



Standard Guide for Analytical Data Interchange Protocol for Chromatographic Data¹

This standard is issued under the fixed designation E 1948; the number immediately following the designation indicates the year of original adoption or, in the case of revision, the year of last revision. A number in parentheses indicates the year of last reapproval. A superscript epsilon (ϵ) indicates an editorial change since the last revision or reapproval.

1. Scope

1.1 This guide covers the implementation of the Chromatographic Data Protocol in analytical software applications. Implementation of this protocol requires:

1.1.1 Specification E 1947, which contains the full set of data definitions. The chromatographic data protocol is not based upon any specific implementation; it is designed to be independent of any particular implementation, so that implementations can change as technology evolves. The protocol is implemented in stages, to speed its acceptance through actual use.

1.1.2 Specification E 1947 contains a full description of the contents of the data communications protocol, including the analytical information categories with data elements and their attributes for most aspects of chromatographic tests.

1.2 The Analytical Information Categories are a practical convenience for breaking down the standardization process into smaller, more manageable pieces. It is easier for developers to build consensus and produce working systems based on smaller information sets, without the burden and complexity of the hundreds of data elements contained in all the categories. The categories also assist vendors and end users in using the guide in their computing environments.

1.3 The NetCDF Data Interchange System is the container used to communicate data between applications in a way that is independent of both computer architectures and end-user applications. In essence, it is a special type of application designed for data interchange.

1.4 The Common Data Language (CDL) Template for Chromatography is a language specification of the chromatography dataset being interchanged. With the use of the NetCDF utilities, this human-readable template can be used to generate an equivalent binary file and the software subroutine calls needed for input and output of data in analytical applications.

2. Referenced Documents

2.1 *ASTM Standards:*²

E 1947 Specification for Analytical Data Interchange Protocol for Chromatographic Data

2.2 *Other Standard:*

NetCDF User's Guide³

3. Features of the Chromatographic Data Interchange Implementation Guide

3.1 The chromatographic data protocol consists of: (1) the data contents (what is being transferred), and (2) the data container (how it is being transferred).

3.2 *Analytical Information Categories*—To make the standardization process more manageable, five Analytical Information Categories were defined:

1. Raw Data
2. Final Results
3. Full Data Processing Method
4. Full Chemical Method
5. Good Laboratory Practices Information

3.2.1 This guide covers implementation of Raw Data (Category 1) and Final Results (Category 2). The development committee has completed implementation and testing of information in these categories.

3.3 *Chromatography Data Elements*—Each data element specifies a piece of information that is to be stored or transferred to another system. A data element has a name, a definition, and may have some default attributes, such as its datatype, the analytical information category to which it belongs, and whether it is mandatory or recommended. A data element may also reference other data elements.

3.4 *NetCDF Data Interchange System*—NetCDF is a data interchange system designed specifically for the requirements of scientific data. It was developed to meet the requirements of

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² For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at service@astm.org. For *Annual Book of ASTM Standards* volume information, refer to the standard's Document Summary page on the ASTM website.

³ Available for Russell K. Rew, Unidata Program Center, University Corporation for Atmospheric Research, P. O. Box 3000, Boulder, CO 80307-3000.

high-performance, large dataset interchange over networks, in a machine- and network-independent way. NetCDF is implemented in the portable C programming language, and is made to be available on all popular scientific computers and operating systems. It achieves platform-independence through the use of the XDR protocol for byte-stream encoding. XDR (eXternal Data Representation) was developed by Sun Microsystems as part of the Network File System (NFS) and is now in the public domain. NetCDF has a particular advantage over many other approaches, such as ASCII flat files or structured ASCII files, because it uses the abstraction of a data access interface or abstract Application Programming Interface (API). This API lets the application developer deal with data more in terms of logical entities, instead of having to think about specific details of encoding and decoding data. This guide specifies the use of version 2.0 or later, and the exchange software should determine only that the version of NetCDF is version 2.0 or greater.

3.5 *Common Data Language*—A NetCDF data file has both binary and human-readable representations. The human-readable form is an ASCII-encoded file written in the NetCDF system’s Common Data Language (CDL). CDL is a data description language, which is a simplified type of programming language without any action statements. The main purpose of a data description language is the specification of data structures. CDL is a generic data description language for scientific data; it is not application-specific. The basic construct in CDL is that of a multidimensional array, to which many types of metadata can be attached. Metadata is additional data about the data, that is, attributes about the data that help turn it into useful information.

3.5.1 For the protocol, a template for chromatography written in CDL is supplied to developers. The CDL template provides an easily comprehended text version of the structure and contents of a binary NetCDF file. Developers use this template with certain NetCDF utilities to generate the software code (subroutine calls) needed to use the NetCDF toolkit. The generated code can then be cut and pasted in the end-user application.

3.5.2 For a full description of the Common Data Language, see the *NetCDF User’s Guide*.

3.6 *NetCDF Utilities*— The NetCDF data interchange system has two very powerful utilities called “ncgen” and “ncdump” that eliminate much work for developers.

3.6.1 After the template for a particular analytical technique dataset (in this case chromatography) is written in CDL, it is fed into ncgen. There are several options with ncgen, the first of which creates the binary file and populates it with fields specified by the CDL template. Other ncgen options generate code for the C language subroutines required for an application to read and write data in the NetCDF file, and the FORTRAN wrapper code needed for FORTRAN programs. The ncdump utility reads the binary files generated by ncgen (which were then populated by the application using the NetCDF libraries), and then generates their ASCII-encoded, human-readable representation. ncdump is routinely used to check the contents of NetCDF files, and debug applications that use NetCDF. It can

also be used to export data to reporting programs. Fig. 1 illustrates the usage of the NetCDF utilities.

3.6.2 Developers typically start with the CDL source code description of the data file they wish to interchange. Then the ncgen utility with the -n option generates the NetCDF binary file. Developers then use ncgen with the -c option to generate the C source code needed for the subroutines to read and write this particular dataset (specified by the CDL template.) These read/write subroutines make up the NetCDF Application Programming Interface (API). These routines are simply cut and pasted into the target application(s). The programmer then uses the API to read or write data in NetCDF datasets.

3.6.3 If FORTRAN is the programming language being used, the ncgen utility with the -f option can generate the FORTRAN source code wrappers needed to embed the NetCDF library code into FORTRAN applications.

3.6.4 Once the application populates the data file with its data, that file can be transferred to another system. If a developer or end user needs a human-readable version of the binary dataset, the binary file can be fed into the ncdump utility, which regenerates the original CDL source code. The ncdump utility has an option switch to regenerate the CDL code with or without the actual experimental data. Some datasets can be enormous and there may be no reason to view the entire dataset. In this way, data are easily encoded into NetCDF files and the required routines to read and write those data are incorporated into applications.

3.6.5 There is another way to build NetCDF data files. Using the NetCDF Application Programming Interface, it is possible to build NetCDF binary files one data element at a time. However, that method is generally more time-consuming, and it is not recommended except for those situations where applications can fully automate the process.

4. Chromatographic Data Protocol Distribution Kit

4.1 Information on how to obtain the kit will be posted on the ASTM web site (www.astm.org) under Committee E-49.

4.2 The Analytical Data Interchange Protocol for Chromatographic Data Distribution Kit contains:

4.2.1 *Software Disks*— NetCDF distribution kit from Uni-data (with the modified makefile needed to make the kit compile out of the box), and the Chromatographic Data Protocol CDL Template.

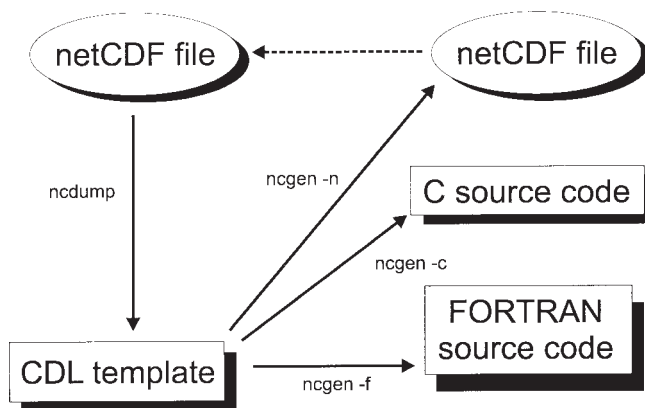


FIG. 1 Usage of NetCDF Utilities

4.2.2 *NetCDF User's Guide*—supplied by Unidata Program Center.

4.2.3 *Specification E 1947*.

4.2.4 *Guide E 1948*.

4.2.5 *Registration Card*, used for tracking and user updates.

5. Hardware and Software

5.1 This section describes the hardware and software configurations used for testing. In general, the NetCDF system puts very few requirements on the hardware, because most routines are left on disk. Only routines being used at any particular time are kept in memory. Any limitations found were typically those not imposed by NetCDF but ones imposed by the operating system or environment.

5.1.1 *Hardware (Personal Computers)*—The personal computer system hardware used for testing was:

5.1.1.1 Intel 80286 processor,

5.1.1.2 640K minimum,

5.1.1.3 monochrome, EGA, VGA graphics,

5.1.1.4 20 megabyte minimum, 80 megabyte hard-disk is typical, and

5.1.1.5 a mouse (optional).

5.1.1.6 NetCDF works well on AT-class machines and higher. NetCDF does not have the items in 5.1.1.1-5.1.1.5 as requirements. These are just the minimum, base-level systems that were used.

5.1.2 *Software*—NetCDF runs on MS-DOS, OS/2, Macintosh, Windows 95, and Windows NT operating systems for personal computers. NetCDF was originally ported from UNIX to DOS running on an IBM-PS/2 Model 80. It was recently ported to the Macintosh OS. NetCDF is written in the C programming language, and there are FORTRAN jackets available for applications that want to use FORTRAN calls. The personal computer software employed for testing and developing NetCDF applications were:

5.1.2.1 Microsoft DOS V3.3 or above,

5.1.2.2 Microsoft C Compiler V6.0,

5.1.2.3 Microsoft Windows V3.0,

5.1.2.4 Microsoft Windows SDK, and

5.1.2.5 NetCDF Version 2.0.1.

5.1.3 *Workstations and Servers*—NetCDF runs easily on UNIX workstations such as Sun 3, Sun 4, VAXstations, DECstation 3100, VAXstation II running ULTRIX or VMS, and IBM RS/6000. There are no particular hardware requirements for workstation class machines, since all workstations have the minimum hardware outlined for personal computers in 5.1.1.

6. Significance and Use

6.1 *General Coding Guidelines*—The NetCDF libraries are supplied to developers as source code. End users receive the libraries in compiled binary form as part of a vendor's application.

6.1.1 Some vendors found that compilation using the huge memory model under Microsoft Windows on MS-DOS was needed because of an array pointer passing its boundary. Many vendors chose to write a conversion program as a stand-alone DOS application, whereby the conversion program used only

text-mode screen I/O. Some vendors are now using it in their MS-Windows applications.

6.1.2 Developers setting out to write a program to convert their data files to the Chromatographic Data Protocol should use the NetCDF utilities *ncgen* and *ncdump*. Applying the *ncgen* utility to the CHROMSTD.CDL template will generate the skeletal code needed to create the NetCDF file. The programmer can then modify the code to create a program that reads the netDCF file from another vendor. After developers create the NetCDF file they should use the *ncdump* program to generate the ASCII representation of the data files, and examine it to ensure the data is being correctly put into the file.

6.2 *Make Files for NetCDF Libraries and Utilities*—In general the compilation is straightforward. The make files were modified after they were received from the Unidata Corporation, because they did not compile the first time on PCs. The changes needed to get the Unidata distribution to run on DOS are (1) rename the file MAKEFILE to UNIX.MK, and (2) rename MSOFT.MK to MAKEFILE, and then run NMAKE. The default switches in the Unidata distribution use the switches for the floating point coprocessor and Microsoft Windows options.

6.2.1 The protocol contain some complete makefile examples for Microsoft C V6.0 running on DOS. The Microsoft C V6.0 compiler manual should be consulted for the exact meaning of the compiler and linker options.

6.2.2 The VMS and SunOS compilation instructions are in directories for those operating systems.

6.3 *NetCDF Library Build Order*—The NetCDF libraries must be built in a specific order. The correct order to build the NetCDF directories is:

```
UTIL
XDR
SRC
NCDUMP
NCGEN
NCTEST
```

6.3.1 The UTIL and XDR makefiles work as distributed using NMAKE with Microsoft C V6.0.

7. CDL Template Structure

7.1 A NetCDF template is built from CDL statements and structured into three sections: (1) dimension declarations, (2) variable declarations, and (3) the data section.

7.2 A few points of clarification about the CDL language are given here to facilitate understanding of the CDL Generic Chromatography Template given in Fig. 2. For more in-depth information on CDL, please consult the *NetCDF User's Guide*.

7.2.1 A NetCDF template starts with the word "NetCDF" followed by the dataset name.

7.2.2 CDL comments are indicated by two forward slash characters (//).

7.2.3 Section indicators (dimensions:, variables:, and data:) end with a colon character (:). these are the only tokens that end with a colon character.

7.2.4 Statements within sections end with the semicolon character (;).

7.2.5 Variable names beginning with numbers must be preceded by an underline character (_). Otherwise the *ncgen* parser will flag an error.

```

netCDF AIA_chromatography_template

{

// File: chromstd12.cdl.      Date: 1-26-92
// CDL code for AIA Chrom netCDF file.  AIA Specification Version 1.0.
// AIA Specification Version 1.0 only includes Analytical Information
// Categories 1 & 2.
// It requires Version 2.0 of netCDF

// Copyright 1991 - The Analytical Instrument Association

dimensions:

// * The following dimensions predefine some useful character array
// * default string lengths.  The actual size of a character dimension is
// * arbitrary.  These were picked by considering computer string length
// * storage efficiency.  These dimension values should
// * not be changed by your data source program.

    _2_byte_string = 2;      // 2 character string length
    _4_byte_string = 4;      // 4 character string length
    _8_byte_string = 8;      // 8 character string length
    _16_byte_string = 16;    // 16 character string length
    _32_byte_string = 32;    // 32 character string length
    _64_byte_string = 64;    // 64 character string length
    _128_byte_string = 128;  // 128 character string length
    _255_byte_string = 255;  // 255 character string length

// * The following variables need to be set by your program.  The values below
// * are given just as examples.  The actual values will be
// * set by the data source program after it
// * does the peak integrations and computes how many peaks and components
// * there are.  This should be done before the netCDF file is written out
// * from your application.

    point_number = 7;        // point number dimension for raw data    M1
    peak_number = 1;         // number of peaks                M2
    error_number = 1;

variables:

// Administrative Information Class - Category 1,2 Data Elements
    :dataset_completeness = "C1+C2"; // M12345
// Categories 1&2, raw & result data
    :aia_template_revision = "1.0"; // M12345
    :netcdf_revision = "2.0"; // M12345

```

FIG. 2 Template

7.2.5.1 Underline characters were chosen for this protocol over hyphen characters, because some compilers may interpret hyphens as subtraction operators. The feature of CDL that allows implicit numerical datatyping of attributes in not being used in the first version of the template. Instead, all floating

point attributes are being handled as strings. This forces programmers to explicitly type variables, thereby encouraging more deliberate programming styles. For example:

```
:aia_template_revision = "0.8"; //M12345
```

```

:languages                                = "English-only for now";
:administrative_comments                   = "none for now!";
:dataset_origin                            = "AIA member company name"; // M5
:dataset_owner                             = "AIA Companies";
:dataset_date_time_stamp                   = "19910901123030-0500";
:injection_date_time_stamp                 = "19910901123030-0500";      // M12345
:experiment_title                          = "Working Demonstrations";
:operator_name                             = "Joe Scibler";           // M5
:separation_experiment_type                 = "liquid chromatography";
:company_method_name                       = "sandope analysis XYZ";
:company_method_id                         = "SAXYZ";
:pre_experiment_program_name               = "setup";
:post_experiment_program_name               = "response calibration";
:source_file_reference                     = "IODINE::dka100:[aia]test.cdl";
char error_log(error_number, _64_byte_string);

// SAMPLE-DESCRIPTION Information Class - Category 1 Data Elements

:sample_id_comments                        = "none";
:sample_id                                 = "JOU812";
:sample_name                               = "test sample";
:sample_type                               = "control";
:sample_injection_volume                   = "2.0";
:sample_amount                             = "2.0";

// DETECTION-METHOD Information Class - Category 1 Data Elements

:detection_method_table_name               = "test 1";
:detector_method_comments                  = "An optical detector";
:detection_method_name                     = " ";
:detector_name                             = "variable wavelength detector";

// the following information is used for proper raw data scaling

float detector_maximum_value;              // M1
float detector_minimum_value;              // M1
:detector_unit                             = "Absorbance Unit";
// M1, e.g., A.U.

// RAW-DATA Information Class - Category 1 Data Elements

:raw_data_table_name                       = "test raw data set";
:retention_unit                            = "time in seconds";
// M12
float actual_run_time_length;              // M12, in retention_unit
float actual_sampling_interval;            // M12, in retention_unit
float actual_delay_time;                   // M12, in retention_unit

float ordinate_values (point_number);      // M1
ordinate_values:uniform_sampling_flag = "Y"; // M1
ordinate_values:autosampler_position = "1.01";
// real_time autosampler position
float raw_data_retention (point_number);
// M1 only if uniform_sampling_flag = "N"

```

FIG. 2 Template (continued)

```
// PEAK-PROCESSING-RESULTS Information Class - Category 2 Elements

:peak_processing_results_table_name      = "Impl. Guide Demo Code";
:peak_processing_results_comments       = "level 1 calibration
                                         results";

:peak_processing_method_name            = "test processing method";
:peak_processing_date_time_stamp        = "19910901123030-0500";
float peak_retention_time(peak_number); // M2
char peak_name(peak_number, _32_byte_string);
float peak_amount(peak_number);        // M3
:peak_amount_unit                       = "grams";
float peak_start_time(peak_number);     // unit = retention_unit
float peak_end_time(peak_number);       // unit = retention_unit
float peak_width(peak_number);
float peak_area(peak_number);           // M2
float peak_area_percent(peak_number);
float peak_height(peak_number);        // M2
float peak_height_percent(peak_number);
float baseline_start_time(peak_number);
float baseline_start_value(peak_number);
float baseline_stop_time(peak_number);
float baseline_stop_value(peak_number);
char peak_start_detection_code(peak_number, _2_byte_string);
                                         //baseline type
char peak_stop_detection_code(peak_number, _2_byte_string);
                                         //baseline type

float retention_index(peak_number);
float migration_time(peak_number);
float peak_asymmetry(peak_number);
float peak_efficiency(peak_number);
float mass_on_column(peak_number);
short manually_reintegrated_peaks(peak_number); // 0=no, 1=yes

data:

// * Data will be inserted here by your program.
// * There will be a great deal of data for our netCDF!
// * The following is an example for part of the data.

ordinate_values = 998760, 997650, 1002340, 1102340, 1203450, 1145670, 1000000;
peak_retention_time = 105;
peak_name = "Peak A";
}
```

FIG. 2 Template (continued)

```
:netcdf_revision = "2.0.1"; //M12345
```

Consult the *NetCDF User's Guide* for more complete information on CDL syntax and usage.

7.2.6 Underline characters only can be used as separators between words within variable names, like:

```
aia-template-revision, or aia_template_revision.
```

7.2.7 Numerical data types for single values can be declared implicitly by putting numbers on the right side of an assignment statement, like:

```
peak_number=2; //number of peaks
```

These numerical datatypes can be floating point or integer values, and can be implicitly datatyped as such.

```
:floating_point_attribute = 1.11; //M12345
```

7.2.8 Numerical data types can be declared explicitly by preceding the variable definition by its data type. Datatype assignments can be for either single value variable definitions or for array variable definitions, for example:

```
float detector_maximum_value;
float ordinate_values(point_number);
```

There is also a way to explicitly declare datatypes on the right

side of an assignment operator. Please consult the NetCDF User's Guide for details.

7.2.9 Metadata are associated with a particular variable by attaching it to that variable with a colon character, for example:

```
ordinate_values:uniform_sampling_flag= "Y";
ordinate_values:autosampler_position= "1.01";
```

7.2.10 Global metadata can be declared simply by not attaching it to any variable, for example:

```
:aia_template_revision= "0.8"; //M12345
```

7.2.11 String attributes can be as long as needed, and are declared by enclosing the strings in quotation marks, for example:

```
:retention_unit= "time in seconds";
```

7.3 *Notes About the Chromatographic Data Protocol Usage of CDL*—Some Mandatory indicator codes (M-codes) for data elements such as M1234, M1, etc., are given in some comment fields of the Protocol template. These are *not* part of CDL syntax. These refer to whether a given data element is mandatory for particular information categories, for example, M1234 specifies that the data element is mandatory for Categories 1, 2, 3, and 4. These M-codes are also given in the Specification E 1947.

8. The Analytical Data Interchange Protocol for Chromatographic Data Template

8.1 The template in Fig. 2 is mostly for Analytical Information Categories 1 and 2 only. Please refer to Specification E 1947 for a complete description of each data element and its allowed value.

9. Other Usage Tips

9.1 *Filename Extensions*—The recommended filename extension is “.cdf,” so that the full name for a NetCDF file would

be “filename.cdf.” This is used so that parsers used to select files can parse filenames based on the “.cdf” extension rather than some other non-standard file extension.

9.2 *Handling of Missing Variables*—The absence of a variable implies that it is not in the file. For example, if a get operation returns an error condition, this implies that the variable does not exist in that file.

9.3 *Performance Tip For Data Value Access*—The point_number dimension was originally declared as “unlimited,” however, this was changed to a finite value because this change allows getting and putting of an entire array at once. This change is minor and will not affect programs, however, it greatly improves performance. Using point_num as unlimited restricts get/put operations to single values at a time, that is, they are treated as records.

9.4 *Getting Valid Date Time Stamps*—In order to get the correct date-time-stamp values in datasets originating from DOS and OS/2 systems, the environment variable for time zone shall be set correctly. The recommended procedure is to set the offset from Greenwich Mean Time *at product installation time*. Some examples of how to set the time-zone environment variable are as follows:

9.4.1 The command “DOS-PROMPT>tz pst 8 pdt” sets the time-zone variable to have a GMT offset of Pacific Standard Time (pst), with a value of 8 h offset from GMT, at Pacific Daylight Time (pdt).

9.4.2 The command “DOS-PROMPT>tz est 5 edt” sets the time-zone variable to have a GMT offset of Eastern Standard Time (est), with a value of 5 h offset for GMT, at Eastern Daylight Time (edt).

10. Keywords

10.1 analytical; chromatographic data; data; data communications; good laboratory practices; method; netCDF; raw data; results

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