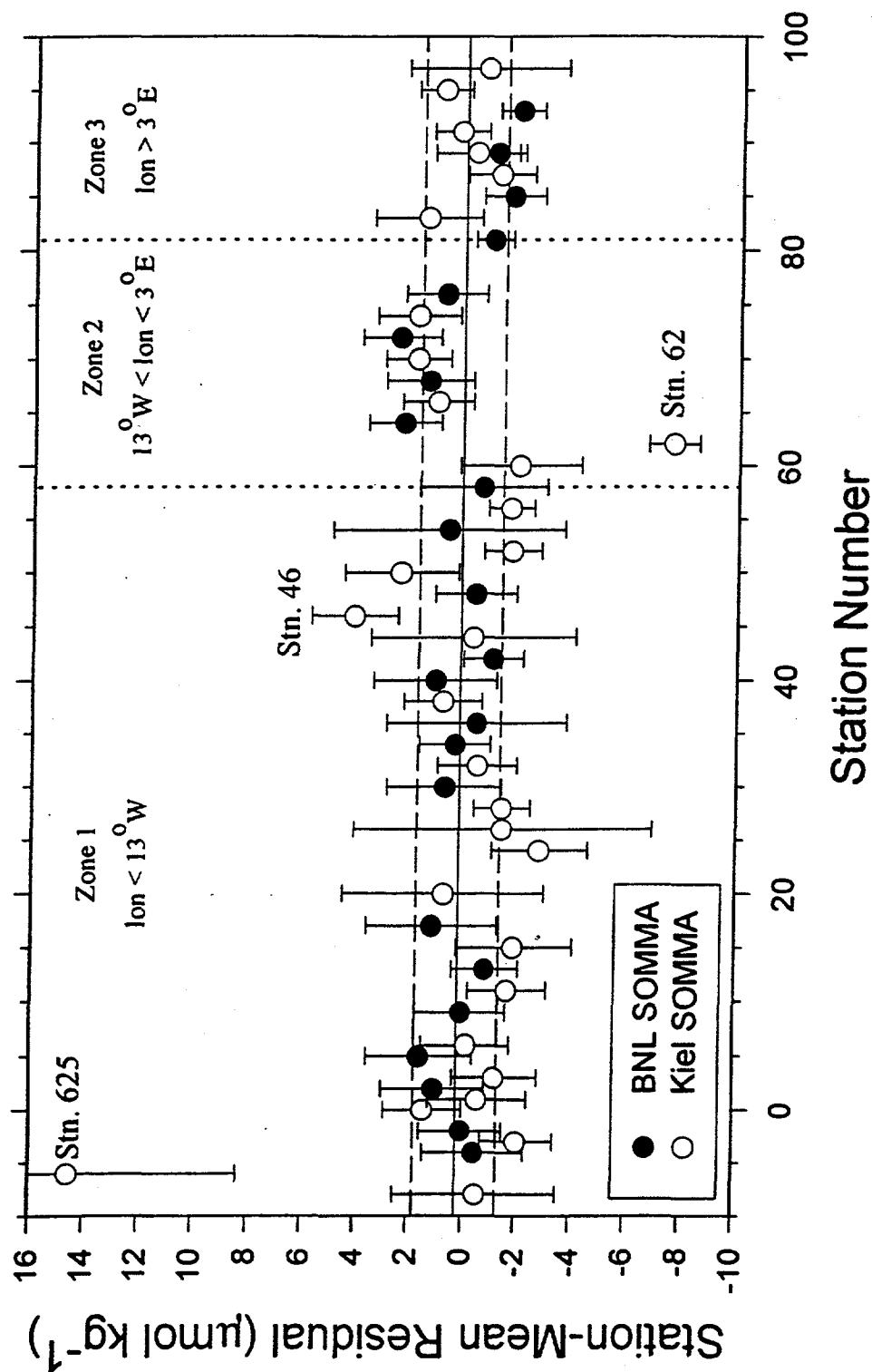


Figure 6. The mean TCO_2 residual (observed - predicted) for each station occupied during R/V *Meteor* Cruise 22/5; residuals were evaluated using a section-wide multiple linear regression (coefficients given for regression no. 0 in Table 5). The error bars denote the 95% confidence interval of the station-mean residual. Filled symbols reflect stations analyzed using the BNL SOMMA, open symbols reflect analyses using the IfMK (Kiel) SOMMA. The demarcation of three geographical zones reflecting three separate ocean basins sampled during the cruise is denoted with the vertical lines. The horizontal lines represent the cruise-wide mean and 95% confidence intervals calculated for the station-mean residuals using the BNL SOMMA results only.

Given the approach used to calibrate the IfMK SOMMA, such deviations could arise from a single incorrect analysis of a CRM which would cause the correction factor (CF_{crm} in Eq. 6) for an entire station to shift based on an incorrect analysis. The BNL SOMMA analyses were less prone to such errors because the primary calibration was based on analyses of pure CO_2 (gas calibration) with the CRM analyses being used as an independent cross-check on this primary calibration. With this approach, any calibration errors that may lead to systematic errors for an entire station are more likely to be identified and corrected.

The residual intercomparison confirms that the overall quality of the combined BNL and IfMK data set is very high. However, three anomalous stations were identified. The TCO_2 results at station 62 appear to be low by 5–7 $\mu\text{mol/kg}$. This station was also sampled for onshore manometric analyses, and the results in Table 3 confirm that, whereas other stations had a mean (ship-shore) difference of $-3.54 \mu\text{mol/kg}$ ($\pm 1.71 \mu\text{mol/kg}$), the shipboard analyses from station 62 were 8–12 $\mu\text{mol/kg}$ lower than the shore-based results. Therefore it was concluded from these two independent lines of evidence that the station 62 results are too low, and they have been flagged as incorrect in the original data file. On the basis of the residual analysis, the TCO_2 results at station 46 may also be high by $\sim 2\text{--}4 \mu\text{mol/kg}$, and TCO_2 results from station 625 may be high by as much as 14 $\mu\text{mol/kg}$. However, there is no independent way to assess the data from these stations, and the anomalous residual could be "real" as a result of error in the predictor variables. Therefore, the TCO_2 data from these stations have been flagged as "questionable." Only these three stations have been flagged; the data collected at the remaining 51 stations that were sampled for TCO_2 appear to be internally consistent.

3.3 TALK Measurements

TALK measurements were collected in 500 mL bottles for 665 samples from 26 stations with the same precautions as for TCO_2 . The bottles were stored in the dark at 4°C and analyzed within 24 hours. The samples were transferred into a closed titration cell with a volume of approximately 120 mL and titrated at $25 \pm 0.1^\circ\text{C}$ with 0.1 M HCl containing 0.6 M NaCl. The titration cell was based on the systems described by Bradshaw and Brewer (1988) and Millero et al. (1993). The potential was followed with an electrode pair consisting of a ROSS (Orion Inc.) glass pH electrode and a ROSS AgCl reference electrode connected to a high precision digital voltmeter. The titration was controlled by a computer that waited for stable emf-readings before adding the next acid increment. The titration curve was analyzed through a modified GRAN-plot method described by Stoll et al. (1993) that used the carbonic acid constants of Goyet and Poisson (1989) and that takes into account the silicate and phosphate concentrations of the sample to obtain the titration alkalinity. The precision of the method was $\pm 2.0 \mu\text{mol/kg}$ as determined by replicate analyses of samples. Standardization was accomplished with NaCO_3 standards in NaCl solutions corrected for the blank arising from impurities in the salt. No reference materials were analyzed for alkalinity during this cruise.

3.4 Underway pCO_2 Measurements

Underway pCO_2 was measured by the method of Schneider et al. (1992). Surface seawater was continuously pumped at a rate of 200–300 mL/min into a glass equilibrator with a volume of approximately 300 mL. The seawater was equilibrated with continuously circulating air entering the bottom of the equilibrator through a frit from a closed loop system. The latter included a heat exchanger to keep the air at the sample temperature, a filter and water trap, and an infrared analyzer (Siemens, Ultramat 5F) to determine the CO_2 content of the equilibrated air. The IR

analyzer and the equilibrator temperature sensor were connected to a PC and to an analog recorder for data display and preservation. The time constant for the equilibration was about 3 min, which corresponded to a spatial resolution of 0.5 mi with the ship speed at 10 kn. Atmospheric air was periodically measured, and the system was calibrated every 12 h using calibration gases with CO₂ mixing ratios of 252.5 and 412.8 ppm. Pressure corrections were made for the effect of water vapor and the pressure at the inlet of the IR analyzer, while the correction for the small difference between in situ and measuring temperature (<10°C) was made according to Gordon and Jones (1973). Fig. 7 shows the plot of underway measurements of temperature, salinity, sea surface pCO₂, and air pCO₂ during the R/V *Meteor* Cruise 22/5 in South Atlantic Ocean.

3.5 Secchi Disk Readings

Between December 30, 1992, and January 28, 1993, as the ship moved eastward, Secchi disk readings were made during daylight hours when the opportunity arose. These data are given in Table 6.

Table 6. Secchi Disk readings made during R/V *Meteor* Cruise 22/5

Date	Local time	Latitude (°S)	Longitude (-°W, +°E)	Conditions	Depth (m)
12/30/1992	15:30	27°55'	-46°40'	clear	25
12/31/1992	12:00	28°05'	-45°56'	clear	30
01/01/1993	17:00	28°50'	-43°35'	cloudy	19
01/03/1993	12:30	30°00'	-40°00'	cloudy	26
01/05/1993	15:30	30°00'	-36°10'	clear	33
01/06/1993	13:00	30°00'	-34°00'	partly cloudy	30
01/07/1993	13:00	30°00'	-32°00'	hazy	19
01/09/1993	16:00	30°00'	-27°00'	hazy	29
01/15/1993	18:00	30°00'	-13°40'	clear	31
01/16/1993	13:00	30°00'	-11°40'	clear	42
01/18/1993	13:00	30°00'	-07°00'	clear	42
01/20/1993	13:00	30°00'	-01°00'	clear	37
01/23/1993	16:30	29°45'	05°06'	sunny	32
01/28/1993	14:00	28°37'	14°41'	sunny	17
01/28/1993	17:00	28°30'	15°00'	sunny	7

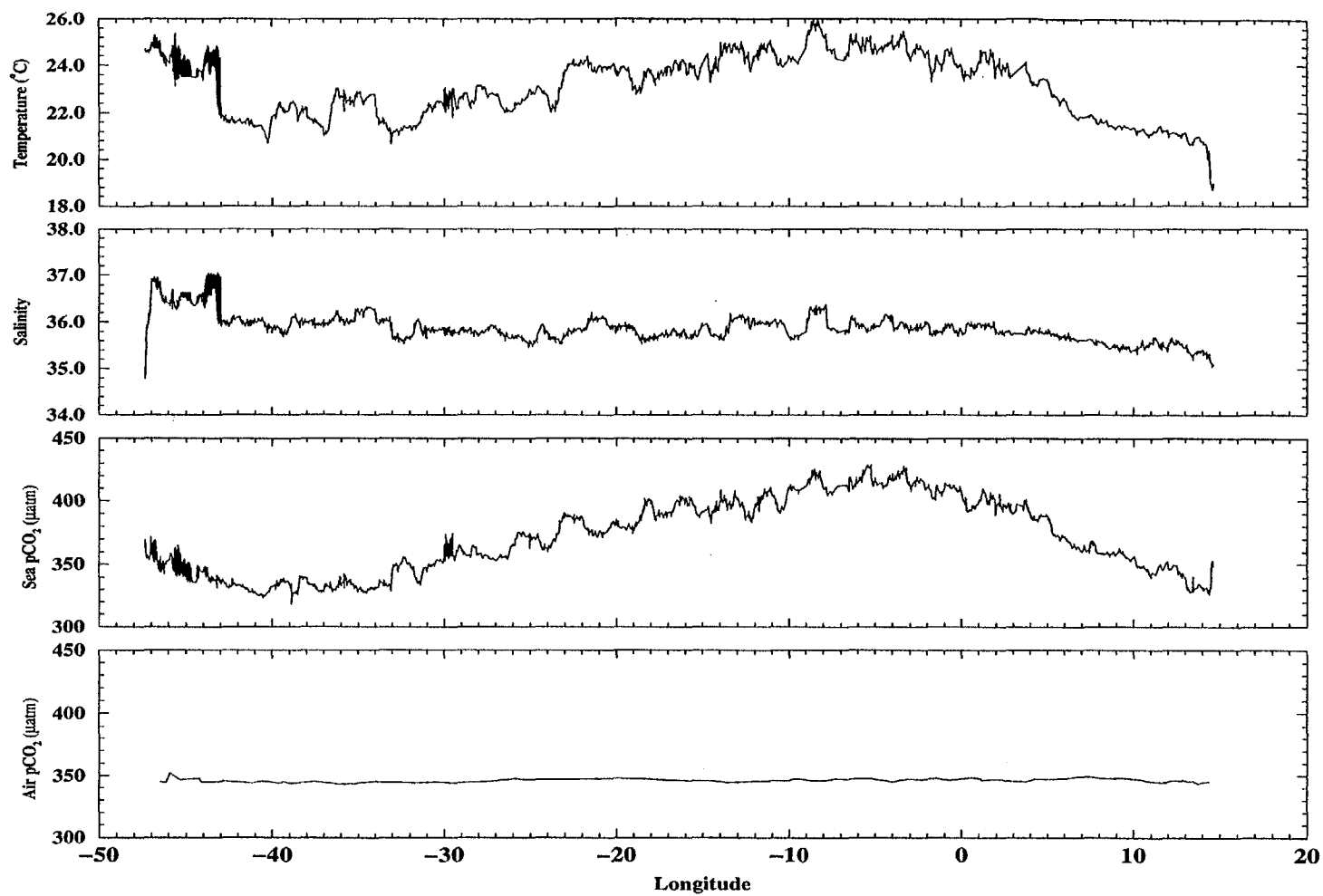


Figure 7. Plot of underway measurements of temperature, salinity, sea surface pCO₂, and air pCO₂ during R/V *Meteor* Cruise 22/5.

4. DATA CHECKS AND PROCESSING PERFORMED BY CDIAC

An important part of the NDP process at the Carbon Dioxide Information Analysis Center (CDIAC) involves the quality assurance (QA) of data before distribution. Data received at CDIAC are rarely in a condition that would permit immediate distribution, regardless of the source. To guarantee data of the highest possible quality, CDIAC conducts extensive QA reviews that involve examining the data for completeness, reasonableness, and accuracy. Although they have common objectives, these reviews are tailored to each data set, often requiring extensive programming efforts. In short, the QA process is a critical component in the value-added concept of supplying accurate, usable data for researchers.

The following information summarizes the data-processing and QA checks performed by CDIAC on the data obtained during the R/V *Meteor* Cruise 22/5 in the South Atlantic Ocean.

1. Carbon-related data and preliminary hydrographic measurements were provided to CDIAC by K. M. Johnson and D. W. R. Wallace of BNL. The final hydrographic and chemical measurements and the station information files were provided by the WOCE Hydrographic Program Office (WHPO) after quality evaluation. A FORTRAN 77 retrieval code was written and used to merge and reformat all data files.
2. To check for obvious outliers, all data were plotted by use of a PLOTNEST.C program written by Stewart C. Sutherland (Lamont-Doherty Earth Observatory). The program plots a series of nested profiles, using the station number as an offset; the first station is defined at the beginning, and subsequent stations are offset by a fixed interval (Figs. 8, 9).
3. To identify "noisy" data and possible systematic, methodological errors, property-property plots for all parameters were generated (Fig. 10), carefully examined, and compared with plots from previous expeditions in the South Atlantic Ocean.
4. All variables were checked for values exceeding physical limits, such as sampling depth values that are greater than the given bottom depths.
5. Dates, times, and coordinates were checked for bogus values (e.g., values of MONTH < 1 or > 12; DAY < 1 or > 31; YEAR < 1992 or > 1993; TIME < 0000 or > 2400; LAT < -25.000 or > 17.000; and LONG < -60.000 or > 20.000).
6. Station locations (latitudes and longitudes) and sampling times were examined for consistency with maps and cruise information supplied by K. M. Johnson and D. W. R. Wallace, BNL.
7. The designation for missing values, given as -9.0 in the original files, was changed to -999.9.

WOCE Section A10
Total CO₂ (all stations)

Profiles which exist in this Pressure (dbar) range are ordered on Station No.
Plotted parameter ranges from 1900 to 2400

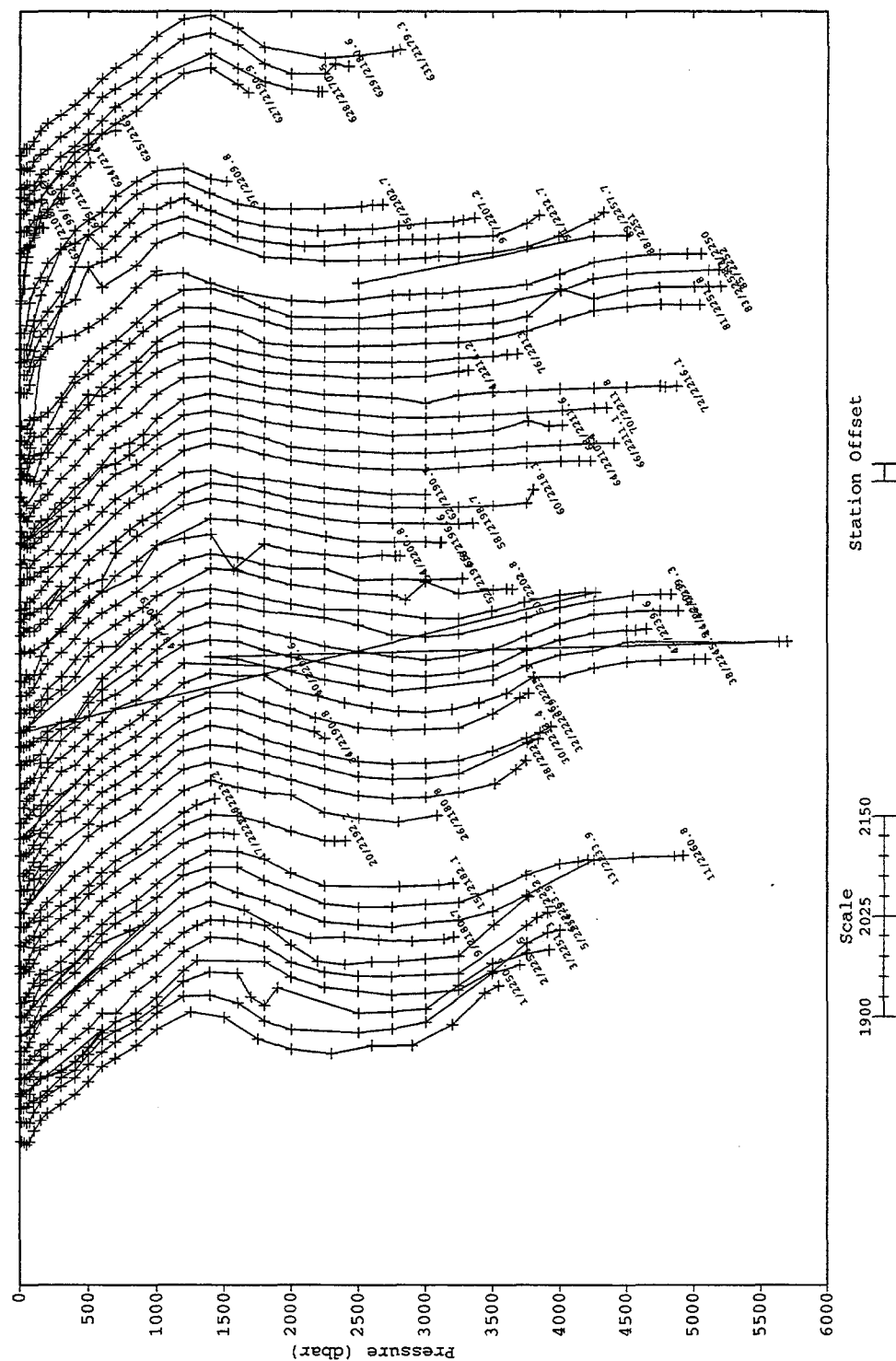


Figure 8. Nested profiles: total carbon dioxide ($\mu\text{mol/kg}$) vs pressure for all stations of WOCE Section A10.

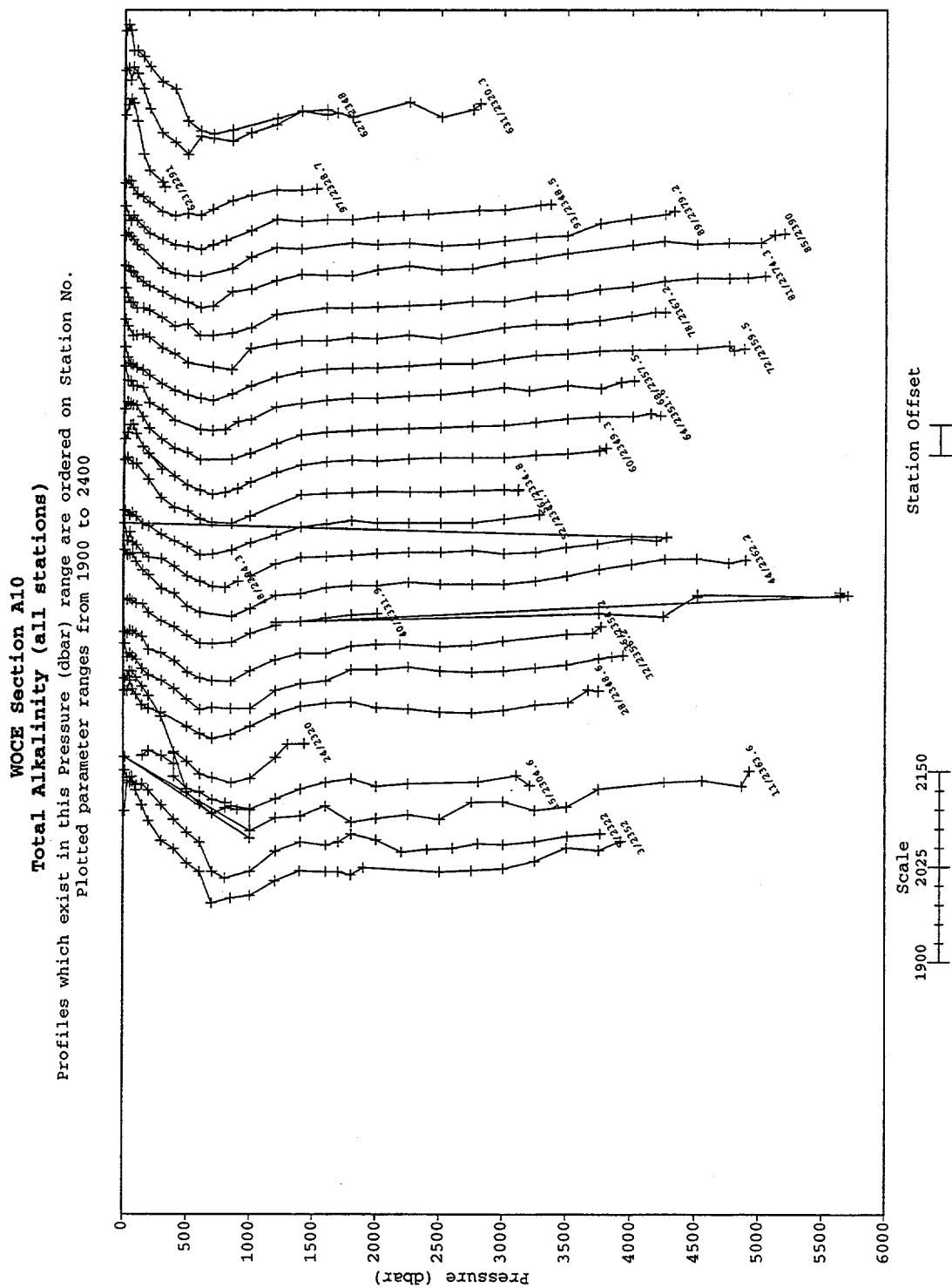


Figure 9. Nested profiles: total alkalinity ($\mu\text{mol/kg}$) vs pressure for all stations of WOCE Section A10.

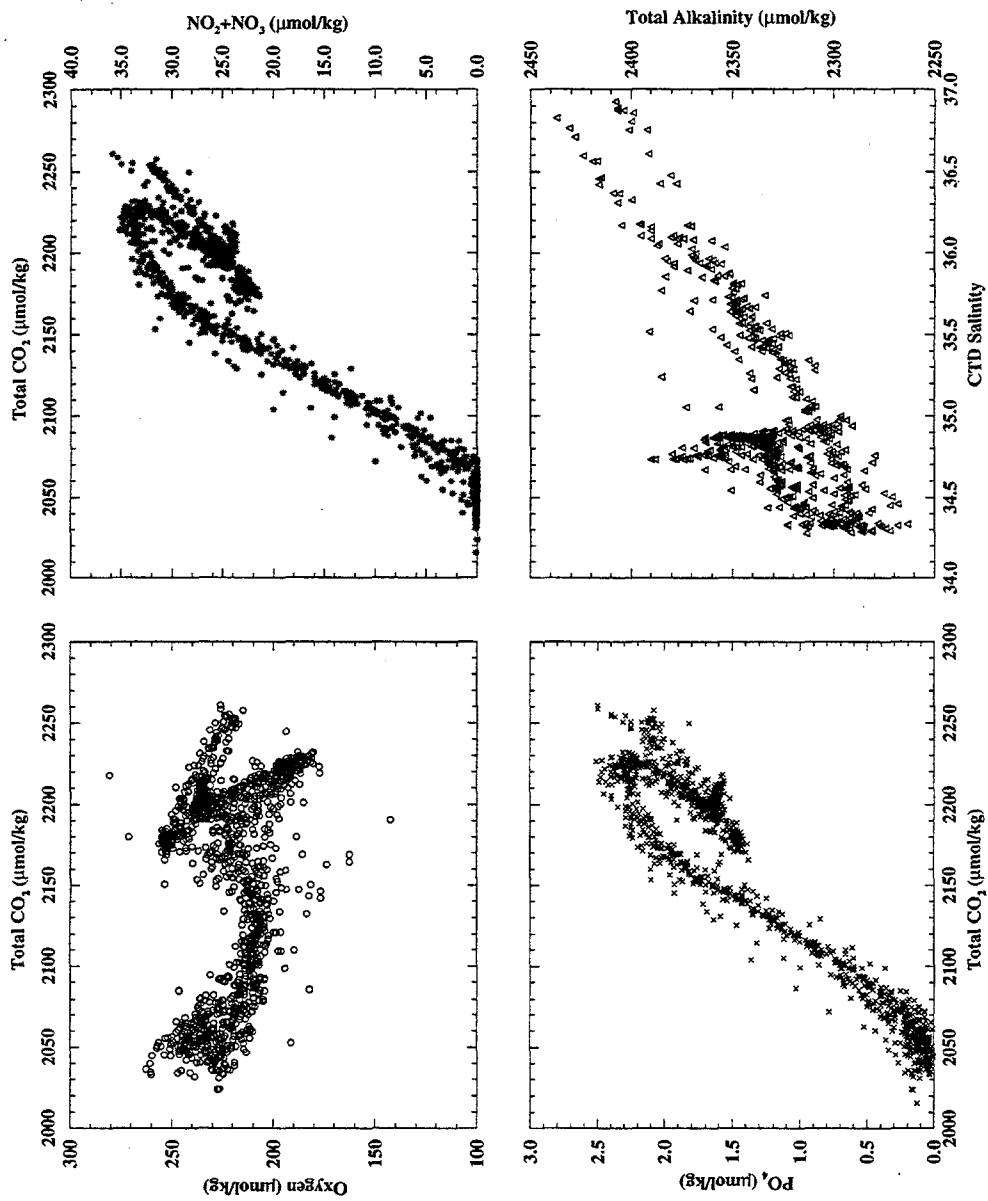


Figure 10. Property-property plots for all stations occupied during R/V Meteor Cruise 22/5.

5. HOW TO OBTAIN THE DATA AND DOCUMENTATION

This database (NDP-066) is available free of charge from CDIAC. The data are available from CDIAC's anonymous file transfer protocol (FTP) area via the Internet. Please note: your computer needs to have FTP software loaded on it (this is built in to most newer operating systems). Commands used to obtain the database are

```
>ftp cdiac.esd.ornl.gov or >ftp 128.219.24.36
Login: anonymous or ftp
Password: YOU@your internet address
Guest login ok, access restrictions apply.
ftp> cd pub/ndp066/
ftp> dir
ftp> mget (files)
ftp> quit
```

The complete documentation and data may also be obtained from CDIAC's Web site at the following URL: <http://cdiac.esd.ornl.gov/oceans/doc.html>

For non-FTP data acquisitions (e.g., floppy diskette, 8-mm tape, CD-ROM, etc.), users may order through CDIAC's online ordering system (http://cdiac.esd.ornl.gov/pns/how_order.html) or contact CDIAC directly to request the data and choice of media.

For additional information, contact CDIAC.

Address: Carbon Dioxide Information Analysis Center
Oak Ridge National Laboratory
P.O. Box 2008
Oak Ridge, Tennessee 37831-6335
U.S.A.

Telephone: (423) 574-3645 (Voice)
(423) 574-2232 (Fax)

Electronic mail: cdiac@ornl.gov

URL: <http://cdiac.esd.ornl.gov/>

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PART 2:
CONTENT AND FORMAT OF DATA FILES

7. FILE DESCRIPTIONS

This section describes the content and format of each of the seven files that comprise this NDP (see Table 7). Because CDIAC distributes the data set in several ways (e.g., via anonymous FTP and on floppy diskette), each of the seven files is referenced by both an ASCII file name, which is given in lower-case, bold-faced type (e.g., **ndp066.txt**) and a file number. The remainder of this section describes (or lists, where appropriate) the contents of each file.

Table 7. Content, size, and format of data files

File number, name, and description	Logical records	File size in bytes
1. ndp066.txt : a detailed description of the cruise network, the three FORTRAN 77 data-retrieval routines, and the three oceanographic data files	1,836	115,581
2. stainv.for : a FORTRAN 77 data-retrieval routine to read and print a10sta.txt (File 5)	45	1,342
3. a10dat.for : a FORTRAN 77 data-retrieval routine to read and print a10dat.txt (File 6)	55	2,075
4. a10pco2.for : a FORTRAN 77 data retrieval routine to read and print a10pco2.txt (File 7)	49	1,743
5. a10sta.txt : a listing of the station locations, sampling dates, and sounding bottom depths for each of the 112 stations	196	14,850
6. a10dat.txt : hydrographic, carbon dioxide, and chemical data from 112 stations	3,422	522,873

Table 7. (continued)

File number, name, and description	Logical records	File size in bytes
7. a10pco2.txt: underway measurements of pCO ₂ along the cruise track	2,822	304,320
Total	6,425	962,784

7.1 ndp066.txt (File 1)

This file contains a detailed description of the data set, the three FORTRAN 77 data retrieval routines, and the three oceanographic data files. It exists primarily for the benefit of individuals who acquire this database as machine-readable data files from CDIAC.

7.2 stainv.for (File 2)

This file contains a FORTRAN 77 data-retrieval routine to read and print **a10sta.txt** (File 5). The following is a listing of this program. For additional information regarding variable definitions, variable lengths, variable types, units, and codes, please see the description for **a10sta.txt**.

```

c*****
c* FORTRAN 77 data retrieval routine to read and print the file      *
c* named "a10sta.txt" (File 5).                                     *
c*****

c*Defines variables*

      INTEGER stat, cast, depth
      REAL latdcm, londcm
      CHARACTER expo*10, sect*9, date*10, time*4
      OPEN (unit=1, file='a10sta.txt')
      OPEN (unit=2, file='a10stat.txt')
      write (2, 5)

c*Writes out column labels*

5      format (1X, 'STATION INVENTORY: R/V Meteor Cruise 22/5',/,
1      1X, 'EXPOCODE', 3X, 'SECT', 1X, 'STNBR', 2X, 'CAST', 9X,
2      'DATE', 2X, 'TIME', 2X, 'LATITUDE', 2X, 'LONGITUDE', 2X,
3      'DEPTH', /)

```

c*Sets up a loop to read and format all the data in the file*

```

      read (1, 6)
6      format (/////////)

7      CONTINUE
      read (1, 10, end=999) expo, sect, stat, cast, date, time,
1 latdcm, londcm, depth

10     format (A9, 4X, A3, 3X, I3, 5X, I1, 3X, A10, 2X, A4, 3X,
1 F7.3, 3X, F8.3, 3X, I4)

      write (2, 20) expo, sect, stat, cast, date, time,
1 latdcm, londcm, depth

20     format (A9, 4X, A3, 3X, I3, 5X, I1, 3X, A10, 2X, A4, 3X,
1 F7.3, 3X, F8.3, 3X, I4)

      GOTO 7
999    close(unit=5)
      close(unit=2)
      stop
      end

```

7.3 a10dat.for (File 3)

This file contains a FORTRAN 77 data retrieval routine to read and print **a10dat.txt** (File 6). The following is a listing of this program. For additional information regarding variable definitions, variable lengths, variable types, units, and codes, please see the description for **a10dat.txt**.

```

c*****
c* FORTRAN 77 data retrieval routine to read and print the file      *
c* named "a10dat.txt" (File 6).                                     *
c*****
      CHARACTER qual*11
      INTEGER sta, cast, bot, pre, som
      REAL ctdtmp, ctdsal, theta, sal, oxy, silca
      REAL phspht, nitrx, cfc11, cfc12, tcarb, talk
      OPEN (unit=1, file='a10dat.txt')
      OPEN (unit=2, file='a10data.txt')
      write (2, 5)

c*Writes out column labels*

5      format (2X,'STNNBR',2X,'CASTNO',2X,'BTLNBR',2X,
1 'CTDPRS',2X,'CTDTMP',2X,'CTDSAL',3X,'THETA',4X,
2 'SALNTY',2X,'OXYGEN',2X,'SILCAT',2X,'PHSPHT',1X,
3 'NO2+NO3',3X,'CFC-11',3X,'CFC-12',2X,'TCARB',1X,
4 'SOMMA#',2X,'ALKALI',8X,'QUALT',/,
5 28X,'DBAR',2X,'ITS-90',2X,'PSS-78',2X,'ITS-90',
6 4X,'PSS-78',1X,4('UMOL/KG',1X),1X,'PMOL/KG',2X,'PMOL/KG',
7 1X,'UMOL/KG',8X,'UMOL/KG',12X,'*',/,
8 17X,'*****',17X,'*****',11X,5('*****',1X),1X,
9 '*****',2X,2('*****',1X),7X,'*****'12X,'*')

c*Sets up a loop to read and format all the data in the file*

```

```

        read (1, 6)
6      format (//////////)

7      CONTINUE
        read (1, 10, end=999) sta, cast, bot, pre, ctdtmp,
1      ctdsal, theta, sal, oxy, silca, phspht, nitrate,
2      cfc11, cfc12, tcarb, som, talk, qual

10     format (5X, I3, 7X, I1, 5X, I3, 4X, I4, 1X, F7.4,
1      1X, F7.4, 1X, F7.4, 1X, F9.4, 1X, F7.1, 1X, F7.2,
2      1X, F7.2, 1X, F7.2, 1X, F8.3, 1X, F8.3, 1X, F7.1,
3      5X, I2, 1X, F7.1, 2X, A11)

        write (2, 20) sta, cast, bot, pre, ctdtmp,
1      ctdsal, theta, sal, oxy, silca, phspht, nitrate,
2      cfc11, cfc12, tcarb, som, talk, qual

20     format (5X, I3, 7X, I1, 5X, I3, 4X, I4, 1X, F7.4,
1      1X, F7.4, 1X, F7.4, 1X, F9.4, 1X, F7.1, 1X, F7.2,
2      1X, F7.2, 1X, F7.2, 1X, F8.3, 1X, F8.3, 1X, F7.1,
3      5X, I2, 1X, F7.1, 2X, A11)

        GOTO 7
999    close(unit=1)
        close(unit=2)
        stop
        end

```

7.4 a10pco2.for (File 4)

This file contains a FORTRAN 77 data retrieval routine to read and print **a10pco2.txt** (File 7). The following is a listing of this program. For additional information regarding variable definitions, variable lengths, variable types, units, and codes, please see the description for **a10pco2.txt**.

```

c*****
c* FORTRAN 77 data retrieval routine to read and print the file      *
c* named "a10pco2.txt" (File 7).                                     *
c*****

c*Defines variables*

      REAL jday, latdcm, londcm, sst, salt, airpre, airxco2
      REAL airpco2, waterpco2
      CHARACTER sect*3, date*10, time*5
      OPEN (unit=1, file='a10pco2.txt')
      OPEN (unit=2, file='a10pco2.dat')
      write (2, 5)

c*Writes out column labels*

5      format (2X, 'UNDERWAY MEASUREMENTS R/V METEOR CRUISE 22/5',/,
1      2X, 'SECT', 9X, 'DATE', 4X, 'TIME', 5X, 'JULIAN', 3X, 'LATIT', 2X,
2      'LONGIT', 3X, 'SSTMP', 2X, 'SALNTY', 2X, 'ATMPRS', 4X, 'XCO2', 5X,
3      'PCO2AIR', 1X, 'PCO2WATER', /, 8X, 'DAY/MO/YEAR', 5X, 'GMT', 7X,
4      'DATE', 5X, 'DCM', 5X, 'DCM', 3X, 'DEG_C', 5X, 'PSS', 4X, 'ATM', 2X,
5      'DRY_AIR_PPM', 3X, 'UATM', 6X, 'UATM', /)

c*Sets up a loop to read and format all the data in the file*

```

```

        read (1, 6)
6       format (/////////)

7       CONTINUE
        read (1, 10, end=999) sect, date, time, jday, latdcm, londcm,
1 sst, salt, airpre, airxco2, airpco2, waterpco2

10      format (3X, A3, 3X, A10, 3X, A5, 2X, F9.3, 1X, F7.2, 1X,
1 F7.2, 1X, F7.2, 1X, F7.2, 1X, F7.1, 2X, F7.2, 3X, F7.1,
2 3X, F7.1)

        write (2, 20) sect, date, time, jday, latdcm, londcm, sst,
1 salt, airpre, airxco2, airpco2, waterpco2

20      format (3X, A3, 3X, A10, 3X, A5, 2X, F9.3, 1X, F7.2, 1X,
1 F7.2, 1X, F7.2, 1X, F7.2, 1X, F7.1, 2X, F7.2, 3X, F7.1,
2 3X, F7.1)

        GOTO 7
999     close(unit=1)
        close(unit=2)
        stop
        end

```

7.5 a10sta.txt (File 5)

This file provides station inventory information for each of the 112 stations occupied during R/V *Meteor* Cruise 22/5. Each line of the file contains an expocode, section number, station number, cast number, sampling date (month/date/year), sampling time, latitude, longitude, and sounding depth. The file is sorted by station number and can be read by using the following FORTRAN 77 code (contained in *stainv.for*, File 2):

```

        INTEGER stat, cast, depth
        CHARACTER expo*10, sect*3, date*10, time*4
        REAL latdcm, londcm

        read (1, 10, end=999) expo, sect, stat, cast, date, time,
1 latdcm, londcm, depth

10      format (A9, 4X, A3, 3X, I3, 5X, I1, 3X, A10, 2X, A4, 3X,
1 F7.3, 3X, F8.3, 3X, I4)

```

Stated in tabular form, the contents include the following:

Variable	Variable type	Variable width	Starting column	Ending column
expo	Character	10	1	10
sect	Character	3	14	16
stat	Numeric	3	20	22
cast	Numeric	1	28	28
date	Character	10	32	41
time	Character	4	44	47
latdcm	Numeric	7	51	57
londcm	Numeric	8	61	68
depth	Numeric	4	72	75

The variables are defined as follows:

expo is the expocode of the cruise;

sect is the WOCE section number;

stat is the station number;

cast is the cast number;

date is the sampling date (month/day/year);

time is the sampling time [Greenwich mean time (GMT)];

latdcm is the latitude of the station (in decimal degrees; negative values indicate the Southern Hemisphere);

londcm is the longitude of the station (in decimal degrees; negative values indicate the Western Hemisphere);

depth is the sounding depth of the station (in meters).

7.6 a10dat.txt (File 6)

This file provides hydrographic, carbon dioxide, and chemical data for the 112 stations occupied during R/V *Meteor* Cruise 22/5. Each line consists of a station number, cast number, bottle number, CTD pressure, CTD temperature, CTD salinity, potential temperature, bottle salinity,

oxygen, silicate, phosphate, nitrate plus nitrite, CFC-11, CFC-12, total CO₂, SOMMA number, total alkalinity, and data-quality flags. The file is sorted by station number and pressure and can be read by using the following FORTRAN 77 code (contained in **10dat.for**, File 3):

```

      CHARACTER qual*11
      INTEGER sta, cast, bot, pre, som
      REAL ctdtmp, ctdsal, theta, sal, oxy, silca
      REAL phspht, nitr, cfc11, cfc12, tcarb, talk

      read (1, 10, end=999) sta, cast, bot, pre, ctdtmp,
1 ctdsal, theta, sal, oxy, silca, phspht, nitr,
2 cfc11, cfc12, tcarb, som, talk, qual

10    format (5X, I3, 7X, I1, 5X, I3, 4X, I4, 1X, F7.4,
1 1X, F7.4, 1X, F7.4, 1X, F9.4, 1X, F7.1, 1X, F7.2,
2 1X, F7.2, 1X, F7.2, 1X, F8.3, 1X, F8.3, 1X, F7.1,
3 5X, I2, 1X, F7.1, 2X, A11)

```

Stated in tabular form, the contents include the following:

Variable	Variable type	Variable width	Starting column	Ending column
sta	Numeric	3	6	8
cast	Numeric	1	16	16
bot	Numeric	3	22	24
pre	Numeric	4	29	32
ctdtmp	Numeric	7	34	40
ctdsal	Numeric	7	42	48
theta	Numeric	7	50	56
sal	Numeric	9	58	66
oxy	Numeric	7	68	74
silca	Numeric	7	76	82
phspht	Numeric	7	84	90
nitr	Numeric	7	92	98
cfc11	Numeric	8	100	107
cfc12	Numeric	8	109	116
tcarb	Numeric	7	118	124
som	Numeric	2	130	131
talk	Numeric	7	133	139
qualt	Character	11	142	152

The variables defined as follows:

sta is the station number;

cast is the cast number;

bot^a	is the bottle number;
pre	is the CTD pressure (in dbar);
ctdtmp	is the CTD temperature (in °C);
ctdsal^a	is the CTD salinity [on the Practical Salinity Scale (PSS)];
theta	is the potential temperature (in °C);
sal^a	is the bottle salinity (on the PSS);
oxy^a	is the oxygen concentration (in $\mu\text{mol/kg}$);
silca^a	is the silicate concentration (in $\mu\text{mol/kg}$);
phspht^a	is the phosphate concentration (in $\mu\text{mol/kg}$);
nitr^a	is the nitrate plus nitrite concentration (in $\mu\text{mol/kg}$);
cfc11^a	is the trichlorofluoromethane-11 concentration (CCl_3F) (in pmol/kg);
cfc12^a	is the dichlorodifluoromethane-12 concentration (CCl_2F_2) (in pmol/kg);
tcarb^a	is the total carbon dioxide concentration (in $\mu\text{mol/kg}$);
som	is the SOMMA number;
talk	is the total alkalinity concentration (in $\mu\text{mol/kg}$);
qualt	is a 11-digit character variable that contains data-quality flag codes for parameters underlined with asterisks ("*****") in the file header.

^aVariables that are underlined with asterisks in the data file's header indicate they have a data-quality flag. Data-quality flags are defined as follows:

- 1 = sample for this measurement was drawn from water bottle but analysis was not received;
- 2 = acceptable measurement;
- 3 = questionable measurement;
- 4 = bad measurement;
- 5 = not reported;
- 6 = mean of replicate measurements;
- 7 = manual chromatographic peak measurement;
- 8 = irregular digital chromatographic peak integration;
- 9 = sample not drawn for this measurement from this bottle.

7.7 a10pco2.txt (File 7)

This file provides underway measurements of $p\text{CO}_2$ during R/V *Meteor* Cruise 22/5. Each line of the file contains a sampling date (month/date/year), latitude, longitude, underway measurements of sea surface temperature, salinity, air $p\text{CO}_2$, and water $p\text{CO}_2$. The file is sorted by longitude and can be read by using the following FORTRAN 77 code (contained in a10pco2.for, File 4):

```

REAL jday, latdcm, londcm, sst, salt, airpre, airxco2
REAL airpco2, waterpco2
CHARACTER sect*3, date*10, time*5

read (1, 10, end=999) sect, date, time, jul, latdcm, londcm,
1 sst, salt, pre, xco2, pco2a, pco2w

10  format (3X, A3, 3X, A10, 3X, A5, 2X, F9.3, 1X, F7.2, 1X,
1 F7.2, 1X, F7.2, 1X, F7.2, 1X, F7.1, 2X, F7.2, 3X, F7.1,
2 3X, F7.1)

```

Stated in tabular form, the contents include the following:

Variable	Variable type	Variable width	Starting column	Ending column
sect	Character	3	4	6
date	Character	10	10	19
time	Character	5	23	27
jday	Numeric	9	30	38
latdcm	Numeric	7	40	46
londcm	Numeric	7	48	54
sst	Numeric	7	56	62
salt	Numeric	7	64	70
airpre	Numeric	7	72	78
airxco2	Numeric	7	81	87
airpco2	Numeric	7	91	97
waterpco2	Numeric	7	101	107

The variables are defined as follows:

sect is the WOCE Section number;

date is the sampling date (day/month/year);

time is the sampling time (GMT);

jday	is the julian day of the century relative to 1900 with time of the day represented as a fractional day (i.e., noon on 1/1/1900 = 0.5);
latdcm	is the latitude of the sampling (in decimal degrees; negative values indicate the Southern Hemisphere);
londcm	is the longitude of the sampling (in decimal degrees; negative values indicate the Western Hemisphere);
sst	is the sea surface temperature (in °C);
salt	is the sea surface salinity (on the PSS);
airpre	is the atmospheric pressure (in atm);
airxco2	is the observed mole fraction of pCO ₂ in air [in ppm (dry air)];
airpco2	is the air pCO ₂ (in μatm);
waterpco2	is the sea surface water pCO ₂ (in μatm).

8. VERIFICATION OF DATA TRANSPORT

The data files contained in this numeric data package can be read by using the FORTRAN 77 data retrieval programs provided. Users should visually examine each data file to verify that the data were correctly transported to their systems. To facilitate the visual inspection process, partial listings of each data file are provided in Tables 8, 9, and 10. Each of these tables contains the first and last twenty five lines of a data file.

Table 8. Partial listing of a10sta.txt (File 5)

First twenty-five lines of the file:

```
*****
* Source: K. Johnson                      B. Schneider                      *
*           D. Wallace                    Baltic Sea Research Institute      *
*                                           Warnemunde, Germany              *
* Brookhaven National Laboratory          L. Mintrop                      *
* Apton, New York, USA                   Institute for Marine Sciences    *
* CDIAC NDP-066, September 1998         Kiel, Germany                  *
*****
*STATION INVENTORY: R/V METEOR
*EXPCODE SECT STNBR CAST DATE TIME LATITUDE LONGITUDE DEPTH
06MT22/5 A10 620 1 12/28/1992 1506 -25.645 -42.175 2290
06MT22/5 A10 622 1 12/30/1992 0626 -27.730 -47.385 177
06MT22/5 A10 623 1 12/30/1992 0914 -27.774 -47.207 326
06MT22/5 A10 624 1 12/30/1992 1129 -27.816 -47.026 536
06MT22/5 A10 625 1 12/30/1992 1348 -27.860 -46.847 758
06MT22/5 A10 626 1 12/30/1992 1755 -27.905 -46.670 1250
06MT22/5 A10 627 1 12/30/1992 2212 -27.950 -46.485 1693
06MT22/5 A10 628 1 12/31/1992 0224 -27.992 -46.308 2217
06MT22/5 A10 629 1 12/31/1992 0650 -28.039 -46.129 2408
06MT22/5 A10 630 1 12/31/1992 1140 -28.088 -45.939 2596
06MT22/5 A10 631 1 12/31/1992 1712 -28.153 -45.674 2782
06MT22/5 A10 632 1 12/31/1992 2240 -28.224 -45.408 2965
06MT22/5 A10 1 2 01/01/1993 0726 -28.418 -44.768 3509
06MT22/5 A10 2 1 01/01/1993 1315 -28.615 -44.222 3694
```

Last twenty-five lines of the file:

```
06MT22/5 A10 85 2 01/24/1993 1622 -29.746 7.621 5200
06MT22/5 A10 85 3 01/24/1993 1816 -29.736 7.627 5203
06MT22/5 A10 86 1 01/25/1993 0110 -29.746 8.465 5091
06MT22/5 A10 86 2 01/25/1993 0247 -29.744 8.465 5086
06MT22/5 A10 87 2 01/25/1993 1036 -29.750 9.297 5032
06MT22/5 A10 87 3 01/25/1993 1235 -29.755 9.307 4986
06MT22/5 A10 88 1 01/25/1993 1925 -29.748 10.148 4900
06MT22/5 A10 88 2 01/25/1993 2141 -29.747 10.185 4853
06MT22/5 A10 89 2 01/26/1993 0527 -29.749 10.980 4296
06MT22/5 A10 89 3 01/26/1993 0700 -29.748 10.981 4271
06MT22/5 A10 90 2 01/26/1993 1419 -29.752 11.830 4002
06MT22/5 A10 90 3 01/26/1993 1615 -29.755 11.852 3994
06MT22/5 A10 91 2 01/26/1993 2025 -29.622 12.171 3818
06MT22/5 A10 91 3 01/26/1993 2150 -29.621 12.187 3812
06MT22/5 A10 92 2 01/27/1993 0248 -29.497 12.467 3657
06MT22/5 A10 92 3 01/27/1993 0436 -29.489 12.474 3652
06MT22/5 A10 93 2 01/27/1993 0944 -29.373 12.795 3358
06MT22/5 A10 93 3 01/27/1993 1104 -29.372 12.815 3341
06MT22/5 A10 94 2 01/27/1993 1620 -29.243 13.114 3098
06MT22/5 A10 95 2 01/27/1993 2135 -29.119 13.428 2640
06MT22/5 A10 96 2 01/28/1993 0248 -29.001 13.735 2136
06MT22/5 A10 97 2 01/28/1993 0645 -28.879 14.045 1490
06MT22/5 A10 98 2 01/28/1993 0955 -28.752 14.366 461
06MT22/5 A10 99 2 01/28/1993 1240 -28.622 14.687 160
06MT22/5 A10 100 2 01/28/1993 1457 -28.503 14.999 174
```

Table 9. Partial listing of a10dat.txt (File 6)

First twenty-five lines of the file

*	Source: K. Johnson				B. Schneider							*						
*	D. Wallace				Baltic Sea Research Institute							*						
*					Warnemunde, Germany							*						
*	Brookhaven National Laboratory											*						
*	Apton, New York, USA				L. Mintrop							*						
*					Institute for Marine Sciences							*						
*	CDIAC NDP-066, September 1998				Kiel, Germany							*						

*	EXPOCODE 06MT22/5 R/V METEOR 22 LEG 5 WOCE SECTION A10															*		
* STNNBR	CASN0	BTLNBR	CTDPNS	CTDTMP	CTDSAL	THETA	SALTNY	OXYGEN	SILCAT	PHPSPH	NO2+NO3	CFC-11	CFC-12	TCARBN	SOMMA#	ALKALI	QUALT	
*			DBAR	ITS-90	PSS-78	ITS-90	PSS-78	UMOL/KG	UMOL/KG	UMOL/KG	UMOL/KG	PMOL/KG	PMOL/KG	UMOL/KG		UMOL/KG		
*	*****																	
*	620	1	324	2271	3.0620	34.9410	2.8832	34.9400	-999.9	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	222999999999	
*	620	1	323	2271	3.0610	34.9410	2.8822	34.9400	-999.9	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	222999999999	
*	620	1	322	2270	3.0600	34.9410	2.8813	34.9380	-999.9	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	222999999999	
*	620	1	321	2270	3.0590	34.9420	2.8803	34.9370	-999.9	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	222999999999	
*	620	1	320	2270	3.0590	34.9420	2.8803	34.9370	-999.9	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	222999999999	
*	620	1	319	2270	3.0610	34.9420	2.8823	34.9370	-999.9	-999.90	-999.90	-999.900	0.019	0.180	-999.9	-9	-999.9	222999924999
*	620	1	318	2269	3.0610	34.9410	2.8824	34.9380	-999.9	-999.90	-999.90	-999.900	0.055	0.273	-999.9	-9	-999.9	222999924999
*	620	1	317	2269	3.0580	34.9420	2.8794	34.9400	-999.9	-999.90	-999.90	-999.900	0.221	0.228	-999.9	-9	-999.9	222999944999
*	620	1	316	2268	3.0630	34.9420	2.8844	34.9400	-999.9	-999.90	-999.90	-999.900	0.009	0.161	-999.9	-9	-999.9	222999924999
*	620	1	315	2268	3.0610	34.9410	2.8825	34.9400	-999.9	-999.90	-999.90	-999.900	0.032	0.040	-999.9	-9	-999.9	222999922999
*	620	1	314	2268	3.0590	34.9410	2.8805	34.9380	-999.9	-999.90	-999.90	-999.900	0.009	0.095	-999.9	-9	-999.9	222999922999
*	620	1	313	2267	3.0620	34.9420	2.8835	34.9400	-999.9	-999.90	-999.90	-999.900	0.016	0.113	-999.9	-9	-999.9	222999922999

Last twenty-five lines of the file

99	2	315	30	17.6020	35.2190	17.5970	-999.9000	226.2	3.24	0.36	3.40	-999.900	-999.900	2054.9	1	-999.9	22922229929
99	2	314	30	17.5930	35.2190	17.5880	-999.9000	-999.9	-999.90	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	22999999999
99	2	313	30	17.5930	35.2200	17.5880	-999.9000	-999.9	-999.90	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	22999999999
99	2	312	49	14.5570	35.2530	14.5498	-999.9000	210.0	4.75	0.67	8.74	-999.900	-999.900	2087.8	1	-999.9	22922229929
99	2	311	49	14.5320	35.2540	14.5248	-999.9000	-999.9	-999.90	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	22999999999
99	2	310	49	14.5280	35.2560	14.5208	-999.9000	-999.9	-999.90	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	22999999999
99	2	309	70	12.7110	35.1020	12.7016	-999.9000	204.3	5.67	0.88	12.55	-999.900	-999.900	2108.5	1	-999.9	22922229929
99	2	308	70	12.7200	35.1030	12.7106	-999.9000	-999.9	-999.90	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	22999999999
99	2	307	70	12.7170	35.1030	12.7076	-999.9000	-999.9	-999.90	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	22999999999
99	2	306	101	11.7430	34.9880	11.7301	-999.9000	200.0	7.09	1.06	15.15	-999.900	-999.900	2121.1	1	-999.9	22922229929
99	2	305	101	11.7490	34.9890	11.7361	-999.9000	-999.9	-999.90	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	22999999999
99	2	304	100	11.7540	34.9900	11.7412	-999.9000	-999.9	-999.90	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	22999999999
99	2	303	157	9.8380	34.7720	9.8201	-999.9000	162.6	13.43	1.59	21.95	-999.900	-999.900	2164.5	1	-999.9	22922229929
99	2	302	156	9.8350	34.7710	9.8172	-999.9000	-999.9	-999.90	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	22999999999
99	2	301	156	9.8320	34.7710	9.8140	-999.9000	-999.9	-999.90	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	22999999999
100	2	324	13	18.3230	34.9820	18.3207	-999.9000	-999.9	-999.90	-999.90	-999.90	-999.900	-999.900	-999.9	-9	-999.9	22999999999
100	2	315	13	18.2810	34.9810	18.2787	-999.9000	248.8	3.02	0.35	1.27	-999.900	-999.900	-999.9	-9	-999.9	22922229999
100	2	313	30	16.7330	34.9500	16.7281	-999.9000	236.1	3.19	0.52	3.83	-999.900	-999.900	-999.9	-9	-999.9	22922229999
100	2	311	50	14.9330	35.0500	14.9255	-999.9000	213.9	4.34	0.70	8.33	-999.900	-999.900	-999.9	-9	-999.9	22922229999
100	2	309	70	12.3020	35.0480	12.2928	-999.9000	192.6	6.12	1.01	14.71	-999.900	-999.900	-999.9	-9	-999.9	22922229999
100	2	307	99	10.9840	34.8930	10.9718	-999.9000	185.6	8.40	1.23	17.81	-999.900	-999.900	-999.9	-9	-999.9	22922229999
100	2	306	100	11.0030	34.8980	10.9907	-999.9000	-999.9	-999.90	-999.90	-999.90	-999.900	-999.900				

Table 10. Partial listing of a10pco2.txt (File 7)

First twenty-five lines of the file:

```
*****
* Source: K. Johnson                      B. Schneider *
*         D. Wallace                      Baltic Sea Research Institute *
*                                         Warnemunde, Germany *
* Brookhaven National Laboratory *
* Apton, New York, USA *
*                                         L. Mintrop *
*                                         Institute for Marine Sciences *
* CDIAC NDP-066, September 1998 *
*                                         Kiel, Germany *
*****
*UNDERWAY MEASUREMENTS: R/V METEOR CRUISE 22/5
* DATE LATITUDE LONGITUDE TEMP SALT AIR_PCO2 WAT_PCO2
30/Dec/1992 -27.731 -47.385 24.70 34.83 -999.9 369.2
30/Dec/1992 -27.730 -47.384 24.67 34.83 -999.9 365.3
30/Dec/1992 -27.731 -47.384 24.67 34.84 -999.9 368.6
30/Dec/1992 -27.728 -47.383 24.68 34.82 -999.9 363.3
30/Dec/1992 -27.727 -47.383 24.69 34.78 -999.9 363.0
30/Dec/1992 -27.739 -47.342 24.59 35.19 -999.9 364.9
30/Dec/1992 -27.747 -47.311 24.61 35.45 -999.9 357.2
30/Dec/1992 -27.755 -47.280 24.61 35.68 -999.9 356.2
30/Dec/1992 -27.763 -47.249 24.56 35.94 -999.9 354.9
30/Dec/1992 -27.771 -47.218 24.66 35.88 -999.9 357.1
30/Dec/1992 -27.778 -47.185 24.61 35.99 -999.9 355.2
30/Dec/1992 -27.787 -47.153 24.54 36.19 -999.9 353.8
30/Dec/1992 -27.795 -47.123 24.67 36.10 -999.9 356.0
30/Dec/1992 -27.805 -47.092 24.64 36.43 -999.9 356.3
```

Last twenty-five lines of the file:

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28/Jan/1993 -28.910 13.980 20.78 35.41 -999.9 331.9
28/Jan/1993 -28.890 14.010 20.77 35.38 -999.9 331.0
28/Jan/1993 -28.890 14.030 20.76 35.38 -999.9 329.4
28/Jan/1993 -28.880 14.040 20.76 35.39 -999.9 329.6
28/Jan/1993 -28.880 14.050 20.76 35.36 344.8 -999.9
28/Jan/1993 -28.890 14.050 20.76 35.34 -999.9 331.7
28/Jan/1993 -28.880 14.060 20.64 35.32 -999.9 331.4
28/Jan/1993 -28.870 14.090 20.62 35.35 -999.9 331.4
28/Jan/1993 -28.860 14.120 20.62 35.34 -999.9 330.5
28/Jan/1993 -28.840 14.150 20.66 35.32 -999.9 331.3
28/Jan/1993 -28.830 14.180 20.65 35.34 -999.9 331.5
28/Jan/1993 -28.810 14.210 20.61 35.36 -999.9 331.6
28/Jan/1993 -28.800 14.240 20.35 35.25 -999.9 330.3
28/Jan/1993 -28.790 14.270 20.16 35.31 -999.9 329.6
28/Jan/1993 -28.770 14.300 19.97 35.27 -999.9 327.6
28/Jan/1993 -28.760 14.330 20.34 35.26 -999.9 330.3
28/Jan/1993 -28.760 14.350 20.35 35.24 -999.9 329.9
28/Jan/1993 -28.750 14.380 20.21 35.34 345.2 -999.9
28/Jan/1993 -28.740 14.400 19.70 35.28 -999.9 327.5
28/Jan/1993 -28.730 14.430 19.94 35.21 -999.9 330.6
28/Jan/1993 -28.710 14.460 19.06 35.14 -999.9 331.7
28/Jan/1993 -28.700 14.500 18.87 35.12 -999.9 346.5
28/Jan/1993 -28.690 14.530 18.71 35.06 -999.9 350.6
28/Jan/1993 -28.680 14.560 18.69 35.10 -999.9 352.9
28/Jan/1993 -28.660 14.580 18.76 35.09 -999.9 352.6
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APPENDIX A:
STATION INVENTORY

APPENDIX A: STATION INVENTORY

This appendix lists station inventory information for the 112 sites occupied during R/V *Meteor* Cruise 22/5 in the South Atlantic Ocean. The meanings of the column headings in Table A-1 are as follows.

EXPOCODE	is the expocode of the cruise;
SECT	is the WOCE section number;
STNNBR	is the station number;
CASTNO	is the cast number;
DATE	is the sampling date (month/day/year);
TIME	is the sampling time (GMT);
LATDCM	is the latitude of the station (in decimal degrees). Stations in the Southern Hemisphere have negative latitudes;
LONDCM	is the longitude of the station (in decimal degrees). Stations in the Western Hemisphere have negative longitudes;
DEPTH	is the sounding bottom depth of each station (in meters).

Table A.1 Station inventory information for the 112 sites occupied
during R/V Meteor Cruise 22/5

*	Source: K. Johnson			B. Schneider			*	
*	D. Wallace			Baltic Sea Research Institute			*	
*				Warnemunde, Germany			*	
*	Brookhaven National Laboratory						*	
*	Apton, New York, USA			L. Mintrop			*	
*				Institute for Marine Sciences			*	
*	CDIAC NDP-066, September 1998			Kiel, Germany			*	

*STATION INVENTORY: R/V METEOR								
*EXPCODE	SECT	STNBR	CAST	DATE	TIME	LATITUDE	LONGITUDE	DEPTH
06MT22/5	A10	620	1	12/28/1992	1506	-25.645	-42.175	2290
06MT22/5	A10	622	1	12/30/1992	0626	-27.730	-47.385	177
06MT22/5	A10	623	1	12/30/1992	0914	-27.774	-47.207	326
06MT22/5	A10	624	1	12/30/1992	1129	-27.816	-47.026	536
06MT22/5	A10	625	1	12/30/1992	1348	-27.860	-46.847	758
06MT22/5	A10	626	1	12/30/1992	1755	-27.905	-46.670	1250
06MT22/5	A10	627	1	12/30/1992	2212	-27.950	-46.485	1693
06MT22/5	A10	628	1	12/31/1992	0224	-27.992	-46.308	2217
06MT22/5	A10	629	1	12/31/1992	0650	-28.039	-46.129	2408
06MT22/5	A10	630	1	12/31/1992	1140	-28.088	-45.939	2596
06MT22/5	A10	631	1	12/31/1992	1712	-28.153	-45.674	2782
06MT22/5	A10	632	1	12/31/1992	2240	-28.224	-45.408	2965
06MT22/5	A10	1	2	01/01/1993	0726	-28.418	-44.768	3509
06MT22/5	A10	2	1	01/01/1993	1315	-28.615	-44.222	3694
06MT22/5	A10	3	1	01/01/1993	1902	-28.833	-43.585	3897
06MT22/5	A10	3	2	01/01/1993	2032	-28.833	-43.588	3882
06MT22/5	A10	4	2	01/02/1993	0315	-29.036	-42.910	4009
06MT22/5	A10	4	3	01/02/1993	0432	-29.031	-42.909	4009
06MT22/5	A10	5	2	01/02/1993	1015	-29.232	-42.329	4001
06MT22/5	A10	5	3	01/02/1993	1150	-29.228	-42.332	4000
06MT22/5	A10	6	2	01/02/1993	1720	-29.422	-41.737	3874
06MT22/5	A10	6	3	01/02/1993	1836	-29.413	-41.731	3847
06MT22/5	A10	7	2	01/03/1993	0013	-29.604	-41.158	3782
06MT22/5	A10	7	3	01/03/1993	0142	-29.596	-41.157	3792
06MT22/5	A10	8	2	01/03/1993	0723	-29.807	-40.587	3786
06MT22/5	A10	8	3	01/03/1993	0858	-29.802	-40.586	3781
06MT22/5	A10	9	2	01/03/1993	1450	-30.006	-39.998	3187
06MT22/5	A10	10	1	01/03/1993	1902	-30.003	-39.530	3972
06MT22/5	A10	10	2	01/03/1993	2041	-30.002	-39.528	4017
06MT22/5	A10	11	2	01/04/1993	0048	-29.998	-39.381	4902
06MT22/5	A10	11	3	01/04/1993	0245	-30.002	-39.377	4901
06MT22/5	A10	12	1	01/04/1993	0712	-30.004	-38.917	4282
06MT22/5	A10	12	2	01/04/1993	0913	-30.010	-38.917	4277
06MT22/5	A10	13	2	01/04/1993	1333	-30.004	-38.494	4226
06MT22/5	A10	13	3	01/04/1993	1503	-30.015	-38.490	4203
06MT22/5	A10	14	1	01/04/1993	1916	-30.003	-38.003	3856
06MT22/5	A10	14	2	01/04/1993	2037	-30.005	-38.008	3868
06MT22/5	A10	15	1	01/05/1993	0155	-29.999	-37.500	3204
06MT22/5	A10	16	1	01/05/1993	0550	-30.002	-37.169	2334
06MT22/5	A10	17	2	01/05/1993	0943	-30.005	-36.824	1573
06MT22/5	A10	18	1	01/05/1993	1300	-30.006	-36.500	1803
06MT22/5	A10	19	2	01/05/1993	1610	-30.000	-36.163	1268
06MT22/5	A10	20	1	01/05/1993	1926	-30.004	-35.836	2380
06MT22/5	A10	21	2	01/05/1993	2328	-30.004	-35.485	2325
06MT22/5	A10	22	1	01/06/1993	0300	-29.996	-35.167	2170
06MT22/5	A10	23	2	01/06/1993	0650	-30.009	-34.823	1637
06MT22/5	A10	24	1	01/06/1993	1012	-29.997	-34.501	1430
06MT22/5	A10	25	2	01/06/1993	1437	-29.989	-34.004	2046
06MT22/5	A10	26	1	01/06/1993	1908	-29.992	-33.509	3175
06MT22/5	A10	27	2	01/06/1993	2352	-29.993	-33.000	3528
06MT22/5	A10	27	3	01/07/1993	0115	-29.981	-33.009	3529
06MT22/5	A10	28	1	01/07/1993	0543	-29.997	-32.499	3747
06MT22/5	A10	28	2	01/07/1993	0712	-29.986	-32.503	3747

Table A.1 (continued)

06MT22/5	A10	29	2	01/07/1993	1205	-29.998	-32.000	3853
06MT22/5	A10	29	3	01/07/1993	1334	-29.986	-31.990	3867
06MT22/5	A10	30	1	01/07/1993	1751	-30.000	-31.504	3858
06MT22/5	A10	30	2	01/07/1993	1925	-30.001	-31.502	3814
06MT22/5	A10	31	2	01/07/1993	2347	-30.000	-30.996	4087
06MT22/5	A10	31	3	01/08/1993	0120	-30.013	-31.001	4090
06MT22/5	A10	32	1	01/08/1993	0530	-30.000	-30.501	3946
06MT22/5	A10	32	2	01/08/1993	0701	-30.002	-30.508	3937
06MT22/5	A10	33	2	01/08/1993	1130	-30.001	-29.991	3528
06MT22/5	A10	33	3	01/08/1993	1256	-29.997	-29.997	3485
06MT22/5	A10	34	1	01/08/1993	1756	-29.998	-29.509	2238
06MT22/5	A10	35	2	01/09/1993	0558	-29.997	-28.997	3202
06MT22/5	A10	36	1	01/09/1993	1025	-29.998	-28.420	3761
06MT22/5	A10	36	2	01/09/1993	1154	-29.999	-28.430	3763
06MT22/5	A10	37	2	01/09/1993	1800	-30.000	-27.568	4881
06MT22/5	A10	37	3	01/09/1993	1948	-30.007	-27.579	4883
06MT22/5	A10	38	1	01/10/1993	0200	-29.998	-26.719	5323
06MT22/5	A10	38	2	01/10/1993	0345	-29.997	-26.731	4920
06MT22/5	A10	39	2	01/10/1993	0955	-29.999	-25.862	4489
06MT22/5	A10	39	3	01/10/1993	1125	-30.001	-25.872	4474
06MT22/5	A10	40	1	01/10/1993	1732	-29.996	-25.018	5592
06MT22/5	A10	40	2	01/10/1993	1958	-30.014	-25.025	5673
06MT22/5	A10	40	3	01/10/1993	2238	-30.028	-25.038	5705
06MT22/5	A10	41	2	01/11/1993	0425	-30.000	-24.170	4805
06MT22/5	A10	41	3	01/11/1993	0600	-29.999	-24.180	4786
06MT22/5	A10	42	1	01/11/1993	1157	-30.000	-23.315	4627
06MT22/5	A10	42	2	01/11/1993	1327	-29.998	-23.317	4658
06MT22/5	A10	43	2	01/11/1993	1940	-29.998	-22.468	4692
06MT22/5	A10	43	3	01/11/1993	2120	-30.006	-22.479	4678
06MT22/5	A10	44	1	01/12/1993	0325	-29.998	-21.622	4874
06MT22/5	A10	44	2	01/12/1993	0505	-29.992	-21.621	4897
06MT22/5	A10	45	2	01/12/1993	1145	-30.002	-20.762	4770
06MT22/5	A10	45	3	01/12/1993	1330	-30.001	-20.767	4782
06MT22/5	A10	46	1	01/12/1993	1929	-29.999	-19.919	4848
06MT22/5	A10	46	2	01/12/1993	2114	-30.003	-19.933	4748
06MT22/5	A10	47	2	01/13/1993	0400	-29.988	-19.067	4064
06MT22/5	A10	47	3	01/13/1993	0550	-30.000	-19.055	4136
06MT22/5	A10	47	4	01/13/1993	0719	-29.979	-19.052	4250
06MT22/5	A10	48	1	01/13/1993	1240	-30.008	-18.385	4203
06MT22/5	A10	48	2	01/13/1993	1422	-30.020	-18.392	4273
06MT22/5	A10	49	2	01/14/1993	0436	-30.001	-17.698	3989
06MT22/5	A10	49	3	01/14/1993	0608	-30.003	-17.690	3949
06MT22/5	A10	50	1	01/14/1993	1101	-29.999	-17.013	3677
06MT22/5	A10	50	2	01/14/1993	1210	-30.014	-17.003	3566
06MT22/5	A10	51	2	01/14/1993	1703	-30.002	-16.333	3698
06MT22/5	A10	51	3	01/14/1993	1824	-30.007	-16.329	3680
06MT22/5	A10	52	1	01/15/1993	0002	-30.000	-15.663	3283
06MT22/5	A10	53	2	01/15/1993	0507	-30.003	-15.000	3834
06MT22/5	A10	53	3	01/15/1993	0652	-30.004	-15.003	3835
06MT22/5	A10	54	1	01/15/1993	1245	-30.006	-14.335	2753
06MT22/5	A10	55	2	01/15/1993	1810	-30.009	-13.661	2297
06MT22/5	A10	56	1	01/15/1993	2331	-30.013	-12.989	3096
06MT22/5	A10	57	2	01/16/1993	0444	-29.997	-12.333	3151
06MT22/5	A10	57	3	01/16/1993	0608	-29.990	-12.338	3131
06MT22/5	A10	58	1	01/16/1993	1115	-30.002	-11.676	3434
06MT22/5	A10	58	2	01/16/1993	1230	-30.018	-11.678	3323
06MT22/5	A10	59	2	01/16/1993	1726	-30.003	-10.994	3760
06MT22/5	A10	59	3	01/16/1993	1839	-30.010	-10.985	3632
06MT22/5	A10	60	1	01/16/1993	2327	-30.001	-10.330	3781
06MT22/5	A10	60	2	01/17/1993	0041	-30.005	-10.330	3791
06MT22/5	A10	61	2	01/17/1993	0537	-30.001	-9.661	3727
06MT22/5	A10	61	3	01/17/1993	0700	-30.006	-9.650	3730
06MT22/5	A10	62	1	01/17/1993	1153	-30.000	-8.998	3972
06MT22/5	A10	62	2	01/17/1993	1310	-30.024	-8.992	3984
06MT22/5	A10	63	2	01/17/1993	1845	-30.003	-8.162	3948
06MT22/5	A10	63	3	01/17/1993	2014	-30.015	-8.154	4025
06MT22/5	A10	64	1	01/18/1993	0210	-30.002	-7.332	4177

Table A.1 (continued)

06MT22/5	A10	64	2	01/18/1993	0334	-30.005	-7.323	4246
06MT22/5	A10	65	2	01/18/1993	0924	-30.007	-6.498	4556
06MT22/5	A10	65	3	01/18/1993	1055	-30.012	-6.506	4135
06MT22/5	A10	65	4	01/18/1993	1341	-30.029	-6.519	3635
06MT22/5	A10	66	1	01/18/1993	1914	-29.999	-5.668	4373
06MT22/5	A10	66	2	01/18/1993	2058	-29.998	-5.664	4390
06MT22/5	A10	67	2	01/19/1993	0241	-30.000	-4.827	4315
06MT22/5	A10	67	3	01/19/1993	0416	-30.006	-4.813	4444
06MT22/5	A10	68	1	01/19/1993	1002	-29.997	-4.001	3990
06MT22/5	A10	68	2	01/19/1993	1117	-29.999	-3.993	3977
06MT22/5	A10	69	2	01/19/1993	1719	-30.010	-3.171	4401
06MT22/5	A10	69	3	01/19/1993	1903	-30.012	-3.172	4369
06MT22/5	A10	70	2	01/20/1993	0055	-29.995	-2.323	4302
06MT22/5	A10	70	3	01/20/1993	0225	-29.993	-2.303	4323
06MT22/5	A10	71	2	01/20/1993	0822	-30.002	-1.498	4779
06MT22/5	A10	71	3	01/20/1993	1013	-29.984	-1.477	4806
06MT22/5	A10	72	1	01/20/1993	1622	-30.016	-0.730	4874
06MT22/5	A10	72	2	01/20/1993	1806	-30.015	-0.726	4875
06MT22/5	A10	73	2	01/20/1993	2340	-29.997	0.002	4008
06MT22/5	A10	73	3	01/21/1993	0056	-29.993	0.007	3962
06MT22/5	A10	74	1	01/21/1993	0617	-29.858	0.567	3329
06MT22/5	A10	75	2	01/21/1993	1058	-29.731	1.133	3739
06MT22/5	A10	75	3	01/21/1993	1215	-29.723	1.138	3735
06MT22/5	A10	76	1	01/21/1993	1641	-29.604	1.701	3665
06MT22/5	A10	76	2	01/21/1993	1802	-29.604	1.701	3647
06MT22/5	A10	77	2	01/21/1993	2255	-29.475	2.267	2712
06MT22/5	A10	78	1	01/22/1993	0332	-29.337	2.830	4251
06MT22/5	A10	78	2	01/22/1993	0510	-29.348	2.826	4241
06MT22/5	A10	79	2	01/22/1993	0938	-29.474	3.304	4777
06MT22/5	A10	79	3	01/22/1993	1116	-29.476	3.312	4770
06MT22/5	A10	80	1	01/22/1993	1538	-29.608	3.778	4944
06MT22/5	A10	80	2	01/22/1993	1723	-29.604	3.772	4933
06MT22/5	A10	81	2	01/23/1993	0730	-29.747	4.239	4971
06MT22/5	A10	81	3	01/23/1993	0902	-29.744	4.243	4985
06MT22/5	A10	82	1	01/23/1993	1524	-29.752	5.100	5253
06MT22/5	A10	82	2	01/23/1993	1719	-29.746	5.107	5262
06MT22/5	A10	83	2	01/23/1993	2349	-29.745	5.932	5198
06MT22/5	A10	83	3	01/24/1993	0138	-29.742	5.945	5192
06MT22/5	A10	84	1	01/24/1993	0817	-29.754	6.780	5207
06MT22/5	A10	84	2	01/24/1993	1007	-29.739	6.780	5209
06MT22/5	A10	85	2	01/24/1993	1622	-29.746	7.621	5200
06MT22/5	A10	85	3	01/24/1993	1816	-29.736	7.627	5203
06MT22/5	A10	86	1	01/25/1993	0110	-29.746	8.465	5091
06MT22/5	A10	86	2	01/25/1993	0247	-29.744	8.465	5086
06MT22/5	A10	87	2	01/25/1993	1036	-29.750	9.297	5032
06MT22/5	A10	87	3	01/25/1993	1235	-29.755	9.307	4986
06MT22/5	A10	88	1	01/25/1993	1925	-29.748	10.148	4900
06MT22/5	A10	88	2	01/25/1993	2141	-29.747	10.185	4853
06MT22/5	A10	89	2	01/26/1993	0527	-29.749	10.980	4296
06MT22/5	A10	89	3	01/26/1993	0700	-29.748	10.981	4271
06MT22/5	A10	90	2	01/26/1993	1419	-29.752	11.830	4002
06MT22/5	A10	90	3	01/26/1993	1615	-29.755	11.852	3994
06MT22/5	A10	91	2	01/26/1993	2025	-29.622	12.171	3818
06MT22/5	A10	91	3	01/26/1993	2150	-29.621	12.187	3812
06MT22/5	A10	92	2	01/27/1993	0248	-29.497	12.467	3657
06MT22/5	A10	92	3	01/27/1993	0436	-29.489	12.474	3652
06MT22/5	A10	93	2	01/27/1993	0944	-29.373	12.795	3358
06MT22/5	A10	93	3	01/27/1993	1104	-29.372	12.815	3341
06MT22/5	A10	94	2	01/27/1993	1620	-29.243	13.114	3098
06MT22/5	A10	95	2	01/27/1993	2135	-29.119	13.428	2640
06MT22/5	A10	96	2	01/28/1993	0248	-29.001	13.735	2136
06MT22/5	A10	97	2	01/28/1993	0645	-28.879	14.045	1490
06MT22/5	A10	98	2	01/28/1993	0955	-28.752	14.366	461
06MT22/5	A10	99	2	01/28/1993	1240	-28.622	14.687	160
06MT22/5	A10	100	2	01/28/1993	1457	-28.503	14.999	174

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