## NCO User's Guide

A suite of netCDF operators<br>Edition 2.8.1, for NCO Version 2.8.1<br>August 2003

by Charlie Zender
Department of Earth System Science
University of California at Irvine

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This is the first edition of the NCO User's Guide, and is consistent with version 2 of 'texinfo.tex'.

Published by Charlie Zender
Department of Earth System Science
University of California at Irvine
Irvine, CA 92697-3100 USA

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The original author of this software, Charlie Zender, wants to improve it with the help of your suggestions, improvements, bug-reports, and patches.
Charlie Zender <zender at uci dot edu>
Department of Earth System Science
University of California at Irvine
Irvine, CA 92697-3100

## Foreword

NCO is the result of software needs that arose while I worked on projects funded by NCAR, NASA, and ARM. Thinking they might prove useful as tools or templates to others, it is my pleasure to provide them freely to the scientific community. Many users (most of whom I have never met) have encouraged the development of NCO. Thanks espcially to Jan Polcher, Keith Lindsay, Arlindo da Silva, John Sheldon, and William Weibel for stimulating suggestions and correspondence. Your encouragment motivated me to complete the NCO User's Guide. So if you like NCO, send me a note! I should mention that NCO is not connected to or officially endorsed by Unidata, ACD, ASP, CGD, or Nike.

Charlie Zender
May 1997
Boulder, Colorado

Major feature improvements entitle me to write another Foreword. In the last five years a lot of work has been done refining NCO. NCO is now an honest-to-goodness open source project. It appears to be much healthier for it. The list of illustrious institutions which do not endorse NCO continues to grow, and now includes UCI.

Charlie Zender
October 2000
Irvine, California

The most remarkable advances in NCO capabilities in the last few years are due to contributions from the Open Source community. Especially noteworthy are the contributions of Henry Butowsky and Rorik Peterson.

Charlie Zender
January 2003
Irvine, California

## Summary

This manual describes NCO, which stands for netCDF Operators. NCO is a suite of programs known as operators. Each operator is a standalone, command line program which is executed at the UNIX (or NT) shell-level like, e.g., ls or mkdir. The operators take netCDF file(s) (or HDF4 files) as input, perform an operation (e.g., averaging or hyperslabbing), and produce a netCDF file as output. The operators are primarily designed to aid manipulation and analysis of data. The examples in this documentation are typical applications of the operators for processing climate model output. This reflects their origin, but the operators are as general as netCDF itself.

## 1 Introduction

### 1.1 Availability

The complete NCO source distribution is currently distributed as a compressed tarfile from http://sourceforge.net/projects/nco and from http://dust.ess.uci.edu/nco/nco.tar.gz. The compressed tarfile must be uncompressed and untarred before building NCO. Uncompress the file with 'gunzip nco.tar.gz'. Extract the source files from the resulting tarfile with 'tar -xvf nco.tar'. GNU tar lets you perform both operations in one step with 'tar -xvzf nco.tar.gz'.

The documentation for nco is called the NCO User's Guide. The User's Guide is available in Postscript, html, Dvi, TEXinfo, and Info formats. These formats are included in the source distribution in the files 'nco.ps', 'nco.html', 'nco.dvi', 'nco.texi', and 'nco.info*', respectively. All the documentation descends from a single source file, 'nco.texi'1. Hence the documentation in every format is very similar. However, some of the complex mathematical expressions needed to describe ncwa can only be displayed in the Postscript and DVI formats.

If you want to quickly see what the latest improvements in NCO are (without downloading the entire source distribution), visit the NCO homepage at http://nco.sourceforge.net. The HTML version of the User's Guide is also available online through the World Wide Web at URL http://nco.sourceforge.net/nco.html.
To build and use nco, you must have netCDF installed. The netCDF homepage is http://www.unidata.ucar.edu/packages/netcdf.

New NCO releases are announced on the netCDF list and on the nco-announce mailing list http://lists.sourceforge.net/mailman/listinfo/nco-announce.

### 1.2 Operating systems compatible with NCO

NCO has been successfully ported and tested and is known to work on the following 32 and 64 bit platforms: ibm aix 4.x, 5.x, FreeBSD 4.x, Gnu/Linux 2.x, LinuxPPC, LinuxAlpha, LinuxSparc64, SGI IRIX 5.x and 6.x, MacOS X 10.x, NEC Super-UX 10.x, DEC osf, Sun SunOS 4.1.x, Solaris 2.x, CRAY Unicos 8.x-10.x, all MS Windows. If you port the code to a new operating system, please send me a note and any patches you required.

The major prerequisite for installing NCO on a particular platform is the successful, prior installation of the netCDF library (and, as of 2003, the UDUnits library). Unidata has shown a commitment to maintaining netCDF and UDUnits on all popular Unix platforms, and is moving towards full support for the Microsoft Windows operating system (OS). Given this, the only difficulty in implementing NCO on a particular platform

[^0]is standardization of various C and Fortran interface and system calls. nCo code is tested for ANSI compliance by compiling with C compilers including those from GNU ('gcc -std=c99 -pedantic -D_BSD_SOURCE' -Wall) ${ }^{2}$, Comeau Computing ('como --c99'), Cray ('cc'), HP/Compaq/DEC ('cc'), IBM ('xlc -c -qlanglvl=extended'), Intel ('icc'), NEC ('cc'), SGI ('cc-LANG:std'), and Sun ('cc'). NCO (all commands and the libnco library) and the C++ interface to netCDF (called libnco_c++) comply with the ISO C++ standards as implemented by Comeau Computing ('como'), Cray ('CC'), GNU ('g++-Wall'), HP/Compaq/DEC ('cxx'), IBM ('xlC'), Intel ('icc'), NEC ('c++'), SGI ('CC -LANG:std'), and Sun ('CC -LANG:std'). See 'nco/bld/Makefile' and 'nco/src/nco_c++/Makefile.old' for more details.

Until recently (and not even yet), ANsi-compliant has meant compliance with the 1989 ISO C-standard, usually called C89 (with minor revisions made in 1994 and 1995). C89 does not allow variable-size arrays nor use of the ' $\%$ z' format for printf. These are nice features of the 1999 ISO C-standard called C99. NCO is C99-compliant where possible and C89-compliant where necessary. Certain branches in the code are required to satisfy the native SGI and SunOS C compilers, which are strictly ANSI C89 compliant, and cannot benefit from C99 features. However, C99 features are fully supported by the GNu, unicos, Solaris, and AIX compilers.

The most time-intensive portion of NCO execution is spent in arithmetic operations, e.g., multiplication, averaging, subtraction. Until August, 1999, these operations were performed in Fortran by default. This was a design decision made in late 1994 based on the speed of Fortran-based object code vs. C-based object code. Since 1994 native C compilers have improved their vectorization capabilities and it has become advantageous to replace all Fortran subroutines with C subroutines. Furthermore, this greatly simplifies the task of compiling on nominally unsupported platforms. As of August 1999, NCO is built entirely in C by default. This allows NCO to compile on any machine with an Ansi C compiler. Furthermore, NCO automatically takes advantage of extensions to ANSI C when compiled with the GNU compiler collection, GCC.

As of July 2000 and NCO version 1.2, NCO no longer supports performing arithmetic operations in Fortran. We decided to sacrifice executable speed for code maintainability Since no objective statistics were ever performed to quantify the difference in speed between the Fortran and C code, the performance penalty incurred by this decision is unknown. Supporting Fortran involves maintaining two sets of routines for every arithmetic operation. The USE_FORTRAN_ARITHMETIC flag is still retained in the 'Makefile'. The file containing the Fortran code, 'nco_fortran.F', has been deprecated but can be resurrected if a volunteer comes forward. If you would like to volunteer to maintain 'nco_fortran. F' please contact me.

### 1.2.1 Compiling NCO for Microsoft Windows OS

NCO has been successfully ported and tested on the Microsoft Windows (98/NT) operating systems. The switches necessary to accomplish this are included in the standard distribution

[^1]of NCO. Using the freely available Cygwin (formerly gnu-win32) development environment ${ }^{3}$, the compilation process is very similar to installing NCO on a uniX system. The preprocessor token PVM_ARCH should be set to WIN32. Note that defining WIN32 has the side effect of disabling Internet features of NCO (see below). Unless you have a Fortran compiler (like g77 or f90) available, no other tokens are required. Users with fast Fortran compilers may wish to activate the Fortran arithmetic routines. To do this, define the preprocessor token USE_FORTRAN_ARITHMETIC in the makefile which comes with NCO, 'Makefile', or in the compilation shell.

The least portable section of the code is the use of standard Unix and Internet protocols (e.g., ftp, rcp, scp, getuid, gethostname, and header files '<arpa/nameser.h>' and '<resolv.h>'). Fortunately, these Unixy calls are only invoked by the single NCO subroutine which is responsible for retrieving files stored on remote systems (see Section 3.3 [Remote storage], page 20). In order to support NCO on the Microsoft Windows platforms, this single feature was disabled (on Windows os only). This was required by Cygwin 18.x-newer versions of Cygwin may support these protocols (let me know if this is the case). The nco operators should behave identically on Windows and UNIX platforms in all other respects.

### 1.3 Libraries

Like all executables, the NCO operators can be built using dynamic linking. This reduces the size of the executable and can result in significant performance enhancements on multiuser systems. Unfortunately, if your library search path (usually the LD_LIBRARY_PATH environment variable) is not set correctly, or if the system libraries have been moved, renamed, or deleted since NCO was installed, it is possible an NCO operator will fail with a message that it cannot find a dynamically loaded (aka shared object or '. so') library. This usually produces a distinctive error message, such as 'ld.so.1:/usr/local/bin/ncea: fatal: libsunmath.so.1: can't open file: errno=2'. If you received an error message like this, ask your system administrator to diagnose whether the library is truly missing ${ }^{4}$, or whether you simply need to alter your library search path. As a final remedy, you can reinstall NCO with all operators statically linked.

## 1.4 netCDF 2.x vs. 3.x

netCDF version 2.x was released in 1993. NCO (specifically ncks) began with netCDF 2.x in 1994. netCDF 3.0 was released in 1996, and we were eager to reap the performance advantages of the newer netCDF implementation. One netCDF 3.x interface call (nc_inq_ libvers) was added to NCO in January, 1998, to aid in maintainance and debugging. In March, 2001, the final conversion of NCO to netCDF 3.x was completed (coincidentally on

[^2]the same day netCDF 3.5 was released). NCO versions 2.0 and higher are built with the -DNO_NETCDF_2 flag to ensure no netCDF 2.x interface calls are used.

However, the ability to compile NCO with only netCDF 2.x calls is worth maintaining because HDF version $4^{5}$ (available from HDF) supports only the netCDF 2.x library calls (see http://hdf.ncsa.uiuc.edu/UG41r3_html/SDS_SD.fm12.html\#47784). Note that there are multiple versions of HDF. Currently HDF version 4.x supports netCDF 2.x and thus NCO version 1.2.x. If NCO version 1.2.x (or earlier) is built with only netCDF 2.x calls then all NCO operators should work with HDF4 files as well as netCDF files ${ }^{6}$. The preprocessor token NETCDF2_ONLY exists in NCO version 1.2.x to eliminate all netCDF 3.x calls. Only versions of NCO numbered 1.2.x and earlier have this capability. The NCO 1.2.x branch will be maintained with bugfixes only (no new features) until HDF begins to fully support the netCDF 3.x interface (which is employed by NCO 2.x). If, at compilation time, NETCDF2_ ONLY is defined, then NCO version 1.2.x will not use any netCDF 3.x calls and, if linked properly, the resulting NCO operators will work with HDF4 files. The 'Makefile' supplied with NCO 1.2.x has been written to simplify building in this HDF capability. When NCO is built with make HDF4=Y, the 'Makefile' will set all required preprocessor flags and library links to build with the HDF4 libraries (which are assumed to reside under /usr/local/hdf4, edit the 'Makefile' to suit your installation).

HDF version 5.x became available in 1999, but did not support netCDF (or, for that matter, Fortran) as of December 1999. By early 2001, hDF version 5.x did support Fortran90. However, support for netCDF 3.x in HDF 5.x is incomplete. Much of the HDF5-netCDF3 interface is complete, however, and it may be separately downloaded from the HDF5-netCDF website. Now that NCO uses only netCDF 3.x system calls we are eager for HDF5 to complete their netCDF 3.x support.

### 1.5 Help and Bug reports

We generally receive three categories of mail from user's: requests for help, bug reports, and requests for new features. Notes saying the equivalent of "Hey, NCO continues to work great and it saves me more time everyday than it took to write this note" are a distant fourth. There is a different protocol for each type of request. Our request is that you communicate with the project via NCO Project Forums. Before posting to the NCO forums described below, you might first register your name and email address with SourceForge.org or else all of your postings will be attributed to "nobody". Once registered you may choose to "monitor" any forum and to receive (or not) email when there are any postings.

If you would like NCO to include a new feature, first check to see if that feature is already on the TODO list. If it is, please consider implementing that feature yourself and sending us the patch! If the feature is not yet on the list then send a note to the NCO Discussion forum.

[^3]Please read the manual before reporting a bug or posting a request for help. Sending questions whose answers are not in the manual is the best way to motivate us to write more documentation. We would also like to accentuate the contrapositive of this statement. If you think you have found a real bug the most helpful thing you can do is simplify the problem to a manageable size and report it. The first thing to do is to make sure you are running the latest publicly released version of NCO.

Once you have read the manual, if you are still unable to get NCO to perform a documented function, write help request. Follow the same procedure as described below for reporting bugs (after all, it might be a bug). That is, describe what you are trying to do, and include the complete commands (with '-D 5'), error messages, and version of NCO. Post your help request to the NCO Help forum.

If you think you are using the right command, but NCO is misbehaving, then you might have found a bug. A core dump, sementation violation, or incorrect numerical answers is always considered a high priority bug. How do you simplify a problem that may be revealing a bug? Cut out extraneous variables, dimensions, and metadata from the offending files and re-run the command until it no longer breaks. Then back up one step and report the problem. Usually the file(s) will be very small, i.e., one variable with one or two small dimensions ought to suffice. Include in the report your run-time environment, the exact error messages (and run the operator with '-D 5' to increase the verbosity of the debugging output), and a copy, or the publically accessible location, of the file(s). Post the bug report to the NCO Project buglist.

## 2 Operator Strategies

### 2.1 NCO operator philosophy

The main design goal has been to produce operators that can be invoked from the command line to perform useful operations on netCDF files. Many scientists work with models and observations which produce too much data to analyze in tabular format. Thus, it is often natural to reduce and massage this raw or primary level data into summary, or second level data, e.g., temporal or spatial averages. These second level data may become the inputs to graphical and statistical packages, and are often more suitable for archival and dissemination to the scientific community. NCO performs a suite of operations useful in manipulating data from the primary to the second level state. Higher level interpretive languages (e.g., IDL, Yorick, Matlab, NCL, Perl, Python), and lower level compiled languages (e.g., C, Fortran) can always perform any task performed by NCO, but often with more overhead. NCO, on the other hand, is limited to a much smaller set of arithmetic and metadata operations than these full blown languages.

Another goal has been to implement enough command line switches so that frequently used sequences of these operators can be executed from a shell script or batch file. Finally, NCO was written to consume the absolute minimum amount of system memory required to perform a given job. The arithmetic operators are extremely efficient; their exact memory usage is detailed in Section 2.9 [Memory usage], page 15.

### 2.2 Climate model paradigm

NCO was developed at NCAR to aid analysis and manipulation of datasets produced by General Circulation Models (GCMS). Datasets produced by GCMs share many features with all gridded scientific datasets and so provide a useful paradigm for the explication of the NCO operator set. Examples in this manual use a GCM paradigm because latitude, longitude, time, temperature and other fields related to our natural environment are as easy to visualize for the layman as the expert.

### 2.3 Temporary output files

NCO operators are designed to be reasonably fault tolerant, so that if there is a system failure or the user aborts the operation (e.g., with $C-c$ ), then no data are lost. The userspecified output-file is only created upon successful completion of the operation ${ }^{1}$. This is accomplished by performing all operations in a temporary copy of output-file. The name of the temporary output file is constructed by appending .pid<process ID>.<operator name>.tmp to the user-specified output-file name. When the operator completes its task with no fatal errors, the temporary output file is moved to the user-specified output-file.

[^4]Note the construction of a temporary output file uses more disk space than just overwriting existing files "in place" (because there may be two copies of the same file on disk until the NCO operation successfully concludes and the temporary output file overwrites the existing output-file). Also, note this feature increases the execution time of the operator by approximately the time it takes to copy the output-file. Finally, note this feature allows the output-file to be the same as the input-file without any danger of "overlap".

Other safeguards exist to protect the user from inadvertently overwriting data. If the output-file specified for a command is a pre-existing file, then the operator will prompt the user whether to overwrite (erase) the existing output-file, attempt to append to it, or abort the operation. However, in processing large amounts of data, too many interactive questions can be a curse to productivity. Therefore NCO also implements two ways to override its own safety features, the ' -0 ' and ' -A ' switches. Specifying ' -0 ' tells the operator to overwrite any existing output-file without prompting the user interactively. Specifying '-A' tells the operator to attempt to append to any existing output-file without prompting the user interactively. These switches are useful in batch environments because they suppress interactive keyboard input.

### 2.4 Appending variables to a file

A frequently useful operation is adding variables from one file to another. This is referred to as appending, although some prefer the terminology merging ${ }^{2}$ or pasting. Appending is often confused with what NCO calls concatenation. In NCO, concatenation refers to splicing a variable along the record dimension. Appending, on the other hand, refers to adding variables from one file to another ${ }^{3}$. In this sense, ncks can append variables from one file to another file. This capability is invoked by naming two files on the command line, input-file and output-file. When output-file already exists, the user is prompted whether to overwrite, append/replace, or exit from the command. Selecting overwrite tells the operator to erase the existing output-file and replace it with the results of the operation. Selecting exit causes the operator to exit - the output-file will not be touched in this case. Selecting append/replace causes the operator to attempt to place the results of the operation in the existing output-file, See Section 4.7 [ncks netCDF Kitchen Sink], page 61.

### 2.5 Addition Subtraction Division Multiplication and Interpolation

Users comfortable with NCO semantics may find it easier to perform some simple mathematical operations in NCO rather than higher level languages. ncbo (see Section 4.3 [ncbo netCDF Binary Operator], page 52) does file addition, subtraction, multiplication, division, and broadcasting. ncflint (see Section 4.6 [ncflint netCDF File Interpolator], page 59) does

2 The terminology merging is reserved for an (unwritten) operator which replaces hyperslabs of a variable in one file with hyperslabs of the same variable from another file
${ }^{3}$ Yes, the terminology is confusing. By all means mail me if you think of a better nomenclature. Should nCO use paste instead of append?
file addition, subtraction, multiplication and interpolation. Sequences of these commands can accomplish simple but powerful operations from the command line.

### 2.6 Averagers vs. Concatenators

The most frequently used operators of NCO are probably the averagers and concatenators. Because there are so many permutations of averaging (e.g., across files, within a file, over the record dimension, over other dimensions, with or without weights and masks) and of concatenating (across files, along the record dimension, along other dimensions), there are currently no fewer than five operators which tackle these two purposes: ncra, ncea, ncwa, ncrcat, and ncecat. These operators do share many capabilities ${ }^{4}$, but each has its unique specialty. Two of these operators, ncrcat and ncecat, are for concatenating hyperslabs across files. The other two operators, ncra and ncea, are for averaging hyperslabs across files ${ }^{5}$. First, let's describe the concatenators, then the averagers.

### 2.6.1 Concatenators ncreat and ncecat

Joining independent files together along a record coordinate is called concatenation. ncrcat is designed for concatenating record variables, while ncecat is designed for concatenating fixed length variables. Consider five files, '85.nc', '86.nc', ... '89.nc' each containing a year's worth of data. Say you wish to create from them a single file, '8589.nc' containing all the data, i.e., spanning all five years. If the annual files make use of the same record variable, then ncrcat will do the job nicely with, e.g., ncrcat 8?.nc 8589.nc. The number of records in the input files is arbitrary and can vary from file to file. See Section 4.9 [ncreat netCDF Record Concatenator], page 68, for a complete description of ncrcat.

However, suppose the annual files have no record variable, and thus their data are all fixed length. For example, the files may not be conceptually sequential, but rather members of the same group, or ensemble. Members of an ensemble may have no reason to contain a record dimension. ncecat will create a new record dimension (named record by default) with which to glue together the individual files into the single ensemble file. If ncecat is used on files which contain an existing record dimension, that record dimension will be converted into a fixed length dimension of the same name and a new record dimension will be created. Consider five realizations, '85a.nc', '85b.nc', ... '85e.nc' of 1985 predictions from the same climate model. Then ncecat 85 ?.nc 85 _ens.nc glues the individual realizations together into the single file, '85_ens.nc'. If an input variable was dimensioned [lat,lon], it will have dimensions [record,lat,lon] in the output file. A restriction of ncecat is that the hyperslabs of the processed variables must be the same from file to file. Normally this means all the input files are the same size, and contain data on different realizations of the
${ }^{4}$ Currently ncea and ncrcat are symbolically linked to the ncra executable, which behaves slightly differently based on its invocation name (i.e., 'argv[0]'). These three operators share the same source code, but merely have different inner loops.
5 The third averaging operator, ncwa, is the most sophisticated averager in Nco. However, ncwa is in a different class than ncra and ncea because it can only operate on a single file per invocation (as opposed to multiple files). On that single file, however, ncwa provides a richer set of averaging options-including weighting, masking, and broadcasting.
same variables. See Section 4.5 [ncecat netCDF Ensemble Concatenator], page 58, for a complete description of ncecat.

Note that ncreat cannot concatenate fixed-length variables, whereas ncecat can concatenate both fixed-length and record variables. To conserve system memory, use ncrcat rather than ncecat when concatenating record variables.

### 2.6.2 Averagers ncea, ncra, and ncwa

The differences between the averagers ncra and ncea are analogous to the differences between the concatenators. ncra is designed for averaging record variables from at least one file, while ncea is designed for averaging fixed length variables from multiple files. ncra performs a simple arithmetic average over the record dimension of all the input files, with each record having an equal weight in the average. ncea performs a simple arithmetic average of all the input files, with each file having an equal weight in the average. Note that ncra cannot average fixed-length variables, but ncea can average both fixed-length and record variables. To conserve system memory, use ncra rather than ncea where possible (e.g., if each input-file is one record long). The file output from ncea will have the same dimensions (meaning dimension names as well as sizes) as the input hyperslabs (see Section 4.4 [ncea netCDF Ensemble Averager], page 56, for a complete description of ncea). The file output from ncra will have the same dimensions as the input hyperslabs except for the record dimension, which will have a size of 1 (see Section 4.8 [ncra netCDF Record Averager], page 66 , for a complete description of ncra).

### 2.6.3 Interpolator ncflint

ncflint can interpolate data between or two files. Since no other operators have this ability, the description of interpolation is given fully on the ncflint reference page (see Section 4.6 [ncflint netCDF File Interpolator], page 59). Note that this capability also allows ncflint to linearly rescale any data in a netCDF file, e.g., to convert between differing units.

### 2.7 Working with large numbers of input files

Occasionally one desires to digest (i.e., concatenate or average) hundreds or thousands of input files. One brave user, for example, recently created a five year time-series of satellite observations by using ncecat to join thousands of daily data files together. Unfotunately, data archives (e.g., NASA EOSDIS) are unlikely to distribute netCDF files conveniently named in a format the '-n loop' switch (which automatically generates arbitrary numbers of input filenames) understands. If there is not a simple, arithmetic pattern to the input filenames (e.g., 'h00001.nc', 'h00002.nc', ... 'h90210.nc') then the ' -n loop' switch is useless. Moreover, when the input files are so numerous that the input filenames are too lengthy (when strung together as a single argument) to be passed by the calling shell to the NCO operator ${ }^{6}$, then the following strategy has proven useful to specify the input filenames

[^5]to NCO. Write a script that creates symbolic links between the irregular input filenames and a set of regular, arithmetic filenames that '-n loop' switch understands. The NCO operator will then succeed at automatically generating the filnames with the '-n loop' option (which circumvents any OS and shell limits on command line size). You can remove the symbolic links once the operator completes its task.

### 2.8 Working with large files

Large files are those files that are comparable in size to the amount of memory (RAM) in your computer. Many users of NCO work with files larger than 100 MB . Files this large not only push the current edge of storage technology, they present special problems for programs which attempt to access the entire file at once, such as ncea, and ncecat. If you need to work with a 300 MB file on a machine with only 32 MB of memory then you will need large amounts of swap space (virtual memory on disk) and NCO will work slowly, or else NCO will fail. There is no easy solution for this and the best strategy is to work on a machine with massive amounts of memory and swap space. That is, if your local machine has problems working with large files, try running NCO from a more powerful machine, such as a network server. Certain machine architectures, e.g., Cray unicos, have special commands which allow one to increase the amount of interactive memory. If you get a core dump on a Cray system (e.g., 'Error exit (core dumped)'), try increasing the available memory by using the ilimit command.

The speed of the NCO operators also depends on file size. When processing large files the operators may appear to hang, or do nothing, for large periods of time. In order to see what the operator is actually doing, it is useful to activate a more verbose output mode. This is accomplished by supplying a number greater than 0 to the '-D debug-level' (or '--debug-level', or '--dbg_lvl') switch. When the debug-level is nonzero, the operators report their current status to the terminal through the stderr facility. Using '-D' does not slow the operators down. Choose a debug-level between 1 and 3 for most situations, e.g., ncea -D $285 . n c 86 . n c 8586 . n c$. A full description of how to estimate the actual amount of memory the multi-file NCO operators consume is given in Section 2.9 [Memory usage], page 15 .

### 2.9 Approximate NCO memory requirements

The multi-file operators currently comprise the record operators, ncra and ncrcat, and the ensemble operators, ncea and ncecat. The record operators require much less memory than the ensemble operators. This is because the record operators are designed to operate on a single record of a file at a time, while the ensemble operators must retrieve an entire variable at a time into memory. Let $M S$ be the peak sustained memory demand of an operator, $F T$ be the memory required to store the entire contents of all the variables to be processed in an input file, $F R$ be the memory required to store the entire contents of a single

[^6]record of each of the variables to be processed in an input file, $V R$ be the memory required to store a single record of the largest record variable to be processed in an input file, $V T$ be the memory required to store the largest variable to be processed in an input file, $V I$ be the memory required to store the largest variable which is not processed, but is copied from the initial file to the output file. All operators require $M I=V I$ during the initial copying of variables from the first input file to the output file. This is the initial (and transient) memory demand. The sustained memory demand is that memory required by the operators during the processing (i.e., averaging, concatenation) phase which lasts until all the input files have been processed. The operators have the following memory requirements: ncrcat requires $M S<=V R$. ncecat requires $M S<=V T$. ncra requires $M S=2 F R+V R$. ncea requires $M S=2 F T+V T$. ncbo requires $M S<=2 V T$. ncflint requires $M S<=2 V T$. Note that only variables which are processed, i.e., averaged or concatenated, contribute to MS. Memory is never allocated to hold variables which do not appear in the output file (see Section 3.5 [Variable subsetting], page 22).

### 2.10 Performance limitations of the operators

1. No buffering of data is performed during ncvarget and ncvarput operations. Hyperslabs too large too hold in core memory will suffer substantial performance penalties because of this.
2. Since coordinate variables are assumed to be monotonic, the search for bracketing the user-specified limits should employ a quicker algorithm, like bisection, than the twosided incremental search currently implemented.
3. C_format, FORTRAN_format, signedness, scale_format and add_offset attributes are ignored by ncks when printing variables to screen.
4. Some random access operations on large files on certain architectures (e.g., 400 MB on unicos) are much slower with these operators than with similar operations performed using languages that bypass the netCDF interface (e.g., Yorick). The cause for this is not understood at present.

## 3 Features common to most operators

Many features have been implemented in more than one operator and are described here for brevity. The description of each feature is preceded by a box listing the operators for which the feature is implemented. Command line switches for a given feature are consistent across all operators wherever possible. If no "key switches" are listed for a feature, then that particular feature is automatic and cannot be controlled by the user.

### 3.1 Command line options

Availability: All operators
Short options: All
Long options: All

NCO achieves flexibility by using command line options. These options are implemented in all traditional UnIX commands as single letter switches, e.g., 'ls -l'. For many years NCo used only single letter option names. In late 2002, we implemented GNU/POSIX extended or long option names for all options. This was done in a backward compatible way such that the full functionality of NCO is still available through the familiar single letter options. In the future, however, some features of NCO may require the use of long options, simply because we have nearly run out of single letter options. More importantly, mnemonics for single letter options are often non-intuitive so that long options provide a more natural way of expressing intent.

Extended options are implemented using the system-supplied 'getopt.h' header file, if possible. This provides the getopt_long function to $\mathrm{NCO}^{1}$.

The syntax of short options (single letter options) is -key value (dash-key-space-value). Here, key is the single letter option name, e.g., '-D 2'.

The syntax of long options (multi-letter options) is --long_name value (dash-dash-key-space-value), e.g., '--dbg_lvl 2' or --long_name=value (dash-dash-key-equal-value), e.g., '--dbg_lvl=2'. Thus the following are all valid for the '-D' (short version) or '--dbg_lvl' (long version) command line option.

```
ncks -D 3 in.nc # Short option
ncks --dbg_lvl=3 in.nc # Long option, preferred form
ncks --dbg_lvl 3 in.nc # Long option, alternate form
```

The last example is preferred for two reasons. First, '--dbg_lvl' is more specific and less ambiguous than '-D'. The long option form makes scripts more self documenting and less error prone. Often long options are named after the source code variable whose value they carry. Second, the equals sign $=$ joins the key (i.e., long_name) to the value in an uninterruptible text block. Experience shows that users are less likely to mis-parse commands when restricted to this form.

[^7]GNU implements a superset of the POSIX standard which allows any unambiguous truncation of a valid option to be used.

```
ncks -D 3 in.nc # Short option
ncks --dbg_lvl=3 in.nc # Long option, full form
ncks --dbg=3 in.nc # Long option, unambiguous truncation
ncks --db=3 in.nc # Long option, unambiguous truncation
ncks --d=3 in.nc # Long option, ambiguous truncation
```

The first four examples are equivalent and will work as expected. The final example will exit with an error since ncks cannot disambiguate whether ' - - d ' is intended as a truncation of '--dbg_lvl', of '--dimension', or of some other long option.

NCO provides many long options for common switches. For example, the debugging level may be set in all operators with any of the switches '-D', '--debug-level', or '--dbg_lvl'. This flexibility allows users to choose their favorite mnemonic. For some, it will be '--debug' (an unambiguous truncation of '--debug-level', and other will prefer '--dbg'. Interactive users usually prefer the minimal amount of typing, i.e., '-D'. We recommend that scripts which are re-usable employ some form of the long options for future maintainability.

This manual generally uses the short option syntax. This is for historical reasons and to conserve space. The remainder of this manual specifies the full long_name of each option. Users are expected to pick the unambiguous truncation of each option name that most suits their taste.

### 3.2 Specifying input files

```
Availability: All operators
Short options: ' -n ', '-p'
Long options: '--nintap', '--pth', '--path'
```

It is important that the user be able to specify multiple input files without tediously typing in each by its full name. There are four different ways of specifying input files to NCO: explicitly typing each, using UNIX shell wildcards, and using the NCO ' -n ' and ' -p ' switches (or their long option equivalents, '--nintap' or '--pth' and '--path', respectively). To illustrate these methods, consider the simple problem of using ncra to average five input files, '85.nc', '86.nc', ... '89.nc', and store the results in '8589.nc'. Here are the four methods in order. They produce identical answers.

```
ncra 85.nc 86.nc 87.nc 88.nc 89.nc 8589.nc
ncra 8[56789].nc 8589.nc
ncra -p input-path 85.nc 86.nc 87.nc 88.nc 89.nc 8589.nc
ncra -n 5,2,1 85.nc 8589.nc
```

The first method (explicitly specifying all filenames) works by brute force. The second method relies on the operating system shell to glob (expand) the regular expression 8 [56789].nc. The shell passes valid filenames which match the expansion to ncra. The third method uses the '-p input-path' argument to specify the directory where all the input files reside. NCO prepends input-path (e.g., '/data/usrname/model') to all input-files
(but not to output-file). Thus, using ' -p ', the path to any number of input files need only be specified once. Note input-path need not end with '/'; the ' $/$ ' is automatically generated if necessary.

The last method passes (with '-n') syntax concisely describing the entire set of filenames ${ }^{2}$. This option is only available with the multi-file operators: ncra, ncreat, ncea, and ncecat. By definition, multi-file operators are able to process an arbitrary number of input-files. This option is very useful for abbreviating lists of filenames representable as alphanumeric_prefix+numeric_suffix+'.'+filetype where alphanumeric_prefix is a string of arbitrary length and composition, numeric_suffix is a fixed width field of digits, and filetype is a standard filetype indicator. For example, in the file 'ccm3_h0001.nc', we have alphanumeric_prefix $=$ 'ccm3_h', numeric_suffix $=$ '0001', and filetype $=$ ' nc '.

NCO is able to decode lists of such filenames encoded using the '-n' option. The simpler (3-argument) '-n' usage takes the form -n file_number, digit_number, numeric_ increment where file_number is the number of files, digit_number is the fixed number of numeric digits comprising the numeric_suffix, and numeric_increment is the constant, integer-valued difference between the numeric_suffix of any two consecutive files. The value of alphanumeric_prefix is taken from the input file, which serves as a template for decoding the filenames. In the example above, the encoding -n 5,2,1 along with the input file name '85.nc' tells NCO to construct five (5) filenames identical to the template '85.nc' except that the final two (2) digits are a numeric suffix to be incremented by one (1) for each successive file. Currently filetype may be either be empty, 'nc', 'cdf', 'hdf', or 'hd5'. If present, these filetype suffixes (and the preceding '.') are ignored by NCO as it uses the ' -n ' arguments to locate, evaluate, and compute the numeric_suffix component of filenames.

Recently the ' -n ' option has been extended to allow convenient specification of filenames with "circular" characteristics. This means it is now possible for NCO to automatically generate filenames which increment regularly until a specified maximum value, and then wrap back to begin again at a specified minimum value. The corresponding ' -n ' usage becomes more complex, taking one or two additional arguments for a total of four or five, respectively: -n file_number, digit_number, numeric_increment[,numeric_ $\max [$, numeric_min]] where numeric_max, if present, is the maximum integer-value of numeric_suffix and numeric_min, if present, is the minimum integer-value of numeric_suffix. Consider, for example, the problem of specifying non-consecutive input files where the filename suffixes end with the month index. In climate modeling it is common to create summertime and wintertime averages which contain the averages of the months June-July-August, and December-January-February, respectively:

```
ncra -n 3,2,1 85_06.nc 85_0608.nc
ncra -n 3,2,1,12 85_12.nc 85_1202.nc
ncra -n 3,2,1,12,1 85_12.nc 85_1202.nc
```

The first example shows that three arguments to the ' -n ' option suffice to specify consecutive months $(06,07,08)$ which do not "wrap" back to a minimum value. The second example shows how to use the optional fourth and fifth elements of the ' -n ' option to specify a wrap value to NCO. The fourth argument to ' -n ', if present, specifies the maximum integer value of numeric_suffix. In this case the maximum value is 12 , and will be formatted as ' 12 '

[^8]in the filename string. The fifth argument to ' -n ', if present, specifies the minimum integer value of numeric_suffix. The default minimum filename suffix is 1 , which is formatted as ' 01 ' in this case. Thus the second and third examples have the same effect, that is, they automatically generate, in order, the filenames '85_12.nc', '85_01.nc', and '85_02.nc' as input to NCO.

### 3.3 Accessing files stored remotely

## Availability: All operators

Short options: ' -p ', ' -1 '
Long options: '---pth', '--path', '--lcl', '--local'
All NCO operators can retrieve files from remote sites as well as from the local file system. A remote site can be an anonymous FTP server, a machine on which the user has rcp or scp privileges, or NCAR's Mass Storage System (MSS). To access a file via an anonymous FTP server, supply the remote file's URL. To access a file using rcp or scp, specify the Internet address of the remote file. Of course in this case you must have rcp or scp privileges which allow transparent (no password entry required) access to the remote machine. This means that ' / /.rhosts' or '~/ssh/authorized_keys' must be set accordingly on both local and remote machines.

To access a file on NCAR's MSS, specify the full MSS pathname of the remote file. NCO will attempt to detect whether the local machine has direct (synchronous) MSS access. In this case, NCO attempts to use the NCAR msrcp command ${ }^{3}$, or, failing that, /usr/local/bin/msread. Otherwise NCO attempts to retrieve the MSS file through the (asynchronous) Masnet Interface Gateway System (Migs) using the nrnet command.

The following examples show how one might analyze files stored on remote systems.

```
ncks -H -l ./ ftp://dust.ess.uci.edu/pub/zender/nco/in.nc
ncks -H -l ./ dust.ess.uci.edu:/home/zender/nco/in.nc
ncks -H -l ./ /ZENDER/nco/in.nc
ncks -H -l ./ mss:/ZENDER/nco/in.nc
ncks -H -l ./ -p http://www.cdc.noaa.gov/cgi-bin/nph-nc/Datasets/\
ncep.reanalysis.dailyavgs/surface air.sig995.1975.nc
```

The first example will work verbatim on your system if your system is connected to the Internet and is not behind a firewall. The second example will work on your system if you have rcp or scp access to the machine dust.ess.uci.edu. The third example will work from NCAR computers with local access to the msrcp, msread, or nrnet commands. The fourth command will work if your local version of NCO was built with DODS capability (see Section 3.3.1 [DODS], page 21). The above commands can be rewritten using the ' -p input-path' option as follows:

```
ncks -H -p ftp://dust.ess.uci.edu/pub/zender/nco -l ./ in.nc
ncks -H -p dust.ess.uci.edu:/home/zender/nco -l ./ in.nc
```

[^9]```
ncks -H -p /ZENDER/nco -l ./ in.nc
ncks -H -p mss:/ZENDER/nco -l ./ in.nc
```

Using ' -p ' is recommended because it clearly separates the input-path from the filename itself, sometimes called the stub. When input-path is not explicitly specified using ' -p ', NCO internally generates an input-path from the first input filename. The automatically generated input-path is constructed by stripping the input filename of everything following the final '/' character (i.e., removing the stub). The '-l output-path' option tells NCO where to store the remotely retrieved file and the output file. Often the path to a remotely retrieved file is quite different than the path on the local machine where you would like to store the file. If ' -1 ' is not specified then NCO internally generates an output-path by simply setting output-path equal to input-path stripped of any machine names. If '-l' is not specified and the remote file resides on the NCAR MSS system, then the leading character of input-path, ' $/$ ', is also stripped from output-path. Specifying output-path as '-1 ./' tells NCO to store the remotely retrieved file and the output file in the current directory. Note that ' -1 .' is equivalent to ' $-1 . /$ ' though the latter is recommended as it is syntactically more clear.

### 3.3.1 DODS

The Distributed Oceanographic Data System (DODS) provides replacements for common data interface libraries like netCDF. The DODS versions of these libraries implement network transparent access to data using the HTTP protocol. NCO may be DODS-enabled by linking nCO to the DODS libraries. Examples of how to do this are given in the DODS documentation and in the 'Makefile' distributed with NCO. Building NCO with make DODS=Y adds the (nonintuitive) commands to link to the DODS libraries installed in the \$DODS_ROOT directory. You will probably need to visit the DODS Homepage to learn which libraries to obtain and link to for the DODS-enabled NCO executables.

Once NCO is DODS-enabled the operators are DODS clients. All DODS clients have network transparent access to any files controlled by a DODS server. Simply specify the path to the file in URL notation

```
ncks -C -d lon,0 -v lon -l ./ -p http://www.cdc.noaa.gov/cgi-bin/nph-nc/
Datasets/ncep.reanalysis.dailyavgs/surface air.sig995.1975.nc foo.nc
```

NCO operates on these remote files without having to transfer the files to the local disk. dods causes all the I/O to appear to NCO as if the files were local. Only the required data (e.g., the variable or hyperslab specified) are transferred over the network. The advantages of this are obvious if you are examining small parts of large files stored at remote locations.

Note that the remote retrieval features of NCO can be used to retrieve any file, including non-netCDF files, via SSH, anonymous FTP, or msrcp. Often this method is quicker than using a browser, or running an FTP session from a shell window yourself. For example, say you want to obtain a JPEG file from a weather server.

```
ncks -p ftp://weather.edu/pub/pix/jpeg -l ./ storm.jpg
```

In this example, ncks automatically performs an anonymous FTP login to the remote machine and retrieves the specified file. When ncks attempts to read the local copy of
'storm.jpg' as a netCDF file, it fails and exits, leaving 'storm.jpg' in the current directory.

### 3.4 Retention of remotely retrieved files

Availability: All operators
Short options: ' - R'
Long options: '--rtn', '--retain'

In order to conserve local file system space, files retrieved from remote locations are automatically deleted from the local file system once they have been processed. Many nco operators were constructed to work with numerous large (e.g., 200 MB ) files. Retrieval of multiple files from remote locations is done serially. Each file is retrieved, processed, then deleted before the cycle repeats. In cases where it is useful to keep the remotely-retrieved files on the local file system after processing, the automatic removal feature may be disabled by specifying ' $-R$ ' on the command line.

### 3.5 Including/Excluding specific variables

Availability: (ncap), ncbo, ncea, ncecat, ncflint, ncks, ncra, ncrcat, ncwa
Short options: ' -v ', ' -x '
Long options: '--variable', '--exclude' or '--xcl'
Variable subsetting is implemented with the ' $-v \operatorname{var}[, \ldots]$ ' and ' -x ' options. A list of variables to extract is specified following the '-v' option, e.g., '-v time,lat,lon'. Not using the ' $-v$ ' option is equivalent to specifying all variables. The ' -x ' option causes the list of variables specified with '-v' to be excluded rather than extracted. Thus ' -x ' saves typing when you only want to extract fewer than half of the variables in a file. Remember, if averaging or concatenating large files stresses your systems memory or disk resources, then the easiest solution is often to use the ' -v ' option to retain only the variables you really need (see Section 2.9 [Memory usage], page 15). Note that, due to its special capabilities, ncap interprets the ' -v ' switch differently (see Section 4.1 [ncap netCDF Arithmetic Processor], page 42). For ncap, the ' $-v$ ' switch takes no arguments and indicates that only user-defined variables should be output. ncap neither accepts nor understands the $-x$ switch.

As of NCO 2.8.1 (August, 2003), variable name arguments of the '-v' switch may contain extended regular expressions. For example, '-v ' 'DST' ' selects all variables beginning with the string DST. Extended regular expressions are defined by the GNU egrep command. The meta-characters used to express pattern matching operations are ' $\$+$ +?.*[]\{\}|'. If the regular expression pattern matches any part of a variable name then that variable is selected. This capability is called wildcarding, and is very useful for sub-setting large data files.

Because of its wide availability, NCO uses the posix regular expression library regex. Regular expressions of arbitary complexity may be used. Since netCDF variable names are
relatively simple constructs, only a few varieties of variable wildcards are likely to be useful. Consider the variables Q01-Q99, Q100, Q_H2O, X_H2O, Q_CO2, X_CO2.

```
ncks -v 'Q+' in.nc # Select variables that start with Q
ncks -v 'Q??' in.nc # Select Q01--Q99, QAA--QZZ
ncks -v 'Q[0-9][0-9]' in.nc # Select Q01--Q99
ncks -v 'H2O$' in.nc # Select Q_H2O, X_H2O
ncks -v '^Q[0-9][0-9]' in.nc # Select Q01--Q99, Q100
ncks -v '^Q[0-9][0-9]$' in.nc # Select Q01--Q99
ncks -v '^[a-z]_[a-z]{3}$' in.nc # Select Q_H2O, X_H2O, Q_CO2, X_CO2
```

Beware - the repetition patter matching operator ' $*$ ' matches zero or more occurences of a regular expression. Thus '^o*', ‘^t*', and ‘^[a-z]*' select all variables. The documentation for the UNIX egrep command contains the detailed description of the extended regular expressions that NCO supports.

One must be careful to protect any special characters in the regular expression specification from being interpreted (globbed) by the shell. This is accomplish by enclosing special characters within single or double quotes

```
ncra -v Q?? in.nc out.nc # Error: Shell attempts to glob wildcards
ncra -v 'Q??' in.nc out.nc # Correct: NCO interprets wildcards
ncra -v 'Q??' in*.nc out.nc
```

The final example shows that commands may use a combination of variable wildcarding and shell filename expansion (globbing).

### 3.6 Including/Excluding coordinate variables

Availability: ncap, ncbo, ncea, ncecat, ncflint, ncks, ncra, ncrcat, ncwa
Short options: '-C', '-c'
Long options: '--no-coords', '--no-crd', '--crd', '--coords'

By default, coordinates variables associated with any variable appearing in the outputfile will also appear in the output-file, even if they are not explicitly specified, e.g., with the '-v' switch. Thus variables with a latitude coordinate lat always carry the values of lat with them into the output-file. This feature can be disabled with '-C', which causes nco to not automatically add coordinates to the variables appearing in the output-file. However, using ' $-C$ ' does not preclude the user from including some coordinates in the output files simply by explicitly selecting the coordinates with the $-v$ option. The ' $-c$ ' option, on the other hand, is a shorthand way of automatically specifying that all coordinate variables in the input-files should appear in the output-file. Thus ' -c ' allows the user to select all the coordinate variables without having to know their names.

### 3.7 C \& Fortran index conventions

Availability: ncbo, ncea, ncecat, ncflint, ncks, ncra, ncrcat, ncwa Short options: ' -F '
Long options: '--fortran'

By default, NCO uses C-style (0-based) indices for all I/O. The '-F' switch tells NCO to switch to reading and writing with Fortran index conventions. In Fortran, indices begin counting from 1 (rather than 0 ), and dimensions are ordered from fastest varying to slowest varying. Consider a file ' $85 . \mathrm{nc}$ ' containing 12 months of data in the record dimension time. The following hyperslab operations produce identical results, a June-July-August average of the data:

```
ncra -d time,5,7 85.nc 85_JJA.nc
ncra -F -d time,6,8 85.nc 85_JJA.nc
```

Printing variable three_dmn_var in file 'in.nc' first with C indexing conventions, then with Fortran indexing conventions results in the following output formats:

```
% ncks -H -v three_dmn_var in.nc
lat[0]=-90 lev[0]=1000 lon[0]=-180 three_dmn_var[0]=0
% ncks -F -H -v three_dmn_var in.nc
lon(1)=0 lev(1)=100 lat(1)=-90 three_dmn_var(1)=0
```


### 3.8 Hyperslabs

Availability: ncbo, ncea, ncecat, ncflint, ncks, ncra, ncrcat, ncwa Short options: ' -d '
Long options: '--dimension', '--dmn'

A hyperslab is a subset of a variable's data. The coordinates of a hyperslab are specified with the - d dim, [min][,[max]] short option (or with the '---dimension' or '--dmn' long options). The bounds of the hyperslab to be extracted are specified by the associated $\min$ and max values. A half-open range is specified by omitting either the min or max parameter but including the separating comma. The unspecified limit is interpreted as the maximum or minimum value in the unspecified direction. A cross-section at a specific coordinate is extracted by specifying only the min limit and omitting a trailing comma. Dimensions not mentioned are passed with no reduction in range. The dimensionality of variables is not reduced (in the case of a cross-section, the size of the constant dimension will be one). If values of a coordinate-variable are used to specify a range or cross-section, then the coordinate variable must be monotonic (values either increasing or decreasing). In this case, command-line values need not exactly match coordinate values for the specified dimension. Ranges are determined by seeking the first coordinate value to occur in the closed range $[\min , \max ]$ and including all subsequent values until one falls outside the range. The
coordinate value for a cross-section is the coordinate-variable value closest to the specified value and must lie within the range or coordinate-variable values.

Coordinate values should be specified using real notation with a decimal point required in the value, whereas dimension indices are specified using integer notation without a decimal point. This convention serves only to differentiate coordinate values from dimension indices. It is independent of the type of any netCDF coordinate variables. For a given dimension, the specified limits must both be coordinate values (with decimal points) or dimension indices (no decimal points).

User-specified coordinate limits are promoted to double precision values while searching for the indices which bracket the range. Thus, hyperslabs on coordinates of type NC_BYTE and NC_CHAR are computed numerically rather than lexically, so the results are unpredictable.

The relative magnitude of min and max indicate to the operator whether to expect a wrapped coordinate (see Section 3.11 [Wrapped coordinates], page 28), such as longitude. If min $>$ max, the NCO expects the coordinate to be wrapped, and a warning message will be printed. When this occurs, NCO selects all values outside the domain $[\max <\min$ ], i.e., all the values exclusive of the values which would have been selected if min and max were swapped. If this seems confusing, test your command on just the coordinate variables with ncks, and then examine the output to ensure NCO selected the hyperslab you expected (coordinate wrapping is currently only supported by ncks).

Because of the way wrapped coordinates are interpreted, it is very important to make sure you always specify hyperslabs in the monotonically increasing sense, i.e., $\min <\max$ (even if the underlying coordinate variable is monotonically decreasing). The only exception to this is when you are indeed specifying a wrapped coordinate. The distinction is crucial to understand because the points selected by, e.g., -d longitude,50. ,340., are exactly the complement of the points selected by -d longitude, 340.,50..

Not specifying any hyperslab option is equivalent to specifying full ranges of all dimensions. This option may be specified more than once in a single command (each hyperslabed dimension requires its own -d option).

### 3.9 Multislabs

Availability: ncks
Short options: '-d'
Long options: '--dimension', '--dmn'

In late 2002, ncks added support for specifying a multislab for any variable. A multislab is a union of one or more hyperslabs which is specified by chaining together hyperslab commands, i.e., $-d$ options (see Section 3.8 [Hyperslabs], page 24). This allows multislabs to overcome some restraints which limit hyperslabs.

A single -d option can only specify a contiguous and/or regularly spaced multidimensional array of data. Multislabs are constructed from multiple -d options and may therefore have non-regularly spaced arrays. For example, suppose it is desired to operate
on all longitudes from 10.0 to 20.0 and from 80.0 to 90.0 degrees. The combined range of longitudes is not selectable in a single hyperslab specfication of the form '-d lon,min,max' or '-d lon,min,max,stride' because its elements are irregularly spaced in coordinate space (and presumably in index space too). The multislab specification for obtaining these values is simply the union of the hyperslabs specifications that comprise the multislab, i.e.,

```
ncks -d lon,10.,20. -d lon,80.,90. in.nc out.nc
ncks -d lon,10.,15. -d lon,15.,20. -d lon,80.,90. in.nc out.nc
```

Any number of hyperslabs specifications may be chained together to specify the multislab. Multislabs are more efficient than the alternative of sequentially performing hyperslab operations and concatenating the results. This is because nCO employs a novel multislab algorithm to minimize the number of I/O operations when retrieving irregularly spaced data from disk.

Users may specify redundant ranges of indices in a multislab, e.g.,

```
ncks -d lon,0,4 -d lon,2,9,2 in.nc out.nc
```

This command retrieves the first five longitudes, and then every other longitude value up to the tenth. Elements 0, 2, and 4 are specified by both hyperslab arguments (hence this is redundant) but will count only once if an arithmetic operation is being performed. The nCO multislab algorithm retrieves each element from disk once and only once. Thus users may take some shortcuts in specifying multislabs and the algorithm will obtain the intended values. Specifying redundant ranges is not encouraged, but may be useful on occasion and will not result in unintended consequences.

A final example shows the real power of multislabs. Suppose the $Q$ variable contains three dimensional arrays of distinct chemical constituents in no particular order. We are interested in the NOy species in a certain geographic range. Say that NO, NO2, and N2O5 are elements 0,1 , and 5 of the species dimension of $Q$. The multislab specification might look something like

```
ncks -d species,0,1 -d species,5 -d lon,0,4 -d lon,2,9,2 in.nc out.nc
```

Multislabs are powerful because they may be specified for every dimension at the same time. Thus multislabs obsolete the need to execute multiple ncks commands to gather the desired range of data. We envision adding multislab support to all arithmetic operators in the future.

### 3.10 UDUnits Support

Availability: ncbo, ncea, ncecat, ncflint, ncks, ncra, ncrcat, ncwa
Short options: '-d'
Long options: '--dimension', '--dmn'
There is more than one way to hyperslab a cat. The UDUnits package provides a library which, if present, NCO uses to translate user-specified physical dimensions into the physical dimensions of data stored in netCDF files. Unidata provides UDUnits under the same terms
as netCDF, so sites should install both. Compiling NCO with UDUnits support is currently optional but may become required in a future version of NCO.

Two examples suffice to demonstrate the power and convenience of UDUnits support. First, consider extraction of a variable containing non-record coordinates with physical dimensions stored in MKS units. In the following example, the user extracts all wavelengths in the visible portion of the spectrum in terms of the units very frequently used in visible spectroscopy, microns:

```
% ncks -O -C -H -u -v wvl -d wvl,"0.4 micron","0.7 micron" in.nc
wvl[0]=5e-07 meter
```

The hyperslab returns the correct values because the wvl variable is stored on disk with a length dimension that UDUnits recognizes in the units attribute. The automagical algorithm that implements this functionality is worth describing since understanding it helps one avoid some potential pitfalls. First, the user includes the physical units of the hyperslab dimensions she supplies, separated by a simple space from the numerical values of the hyperslab limits. She encloses each coordinate specifications in quotes so that the shell does not break the value-space-unit string into separate arguments before passing them to NCO. Double quotes ("foo") or single quotes ('foo') are equally valid for this purpose. Second, NCO recognizes that units translation is requested because each hyperslab argument contains text characters and non-initial spaces. Third, nco determines whether the wvl is dimensioned with a coordinate variable that has a units attribute. In this case, wvl itself is a coordinate variable. The value of its units attribute is meter. Thus wvl passes this test so UDUnits conversion is attempted. If the coordinate associated with the variable does not contain a units attribute, then NCO aborts. Fourth, NCO passes the specified and desired dimension strings (microns are specified by the user, meters are required by NCO) to the UDUnits library. Fifth, the UDUnits library that these dimension are commensurate and it returns the appropriate linear scaling factors to convert from microns to meters to NCO. If the units are incommensurate (i.e., not expressible in the same fundamental MKS units), or are not listed in the UDUnits database, then NCO aborts since it cannot determine the user's intent. Finally, nco uses the scaling information to convert the userspecified hyperslab limits into the same physical dimensions as those of the corresponding cooridinate variable on disk. At this point, NCO can perform a coordinate hyperslab using the same algorithm as if the user had specified the hyperslab without requesting units conversion.

The translation and dimennterpretation of time coordinates shows a more powerful, and probably more common, application of the UDUnits feature. In this example, the user prints all data between the eighth and ninth of December, 1999, from a variable whose time dimension is hours since the year 1900:

```
% ncks -0 -C -H -u -v time_udunits -d time_udunits,"1999-12-08 \
12:00:0.0","1999-12-09 00:00:0.0",2 in.nc foo2.nc
% time_udunits[1]=876018 hours since 1900-01-01 00:00:0.0
```

Here, the user invokes the stride (see Section 3.12 [Stride], page 29) capability to obtain every other timeslice. This is possible because the UDUnits feature is additive, not exclusive - it works in conjunction with all other hyperslabbing (see Section 3.8 [Hyperslabs], page 24)
options and in all operators which support hyperslabbing. The following example shows how one might average data in a time period spread across multiple input files

```
ncra -0 -d time,"1939-09-09 12:00:0.0","1945-05-08 00:00:0.0" \
in1.nc in2.nc in3.nc out.nc
```

Note that there is no excess whitespace before or after the individual elements of the ' -d ' argument. This is important since, as far as the shell knows, '- d ' takes only one command-line argument. Parsing this argument into its component dim, [min], [max], stride elements (see Section 3.8 [Hyperslabs], page 24) is the job of Nco. When unquoted whitespace is present between these elements, the shell passes NCO arugment fragments which will not parse as intended.

The UDUnits package documentation describes the supported formats of time dimensions. Among the metadata conventions which adhere to these formats are the Climate and Forecast (CF) Conventions and the Cooperative Ocean/Atmosphere Research Data Service (COARDS) Conventions. The following '-d arguments' extract the same data using commonly encountered time dimension formats:

```
-d time,"1918-11-11 11:00:0.0","1939-09-09 00:00:0.0"
```

All of these formats include at least one dash - in a non-leading character position (a dash in a leading character position is a negative sign). NCO assumes that a non-leading dash in a limit string indicates that a UDUnits date conversion is requested.
netCDF variables should always be stored with MKS units, so that application programs may assume MKS dimensions apply to all input variables. The UDUnits feature is intended to alleviate some of the NCO user's pain when handling MKS units. It connects users who think in human-friendly units (e.g., miles, millibars, days) to extract data which are always stored in God's units, MKS (e.g., meters, Pascals, seconds). The feature is not intended to encourage writers to store data in esoteric units (e.g., furlongs, pounds per square inch, fortnights).

### 3.11 Wrapped coordinates

```
Availability: ncks
Short options: '-d'
Long options: '--dimension', '--dmn'
```

A wrapped coordinate is a coordinate whose values increase or decrease monotonically (nothing unusual so far), but which represents a dimension that ends where it begins (i.e., wraps around on itself). Longitude (i.e., degrees on a circle) is a familiar example of a wrapped coordinate. Longitude increases to the East of Greenwich, England, where it is defined to be zero. Halfway around the globe, the longitude is 180 degrees East (or West). Continuing eastward, longitude increases to 360 degrees East at Greenwich. The longitude values of most geophysical data are either in the range $[0,360)$, or $[-180,180)$. In either case, the Westernmost and Easternmost longitudes are numerically separated by 360 degrees, but represent contiguous regions on the globe. For example, the Saharan desert stretches from roughly 340 to 50 degrees East. Extracting the hyperslab of data representing the

Sahara from a global dataset presents special problems when the global dataset is stored consecutively in longitude from 0 to 360 degrees. This is because the data for the Sahara will not be contiguous in the input-file but is expected by the user to be contiguous in the output-file. In this case, ncks must invoke special software routines to assemble the desired output hyperslab from multiple reads of the input-file.

Assume the domain of the monotonically increasing longitude coordinate lon is $0<$ lon $<360$. ncks will extract a hyperslab which crosses the Greenwich meridian simply by specifying the westernmost longitude as min and the easternmost longitude as max. The following commands extract a hyperslab containing the Saharan desert:

```
ncks -d lon,340.,50. in.nc out.nc
ncks -d lon,340.,50. -d lat,10.,35. in.nc out.nc
```

The first example selects data in the same longitude range as the Sahara. The second example further constrains the data to having the same latitude as the Sahara. The coordinate lon in the output-file, 'out.nc', will no longer be monotonic! The values of lon will be, e.g., ' $340,350,0,10,20,30,40,50$ '. This can have serious implications should you run 'out.nc' through another operation which expects the lon coordinate to be monotonically increasing. Fortunately, the chances of this happening are slim, since lon has already been hyperslabbed, there should be no reason to hyperslab lon again. Should you need to hyperslab lon again, be sure to give dimensional indices as the hyperslab arguments, rather than coordinate values (see Section 3.8 [Hyperslabs], page 24).

### 3.12 Stride

```
Availability: ncks, ncra, ncrcat
Short options: '-d'
Long options: '--dimension', '--dmn'
```

ncks offers support for specifying a stride for any hyperslab, while ncra and ncrcat suport the stride argument only for the record dimension. The stride is the spacing between consecutive points in a hyperslab. A stride of 1 means pick all the elements of the hyperslab, but a stride of 2 means skip every other element, etc. Using the stride option with ncra and ncrcat makes it possible, for instance, to average or concatenate regular intervals across multi-file input data sets.

The stride is specified as the optional fourth argument to the '-d' hyperslab specification: -d dim, [min][,[max]][,[stride]]. Specify stride as an integer (i.e., no decimal point) following the third comma in the ' -d ' argument. There is no default value for stride. Thus using '-d time, , ,2' is valid but '-d time, , , 2.0' and '-d time, ,,' are not. When stride is specified but min is not, there is an ambiguity as to whether the extracted hyperslab should begin with (using C-style, 0 -based indexes) element 0 or element 'stride-1'. NCO must resolve this ambiguity and it chooses element 0 as the first element of the hyperslab when min is not specified. Thus '-d time, , ,stride' is syntactically equivalent to ' -d time, 0, ,stride'. This means, for example, that specifying the operation '-d time, , , 2' on the array ' $1,2,3,4,5$ ' selects the hyperslab ' $1,3,5$ '. To obtain the hyperslab ' 2,4 ' instead, simply explicitly specify the starting index as 1 , i.e., '-d time, 1, , $2^{\prime}$.

For example, consider a file '8501_8912.nc' which contains 60 consecutive months of data. Say you wish to obtain just the March data from this file. Using 0 -based subscripts (see Section 3.7 [Fortran indexing], page 24) these data are stored in records $2,14, \ldots 50$ so the desired stride is 12 . Without the stride option, the procedure is very awkward. One could use ncks five times and then use ncreat to concatenate the resulting files together:

```
for idx in 02 14 26 38 50; do # Bourne Shell
ncks -d time,${idx} 8501_8912.nc foo.${idx}
done
foreach idx (02 14 26 38 50) # C Shell
ncks -d time,${idx} 8501_8912.nc foo.${idx}
end
ncrcat foo.?? 8589_03.nc
rm foo.??
```

With the stride option, ncks performs this hyperslab extraction in one operation:

```
ncks -d time,2,,12 8501_8912.nc 8589_03.nc
```

See Section 4.7 [ncks netCDF Kitchen Sink], page 61, for more information on ncks.
The stride option is supported by ncra and ncrcat for the record dimension only. This makes it possible, for instance, to average or concatenate regular intervals across multi-file input data sets.

```
ncra -F -d time,3,,12 85.nc 86.nc 87.nc 88.nc 89.nc 8589_03.nc
ncrcat -F -d time,3,,12 85.nc 86.nc 87.nc 88.nc 89.nc 8503_8903.nc
```


### 3.13 Missing values

Availability: ncap, ncbo, ncea, ncflint, ncra, ncwa
Short options: None

The phrase missing data refers to data points that are missing, invalid, or for any reason not intended to be arithmetically processed in the same fashion as valid data. The nco arithmetic operators attempt to handle missing data in an intelligent fashion. There are four steps in the NCO treatment of missing data:

1. Identifying variables which may contain missing data.

NCO follows the convention that missing data should be stored with the missing_value specified in the variable's missing_value attribute ${ }^{4}$. The only way nco recognizes that a variable may contain missing data is if the variable has a missing_value attribute. In this case, any elements of the variable which are numerically equal to the missing_value are treated as missing data.

[^10]2. Converting the missing_value to the type of the variable, if neccessary.

Consider a variable var of type var_type with a missing_value attribute of type att_type containing the value missing_value. As a guideline, the type of the missing_ value attribute should be the same as the type of the variable it is attached to. If var_type equals att_type then NCO straightforwardly compares each value of var to missing_value to determine which elements of var are to be treated as missing data. If not, then NCO will internally convert att_type to var_type by using the implicit conversion rules of C, or, if att_type is NC_CHAR ${ }^{5}$, by typecasting the results of the C function strtod(missing_value). You may use the NCO operator ncatted to change the missing_value attribute and all data whose data is missing_value to a new value (see Section 4.2 [ncatted netCDF Attribute Editor], page 48).
3. Identifying missing data during arithmetic operations.

When an NCO arithmetic operator is processing a variable var with a missing_value attribute, it compares each value of var to missing_value before performing an operation. Note the missing_value comparison inflicts a performance penalty on the operator. Arithmetic processing of variables which contain the missing_value attribute always incurs this penalty, even when none of the data are missing. Conversely, arithmetic processing of variables which do not contain the missing_value attribute never incurs this penalty. In other words, do not attach a missing_value attribute to a variable which does not contain missing data. This exhortation can usually be obeyed for model generated data, but it may be harder to know in advance whether all observational data will be valid or not.
4. Treatment of any data identified as missing in arithmetic operators.

NCO averagers (ncra, ncea, ncwa) do not count any element with the value miss-ing-value towards the average. ncbo and ncflint define a missing_value result when either of the input values is a missing_value. Sometimes the missing_value may change from file to file in a multi-file operator, e.g., ncra. NCO is written to account for this (it always compares a variable to the missing-value assigned to that variable in the current file). Suffice it to say that, in all known cases, NCO does "the right thing".

### 3.14 Operation Types

```
Availability: ncra, ncea, ncwa
Short options: '-y'
Long options: '--operation', '--op_typ'
```

The '-y op_typ' switch allows specification of many different types of operations Set op_typ to the abbreviated key for the corresponding operation:

```
avg Mean value (default)
sqravg Square of the mean
avgsqr Mean of sum of squares
```

${ }^{5}$ For example, the DOE ARM program often uses att_type $=$ NC_CHAR and missing_value $=$ ' -99999. '

```
max Maximium value
min Minimium value
rms Root-mean-square (normalized by N)
rmssdn Root-mean square normalized by N-1
sqrt Square root of the mean
ttl Sum of values
```

If an operation type is not specified with '-y' then the operator will perform an arithmetic average by default. The mathematical definition of each operation is given below. See Section 4.11 [ncwa netCDF Weighted Averager], page 72, for additional information on masks and normalization. Averaging is the default, and will be described first so the terminology for the other operations is familiar.

The masked, weighted average of a variable $x$ can be generally represented as

$$
\bar{x}_{j}=\frac{\sum_{i=1}^{i=N} \mu_{i} m_{i} w_{i} x_{i}}{\sum_{i=1}^{i=N} \mu_{i} m_{i} w_{i}}
$$

where $\bar{x}_{j}$ is the $j^{\prime}$ 'th element of the output hyperslab, $x_{i}$ is the $i$ 'th element of the input hyperslab, $\mu_{i}$ is 1 unless $x_{i}$ equals the missing value, $m_{i}$ is 1 unless $x_{i}$ is masked, and $w_{i}$ is the weight. This formiddable looking formula represents a simple weighted average whose bells and whistles are all explained below. It is not early to note, however, that when $\mu_{i}=m_{i}=w_{i}=1$, the generic averaging expression above reduces to a simple arithmetic average. Furthermore, $m_{i}=w_{i}=1$ for all operators besides ncwa, but these variables are included in the discussion below for completeness and for possible future use in other operators.

The size $J$ of the output hyperslab for a given variable is the product of all the dimensions of the input variable which are not averaged over. The size $N$ of the input hyperslab contributing to each $\bar{x}_{j}$ is simply the product of the sizes of all dimensions which are averaged over (i.e., dimensions specified with ' -a '). Thus $N$ is the number of input elements which potentially contribute to each output element. An input element $x_{i}$ contributes to the output element $x_{j}$ except in two conditions:

1. $x_{i}$ equals the missing value (see Section 3.13 [Missing values], page 30) for the variable.
2. $x_{i}$ is located at a point where the masking condition (see Section 4.11.1 [Masking condition], page 73) is false.

Points $x_{i}$ in either of these two categories do not contribute to $x_{j}$, they are ignored. We now define these criteria more rigorously.

Each $x_{i}$ has an associated Boolean weight $\mu_{i}$ whose value is 0 or 1 (false or true). The value of $\mu_{i}$ is 1 (true) unless $x_{i}$ equals the missing value (see Section 3.13 [Missing values], page 30) for the variable. Thus, for a variable with no missing_value attribute, $\mu_{i}$ is always 1. All NCO arithmetic operators (ncbo, ncra, ncea, ncflint, ncwa) treat missing values analogously.

Besides (weighted) averaging, ncwa, ncra, and ncea also compute some common nonlinear operations which may be specified with the '-y' switch (see Section 3.14 [Operation

Types], page 31). The other rank-reducing operations are simple variations of the generic weighted mean described above. The total value of $x(-\mathrm{y} t \mathrm{tl})$ is

$$
\bar{x}_{j}=\sum_{i=1}^{i=N} \mu_{i} m_{i} w_{i} x_{i}
$$

Note that the total is the same as the numerator of the mean of $x$, and may also be obtained in ncwa by using the ' $-N$ ' switch (see Section 4.11 [ncwa netCDF Weighted Averager], page 72).

The minimum value of $x(-\mathrm{y} \min )$ is

$$
\bar{x}_{j}=\min \left[\mu_{1} m_{1} w_{1} x_{1}, \mu_{2} m_{2} w_{2} x_{2}, \ldots, \mu_{N} m_{N} w_{N} x_{N}\right]
$$

Analogously, the maximum value of $x$ (-y max) is

$$
\bar{x}_{j}=\max \left[\mu_{1} m_{1} w_{1} x_{1}, \mu_{2} m_{2} w_{2} x_{2}, \ldots, \mu_{N} m_{N} w_{N} x_{N}\right]
$$

Thus the minima and maxima are determined after any weights are applied.
The square of the mean value of $x$ (-y sqravg) is

$$
\bar{x}_{j}=\left(\frac{\sum_{i=1}^{i=N} \mu_{i} m_{i} w_{i} x_{i}}{\sum_{i=1}^{i=N} \mu_{i} m_{i} w_{i}}\right)^{2}
$$

The mean of the sum of squares of $x$ (-y avgsqr) is

$$
\bar{x}_{j}=\frac{\sum_{i=1}^{i=N} \mu_{i} m_{i} w_{i} x_{i}^{2}}{\sum_{i=1}^{i=N} \mu_{i} m_{i} w_{i}}
$$

If $x$ represents a deviation from the mean of another variable, $x_{i}=y_{i}-\bar{y}$ (possibly created by ncbo in a previous step), then applying avgsqr to $x$ computes the approximate variance of $y$. Computing the true variance of $y$ requires subtracting 1 from the denominator, discussed below. For a large sample size however, the two results will be nearly indistinguishable.

The root mean square of $x$ ( -y rms) is

$$
\bar{x}_{j}=\sqrt{\frac{\sum_{i=1}^{i=N} \mu_{i} m_{i} w_{i} x_{i}^{2}}{\sum_{i=1}^{i=N} \mu_{i} m_{i} w_{i}}}
$$

Thus rms simply computes the squareroot of the quantity computed by avgsqr.
The root mean square of $x$ with standard-deviation-like normalization (-y rmssdn) is implemented as follows. When weights are not specified, this function is the same as the root mean square of $x$ except one is subtracted from the sum in the denominator

$$
\bar{x}_{j}=\sqrt{\frac{\sum_{i=1}^{i=N} \mu_{i} m_{i} x_{i}^{2}}{-1+\sum_{i=1}^{i=N} \mu_{i} m_{i}}}
$$

If $x$ represents the deviation from the mean of another variable, $x_{i}=y_{i}-\bar{y}$, then applying rmssdn to $x$ computes the standard deviation of $y$. In this case the -1 in the denominator compensates for the degree of freedom already used in computing $\bar{y}$ in the numerator. Consult a statistics book for more details.

When weights are specified it is unclear how to compensate for this extra degree of freedom. Weighting the numerator and denominator of the above by $w_{i}$ and subtracting one from the denominator is only appropriate when all the weights are 1.0. When the weights are arbitrary (e.g., Gaussian weights), subtracting one from the sum in the denominator does not necessarily remove one degree of freedom. Therefore when -y rmssdn is requested and weights are specified, ncwa actually implements the rms procedure. ncea and ncra, which do not allow weights to be specified, always implement the rmssdn procedure when asked.

The square root of the mean of $x(-\mathrm{y}$ sqrt $)$ is

$$
\bar{x}_{j}=\sqrt{\frac{\sum_{i=1}^{i=N} \mu_{i} m_{i} w_{i} x_{i}}{\sum_{i=1}^{i=N} \mu_{i} m_{i} w_{i}}}
$$

The definitions of some of these operations are not universally useful. Mostly they were chosen to facilitate standard statistical computations within the nco framework. We are open to redefining and or adding to the above. If you are interested in having other statistical quantities defined in NCO please contact the NCO project (see Section 1.5 [Help and Bug reports], page 8).

## EXAMPLES

Suppose you wish to examine the variable prs_sfc(time,lat,lon) which contains a time series of the surface pressure as a function of latitude and longitude. Find the minimium value of prs_sfc over all dimensions:

```
ncwa -y min -v prs_sfc in.nc foo.nc
```

Find the maximum value of prs_sfc at each time interval for each latitude:

```
ncwa -y max -v prs_sfc -a lon in.nc foo.nc
```

Find the root-mean-square value of the time-series of prs_sfc at every gridpoint:

```
ncra -y rms -v prs_sfc in.nc foo.nc
ncwa -y rms -v prs_sfc -a time in.nc foo.nc
```

The previous two commands give the same answer but ncra is preferred because it has a smaller memory footprint. Also, ncra leaves the (degenerate) time dimension in the output file (which is usually useful) whereas ncwa removes the time dimension.

These operations work as expected in multi-file operators. Suppose that prs_sfc is stored in multiple timesteps per file across multiple files, say 'jan.nc', 'feb.nc', 'march.nc'. We can now find the three month maximium surface pressure at every point.

```
ncea -y max -v prs_sfc jan.nc feb.nc march.nc out.nc
```

It is possible to use a combination of these operations to compute the variance and standard deviation of a field stored in a single file or across multiple files. The procedure to compute
the temporal standard deviation of the surface pressure at all points in a single file 'in.nc' involves three steps.

```
ncwa -0 -v prs_sfc -a time in.nc out.nc
ncbo -0 --op_typ=sub -v prs_sfc in.nc out.nc out.nc
ncra -0 -y rmssdn out.nc out.nc
```

First the output file 'out.nc' is contructed containing the temporal mean of prs_sfc. Next 'out.nc' is overwritten with the deviation from the mean. Finally 'out.nc' is overwritten with the root-mean-square of itself. Note the use of '-y rmssdn' (rather than '-y rms') in the final step. This ensures the standard deviation is correctly normalized by one fewer than the number of time samples. The procedure to compute the variance is identical except for the use of '-y var' instead of '-y rmssdn' in the final step.

The procedure to compute the spatial standard deviation of a field in a single file 'in.nc' involves three steps.

```
ncwa -0 -v prs_sfc,gw -a lat,lon -w gw in.nc out.nc
ncbo -0 --op_typ=sub -v prs_sfc,gw in.nc out.nc out.nc
ncwa -0 -y rmssdn -v prs_sfc -a lat,lon -w gw out.nc out.nc
```

First the appropriately weighted (with '-w gw') spatial mean values are written to the output file. This example includes the use of a weighted variable specified with ' -w gw '. When using weights to compute standard deviations one must remember to include the weights in the initial output files so that they may be used again in the final step. The initial output file is then overwritten with the gridpoint deviations from the spatial mean. Finally the root-mean-square of the appropriately weighted spatial deviations is taken.

The procedure to compute the standard deviation of a time-series across multiple files involves one extra step since all the input must first be collected into one file.

```
ncrcat -0 -v tpt in.nc in.nc foo1.nc
ncwa -0 -a time foo1.nc foo2.nc
ncbo -0 --op_typ=sub -v tpt foo1.nc foo2.nc foo2.nc
ncra -0 -y rmssdn foo2.nc out.nc
```

The first step assembles all the data into a single file. This may require a lot of temporary disk space, but is more or less required by the ncbo operation in the third step.

### 3.15 Type conversion

```
Availability: ncap, ncbo, ncea, ncra, ncwa
```

Short options: None
Type conversion refers to the casting of one fundamental data type to another, e.g., converting NC_SHORT ( 2 bytes) to NC_DOUBLE ( 8 bytes). As a general rule, type conversions should be avoided for at least two reasons. First, type conversions are expensive since they require creating (temporary) buffers and casting each element of a variable from the type it was stored at to some other type. Second, the dataset's creator probably had a good reason
for storing data as, say, NC_FLOAT rather than NC_DOUBLE. In a scientific framework there is no reason to store data with more precision than the observations were made. Thus NCO tries to avoid performing type conversions when performing arithmetic.

Type conversion during arithmetic in the languages C and Fortran is performed only when necessary. All operands in an operation are converted to the most precise type before the operation takes place. However, following this parsimonious conversion rule dogmatically results in numerous headaches. For example, the average of the two NC_SHORTs 17000s and 17000s results in garbage since the intermediate value which holds their sum is also of type NC_SHORT and thus cannot represent values greater than $32,767^{6}$. There are valid reasons for expecting this operation to succeed and the NCO philosophy is to make operators do what you want, not what is most pure. Thus, unlike C and Fortran, but like many other higher level interpreted languages, NCO arithmetic operators will perform automatic type conversion when all the following conditions are met ${ }^{7}$ :

1. The operator is ncea, ncra, or ncwa. ncbo is not yet included in this list because subtraction did not benefit from type conversion. This will change in the future
2. The arithmetic operation could benefit from type conversion. Operations that could benefit (e.g., from larger representable sums) include averaging, summation, or any "hard" arithmetic. Type conversion does not benefit searching for minima and maxima ('-y min', or ' -y max').
3. The variable on disk is of type NC_BYTE, NC_CHAR, NC_SHORT, or NC_INT. Type NC_ DOUBLE is not type converted because there is no type of higher precision to convert to. Type NC_FLOAT is not type converted because, in our judgement, the performance penalty of always doing so would outweigh the (extremely rare) potential benefits.

When these criteria are all met, the operator converts the variable in question to type NC_DOUBLE, performs all the arithmetic operations, casts the NC_DOUBLE type back to the original type, and finally writes the result to disk. The result written to disk may not be what you expect, because of incommensurate ranges represented by different types, and because of (lack of) rounding. First, continuing the example given above, the average (e.g., '-y avg') of 17000 s and 17000 s is written to disk as 17000 s. The type conversion feature of NCO makes this possible since the arithmetic and intermediate values are stored as NC_DOUBLEs, i.e., 34000.0 d and only the final result must be represented as an NC_ SHORT. Without the type conversion feature of NCO, the average would have been garbage (albeit predictable garbage near -15768s). Similarly, the total (e.g., '-y ttl') of 17000s and 17000s written to disk is garbage (actually -31536s) since the final result (the true total) of 34000 is outside the range of type NC_SHORT.

Type conversions use the floor function to convert floating point number to integers. Type conversions do not attempt to round floating point numbers to the nearest integer. Thus the average of 1 s and 2 s is computed in double precisions arithmetic as $(1.0 \mathrm{~d}+$ $1.5 \mathrm{~d}) / 2)=1.5 \mathrm{~d}$. This result is converted to NC_SHORT and stored on disk as $\mathrm{floor}(1.5 \mathrm{~d})=$

[^11]$1 \mathrm{~s}^{8}$. Thus no "rounding up" is performed. The type conversion rules of C can be stated as follows: If $n$ is an integer then any floating point value x satisfying $n \leq x<n+1$ will have the value $n$ when converted to an integer.

### 3.16 Suppressing interactive prompts

## Availability: All operators

Short options: ' -0 ', '-A'
Long options: '--ovr', '--overwrite', '--apn', '--append'

If the output-file specified for a command is a pre-existing file, then the operator will prompt the user whether to overwrite (erase) the existing output-file, attempt to append to it, or abort the operation. However, in processing large amounts of data, too many interactive questions can be a curse to productivity. Therefore nco also implements two ways to override its own safety features, the ' -0 ' and ' -A ' switches. Specifying ' -0 ' tells the operator to overwrite any existing output-file without prompting the user interactively. Specifying ' -A ' tells the operator to attempt to append to any existing output-file without prompting the user interactively. These switches are useful in batch environments because they suppress interactive keyboard input.

### 3.17 History attribute

Availability: All operators
Short options: '-h'
Long options: '--hst', '--history'

All operators automatically append a history global attribute to any file they modify or create. The history attribute consists of a timestamp and the full string of the invocation command to the operator, e.g., 'Mon May 26 20:10:24 1997: ncks in.nc foo.nc'. The full contents of an existing history attribute are copied from the first input-file to the outputfile. The timestamps appear in reverse chronological order, with the most recent timestamp appearing first in the history attribute. Since NCO and many other netCDF operators adhere to the history convention, the entire data processing path of a given netCDF file may often be deduced from examination of its history attribute. As of May, 2002, nco is case-insensitive to the spelling of the history attribute name. Thus attributes named History or HISTORY (which are non-standard and not recommended) will be treated as valid history attributes. When more than one global attribute fits the case-insensitive search for "history", the first one found will be used. history attribute To avoid information overkill, all operators have an optional switch ('-h') to override automatically appending the history attribute (see Section 4.2 [ncatted netCDF Attribute Editor], page 48).

[^12]
### 3.18 NCAR CSM Conventions

Availability: ncbo, ncea, ncecat, ncflint, ncra, ncwa
Short options: None

NCO recognizes NCAR CSM history tapes, and treats them specially. If you do not work with NCAR CSM data then you may skip this section. The CSM netCDF convention is described at http://www.cgd.ucar.edu/csm/experiments/output.format.html. Most of the CSM netCDF convention is transparent to $\mathrm{NCO}^{9}$. There are no known pitfalls associated with using any NCO operator on files adhering to this convention ${ }^{10}$. However, to facilitate maximum user friendliness, nco does treat certain variables in some CSM files specially. The special functions are not required by the CSM netCDF convention, but experience has shown they do make life easier.

Currently, NCO determines whether a datafile is a CSM output datafile simply by checking whether value of the global attribute convention (if it exists) equals 'NCAR-CSM'. Should convention equal 'NCAR-CSM' in the (first) input-file, NCO will attempt to treat certain variables specially, because of their meaning in CSM files. NCO will not average the following variables often found in CSM files: ntrm, ntrn, ntrk, ndbase, nsbase, nbdate, nbsec, mdt, mhisf. These variables contain scalar metadata such as the resolution of the host CSM model and it makes no sense to change their values. Furthermore, the ncbo operator does not operate on (i.e., add, subtract, etc.) the following variables: gw, ORO, date, datesec, hyam, hybm, hyai, hybi. These variables represent the Gaussian weights, the orography field, time fields, and hybrid pressure coefficients. These are fields which you want to remain unaltered in the output file $99 \%$ of the time. If you decide you would like any of the above CSM fields processed, you must use ncrename to rename them first.

### 3.19 ARM Conventions

Availability: ncreat
Short options: None
ncrcat has been programmed to recognize ARM (Atmospheric Radiation Measurement Program) data files. If you do not work with ARM data then you may skip this section. ARM data files store time information in two variables, a scalar, base_time, and a record variable, time_offset. Subtle but serious problems can arise when these type of files are just blindly concatenated. Therefore ncrcat has been specially programmed to be able to

[^13]chain together consecutive ARM input-files and produce and an output-file which contains the correct time information. Currently, ncreat determines whether a datafile is an ARM datafile simply by testing for the existence of the variables base_time, time_offset, and the dimension time. If these are found in the input-file then ncrcat will automatically perform two non-standard, but hopefully useful, procedures. First, ncrcat will ensure that values of time_offset appearing in the output-file are relative to the base_time appearing in the first input-file (and presumably, though not necessarily, also appearing in the outputfile). Second, if a coordinate variable named time is not found in the input-files, then ncrcat automatically creates the time coordinate in the output-file. The values of time are defined by the ARM convention time $=$ base_time + time_offset. Thus, if output-file contains the time_offset variable, it will also contain the time coordinate. A short message is added to the history global attribute whenever these ARm-specific procedures are executed.

### 3.20 Operator version

```
Availability: All operators
Short options: ' \(-r\) '
Long options: ‘--revision', ‘--version', or ‘--vrs’
```

All operators can be told to print their internal version number and copyright notice and then quit with the ' $-r$ ' switch. The internal version number varies between operators, and indicates the most recent change to a particular operator's source code. This is useful in making sure you are working with the most recent operators. The version of NCO you are using might be, e.g., 1.2. However using '-r' on, say, ncks, will produce something like ' NCO netCDF Operators version 1.2 Copyright (C) 1995--2000 Charlie Zender ncks version 1.30 (2000/07/31) "Bolivia"'. This tells you ncks contains all patches up to version 1.30, which dates from July 31, 2000.

## 4 Reference manual for all operators

This chapter presents reference pages for each of the operators individually. The operators are presented in alphabetical order. All valid command line switches are included in the syntax statement. Recall that descriptions of many of these command line switches are provided only in Chapter 3 [Common features], page 17, to avoid redundancy. Only options specific to, or most useful with, a particular operator are described in any detail in the sections below.

## 4.1 ncap netCDF Arithmetic Processor

## SYNTAX

```
ncap [-A] [-C] [-c] [-D dbg]
[-d dim,[min][,[max]][,[stride]]] [-F] [-f]
[-l path] [-0] [-p path] [-R] [-r]
[-s algebra] [-S fl.nco] [-v]
input-file [output-file]
DESCRIPTION
```

Note: documentation for ncap is incomplete and evolving. The ncap parser tends to develop fitfully, and the best documentation for recent capabilities is the 'ChangeLog' file.
ncap arithmetically processes a netCDF file. The processing instructions are contained either in the NCO script file 'fl.nco' or in a sequence of command line arguments. The options '-s' (or long options '--spt' or '--script') are used for in-line scripts and '-S' (or long options '--fl_spt' or '--script-file') are used to provide the filename where (usually multiple) scripting commands are pre-stored. ncap was written to perform arbitrary albebraic transformations of data and archive the results as easily as possible. The results of the algebraic manipulations are called derived fields.

Unlike the other operators, ncap does not accept a list of variables to be operated on as an argument to ' -v ' (see Section 3.5 [Variable subsetting], page 22). Rather, the ' -v ' switch takes no arguments and indicates that ncap should output only user-defined variables. ncap does not accept or understand the -x switch.

### 4.1.1 Left hand casting

The following examples demonstrate the utility of the left hand casting ability of ncap. Consider first this simple, artificial, example. If lat and lon are one dimensional coordinates of dimensions lat and lon, respectively, then addition of these two one-dimensional arrays is intrinsically ill-defined because whether lat_lon should be dimensioned lat by lon or lon by lat is ambiguous (assuming that addition is to remain a commutative procedure, i.e., one that does not depend on the order of its arguments). Differing dimensions are said to be orthogonal to one another, and sets of dimensions which are mutually exclusive are orthogonal as a set and any arithmetic operation between variables in orthogonal dimensional spaces is ambiguous without further information.

The ambiguity may be resolved by enumerating the desired dimension ordering of the output expression inside square brackets on the left hand side (LHS) of the equals sign. This is called left hand casting because the user resolves the dimensional ordering of the RHS of the expression by specifying the desired ordering on the LHS.

```
ncap -0 -s "lat_lon[lat,lon]=lat+lon" in.nc out.nc
ncap -0 -s "lon_lat[lon,lat]=lat+lon" in.nc out.nc
```

The explicit list of dimensions on the LHS, [lat, lon] resolves the otherwise ambiguous ordering of dimensions in lat_lon. In effect, the LHS casts its rank properties onto the RHS.

Without LhS casting, the dimensional ordering of lat_lon would be undefined and, hopefully, ncap would print an error message.

Consider now a slightly more complex example. In geophysical models, a coordinate system based on a blend of terrain-following and density-following surfaces is called a hybrid coordinate system. In this coordinate system, four variables must be manipulated to obtain the pressure of the vertical coordinate: $P O$ is the domain-mean surface pressure offset (a scalar), $P S$ is the local (time-varying) surface pressure (usually two horizontal spatial dimensions, i.e, latitude by longitude), hyam is the weight given to surfaces of constant density (one spatial dimension, pressure, which is orthogonal to the horizontal dimensions), and hybm is the weight given to surfaces of constant elevation (also one spatial dimension). This command constructs a four-dimensional pressure prs_mdp from the four input variables of mixed rank and orthogonality:

```
ncap -0 -s "prs_mdp[time,lat,lon,lev]=P0*hyam+PS*hybm" in.nc out.nc
```

Manipulating the four fields which define the pressure in a hybrid coordinate system is easy with left hand casting.

### 4.1.2 Syntax of ncap statements

Mastering ncap is relatively simple. Each valid statement statement consists of standard forward algebraic expression. The 'fl.nco', if present, is simply a list of such statements, whitespace, and comments. The syntax of statements is most like the computer language C. The following characteristics of C are preserved:

Array syntax
Arrays elements are placed within [] characters;
Array indexing
Arrays are 0-based;
Array storage
Last dimension is most rapidly varying;
Assignment statements
A semi-colon ';' indicates the end of an assignment statement.
Comments
Multi-line comments are enclosed within /* */ characters. Single line comments are preceded by // characters.

Nesting Files may be nested in scripts using \#include script. Note that the \#include command is not followed by a semi-colon because it is a pre-processor directive, not an assignment statement. The filename 'script' is interpreted relative to the run directory.

Attribute syntax
The at-sign @ is used to delineate an attribute name from a variable name.

### 4.1.3 Intrinsic functions

ncap contains a small but growing library of intrinsic functions. In addition to the standard mathematical functions (see Section 4.1.4 [Intrinsic mathematical functions], page 45), ncap currently supports packing and unpacking.

## Packing and Unpacking Functions

pack(x) Packing The standard packing algorithm is applied to variable $x$. The packing algorithm is lossy, and produces data with the same dynamic range as the original but which requires no more than half the space to store. The packed variable is stored (usually) as type NC_SHORT with the two attributes required to unpack the variable, scale_factor and add_offset, stored at the original precision of the variable. Let min and max be the minimum and maximum values of $x$.

$$
\begin{aligned}
\text { scale_factor } & =(\max -\min ) / \mathrm{ndrv} \\
\text { add_offset } & =(\min +\max ) / 2 \\
\mathrm{pck} & =(\mathrm{upk}-\text { add_offset }) / \text { scale_factor } \\
& =\frac{\mathrm{ndrv} \times[\mathrm{upk}-(\min +\max ) / 2]}{\max -\min }
\end{aligned}
$$

where ndrv is the number of discrete representable values for given type of packed variable. The theoretical maximum value for ndrv is two raised to the number of bits used to store the packed variable. Thus if the variable is packed into type NC_SHORT, a 2 byte datatype, then there are at most $2^{1} 6=65536$ distinct values representible. In practice, the number of discretely representible values is taken to be one less than the theoretical maximum. This leaves extra space and solves potential problems with rounding which can occur during the unpacking of the variable. Thus for NC_SHORT, $n d r v=65536-1=65535$. Less often, the variable may be packed into type NC_CHAR, where $n d r v=256-1=$ 255 , or type NC_INT where where $n d r v=4294967295-1=4294967294$.
unpack (x)
Unpacking The standard unpacking algorithm is applied to variable $x$. The unpacking algorithm depends on the presence of two attributes, scale_factor and add_offset. If scale_factor is present for a variable, the data are multiplied by the value scale_factor after the data are read. If add_offset is present for a variable, then the add_offset value is added to the data after the data are read. If both scale_factor and add_offset attributes are present, the data are first scaled by scale_factor before the offset add_offset is added.

$$
\begin{aligned}
\text { upk } & =\text { scale_factor } \times \text { pck }+ \text { add_offset } \\
& =\frac{\text { pck } \times(\max -\min )}{\operatorname{ndrv}}+\frac{\min +\max }{2}
\end{aligned}
$$

When scale_factor and add_offset are used for packing, the associated variable (containing the packed data) is typically of type byte or short, whereas the
unpacked values are intended to be of type float or double. An attribute's scale_factor and add_offset should both be of the type intended for the unpacked data, e.g., float or double.

## Type Conversion Functions

```
byte(x) Convert to NC_BYTE Converts x to external type NC_BYTE, a C-type signed
    char.
char(x) Convert to NC_CHAR Converts x to external type NC_CHAR, a C-type unsigned
    char.
double(x)
    Convert to NC_DOUBLE Converts x to external type NC_DOUBLE, a C-type double.
float(x) Convert to NC_FLOAT Converts x to external type NC_FLOAT, a C-type float.
int(x) Convert to NC_INT Converts x to external type NC_INT, a C-type int.
short(x) Convert to NC_SHORT Converts x to external type NC_SHORT, a C-type short.
```


### 4.1.4 Intrinsic mathematical functions

ncap supports the standard mathematical functions supplied with most operating systems. Standard calculator notation is used for addition + , subtraction -, multiplication *, division $/$, exponentiation ^, and modulus \%. The available elementary mathematical functions are:
abs (x) Absolute value Absolute value of x .
$\operatorname{acos}(\mathrm{x}) \quad$ Arc-cosine Arc-cosine of x where x is specified in radians.
acosh ( x ) Hyperbolic arc-cosine Hyperbolic arc-cosine of $x$ where $x$ is specified in radians.
$\operatorname{asin}(x) \quad$ Arc-sine Arc-sine of $x$ where $x$ is specified in radians.
$\operatorname{asinh}(\mathrm{x}) \quad$ Hyperbolic arc-sine Hyperbolic arc-sine of $x$ where $x$ is specified in radians.
$\operatorname{atan}(\mathrm{x}) \quad$ Arc-tangent Arc-tangent of x where x is specified in radians between $-\pi / 2$ and $\pi / 2$.
atanh ( x ) Hyperbolic arc-tangent Hyperbolic arc-tangent of x where x is specified in radians between $-\pi / 2$ and $\pi / 2$.
ceil (x) Ceil Ceiling of $x$.
cerf( x ) Complementary error function Complementary error function of x where x is specified between -1 and 1 .
$\cos (x) \quad$ Cosine Cosine of $x$ where $x$ is specified in radians.
$\cosh (\mathrm{x}) \quad$ Hyperbolic cosine Hyperbolic cosine of $x$ where $x$ is specified in radians.
$\operatorname{erf}(\mathrm{x}) \quad$ Error function Error function of x where x is specified between -1 and 1 .
$\exp (\mathrm{x}) \quad$ Exponential Exponential of $x, e^{x}$.

```
floor(x) Floor Floor of x.
gamma(x) Gamma function Gamma function of x, \Gamma(x).
log(x) Natural Logarithm Natural logarithm of x, ln(x).
log10(x) Base 10 Logarithm Base 10 logarithm of x, 知10}(x)
nearbyint(x)
```

Round inexactly Nearest integer to $x$ is returned in floating point format. No exceptions are raised for inexact conversions.
rint(x) Round exactly Nearest integer to x is returned in floating point format. Exceptions are raised for inexact conversions.
round ( x ) Round Nearest integer to x is returned in floating point format. Round halfway cases away from zero, regardless of current IEEE rounding direction.
$\sin (x) \quad$ Sine Sine of $x$ where $x$ is specified in radians.
$\sinh (\mathrm{x}) \quad$ Hyperbolic sine Hyperbolic sine of x where x is specified in radians.
sqrt (x) Square Root Square Root of $x, \sqrt{x}$.
$\tan (\mathrm{x}) \quad$ Tangent Tangent of $x$ where $x$ is specified in radians.
$\tanh (\mathrm{x}) \quad$ Hyperbolic tangent Hyperbolic tangent of x where x is specified in radians.
trunc (x) Truncate Nearest integer to $x$ is returned in floating point format. Round halfway cases toward zero, regardless of current IEEE rounding direction.

The complete list of mathematical functions supported is platform-specific. Functions mandated by ANSI C are guaranteed to be present and are indicated with an asterisk ${ }^{1}$. and are indicated with an asterisk. Use the ' -f ' (or 'fnc_tbl' or 'prn_fnc_tbl') switch to print a complete list of functions supported on your platform. This prints a list of functions and whether they are supported for netCDF variables of intrinsic type NC_FLOAT and NC_DOUBLE. ${ }^{2}$

## EXAMPLES

Define new attribute new for existing variable one as twice the existing attribute double_att of variable att_var:

```
ncap -0 -s "one@new=2*att_var@double_att" in.nc out.nc
```

Average variables of mixed types (result is of type double):

[^14]```
ncap -0 -s "average=(var_float+var_double+var_int)/3" in.nc out.nc
```

Multiple commands may be given to ncap in three ways. First, the commands may be placed in a script which is executed, e.g., 'tst.nco'. Second, the commands may be individually specified with multiple ' $-s$ ' arguments to the same ncap invocation. Third, the commands may be chained together into a single '-s' argument to ncap. Assuming the file 'tst.nco' contains the commands $a=3 ; b=4 ; c=s q r t\left(a^{\wedge} 2+b\right.$ ^2 $)$; , then the following ncap invocations produce identical results:

```
ncap -0 -v -S tst.nco in.nc out.nc
ncap -0 -v -s "a=3" -s "b=4" -s "c=sqrt(a^2+b^2)" in.nc out.nc
ncap -0 -v -s "a=3;b=4;c=sqrt(a^2+b^2)" in.nc out.nc
```

The second and third examples show that ncap does not require that a trailing semicolon ';' be placed at the end of a ' $-s$ ' argument, although a trailing semi-colon ';' is always allowed. However, semi-colons are required to separate individual assignment statements chained together as a single '-s' argument.

Imagine you wish to create a binary flag based on the value of an array. The flag should have value 1.0 where the array exceeds 1.0 , and a value of 0.0 elsewhere. Assume the array named ORO is in 'in.nc'. The variable ORO_flg in 'out.nc'

```
# Add degenerate "record" dimension to ORO for averaging
ncecat -O -v ORO in.nc foo.nc
# Average degenerate "record" dimension using ORO as mask
ncwa -a record -0 -m ORO -M 1.0 -o gt foo.nc foo.nc
# ORO is either 0.0 or > 1.0 everywhere
# Create ORO_frc in [0.0,1.0) then add 0.99 and convert to int
ncap -0 -s "ORO_frc=ORO-int(ORO)" -s "ORO_flg=int(ORO_frc+0.99)" foo.nc out.nc
# ORO_flg now equals 0 or 1
```

This example uses ncap to compute the covariance of two variables. Let the variables $u$ and $v$ be the horizontal wind components. The covariance of $u$ and $v$ is defined as the time mean product of the deviations of $u$ and $v$ from their respective time means. Symbolically, the covariance $\left[u^{\prime} v^{\prime}\right]=[u v]-[u][v]$ where $[x]$ denotes the time-average of $x$, $[x] \equiv \frac{1}{\tau} \int_{t=0}^{t=\tau} x(t) d t$ and $x^{\prime}$ denotes the deviation from the time-mean. The covariance tells us how much of the correlation of two signals arises from the signal fluctuations versus the mean signals. Sometimes this is called the eddy covariance. We will store the covariance in the variable uprmvprm.

```
ncra -0 -v u,v in.nc foo.nc # Compute time mean of u,v
ncrename -O -v u,uavg -v v,vavg foo.nc # Rename to avoid conflict
ncks -A -v u,v in.nc foo.nc # Place originals with time means
ncap -0 -s "uprmvprm=u*v-uavg*vavg" foo.nc foo.nc # Covariance
ncra -O -v uprmvprm foo.nc out.nc # Time-mean covariance
```

The same answer would be obtained by replacing the step involving ncap with

```
ncap -0 -s "uprmvprm=(u-uavg)*(v-vavg)" foo.nc foo.nc # Covariance
```


## 4.2 ncatted netCDF Attribute Editor

## SYNTAX

```
ncatted [-a att_dsc] [-a ...] [-D dbg] [-h]
[-l path] [-0] [-p path] [-R] [-r]
input-file [output-file]
```


## DESCRIPTION

ncatted edits attributes in a netCDF file. If you are editing attributes then you are spending too much time in the world of metadata, and ncatted was written to get you back out as quickly and painlessly as possible. ncatted can append, create, delete, modify, and overwrite attributes (all explained below). Furthermore, ncatted allows each editing operation to be applied to every variable in a file, thus saving you time when you want to change attribute conventions throughout a file. ncatted interprets character attributes as strings.

Because repeated use of ncatted can considerably increase the size of the history global attribute (see Section 3.17 [History attribute], page 37), the '-h' switch is provided to override automatically appending the command to the history global attribute in the output-file.

When ncatted is used to change the missing_value attribute, it changes the associated missing data self-consistently. If the internal floating point representation of a missing value, e.g., 1.0e36, differs between two machines then netCDF files produced on those machines will have incompatible missing values. This allows ncatted to change the missing values in files from different machines to a single value so that the files may then be concatenated together, e.g., by ncrcat, without losing any information. See Section 3.13 [Missing values], page 30, for more information.

The key to mastering ncatted is understanding the meaning of the structure describing the attribute modification, att_dsc specified by the required option '-a' or '--attribute'. Each att_dsc contains five elements, which makes using ncatted somewhat complicated, but powerful. The att_dsc argument structure contains five arguments in the following order:

$$
\text { att_dsc }=\text { att_nm, var_nm, mode, att_type, att_val }
$$

att_nm Attribute name. Example: units
var_nm Variable name. Example: pressure
mode Edit mode abbreviation. Example: a. See below for complete listing of valid values of mode.
att_type Attribute type abbreviation. Example: c. See below for complete listing of valid values of att_type.
att_val Attribute value. Example: pascal.

There should be no empty space between these five consecutive arguments. The description of these arguments follows in their order of appearance.

The value of att_nm is the name of the attribute you want to edit. This meaning of this should be clear to all users of the ncatted operator. If att_nm is omitted (i.e., left blank) and Delete mode is selected, then all attributes associated with the specified variable will be deleted.

The value of var_nm is the name of the variable containing the attribute (named att_nm) that you want to edit. There are two very important and useful exceptions to this rule. The value of var_nm can also be used to direct ncatted to edit global attributes, or to repeat the editing operation for every variable in a file. A value of var_nm of "global" indicates that att_nm refers to a global attribute, rather than a particular variable's attribute. This is the method ncatted supports for editing global attributes. If var_nm is left blank, on the other hand, then ncatted attempts to perform the editing operation on every variable in the file. This option may be convenient to use if you decide to change the conventions you use for describing the data.

The value of mode is a single character abbreviation ( $\mathrm{a}, \mathrm{c}, \mathrm{d}, \mathrm{m}$, or o) standing for one of five editing modes:
a Append. Append value att_val to current var_nm attribute att_nm value att_val, if any. If var_nm does not have an attribute att_nm, there is no effect.
c Create. Create variable var_nm attribute att_nm with att_val if att_nm does not yet exist. If var_nm already has an attribute att_nm, there is no effect.
d Delete. Delete current var_nm attribute att_nm. If var_nm does not have an attribute att_nm, there is no effect. If att_nm is omitted (left blank), then all attributes associated with the specified variable are automatically deleted. When Delete mode is selected, the att_type and att_val arguments are superfluous and may be left blank.
m
Modify. Change value of current var_nm attribute att_nm to value att_val. If var_nm does not have an attribute att_nm, there is no effect.
o Overwrite. Write attribute att_nm with value att_val to variable var_nm, overwriting existing attribute att_nm, if any. This is the default mode.

The value of att_type is a single character abbreviation (f, d, l, i, s, c, or b) standing for one of the seven primitive netCDF data types:
f Float. Value(s) specified in att_val will be stored as netCDF intrinsic type NC_FLOAT.
d Double. Value(s) specified in att_val will be stored as netCDF intrinsic type NC_DOUBLE.
i Integer. Value(s) specified in att_val will be stored as netCDF intrinsic type NC_INT.

1 Long. Value(s) specified in att_val will be stored as netCDF intrinsic type NC_LONG.
s
Short. Value(s) specified in att_val will be stored as netCDF intrinsic type NC_SHORT.
c Char. Value(s) specified in att_val will be stored as netCDF intrinsic type NC_CHAR.
b Byte. Value(s) specified in att_val will be stored as netCDF intrinsic type NC_BYTE.

The specification of att_type is optional in Delete mode.
The value of att_val is what you want to change attribute att_nm to contain. The specification of att_val is optional in Delete mode. Attribute values for all types besides NC_CHAR must have an attribute length of at least one. Thus att_val may be a single value or one-dimensional array of elements of type att_type. If the att_val is not set or is set to empty space, and the att_type is NC_CHAR, e.g., -a units,T,o,c,"" or -a units, $\mathrm{T}, \mathrm{o}, \mathrm{c}$, , then the corresponding attribute is set to have zero length. When specifying an array of values, it is safest to enclose att_val in single or double quotes, e.g., -a levels, $\mathrm{T}, \mathrm{o}, \mathrm{s}, " 1,2,3,4$ " or -a levels, $\mathrm{T}, \mathrm{o}, \mathrm{s}, ' 1,2,3,4$ '. The quotes are strictly unnecessary around att_val except when att_val contains characters which would confuse the calling shell, such as spaces, commas, and wildcard characters.

NCO processing of NC_CHAR attributes is a bit like Perl in that it attempts to do what you want by default (but this sometimes causes unexpected results if you want unusual data storage). If the att_type is NC_CHAR then the argument is interpreted as a string and it may contain C-language escape sequences, e.g., $\backslash \mathrm{n}$, which NCO will interpret before writing anything to disk. NCO translates valid escape sequences and stores the appropriate ASCII code instead. Since two byte escape sequences, e.g., \n, represent one byte Ascii codes, e.g., ASCII 10 (decimal), the stored string attribute is one byte shorter than the input string length for each embedded escape sequence. The most frequently used C-language escape sequences are $\backslash \mathrm{n}$ (for linefeed) and $\backslash \mathrm{t}$ (for horizontal tab). These sequences in particular allow convenient editing of formatted text attributes. The other valid ASCII codes are $\backslash \mathrm{a}$, $\backslash \mathrm{b}, \backslash \mathrm{f}, \backslash \mathrm{r}, \backslash \mathrm{v}$, and $\backslash \backslash$. See Section 4.7 [ncks netCDF Kitchen Sink], page 61, for more examples of string formatting (with the ncks '-s' option) with special characters.

Analogous to printf, other special characters are also allowed by ncatted if they are "protected" by a backslash. The characters ", ', ?, and \may be input to the shell as \", $\backslash ', \backslash$ ?, and $\backslash \backslash$. NCO simply strips away the leading backslash from these characters before editing the attribute. No other characters require protection by a backslash. Backslashes which precede any other character (e.g., $3, \mathrm{~m}, \$, \mathrm{I}, \&, \varrho, \%$, \{, and \}) will not be filtered and will be included in the attribute.

Note that the NUL character $\backslash 0$ which terminates C language strings is assumed and need not be explicitly specified. If $\backslash 0$ is input, it will not be translated (because it would terminate the string in an additional location). Because of these context-sensitive rules, if wish to use an attribute of type NC_CHAR to store data, rather than text strings, you should use ncatted with care.

## EXAMPLES

Append the string "Data version 2.0. $\backslash \mathrm{n}$ " to the global attribute history:

```
ncatted -0 -a history,global,a,c,"Data version 2.O\n" in.nc
```

Note the use of embedded C language printf()-style escape sequences.
Change the value of the long_name attribute for variable T from whatever it currently is to "temperature":

```
ncatted -0 -a long_name,T,o,c,temperature in.nc
```

Delete all existing units attributes:

```
ncatted -0 -a units,,d,, in.nc
```

The value of var_nm was left blank in order to select all variables in the file. The values of att_type and att_val were left blank because they are superfluous in Delete mode.

Delete all attributes associated with the tpt variable:

```
ncatted -0 -a ,tpt,d,, in.nc
```

The value of att_nm was left blank in order to select all attributes associate with the variable.

Modify all existing units attributes to "meter second-1"

```
ncatted -0 -a units,,m,c,"meter second-1" in.nc
```

Overwrite the quanta attribute of variable energy to an array of four integers.

```
ncatted -0 -a quanta,energy,o,s,"010,101,111,121" in.nc
```

Demonstrate input of C-language escape sequences (e.g., $\backslash \mathrm{n}$ ) and other special characters (e.g., \")
ncatted -h -a special,global,o,c,
'\nDouble quote: \"\nTwo consecutive double quotes: \"\"\n Single quote: Beyond my shell abilities! \nBackslash: <br>\n Two consecutive backslashes: <br><br><br>nQuestion mark: \?\n' in.nc

Note that the entire attribute is protected from the shell by single quotes. These outer single quotes are necessary for interactive use, but may be omitted in batch scripts.

## 4.3 ncbo netCDF Binary Operator

## SYNTAX

```
ncbo [-A] [-C] [-c] [-D dbg]
[-d dim,[min][,[max]]] [-F] [-h] [-l path]
[-0] [-p path] [-R] [-r] [-v var[,...]]
[-x] [-y op_typ] file_1 file_2 file_3
```


## DESCRIPTION

ncbo performs binary operations on variables in file_1 and the corresponding variables (those with the same name) in file_2 and stores the results in file_3. The binary operation operates on the entire files (modulo any excluded variables). One of the four standard arithmetic binary operations currently supported must be selected with the '-y op_typ' switch (or long options '--op_typ' or '--operation'). The valid binary operations for ncbo, their definitions, and corresponding values of the op_typ key are:

Addition Definition: file_3 = file_1 + file_2
Alternate invocation: ncadd
op_typ key values: 'add', '+', 'addition'
Examples: 'ncbo --op_typ=add 1.nc 2.nc 3.nc', 'ncadd 1.nc 2.nc 3.nc'

Subtraction
Definition: file_3 = file_1 - file_2
Alternate invocations: ncdiff, ncsub, ncsubtract
op_typ key values: 'sbt', '-', 'dff', 'diff', 'sub', 'subtract', 'subtraction' Examples: 'ncbo --op_typ=-1.nc 2.nc 3.nc', 'ncdiff 1.nc 2.nc 3.nc'

Multiplication
Definition: file_3 $=$ file_ 1 * file_2
Alternate invocations: ncmult, ncmultiply
op_typ key values: 'mlt', '*', 'mult', 'multiply', 'multiplication'
Examples: 'ncbo --op_typ=mlt 1.nc 2.nc 3.nc', 'ncmult 1.nc 2.nc 3.nc'

Division Definition: file_3 = file_1 / file_2
Alternate invocation: ncdivide
op_typ key values: 'dvd', '/', 'divide', 'division'
Examples: 'ncbo --op_typ=/ 1.nc 2.nc 3.nc', 'ncdivide 1.nc 2.nc 3.nc'

Care should be taken when using the shortest form of key values, i.e., ' + ', ' - ', '*', '/'. Some of these single characters may have special meanings to the shell ${ }^{1}$. They should be

[^15]protected from the shell by placing them in quotes so that the shell does not attempt to interpret (glob) them ${ }^{2}$. For example, the following commands are equivalent

```
ncbo --op_typ=* 1.nc 2.nc 3.nc # Dangerous? (shell may attempt globbing)
ncbo --op_typ='*' 1.nc 2.nc 3.nc # Safe ('*' protected from shell)
ncbo --op_typ="*" 1.nc 2.nc 3.nc # Safe ('*' protected from shell)
ncbo --op_typ=mlt 1.nc 2.nc 3.nc
ncbo --op_typ=mult 1.nc 2.nc 3.nc
ncbo --op_typ=multiply 1.nc 2.nc 3.nc
ncbo --op_typ=multiplication 1.nc 2.nc 3.nc
ncmult 1.nc 2.nc 3.nc # First use ln -s ncbo ncmult
ncmultiply 1.nc 2.nc 3.nc # First use ln -s ncbo ncmult
```

No particular argument or invocation form is preferred. Users are encouraged to use the forms which are most intuitive to them.

Normally, an operation type must be specified with '-y' or ncbo will fail. Exceptions to this rule may be created to suit that tastes of your particular site. For many years, ncdiff was the main binary file operator. As a result, many users prefer to continue invoking ncdiff rather than memorizing a new command ('ncbo -y sbt') which behaves identically to the old ncdiff command. The is much to be said for the simplicity of ncdiff. However, from a software maintenance standpoint, maintaining a distinct executable for each binary operation (e.g., ncadd) is untenable.
ncbo subtracts variables in file_2 from the corresponding variables (those with the same name) in file_1 and stores the results in file_3. Variables in file_2 are broadcast to conform to the corresponding variable in file_1 if necessary. Broadcasting a variable means creating data in non-existing dimensions from the data in existing dimensions. For example, a two dimensional variable in file_2 can be subtracted from a four, three, or two (but not one or zero) dimensional variable (of the same name) in file_1. This functionality allows the user to compute anomalies from the mean. Note that variables in file_1 are not broadcast to conform to the dimensions in file_2. Thus, ncbo, the number of dimensions, or rank, of any processed variable in file_1 must be greater than or equal to the rank of the same variable in file_2. Furthermore, the size of all dimensions common to both file_1 and file_2 must be equal.

When computing anomalies from the mean it is often the case that file_2 was created by applying an averaging operator to a file with initially the same dimensions as file_1 (often file_1 itself). In these cases, creating file_2 with ncra rather than ncwa will cause the ncbo operation to fail. For concreteness say the record dimension in file_1 is time. If file_2 were created by averaging file_1 over the time dimension with the ncra operator rather than with the ncwa operator, then file_2 will have a time dimension of size 1 rather than having no time dimension at all ${ }^{3}$. In this case the input files to ncbo, file_1 and file_2,

[^16]will have unequally sized time dimensions which causes ncbo to fail. To prevent this from occuring, use ncwa to remove the time dimension from file_2. An example is given below.
ncbo will never difference coordinate variables or variables of type NC_CHAR or NC_BYTE. This ensures that coordinates like (e.g., latitude and longitude) are physically meaningful in the output file, file_3. This behavior is hardcoded. ncbo applies special rules to some NCAR CSM fields (e.g., ORO). See Section 3.18 [NCAR CSM Conventions], page 38 for a complete description. Finally, we note that ncflint (see Section 4.6 [ncflint netCDF File Interpolator], page 59) is designed for file interpolation. As such, it also performs file subtraction, addition, multiplication, albeit in a more convoluted way than ncbo.

## EXAMPLES

Say files '85_0112.nc' and '86_0112.nc' each contain 12 months of data. Compute the change in the monthly averages from 1985 to 1986:

```
ncbo -op_typ=sub 86_0112.nc 85_0112.nc 86m85_0112.nc
ncdiff 86_0112.nc 85_0112.nc 86m85_0112.nc
```

The following examples demonstrate the broadcasting feature of ncbo. Say we wish to compute the monthly anomalies of T from the yearly average of T for the year 1985. First we create the 1985 average from the monthly data, which is stored with the record dimension time.

```
ncra 85_0112.nc 85.nc
ncwa -0 -a time 85.nc 85.nc
```

The second command, ncwa, gets rid of the time dimension of size 1 that ncra left in ' $85 . \mathrm{nc}$ '. Now none of the variables in ' $85 . \mathrm{nc}$ ' has a time dimension. A quicker way to accomplish this is to use ncwa from the beginning:

```
ncwa -a time 85_0112.nc 85.nc
```

We are now ready to use ncbo to compute the anomalies for 1985:

```
ncdiff -v T 85_0112.nc 85.nc t_anm_85_0112.nc
```

Each of the 12 records in 't_anm_85_0112.nc' now contains the monthly deviation of $T$ from the annual mean of $T$ for each gridpoint.

Say we wish to compute the monthly gridpoint anomalies from the zonal annual mean. A zonal mean is a quantity that has been averaged over the longitudinal (or $x$ ) direction. First we use ncwa to average over longitudinal direction lon, creating '85_x.nc', the zonal mean of ' $85 . \mathrm{nc}$ '. Then we use ncbo to subtract the zonal annual means from the monthly gridpoint data:
ncwa -a lon 85.nc 85_x.nc
ncdiff 85_0112.nc 85_x.nc tx_anm_85_0112.nc
This examples works assuming '85_0112.nc' has dimensions time and lon, and that '85_x.nc' has no time or lon dimension.

As a final example, say we have five years of monthly data (i.e., 60 months) stored in '8501_8912.nc' and we wish to create a file which contains the twelve month seasonal cycle of the average monthly anomaly from the five-year mean of this data. The following method
is just one permutation of many which will accomplish the same result. First use ncwa to create the file containing the five-year mean:

```
ncwa -a time 8501_8912.nc 8589.nc
```

Next use ncbo to create a file containing the difference of each month's data from the five-year mean:
ncbo 8501_8912.nc 8589.nc t_anm_8501_8912.nc
Now use ncks to group the five January anomalies together in one file, and use ncra to create the average anomaly for all five Januarys. These commands are embedded in a shell loop so they are repeated for all twelve months:

```
for idx in 01 02 03 04 05 06 07 08 09 10 11 12; do # Bourne Shell
ncks -F -d time,${idx},,12 t_anm_8501_8912.nc foo.${idx}
ncra foo.${idx} t_anm_8589_${idx}.nc
done
foreach idx (01 02 03 04 05 06 07 08 09 10 11 12) # C Shell
ncks -F -d time,${idx},,12 t_anm_8501_8912.nc foo.${idx}
ncra foo.${idx} t_anm_8589_${idx}.nc
end
```

Note that ncra understands the stride argument so the two commands inside the loop may be combined into the single command

```
ncra -F -d time,${idx},,12 t_anm_8501_8912.nc foo.${idx}
```

Finally, use ncrcat to concatenate the 12 average monthly anomaly files into one twelverecord file which contains the entire seasonal cycle of the monthly anomalies:
ncrcat t_anm_8589_??.nc t_anm_8589_0112.nc

## 4.4 ncea netCDF Ensemble Averager

## SYNTAX

```
ncea [-A] [-C] [-c] [-D dbg]
[-d dim,[min][,[max]]] [-F] [-h] [-l path]
[-n loop] [-0] [-p path] [-R] [-r] [-v var[,...]]
[-x] [-y op_typ] input-files output-file
```


## DESCRIPTION

ncea performs gridpoint averages of variables across an arbitrary number (an ensemble) of input files, with each file receiving an equal weight in the average. Each variable in the output-file will be the same size as the same variable in any one of the in the inputfiles, and all input-files must be the same size. Whereas ncra only performs averages over the record dimension (e.g., time), and weights each record in the record dimension evenly, ncea averages entire files, and weights each file evenly. All dimensions, including the record dimension, are treated identically and preserved in the output-file. See Section 2.6 [Averaging vs. Concatenating], page 13, for a description of the distinctions between the various averagers and concatenators.

The file is the logical unit of organization for the results of many scientific studies. Often one wishes to generate a file which is the gridpoint average of many separate files. This may be to reduce statistical noise by combining the results of a large number of experiments, or it may simply be a step in a procedure whose goal is to compute anomalies from a mean state. In any case, when one desires to generate a file whose properties are the mean of all the input files, then ncea is the operator to use. ncea assumes coordinate variable are properties common to all of the experiments and so does not average them across files. Instead, ncea copies the values of the coordinate variables from the first input file to the output file.

## EXAMPLES

Consider a model experiment which generated five realizations of one year of data, say 1985. You can imagine that the experimenter slightly perturbs the initial conditions of the problem before generating each new solution. Assume each file contains all twelve months (a seasonal cycle) of data and we want to produce a single file containing the ensemble average (mean) seasonal cycle. Here the numeric filename suffix denotes the experiment number (not the month):

```
ncea 85_01.nc 85_02.nc 85_03.nc 85_04.nc 85_05.nc 85.nc
ncea 85_0[1-5].nc 85.nc
ncea -n 5,2,1 85_01.nc 85.nc
```

These three commands produce identical answers. See Section 3.2 [Specifying input files], page 18, for an explanation of the distinctions between these methods. The output file, ' $85 . \mathrm{nc}$ ', is the same size as the inputs files. It contains 12 months of data (which might or might not be stored in the record dimension, depending on the input files), but each value in the output file is the average of the five values in the input files.

In the previous example, the user could have obtained the ensemble average values in a particular spatio-temporal region by adding a hyperslab argument to the command, e.g.,

```
ncea -d time,0,2 -d lat,-23.5,23.5 85_??.nc 85.nc
```

In this case the output file would contain only three slices of data in the time dimension. These three slices are the average of the first three slices from the input files. Additionally, only data inside the tropics is included.

## 4.5 ncecat netCDF Ensemble Concatenator

## SYNTAX

```
ncecat [-A] [-C] [-c] [-D dbg]
[-d dim,[min][,[max]]] [-F] [-h] [-1 path]
[-n loop] [-0] [-p path] [-R] [-r] [-v var[,...]]
    [-x] input-files output-file
```


## DESCRIPTION

ncecat concatenates an arbitrary number of input files into a single output file. Input files are glued together by creating a record dimension in the output file. Input files must be the same size. Each input file is stored consecutively as a single record in the output file. Thus, the size of the output file is the sum of the sizes of the input files. See Section 2.6 [Averaging vs. Concatenating], page 13, for a description of the distinctions between the various averagers and concatenators.

Consider five realizations, '85a.nc', '85b.nc', ... '85e.nc' of 1985 predictions from the same climate model. Then ncecat 85 ? .nc 85 _ens.nc glues the individual realizations together into the single file, '85_ens.nc'. If an input variable was dimensioned [lat,lon], it will have dimensions [record,lat,lon] in the output file. A restriction of ncecat is that the hyperslabs of the processed variables must be the same from file to file. Normally this means all the input files are the same size, and contain data on different realizations of the same variables.

## EXAMPLES

Consider a model experiment which generated five realizations of one year of data, say 1985. You can imagine that the experimenter slightly perturbs the initial conditions of the problem before generating each new solution. Assume each file contains all twelve months (a seasonal cycle) of data and we want to produce a single file containing all the seasonal cycles. Here the numeric filename suffix denotes the experiment number (not the month):

```
ncecat 85_01.nc 85_02.nc 85_03.nc 85_04.nc 85_05.nc 85.nc
ncecat 85_0[1-5].nc 85.nc
ncecat -n 5,2,1 85_01.nc 85.nc
```

These three commands produce identical answers. See Section 3.2 [Specifying input files], page 18, for an explanation of the distinctions between these methods. The output file, ' $85 . \mathrm{nc}$ ', is five times the size as a single input-file. It contains 60 months of data (which might or might not be stored in the record dimension, depending on the input files).

## 4.6 ncflint netCDF File Interpolator

## SYNTAX

```
ncflint [-A] [-C] [-c] [-D dbg]
[-d dim,[min][,[max]]] [-F] [-h]
[-i var,val3]
[-l path] [-0] [-p path] [-R] [-r] [-v var[,...]]
[-w wgt1[,wgt2]] [-x] file_1 file_2 file_3
```


## DESCRIPTION

ncflint creates an output file that is a linear combination of the input files. This linear combination can be a weighted average, a normalized weighted average, or an interpolation of the input files. Coordinate variables are not acted upon in any case, they are simply copied from file_1.

There are two conceptually distinct methods of using ncflint. The first method is to specify the weight each input file is to have in the output file. In this method, the value val3 of a variable in the output file file_3 is determined from its values vall and val2 in the two input files according to val $3=w g t 1 \times v a l 1+w g t 2 \times v a l 2$. Here at least wgt1, and, optionally, wgt2, are specified on the command line with the '-w' (or '--weight' or '--wgt_var') switch. If only wgt1 is specified then wgt2 is automatically computed as wgt2 $=1$-wgt1. Note that weights larger than 1 are allowed. Thus it is possible to specify wgt1 $=2$ and wgt2 $=-3$. One can use this functionality to multiply all the values in a given file by a constant.

The second method of using ncflint is to specify the interpolation option with '-i' (or with the '--ntp' or '--interpolate' long options). This is really the inverse of the first method in the following sense. When the user specifies the weights directly, ncflint has no work to do besides multiplying the input values by their respective weights and adding the results together to produce the output values. This assumes it is the weights that are known a priori. In another class of cases it is the arrival value (i.e., val3) of a particular variable var that is known a priori. In this case, the implied weights can always be inferred by examining the values of var in the input files. This results in one equation in two unknowns, wgt1 and wgt2: val3 $=w g t 1 \times$ val $1+w g t 2 \times$ val 2 . Unique determination of the weights requires imposing the additional constraint of normalization on the weights: wgt1 + wgt2 $=1$. Thus, to use the interpolation option, the user specifies var and val3 with the '-i' option. ncflint will compute wgt1 and wgt2, and use these weights on all variables to generate the output file. Although var may have any number of dimensions in the input files, it must represent a single, scalar value. Thus any dimensions associated with var must be degenerate, i.e., of size one.

If neither '-i' nor '-w' is specified on the command line, ncflint defaults to weighting each input file equally in the output file. This is equivalent to specifying ' -w 0.5 ' or ' -w $0.5,0.5$ '. Attempting to specify both ' -i ' and ' -w ' methods in the same command is an error.
ncflint is programmed not to interpolate variables of type NC_CHAR and NC_BYTE. This behavior is hardcoded.

## EXAMPLES

Although it has other uses, the interpolation feature was designed to interpolate file_3 to a time between existing files. Consider input files ' $85 . \mathrm{nc}$ ' and ' $87 . \mathrm{nc}$ ' containing variables describing the state of a physical system at times time $=85$ and time $=87$. Assume each file contains its timestamp in the scalar variable time. Then, to linearly interpolate to a file ' $86 . n c$ ' which describes the state of the system at time at time $=86$, we would use

```
ncflint -i time,86 85.nc 87.nc 86.nc
```

Say you have observational data covering January and April 1985 in two files named '85_01.nc' and '85_04.nc', respectively. Then you can estimate the values for February and March by interpolating the existing data as follows. Combine '85_01.nc' and '85_04.nc' in a $2: 1$ ratio to make ' 85 _02.nc':

```
ncflint -w 0.667 85_01.nc 85_04.nc 85_02.nc
ncflint -w 0.667,0.333 85_01.nc 85_04.nc 85_02.nc
```

Multiply ' $85 . n c$ ' by 3 and by -2 and add them together to make 'tst.nc':

```
ncflint -w 3,-2 85.nc 85.nc tst.nc
```

This is an example of a null operation, so 'tst.nc' should be identical (within machine precision) to ' $85 . \mathrm{nc}$ '.

Add '85.nc' to '86.nc' to obtain '85p86.nc', then subtract '86.nc' from '85.nc' to obtain ' $85 \mathrm{~m} 86 . \mathrm{nc}$ '

```
ncflint -w 1,1 85.nc 86.nc 85p86.nc
ncflint -w 1,-1 85.nc 86.nc 85m86.nc
ncdiff 85.nc 86.nc 85m86.nc
```

Thus ncflint can be used to mimic some ncbo operations. However this is not a good idea in practice because ncflint does not broadcast (see Section 4.3 [ncbo netCDF Binary Operator], page 52) conforming variables during arithmetic. Thus the final two commands would produce identical results except that ncflint would fail if any variables needed to be broadcast.

Rescale the dimensional units of the surface pressure prs_sfc from Pascals to hectopascals (millibars)

```
ncflint -0 -C -v prs_sfc -w 0.01,0.0 in.nc in.nc out.nc
ncatted -0 -a units,prs_sfc,o,c,millibar out.nc
```


## 4.7 ncks netCDF Kitchen Sink

## SYNTAX

```
ncks [-A] [-a] [-B] [-b binary-file] [-C] [-c] [-D dbg]
[-d dim,[min][,[max]][,[stride]]]
[-F] [-H] [-h] [-l path] [-M] [-m] [-0] [-p path] [-q]
[-R] [-r] [-s format] [-u] [-v var[,...]] [-x]
input-file [output-file]
```

DESCRIPTION
ncks combines selected features of ncdump, ncextr, and the nccut and ncpaste specifications into one versatile utility. ncks extracts a subset of the data from input-file and prints it as ASCII text to 'stdout', writes it in flat binary format to 'binary-file', and writes (or pastes) it in netCDF format to output-file.
ncks will print netCDF data in ASCII format to stdout, like ncdump, but with these differences: ncks prints data in a tabular format intended to be easy to search for the data you want, one datum per screen line, with all dimension subscripts and coordinate values (if any) preceding the datum. Option '-s' (or lon options '--sng', '--string', '--fmt', or '--format') allows the user the format the data using C-style format strings.
 control the formatted appearance of the data.
ncks will extract (and optionally create a new netCDF file comprised of) only selected variable from the input file, like ncextr but with these differences: Only variables and coordinates may be specifically included or excluded - all global attributes and any attribute associated with an extracted variable will be copied to the screen and/or output netCDF file. Options ' -c ', ' -C ', ' -v ', and ' -x ' (and their long option synomyms) control which variables are extracted.
ncks will extract hyperslabs from the specified variables. In fact ncks implements the nccut specification exactly. Option '-d' controls the hyperslab specification.

Input dimensions that are not associated with any output variable will not appear in the output netCDF. This feature removes superfluous dimensions from a netCDF file.
ncks will append variables and attributes from the input-file to output-file if output-file is a pre-existing netCDF file whose relevant dimensions conform to dimension sizes of inputfile. The append features of ncks are intended to provide a rudimentary means of adding data from one netCDF file to another, conforming, netCDF file. When naming conflicts exists between the two files, data in output-file is usually overwritten by the corresponding data from input-file. Thus it is recommended that the user backup output-file in case valuable data are accidentally overwritten.

If output-file exists, the user will be queried whether to overwrite, append, or exit the ncks call completely. Choosing overwrite destroys the existing output-file and create an entirely new one from the output of the ncks call. Append has differing effects depending on the uniqueness of the variables and attributes output by ncks: If a variable or attribute
extracted from input-file does not have a name conflict with the members of output-file then it will be added to output-file without overwriting any of the existing contents of outputfile. In this case the relevant dimensions must agree (conform) between the two files; new dimensions are created in output-file as required. When a name conflict occurs, a global attribute from input-file will overwrite the corresponding global attribute from output-file. If the name conflict occurs for a non-record variable, then the dimensions and type of the variable (and of its coordinate dimensions, if any) must agree (conform) in both files. Then the variable values (and any coordinate dimension values) from input-file will overwrite the corresponding variable values (and coordinate dimension values, if any) in output-file ${ }^{1}$.

Since there can only be one record dimension in a file, the record dimension must have the same name (but not necessarily the same size) in both files if a record dimension variable is to be appended. If the record dimensions are of differing sizes, the record dimension of output-file will become the greater of the two record dimension sizes, the record variable from input-file will overwrite any counterpart in output-file and fill values will be written to any gaps left in the rest of the record variables (I think). In all cases variable attributes in output-file are superseded by attributes of the same name from input-file, and left alone if there is no name conflict.

Some users may wish to avoid interactive ncks queries about whether to overwrite existing data. For example, batch scripts will fail if ncks does not receive responses to its queries. Options ' -0 ' and ' -A ' are available to force overwriting existing files and variables, respectively.

## Options specific to ncks

The following list provides a short summary of the features unique to ncks. Features common to many operators are described in Chapter 3 [Common features], page 17.

[^17][^18]'-b 'file''
Activate native machine binary output writing to binary file 'file'. Also '--fl_bnr' and '--binary-file'. Writing packed variables in binary format is not supported.
'-d dim,[min][,[max]][,[stride]]'
Add stride argument to hyperslabber. For a complete description of the stride argument, See Section 3.12 [Stride], page 29.
‘-H’ Print data to screen. Also activated using '--print' or '--prn'. Unless otherwise specified (with -s), each element of the data hyperslab is printed on a separate line containing the names, indices, and, values, if any, of all of the variables dimensions. The dimension and variable indices refer to the location of the corresponding data element with respect to the variable as stored on disk (i.e., not the hyperslab).

```
% ncks -H -C -v three_dmn_var in.nc
lat[0]=-90 lev[0]=100 lon[0]=0 three_dmn_var[0]=0
lat[0]=-90 lev[0]=100 lon[1]=90 three_dmn_var[1]=1
lat[0]=-90 lev[0]=100 lon[2]=180 three_dmn_var[2]=2
lat[1]=90 lev[2]=1000 lon[1]=90 three_dmn_var[21]=21
lat[1]=90 lev[2]=1000 lon[2]=180 three_dmn_var[22]=22
lat[1]=90 lev[2]=1000 lon[3]=270 three_dmn_var[23]=23
```

Printing the same variable with the ' $-F$ ' option shows the same variable indexed with Fortran conventions

```
% ncks -F -H -C -v three_dmn_var in.nc
lon(1)=0 lev(1)=100 lat(1)=-90 three_dmn_var(1)=0
lon(2)=90 lev(1)=100 lat(1)=-90 three_dmn_var(2)=1
lon(3)=180 lev(1)=100 lat(1)=-90 three_dmn_var(3)=2
```

Printing a hyperslab does not affect the variable or dimension indices since these indices are relative to the full variable (as stored in the input file), and the input file has not changed. However, if the hypserslab is saved to an output file and those values are printed, the indices will change:

```
% ncks -0 -H -d lat,90.0 -d lev,1000.0 -v three_dmn_var in.nc out.nc
lat[1]=90 lev[2]=1000 lon[0]=0 three_dmn_var[20]=20
lat[1]=90 lev[2]=1000 lon[1]=90 three_dmn_var[21]=21
lat[1]=90 lev[2]=1000 lon[2]=180 three_dmn_var[22]=22
lat[1]=90 lev[2]=1000 lon[3]=270 three_dmn_var[23]=23
% ncks -C -H -v three_dmn_var out.nc
lat[0]=90 lev[0]=1000 lon[0]=0 three_dmn_var[0]=20
lat[0]=90 lev[0]=1000 lon[1]=90 three_dmn_var[1]=21
lat[0]=90 lev[0]=1000 lon[2]=180 three_dmn_var[2]=22
lat[0]=90 lev[0]=1000 lon[3]=270 three_dmn_var[3]=23
'-M' Print to screen the global metadata describing the file. This includes file summary information and global attributes. Also '--Mtd' and '--Metadata'.
```

'-m' Print variable metadata to screen (similar to ncdump -h). This displays all metadata pertaining to each variable, one variable at a time. Also '--mtd' and '--metadata'.
'-q' Toggle printing of dimension indices and coordinate values when printing arrays. The name of each variable will appear flush left in the output. This is useful when trying to locate specific variables when displaying many variables with different dimensions. Also '--quiet'.
'-s format'
String format for text output. Accepts C language escape sequences and printf() formats. Also '--string', '--format', and '--fmt'.
'-u' Accompany the printing of a variable's values with its units attribute, if any. Also '--units'.

## EXAMPLES

View all data in netCDF 'in.nc', printed with Fortran indexing conventions:

```
ncks -H -F in.nc
```

Copy the netCDF file 'in.nc' to file 'out.nc'.
ncks -0 in.nc out.nc
Now the file 'out.nc' contains all the data from 'in.nc'. There are, however, two differences between 'in.nc' and 'out.nc'. First, the history global attribute (see Section 3.17 [History attribute], page 37) will contain the command used to create 'out.nc'. Second, the variables in 'out.nc' will be defined in alphabetical order. Of course the internal storage of variable in a netCDF file should be transparent to the user, but there are cases when alphabetizing a file is useful (see description of -a switch).

Print variable three_dmn_var from file 'in.nc' with default notations. Next print three_dmn_var as an un-annotated text column. Then print three_dmn_var signed with very high precision. Finally, print three_dmn_var as a comma-separated list.

```
% ncks -H -C -v three_dmn_var in.nc
lat[0]=-90 lev[0]=100 lon[0]=0 three_dmn_var[0]=0
lat[0]=-90 lev[0]=100 lon[1]=90 three_dmn_var[1]=1
lat[1]=90 lev[2]=1000 lon[3]=270 three_dmn_var[23]=23
% ncks -s "%f\n" -H -C -v three_dmn_var in.nc
0.000000
1.000000
23.000000
% ncks -s "%+16.10f\n" -H -C -v three_dmn_var in.nc
    +0.0000000000
    +1.0000000000
    +23.0000000000
% ncks -s "%f, " -H -C -v three_dmn_var in.nc
```

$$
0.000000,1.000000, \ldots, 23.000000,
$$

The second and third options are useful when pasting data into text files like reports or papers. See Section 4.2 [ncatted netCDF Attribute Editor], page 48, for more details on string formatting and special characters.

One dimensional arrays of characters stored as netCDF variables are automatically printed as strings, whether or not they are NUL-terminated, e.g.,

```
ncks -v fl_nm in.nc
```

The \%c formatting code is useful for printing multidimensional arrays of characters representing fixed length strings

```
ncks -H -s "%c" -v fl_nm_arr in.nc
```

Using the \%s format code on strings which are not NUL-terminated (and thus not technically strings) is likely to result in a core dump.

Create netCDF 'out.nc' containing all variables, and any associated coordinates, except variable time, from netCDF 'in.nc':

```
ncks -x -v time in.nc out.nc
```

Extract variables time and pressure from netCDF 'in.nc'. If 'out.nc' does not exist it will be created. Otherwise the you will be prompted whether to append to or to overwrite 'out.nc':

```
ncks -v time,pressure in.nc out.nc
ncks -C -v time,pressure in.nc out.nc
```

The first version of the command creates an 'out.nc' which contains time, pressure, and any coordinate variables associated with pressure. The 'out.nc' from the second version is guaranteed to contain only two variables time and pressure.

Create netCDF 'out.nc' containing all variables from file 'in.nc'. Restrict the dimensions of these variables to a hyperslab. Print (with -H) the hyperslabs to the screen for good measure. The specified hyperslab is: the fifth value in dimension time; the half-open range lat $>0$. in coordinate lat; the half-open range lon $<330$. in coordinate lon; the closed interval $0.3<$ band $<0.5$ in coordinate band; and cross-section closest to 1000 . in coordinate lev. Note that limits applied to coordinate values are specified with a decimal point, and limits applied to dimension indices do not have a decimal point See Section 3.8 [Hyperslabs], page 24.

```
ncks -H -d time,5 -d lat,,0.0 -d lon,330.0, -d band,0.3,0.5
-d lev,1000.0 in.nc out.nc
```

Assume the domain of the monotonically increasing longitude coordinate lon is $0<$ lon $<360$. Here, lon is an example of a wrapped coordinate. ncks will extract a hyperslab which crosses the Greenwich meridian simply by specifying the westernmost longitude as min and the easternmost longitude as max, as follows:

```
ncks -d lon,260.0,45.0 in.nc out.nc
```

For more details See Section 3.11 [Wrapped coordinates], page 28.

## 4.8 ncra netCDF Record Averager

## SYNTAX

```
ncra [-A] [-C] [-c] [-D dbg]
[-d dim,[min][,[max]][,[stride]]] [-F] [-h] [-1 path]
[-n loop] [-0] [-p path] [-R] [-r] [-v var[,...]]
[-x] [-y op_typ] input-files output-file
```

DESCRIPTION
ncra averages record variables across an arbitrary number of input files. The record dimension is retained as a degenerate (size 1) dimension in the output variables. See Section 2.6 [Averaging vs. Concatenating], page 13, for a description of the distinctions between the various averagers and concatenators.

Input files may vary in size, but each must have a record dimension. The record coordinate, if any, should be monotonic for (or else non-fatal warnings may be generated). Hyperslabs of the record dimension which include more than one file are handled correctly. ncra supports the stride argument to the '-d' hyperslab option for the record dimension only, stride is not supported for non-record dimensions.
ncra weights each record (e.g., time slice) in the input-files equally. ncra does not attempt to see if, say, the time coordinate is irregularly spaced and thus would require a weighted average in order to be a true time average.

## EXAMPLES

Average files '85.nc', '86.nc', ... '89.nc' along the record dimension, and store the results in '8589.nc':

```
ncra 85.nc 86.nc 87.nc 88.nc 89.nc 8589.nc
ncra 8[56789].nc 8589.nc
ncra -n 5,2,1 85.nc 8589.nc
```

These three methods produce identical answers. See Section 3.2 [Specifying input files], page 18, for an explanation of the distinctions between these methods.

Assume the files '85.nc', '86.nc', .. . '89.nc' each contain a record coordinate time of length 12 defined such that the third record in ' $86 . \mathrm{nc}$ ' contains data from March 1986, etc. nco knows how to hyperslab the record dimension across files. Thus, to average data from December, 1985 through February, 1986:

```
ncra -d time,11,13 85.nc 86.nc 87.nc 8512_8602.nc
ncra -F -d time,12,14 85.nc 86.nc 87.nc 8512_8602.nc
```

The file ' $87 . n c$ ' is superfluous, but does not cause an error. The ' $-F$ ' turns on the Fortran (1-based) indexing convention. The following uses the stride option to average all the March temperature data from multiple input files into a single output file

```
ncra -F -d time,3,,12 -v temperature 85.nc 86.nc 87.nc 858687_03.nc
```

See Section 3.12 [Stride], page 29, for a description of the stride argument.

Assume the time coordinate is incrementally numbered such that January, $1985=1$ and December, $1989=60$. Assuming '??' only expands to the five desired files, the following averages June, 1985-June, 1989:

```
ncra -d time,6.,54. ??.nc 8506_8906.nc
```


## 4.9 ncreat netCDF Record Concatenator

## SYNTAX

```
ncrcat [-A] [-C] [-c] [-D dbg]
[-d dim,[min][,[max]][,[stride]]] [-F] [-h] [-l path]
[-n loop] [-0] [-p path] [-R] [-r] [-v var[,...]]
[-x] input-files output-file
```


## DESCRIPTION

ncreat concatenates record variables across an arbitrary number of input files. The final record dimension is by default the sum of the lengths of the record dimensions in the input files. See Section 2.6 [Averaging vs. Concatenating], page 13, for a description of the distinctions between the various averagers and concatenators.

Input files may vary in size, but each must have a record dimension. The record coordinate, if any, should be monotonic (or else non-fatal warnings may be generated). Hyperslabs of the record dimension which include more than one file are handled correctly. ncra supports the stride argument to the '-d' hyperslab option for the record dimension only, stride is not supported for non-record dimensions.
ncrcat applies special rules to ARM convention time fields (e.g., time_offset). See Section 3.19 [ARM Conventions], page 38 for a complete description.

## EXAMPLES

Concatenate files '85.nc', '86.nc', ... '89.nc' along the record dimension, and store the results in '8589.nc':

```
ncrcat 85.nc 86.nc 87.nc 88.nc 89.nc 8589.nc
ncrcat 8[56789].nc 8589.nc
ncrcat -n 5,2,1 85.nc 8589.nc
```

These three methods produce identical answers. See Section 3.2 [Specifying input files], page 18, for an explanation of the distinctions between these methods.

Assume the files '85.nc', '86.nc', ... '89.nc' each contain a record coordinate time of length 12 defined such that the third record in ' $86 . \mathrm{nc}$ ' contains data from March 1986, etc. nCO knows how to hyperslab the record dimension across files. Thus, to concatenate data from December, 1985-February, 1986:

```
ncrcat -d time,11,13 85.nc 86.nc 87.nc 8512_8602.nc
ncrcat -F -d time,12,14 85.nc 86.nc 87.nc 8512_8602.nc
```

The file ' $87 . n c$ ' is superfluous, but does not cause an error. The ' -F ' turns on the Fortran (1-based) indexing convention.

The following uses the stride option to concatenate all the March temperature data from multiple input files into a single output file

```
ncrcat -F -d time,3,,12 -v temperature 85.nc 86.nc 87.nc 858687_03.nc
```

See Section 3.12 [Stride], page 29, for a description of the stride argument.

Assume the time coordinate is incrementally numbered such that January, $1985=1$ and December, $1989=60$. Assuming ?? only expands to the five desired files, the following concatenates June, 1985-June, 1989:

```
ncrcat -d time,6.,54. ??.nc 8506_8906.nc
```


### 4.10 ncrename netCDF Renamer

## SYNTAX

```
ncrename [-a old_name,new_name] [-a ...] [-D dbg]
[-d old_name,new_name] [-d ...] [-h] [-l path] [-0] [-p path]
[-R] [-r] [-v old_name,new_name] [-v ...]
input-file [output-file]
```


## DESCRIPTION

ncrename renames dimensions, variables, and attributes in a netCDF file. Each object that has a name in the list of old names is renamed using the corresponding name in the list of new names. All the new names must be unique. Every old name must exist in the input file, unless the old name is preceded by the character '.'. The validity of old_name is not checked prior to the renaming. Thus, if old_name is specified without the the '.' prefix and is not present in input-file, ncrename will abort. The new_name should never be prefixed by a '.' (the period will be included as part of the new name). The OPTIONS and EXAMPLES show how to select specific variables whose attributes are to be renamed.
ncrename is the exception to the normal rules that the user will be interactively prompted before an existing file is changed, and that a temporary copy of an output file is constructed during the operation. If only input-file is specified, then ncrename will change the names of the input-file in place without prompting and without creating a temporary copy of inputfile. This is because the renaming operation is considered reversible if the user makes a mistake. The new_name can easily be changed back to old_name by using ncrename one more time.

Note that renaming a dimension to the name of a dependent variable can be used to invert the relationship between an independent coordinate variable and a dependent variable. In this case, the named dependent variable must be one-dimensional and should have no missing values. Such a variable will become a coordinate variable.

According to the netCDF User's Guide, renaming properties in netCDF files does not incur the penalty of recopying the entire file when the new_name is shorter than the old_name.
OPTIONS
'-a old_name,new_name'
Attribute renaming. The old and new names of the attribute are specified by the associated old_name and new_name values. Global attributes are treated no differently than variable attributes. This option may be specified more than once. As mentioned above, all occurrences of the attribute of a given name will be renamed unless the '.' form is used, with one exception. To change the attribute name for a particular variable, specify the old_name in the format old_var_name@old_att_name. The '@' symbol serves to delimit the variable name from the attribute name. If the attribute is uniquely named (no other variables contain the attribute) then the old_var_name@old_att_name syntax is redundant. The var_name@att_name syntax is accepted, but not required, for the new_name.
'-d old_name,new_name'
Dimension renaming. The old and new names of the dimension are specified by the associated old_name and new_name values. This option may be specified more than once.
'-v old_name, new_name'
Variable renaming. The old and new names of the variable are specified by the associated old_name and new_name values. This option may be specified more than once.

## EXAMPLES

Rename the variable $p$ to pressure and $t$ to temperature in netCDF 'in.nc'. In this case p must exist in the input file (or ncrename will abort), but the presence of t is optional:
ncrename $-v$ p,pressure $-v$.t,temperature in.nc
ncrename does not automatically attach dimensions to variables of the same name. If you want to rename a coordinate variable so that it remains a coordinate variable, you must separately rename both the dimension and the variable:
ncrename -d lon,longitude -v lon,longitude in.nc
Create netCDF 'out.nc' identical to 'in.nc' except the attribute _FillValue is changed to missing_value which possess it), the attribute units is renamed to CGS_units (but only in those variables which possess it) and the global attribute Zaire is renamed to Congo:

```
ncrename -a _FillValue,missing_value -a .units,CGS_units \
-a tpt@hieght,height -a prs_sfc@.hieght,height in.nc out.nc
```

The presence and absence of the '.' and '@' features cause this command to execute successfully only if a number of conditions are met. All variables must have a _FillValue attribute and _FillValue must also be a global attribute. The units attribute, on the other hand, will be renamed to CGS_units wherever it is found but need not be present in the file at all (either as a global or a variable attribute). The variable tpt must contain the hieght attribute. The variable prs_sfc need not exist, and need not contain the hieght attribute.

### 4.11 ncwa netCDF Weighted Averager

## SYNTAX

```
ncwa [-A] [-a dim[,...]] [-C] [-c] [-D dbg]
[-d dim,[min][,[max]]] [-F] [-h] [-I] [-l path]
[-M mask_val] [-m mask_var] [-N] [-n] [-0] [-o condition]
[-p path] [-R] [-r] [-v var[,...]] [-W] [-w weight]
[-x] [-y op_typ] input-file output-file
```


## DESCRIPTION

ncwa averages variables in a single file over arbitrary dimensions, with options to specify weights, masks, and normalization. See Section 2.6 [Averaging vs. Concatenating], page 13 , for a description of the distinctions between the various averagers and concatenators. The default behavior of ncwa is to arithmetically average every numerical variable over all dimensions and produce a scalar result. To average variables over only a subset of their dimensions, specify these dimensions in a comma-separated list following '-a', e.g., '-a time, lat,lon'. As with all arithmetic operators, the operation may be restricted to an arbitrary hypserslab by employing the '-d' option (see Section 3.8 [Hyperslabs], page 24). ncwa also handles values matching the variable's missing_value attribute correctly. Moreover, ncwa understands how to manipulate user-specified weights, masks, and normalization options. With these options, ncwa can compute sophisticated averages (and integrals) from the command line.
mask_var and weight, if specified, are broadcast to conform to the variables being averaged. The rank of variables is reduced by the number of dimensions which they are averaged over. Thus arrays which are one dimensional in the input-file and are averaged by ncwa appear in the output-file as scalars. This allows the user to infer which dimensions may have been averaged. Note that that it is impossible for ncwa to make make a weight or mask_var of rank $W$ conform to a var of rank $V$ if $W>V$. This situation often arises when coordinate variables (which, by definition, are one dimensional) are weighted and averaged. ncwa assumes you know this is impossible and so ncwa does not attempt to broadcast weight or mask_var to conform to var in this case, nor does ncwa print a warning message telling you this, because it is so common. Specifying $d b g>2$ does cause ncwa to emit warnings in these situations, however.

Non-coordinate variables are always masked and weighted if specified. Coordinate variables, however, may be treated specially. By default, an averaged coordinate variable, e.g., latitude, appears in output-file averaged the same way as any other variable containing an averaged dimension. In other words, by default ncwa weights and masks coordinate variables like all other variables. This design decision was intended to be helpful but for some applications it may be preferable not to weight or mask coordinate variables just like all other variables. Consider the following arguments to ncwa: -a latitude -w lat_wgt -d latitude, $0 ., 90$. where lat_wgt is a weight in the latitude dimension. Since, by default ncwa weights coordinate variables, the value of latitude in the output-file depends on the weights in lat_wgt and is not likely to be 45.0, the midpoint latitude of the hyperslab. Option '-I' overrides this default behavior and causes ncwa not to weight or mask
coordinate variables ${ }^{1}$. In the above case, this causes the value of latitude in the outputfile to be 45.0, an appealing result. Thus, ' - I' specifies simple arithmetic averages for the coordinate variables. In the case of latitude, '-I' specifies that you prefer to archive the central latitude of the hyperslab over which variables were averaged rather than the area weighted centroid of the hyperslab ${ }^{2}$. The mathematical definition of operations involving rank reduction is given above (see Section 3.14 [Operation Types], page 31).

### 4.11.1 Masking condition

Each $x_{i}$ also has an associated masking weight $m_{i}$ whose value is 0 or 1 (false or true). The value of $m_{i}$ is always 1 unless a mask_var is specified (with '-m'). As noted above, mask_var is broadcast, if possible, to conform to the variable being averaged. In this case, the value of $m_{i}$ depends on the masking condition. As expected, $m_{i}=1$ when the masking condition is true and $m_{i}=0$ otherwise.

The masking condition has the syntax mask_var condition mask_val. Here mask_var is the name of the masking variable (specified with '-m', '--mask-variable', '--mask_variable', '--msk_nm', or '--msk_var'). The truth condition argument (specified with '-o', '--op_rlt', '--cmp', '--compare', or '--op_cmp' may be any one of the six arithmetic comparatives: eq, ne, gt, lt, ge, le. These are the Fortran-style character abbreviations for the logical operations $=, \neq,>,<, \geq, \leq$. The masking condition defaults to eq (equality). The mask_val argument to '-M' (or '--mask-value', or '--msk_val') is the right hand side of the masking condition. Thus for the $i$ 'th element of the hyperslab to be averaged, the masking condition is maski condition mask_val.

Each $x_{i}$ is also associated with an additional weight $w_{i}$ whose value may be user-specified. The value of $w_{i}$ is identically 1 unless the user specifies a weighting variable weight (with '-w', '--weight', or '--wgt_var'). In this case, the value of $w_{i}$ is determined by the weight variable in the input-file. As noted above, weight is broadcast, if possible, to conform to the variable being averaged.
$M$ is the number of input elements $x_{i}$ which actually contribute to output element $x_{j}$. $M$ is also known as the tally and is defined as

$$
M=\sum_{i=1}^{i=N} \mu_{i} m_{i}
$$

$M$ is identical to the denominator of the generic averaging expression except for the omission of the weight $w_{i}$. Thus $M=N$ whenever no input points are missing values or are masked. Whether an element contributes to the output, and thus increments $M$ by one, has more to do with the above two criteria (missing value and masking) than with the numeric value of the element per se. For example, $x_{i}=0.0$ does contribute to $x_{j}$ (assuming the missing_value attribute is not 0.0 and location $i$ is not masked). The value $x_{i}=0.0$ will not change the numerator of the generic averaging expression, but it will change the denominator (unless its weight $w_{i}=0.0$ as well).

[^19]
### 4.11.2 Normalization

ncwa has one switch which controls the normalization of the averages appearing in the output-file. Short option ' -N ' (or long options ' --nmr ' or '--numerator') prevents ncwa from dividing the weighted sum of the variable (the numerator in the averaging expression) by the weighted sum of the weights (the denominator in the averaging expression). Thus ' -N ' tells ncwa to return just the numerator of the arithmetic expression defining the operation (see Section 3.14 [Operation Types], page 31).

## EXAMPLES

Given file '85_0112.nc':

```
netcdf 85_0112 {
dimensions:
        lat = 64 ;
        lev = 18 ;
        lon = 128 ;
        time = UNLIMITED ; // (12 currently)
variables:
        float lat(lat) ;
        float lev(lev) ;
        float lon(lon) ;
        float time(time) ;
        float scalar_var ;
        float three_dmn_var(lat, lev, lon) ;
        float two_dmn_var(lat, lev) ;
        float mask(lat, lon) ;
        float gw(lat) ;
}
```

Average all variables in 'in.nc' over all dimensions and store results in 'out.nc':

```
ncwa in.nc out.nc
```

Every variable in 'in.nc' is reduced to a scalar in 'out.nc' because, by default, averaging is performed over all dimensions.

Store the zonal (longitudinal) mean of 'in.nc' in 'out.nc':

```
ncwa -a lon in.nc out.nc
```

Here the tally is simply the size of lon, or 128 .
Compute the meridional (latitudinal) mean, with values weighted by the corresponding element of $g w^{3}$ :

```
ncwa -w gw -a lat in.nc out.nc
```

Here the tally is simply the size of lat, or 64 . The sum of the Gaussian weights is 2.0 .
Compute the area mean over the tropical Pacific:

[^20]```
ncwa -w gw -a lat,lon -d lat,-20.,20. -d lon,120.,270.
in.nc out.nc
```

Here the tally is $64 \times 128=8192$.
Compute the area mean over the globe, but include only points for which $O R O<0.5^{4}$ :

```
ncwa -m ORO -M 0.5 -o lt -w gw -a lat,lon in.nc out.nc
```

Assuming $70 \%$ of the gridpoints are maritime, then here the tally is $0.70 \times 8192 \approx 5734$.
Compute the global annual mean over the maritime tropical Pacific:

```
ncwa -m ORO -M 0.5 -o lt -w gw -a lat,lon,time
-d lat,-20.0,20.0 -d lon,120.0,270.0 in.nc out.nc
```

Determine the total area of the maritime tropical Pacific, assuming the variable area contains the area of each gridcell

```
ncwa -N -v area -m ORO -M 0.5 -o lt -a lat,lon
-d lat,-20.0,20.0 -d lon,120.0,270.0 in.nc out.nc
```

Weighting area (e.g., by $g w$ ) is not appropriate because area is already area-weighted by definition. Thus the ' -N ' switch, or, equivalently, the ' -y ttl' switch, are all that are needed to correctly integrate the cell areas into a total regional area.

[^21]
## 5 Contributing

We welcome contributions from anyone. The NCO project homepage at https://sf.net/projects/nco contains more information on how to contribute.

Charlie Zender
Concept, design and implementation of NCO from 1995-2000. Since then, mainly packing, NCO library redesign, ncap features, project coordination, code maintenance and porting, documentation, and ncbo.

Henry Butowsky
Non-linear operations and min(), max(), total() support in ncra and ncwa. Type conversion for arithmetic. Migration to netCDF3 API. ncap parser, lexer, and I/O. Multislabbing algorithm. Various hacks.

Rorik Peterson
Autotool build support, long options.
Brian Mays
Packaging for Debian GNU/Linux, nroff man pages.
George Shapovalov
Packaging for Gentoo GNU/Linux.
Bill Kocik Memory management.
Len Makin
NECSX architecture support.
Jim Edwards
AIX architecture support.
Juliana Rew
Compatibility with large PIDs.
Keith Lindsay, Martin Dix
Excellent bug reports.

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[^0]:    1 To produce these formats, 'nco.texi' was simply run through the freely available programs texi2dvi, dvips, texi2html, and makeinfo. Due to a bug in $\mathrm{T}_{\mathrm{E}} \mathrm{X}$, the resulting Postscript file, ' nco . ps', contains the Table of Contents as the final pages. Thus if you print 'nco.ps', remember to insert the Table of Contents after the cover sheet before you staple the manual.

[^1]:    ${ }^{2}$ The '_BSD_SOURCE' token is required on some Linux platforms where gcc dislikes the network header files like 'netinet/in.h').

[^2]:    ${ }^{3}$ The Cygwin package is available from
    http://sourceware.redhat.com/cygwin
    Currently, Cygwin 20.x comes with the gNu C/C++/Fortran compilers (gcc, g++, g77). These GNU compilers may be used to build the netCDF distribution itself.
    ${ }^{4}$ The ldd command, if it is available on your system, will tell you where the executable is looking for each dynamically loaded library. Use, e.g., ldd 'which ncea'.

[^3]:    ${ }^{5}$ The Hierarchical Data Format, or HDF, is another self-describing data format similar to, but more elaborate than, netCDF.
    ${ }^{6}$ One must link the NCO code to the HDF4 MFHDF library instead of the usual netCDF library. However, the MFHDF library only supports netCDF 2.x calls. Thus I will try to keep this capability in NCO as long as it is not too much trouble.

[^4]:    ${ }^{1}$ The ncrename operator is an exception to this rule. See Section 4.10 [ncrename netCDF Renamer], page 70 .

[^5]:    ${ }^{6}$ The exact length which exceeds the operating system internal limit for command line lengths varies from OS to OS and from shell to shell. GNU bash may not have any arbitrary fixed limits to the size of

[^6]:    command line arguments. Many oss cannot handle command line arguments longer than a few thousand characters. When this occurs, the ansi C-standard argc-argv method of passing arguments from the calling shell to a C-program (i.e., an NCO operator) breaks down.

[^7]:    1 If a getopt_long function cannot be found on the system, NCO will use the getopt_long from the my_getopt package by Benjamin Sittler bsittler@iname.com. This is BSD-licensed software available from http://www.geocities.com/ResearchTriangle/Node/9405/\#my_getopt.

[^8]:    ${ }^{2}$ The ' -n ' option is a backward compatible superset of the NINTAP option from the NCAR CCM Processor.

[^9]:    ${ }^{3}$ The msrcp command must be in the user's path and located in one of the following directories: /usr/local/bin, /usr/bin, /opt/local/bin, or /usr/local/dcs/bin.

[^10]:    4 nCO averagers have a bug (TODO 121) which may cause them to behave incorrectly if the missing_value $=$ ' 0.0 ' for a variable to be averaged. The workaround for this bug is to change missing_value to anything besides zero.

[^11]:    ${ }^{6} \quad 32767=2^{15}-1$
    ${ }^{7}$ Operators began performing type conversions before arithmetic in NCO version 1.2, August, 2000. Previous versions never performed unnecessary type conversion for arithmetic.

[^12]:    8 The actual type conversions are handled by intrinsic C-language type conversion, so the floor() function is not explicitly called, but the results are the same as if it were.

[^13]:    9 The exception is appending/altering the attributes $x_{-} o p, y_{-} o p, z_{-} o p$, and $t_{\_} o p$ for variables which have been averaged across space and time dimensions. This feature is scheduled for future inclusion in NCo.
    10 The CSM convention recommends time be stored in the format time since base_time, e.g., the units attribute of time might be 'days since 1992-10-8 15:15:42.5-6:00'. A problem with this format occurs when using ncrcat to concatenate multiple files together, each with a different base_time. That is, any time values from files following the first file to be concatenated should be corrected to the base_time offset specified in the units attribute of time from the first file. The analogous problem has been fixed in ARM files (see Section 3.19 [ARM Conventions], page 38) and could be fixed for CSM files if there is sufficient lobbying, and if Unidata fixes the UDUnits package to build out of the box on Linux.

[^14]:    1 ANSI C compilers are guaranteed to support double precision versions of these functions. These functions normally operate on netCDF variables of type NC_DOUBLE without having to perform intrinsic conversions. For example, ANSI compilers provide sin for the sine of C-type double variables. The ANSI standard does not require, but many compilers provide, an extended set of mathematical functions that apply to single (float) and quadruple (long double) precision variables. Using these functions (e.g., sinf for float, sinl for long double), when available, is more efficient than casting variables to type double, performing the operation, and then recasting. NCO uses the faster intrinsic functions when they are available, and uses the casting method when they are not.
    ${ }^{2}$ Linux supports more of these intrinsic functions than other OSs.

[^15]:    ${ }^{1}$ A naked (i.e., unprotected or unquoted) '*' is a wildcard character. A naked '-' may interpretedconfuse command line parser. ' + ' and '/' are relatively harmless.

[^16]:    2 The widely used shell Bash correctly interprets all these special characters even when they are not quoted. That is, Bash does not prevent NCO from correctly interpreting the intended arithmetic operation when the following arguments are given (without quotes) to ncbo: '--op_typ=+', '--op_typ=-', '--op_typ=*', and '--op_typ=/'
    ${ }^{3}$ This is because ncra collapses the record dimension to a size of 1 (making it a degenerate dimension), but does not remove it, while ncwa removes all dimensions it averages over. In other words, ncra changes the size but not the rank of variables, while ncwa changes both the size and the rank of variables.

[^17]:    '-a' Do not alphabetize extracted fields. By default, the specified output variables are extracted, printed, and written to disk in alphabetical order. This tends to make long output lists easier to search for particular variables. Specifying -a results in the variables being extracted, printed, and written to disk in the order in which they were saved in the input file. Thus -a retains the original ordering of the variables. Also '--abc' and '--alphabetize'.

    ```
    '-B 'file''
    ```

    Activate native machine binary output writing to the default binary file, 'ncks.bnr'. The -B switch is redundant when the -b 'file' option is specified, and native binary output will be directed to the binary file 'file'. Also '--bnr' and '--binary'. Writing packed variables in binary format is not supported.

[^18]:    1 Those familiar with netCDF mechanics might wish to know what is happening here: ncks does not attempt to redefine the variable in output-file to match its definition in input-file, ncks merely copies the values of the variable and its coordinate dimensions, if any, from input-file to output-file.

[^19]:    1 The default behavior of (' ${ }^{-I}$ ') changed on 1998/12/01—before this date the default was not to weight or mask coordinate variables.
    ${ }^{2}$ If lat_wgt contains Gaussian weights then the value of latitude in the output-file will be the areaweighted centroid of the hyperslab. For the example given, this is about 30 degrees.

[^20]:    ${ }^{3}$ gw stands for Gaussian weight in the NCAR climate model.

[^21]:    ${ }^{4}$ ORO stands for Orography in the NCAR climate model. ORO $<0.5$ selects the gridpoints which are covered by ocean.

