Parallel and Large-scale Simulation Enhancements to CGNS

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1 Overview of changes introduced in the HDF5_Parallel branch

Many of the changes discussed in the following sections address the currently (as of version 3.2.1) underperforming parallel capabilities of the CGNS library. For example, the CGNS function cgp_open, which opens a CGNS file for processing, has substantially increasing execution time as the number of processes is increased, Fig. 1 (trunk). The current improvement for cgp_open is substantial at 100-1000 times faster (branch) than the previous implementation (trunk). In fact, for runs with the largest number of processes (>1024) the batch job had a time limit of 5 minutes and not all the processes had completed cgp_open before this limit was reached. Obviously, the previous implementation of cgp_open is a lot worse than reported in the figure.

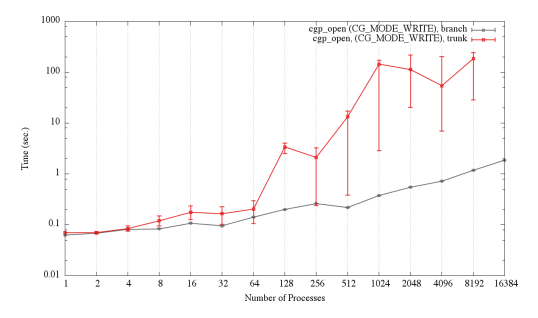


Figure 1: Time for completion of cgp_open in write mode for the original implementation (trunk) and the current implementation (branch), the error bars correspond to the minimum and maximum time over all the processes that had completed before the batch job time limit was reached.

Section 2 lists changes to CGNS that could effect the end user and introduces new functions and specifications. Fortran programmers should take notice of Section 2.2 which highlights changes introduced for better interoperability with the C CGNS library. Section 3 gives example installation guides for GPFS and Lustre hardware. Items listed in blue affect compatibility of older codes when using CGNS v3.2.2. Known problems are highlighted in red. The branch can be checked out from sourceforge at,

http://svn.code.sf.net/p/cgns/code/cgns/branches/HDF5 Parallel

2 General behavior changes and new recommendations for parallel performance

- The flush functions should not be used. Writing and reading immediately avoids IO contention occurring when flush is being used.
- The parallel routines are meant for parallel file systems (GPFS or Lustre).
- The default parallel input/output mode was changed from CGP_INDEPENDENT to CGP_COLLECTIVE.
- An extra argument for passing MPI info to the CGNS library was added to cgp pio mode.

 \mathbf{C}

```
int cgp_pio_mode(CGNS_ENUMT(PIOmode_t) mode, MPI_Info info)
```

Fortran

```
CALL cgp_pio_mode_f(mode, comm_info, ierr)
    INTEGER(KIND(CGP_COLLECTIVE)) :: mode ! Use parameters CGP_INDEPENDENT or CGP_COLLECTIVE
    INTEGER :: comm_info
    INTEGER :: ierr
```

• Functions for parallel reading and writing multi-component datasets using a single call was introduced. The new APIs use new capabilities are tentively to be introduced in version 1.8.15 of the HDF5 library. The new APIs pack multiple datasets into a single buffer and the underlying MPI IO completes the IO request using just one call. The availability of the new functions in the HDF5 library is checked at compile time. The current limitation (due to MPI) is that the size of the sum of the datasets must be less than 2GB. Example usage can be found in benchmark_hdf5.c and benchmark_hdf5_f90.F90 in ptests.

 \mathbf{C}

```
int cgp_coord_multi_read_data(int fn, int B, int Z, int *C,
                              const cgsize_t *rmin, const cgsize_t *rmax,
                              void *coordsX, void *coordsY, void *coordsZ);
int cgp_coord_multi_write_data(int fn, int B, int Z, int *C,
                               const cgsize_t *rmin, const cgsize_t *rmax,
                               const void *coordsX, const void *coordsY, const void *coordsZ);
 int cgp_field_multi_read_data(int fn, int B, int Z, int S, int *F,
                               const cgsize_t *rmin, const cgsize_t *rmax,
                               int nsets, ...);
/* ... nsets of variable arguments, *solution_array, corresponding to the order given by F */
int cgp_field_multi_write_data(int fn, int B, int Z, int S, int *F,
                               const cgsize_t *rmin, const cgsize_t *rmax,
                               int nsets, ...);
/* ... nsets of variable arguments, *solution_array, corresponding to the order given by F */
int cgp_array_multi_write_data(int fn, int *A, const cgsize_t *rmin, const cgsize_t *rmax,
                               int nsets, ...);
/* ... nsets of variable arguments, *field_array, corresponding to the order given by F */
int cgp_array_multi_read_data(int fn, int *A, const cgsize_t *rmin,const cgsize_t *rmax,
                              int nsets, ...);
/* ... nsets of variable arguments, *field_array, corresponding to the order given by F */
```

Fortran

```
CALL cgp_coord_multi_read_data_f(fn, B, Z, C, rmin, rmax, coordsX, coordsY, coordsZ, ier)
  INTEGER :: fn
  INTEGER :: B
  INTEGER :: Z
  INTEGER :: C
  INTEGER(CG_SIZE_T) :: rmin
  INTEGER(CG_SIZE_T) :: rmax
  REAL :: coordsX, coordsY, coordsZ
  INTEGER :: ier
CALL cgp_coord_multi_write_data_f(fn, B, Z, C, rmin, rmax, coordsX, coordsY, coordsZ, ier)
  INTEGER :: fn
  INTEGER :: B
  INTEGER :: Z
  INTEGER :: C
  INTEGER(CG_SIZE_T) :: rmin
  INTEGER(CG_SIZE_T) :: rmax
  REAL :: coordsX, coordsY, coordsZ
  INTEGER :: ier
CALL cgp_field_multi_write_data_f(fn, B, Z, S, F, rmin, rmax, ier, nsets, |...|)
  INTEGER :: fn
  INTEGER :: B
  INTEGER :: Z
  INTEGER :: C
  INTEGER(CG_SIZE_T) :: rmin
  INTEGER(CG_SIZE_T) :: rmax
  INTEGER :: ier
  INTEGER :: nsets
   ... REAL, DIMENSION(*) :: field_array ! entered nsets times
CALL cgp_field_multi_read_data_f(fn, B, Z, S, F, rmin, rmax, ier, nsets, |...|)
   INTEGER :: fn
  INTEGER :: B
  INTEGER :: Z
  INTEGER :: C
  INTEGER(CG_SIZE_T) :: rmin
  INTEGER(CG_SIZE_T) :: rmax
  INTEGER :: ier
  INTEGER :: nsets
   ... REAL, DIMENSION(*) :: field_array ! entered nsets times
```

```
CALL cgp_array_multi_write_data_f(fn, B, Z, S, F, rmin, rmax, ier, nsets, |...)
   INTEGER :: fn
   INTEGER :: B
  INTEGER :: Z
   INTEGER :: C
   INTEGER(CG_SIZE_T) :: rmin
   INTEGER(CG_SIZE_T) :: rmax
   INTEGER :: ier
   INTEGER :: nsets
   ... REAL, DIMENSION(*) :: data_array ! entered nsets times
CALL cgp_array_multi_read_data_f(fn, B, Z, S, F, rmin, rmax, ier, nsets, ...)
   INTEGER :: fn
   INTEGER :: B
   INTEGER :: Z
   INTEGER :: C
   INTEGER(CG_SIZE_T) :: rmin
   INTEGER(CG_SIZE_T) :: rmax
   INTEGER :: ier
   INTEGER :: nsets
   ... REAL, DIMENSION(*) :: data_array ! entered nsets times
```

2.1 New C changes

• A new parallel example benchmark program, benchmark hdf5.c, was added to directory ptests.

2.1.1 Internal library changes

- 1. Fixed issue with autotools putting a blank "-l" in "MPILIBS =" when compiling library using using mpi.
- 2. Replaced the hid_t to double (and vice-versa) utilities to_HDF_ID and to_ADF_ID from a type cast to a function which uses memcpy for the conversion. This is need to for the upcomming release of HDF5 1.10 where hid_t was changed from a 32 bit integer to a 64 bit integer.

2.2 New Fortran changes

All users are **strongly** encouraged to use a Fortran 2003 standard compliant compiler. Using a Fortran 2003 compiler guarantees interoperability with the C APIs via the ISO_C_BINDING module. Many changes were added to the CGNS library in order to take full advantage of the interoperability offered by the ISO_C_BINDING module.

- 1. Configure was changed to check if the Fortran compiler is Fortran 2003 compliant. If it is then the features of ISO_C_BINDING will be used.
- 2. The predefined CGNS constant parameters data types were changed from INTEGER to ENUM, BIND(C) for better C interoperability. The users should use the predefined constants whenever possible and not the numerical value represented by the constants.
- 3. INCLUDE "cgslib" h" was changed in favor of using a module, USE CGNS.
 - (a) This allows defining a KIND type for integers instead of the current way of using the preprocessor dependent *cgsize t*.
 - (b) Backward compatibility might be added before the merge to the trunk.

- 4. The user should be sure to declare the arguments declared *int* in the C APIs as INTEGER in Fortran. The ONLY Fortran arguments declared as type *cgsize_t* should be the arguments which are also declared *cgsize_t* in the C APIs. This is very important when building with option *-enable-64bit*.
- 5. Assuming the rules in step 4 were followed, users should not need to use parameter CG_BUILD_64BIT since Fortran's cgsize_t is now guaranteed to match C's cgsize_t.
- 6. Fortran programs defining CGNS data types with a default INTEGER size of 8 bytes are not currently compatible. This is independent of whether or not -enable-64bit is being used. For clarification, using -enable-64bit allows for data types (i.e. those declared as cgsize_t) to be able to store values which are too large to be stored as 4 byte integers (i.e. numbers greater than 2,147,483,647). It is not necessary, or advisable, to have CGNS INTEGER types (types declared int in C) to be 8 bytes; the variables declared as cgsize_t will automatically handle data types that can not be stored as 4 byte integers when -enable-64bit is being used.
 - (a) CGNS developer's note: A new C data type, cgint_f, was introduced to be interpretable with the C type int. In order to allow for default 8 byte integers in Fortran: (1) The C API wrappers in cg_ftoc.c need to be changed from cgsize_t to cgint_f everywhere the C argument is declared as an int in C, (2) configure needs to detect what size the default integer is in Fortran and find the corresponding size in C in order to set the correct size of cgint_f.
- 7. Two new benchmarking programs were introduced in directory ptests:
 - (a) benchmarking hdf5 f90.F90 uses the conventional Fortran wrappers.
 - (b) benchmarking hdf5 f03.F90 calls the C APIs directly, no Fortran wrappers are used.
- 8. A new Fortran API was added for determining the CGNS data type of a variable which is interoperable with the C data type.

```
Function cg_get_type(var)
  type, INTENT(IN) :: var
  INTEGER(KIND(enumvar)) :: cg_get_type
```

An example of using the new function to automatically specify the CGNS type corresponding to the Fortran data type is,

```
INTEGER, DIMENSION(1:10) :: Array_i

CALL cgp_array_write_f("ArrayI",cg_get_type(Array_i(1)),1,INT(nijk(1),cgsize_t),Ai, err)
```

3 Parallel installation instructions

Two parallel files systems were investigated: GPFS (mira, Argonne National Laboratory) and Lustre (Pleiades NASA). The following descriptions were for those systems, but the overall procedure should be similar on different machines of the same type. Example build scripts for these systems can be found in src/SampleScripts of the CGNS source code. They include scripts for building zlib, hdf5 (assuming the user does not already have them installed system wide) and a script for building CGNS. All the scripts use autotools; cmake remains untested. The next few examples assume all the needed packages are in \${HOME}/packages and all the build scripts are placed in \${HOME}/packages. This information can also be found in the README.txt in the scripts directory.

3.1 Building on IBM Blue Gene (GPFS)

- 1. Building zlib from source: Download and extract the zlib source: http://www.zlib.net/
 - (a) cd into the top level zlib source directory.
 - (b) modify and run the script: ../build zlib
- 2. Building hdf5 from source
 - (a) From the top level of the hdf5 library, change the \${HOME}/packages to where zlib was installed in STEP 1.
 - (b) ../build_hdf5 -without-pthread -disable-shared -enable-parallel -enable-production \ -enable-fortran enable-fortran2003 \ -disable-stream-vfd -disable-direct-vfd \ -with-zlib= ${HOME}/packages/zlib-1.2.8/lib-prefix={HOME}/packages/phdf5-trunk$

where prefix is set for where the hdf5 library will get installed. There should be no need to modify the script.

- 3. Building cgns from source:
 - (a) cd into the cgns/src directory
 - (b) modify and run: <pathto>/build cgns
 - (c) make
 - (d) To make the tests: cd ptests; make;make tests
- 4. Important parameters for good performance on GPFS:
 - (a) The environment variable BGLOCKLESSMPIO_F_TYPE=0x47504653 should be set. For example, this can be set in a batch job using qsub—env BGLOCKLESSMPIO_F_TYPE=0x47504653

3.2 Building on SGI (Lustre)

- 1. Building zlib from source: Download and extract the zlib source: http://www.zlib.net/
 - (a) cd into the top level zlib source directory.
 - (b) modify and run the script: ../build zlib
- 2. Building hdf5 from source:
 - (a) From the top level of the hdf5 library, change the \${HOME}/packages to where zlib was installed in STEP 1.
 - (b) ../build_hdf5
- 3. Building cgns from source:
 - (a) cd into the cgns/src directory
 - (b) modify and run: <pathto>/build cgns
 - (c) make
 - (d) To make the tests: cd ptests; make; make tests
- 4. Important parameters for good performance:
 - (a) The Lustre parameters have not been fully tested.
 - (b) On Pleiades, If setstripe -c 64 -s 0 /nobackupp8/<dir>, has shown good performance.

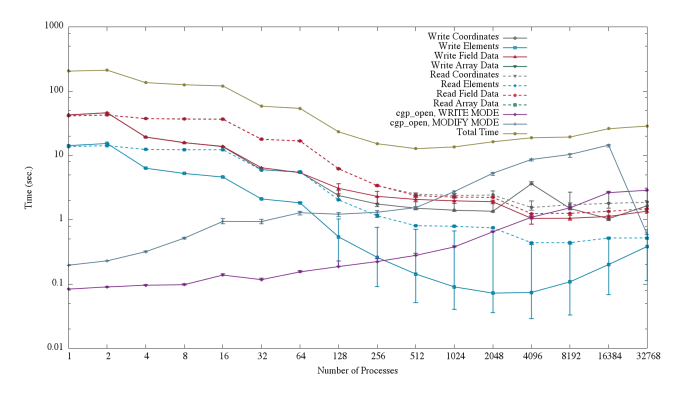


Figure 2: Results for benchmark_hdf5.c on GPFS (cetus, ANL).

3.3 Parallel performace results

The following results are for the benchmark_hdf5* programs found in the ptests directory. The benchmark simulates writing and then reading a ~33.5 million 6-node pentahedra elements mesh with ~201 million nodes. The benchmark results from benchmark_hdf5.c show improvement in cgp_open for up to 32,768 processes, Fig. 2 over the previous implementation. A comparison of benchmark_hdf5_f90.F90 and benchmark_hdf5_f03.F90 also shows negligible performance differences when using the Fortran 90 wrapper routines (benchmark_hdf5_f90.F90) and when calling the C CGNS APIs directly (benchmark_hdf5_f03.F90), Fig. 3. Additionally, negligible difference in performance and scaling exists when calling CGNS from Fortran and C, Fig. 4.

The benchmark results from benchmark_hdf5.c on the Lustre file system (Pleiades, NASA) is presented in Fig. 5. The system defaults for the Lustre file system were used along with a strip count of 64 and a strip size of 4 MB. Overall the Lustre file system appears to be faster at reading a writing, but the GPFS scaled better for the benchmark.

Acknowledgments

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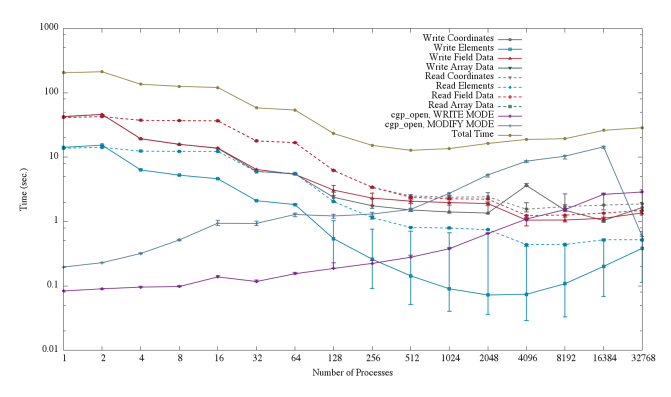


Figure 3: Results for benchmark_hdf5_f03.F90 (shown in color) and benchmark_hdf5_f90.F90 (shown in grey) on GPFS (cetus, ANL).

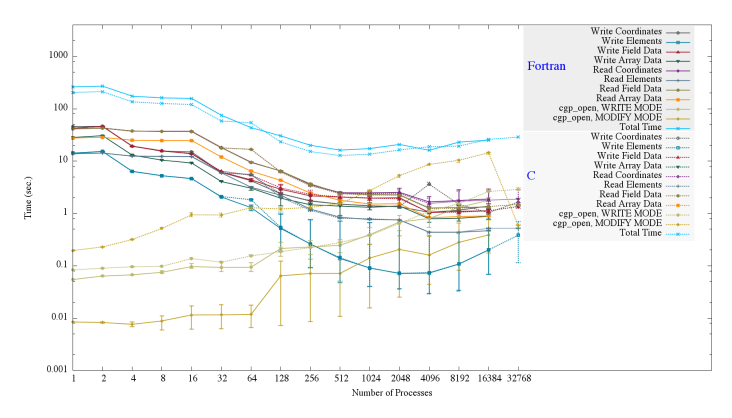


Figure 4: Comparison of Fortran (benchmark_hdf5_f03.F90, solid lines) and C (benchmark_hdf5.c, dashed lines) on GPFS (cetus, ANL).

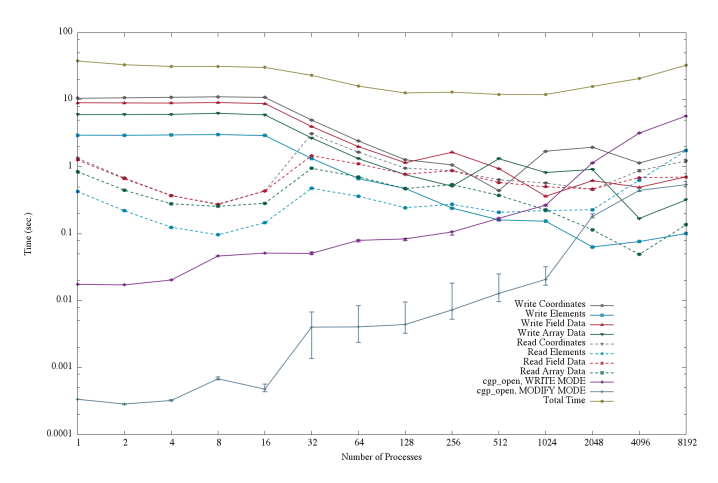


Figure 5: Results for $benchmark_\,hdf5.c$ on Lustre file system (Pleiades, NASA).