Parallel Fast Fourier Transforms

Theory, Methods and Libraries.
A small introduction.

HPC Numerical Libraries
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Massimiliano Guarrasi
m.guarrasi@cineca.it
Agenda

- Introduction to F.T.:
  - Theorems
  - DFT
  - FFT

- Parallel Domain Decomposition
  - Slab Decomposition
  - Pencil Decomposition

- Some Numerical libraries:
  - FFTW
    - Some useful commands
    - Some Examples
  - 2Decomp&FFT
    - Some useful commands
    - Example
  - P3DFFT
    - Some useful commands
    - Example
  - Performance results
Introduction to Fourier Transforms
Fourier Transforms

\[ H(f) = \int_{-\infty}^{\infty} h(t) e^{2\pi jft} \, dt \]

\[ h(t) = \int_{-\infty}^{\infty} H(f) e^{-2\pi jft} \, df \]

Frequency Domain  \leftrightarrow  Time Domain

Real Space  \leftrightarrow  Reciprocal Space
Some useful Theorems

**Convolusion Theorem**

\[ g(t) * h(t) \Leftrightarrow G(f) \cdot H(f) \]

**Correlation Theorem**

\[ Corr(g, h) \equiv \int_{-\infty}^{+\infty} g(\tau)h(t + \tau)d\tau \Leftrightarrow G(f)H^*(f) \]

**Power Spectrum**

\[ Corr(h, h) \equiv \int_{-\infty}^{+\infty} h(\tau)h(t + \tau)d\tau \Leftrightarrow |H(f)|^2 \]
Discrete Fourier Transform (DFT)

In many application contexts the Fourier transform is approximated with a Discrete Fourier Transform (DFT):

\[
H(f_n) = \int_{-\infty}^{\infty} h(t) e^{2\pi i f_n t} \, dt \approx \sum_{k=0}^{N-1} h_k e^{2\pi i f_n k / N} \Delta = \Delta \sum_{k=0}^{N-1} h_k e^{2\pi i f_n k / N}
\]

\[
\begin{align*}
t_k &= \Delta k / N \\
f_n &= n / \Delta
\end{align*}
\]

The last expression is periodic, with period N. It define a between 2 sets of numbers, \( H_n \) & \( h_k \) (\( H(f_n) = \Delta H_n \))
Discrete Fourier Transforms (DFT)

\[ H_n = \sum_{k=0}^{N-1} h_k e^{2\pi i kn/N} \]

\[ h_k = \frac{1}{N} \sum_{n=0}^{N-1} H_n e^{-2\pi i kn/N} \]

frequencies from 0 to \( fc \) (maximum frequency) are mapped in the values with index from 0 to \( N/2-1 \), while negative ones are up to -\( fc \) mapped with index values of \( N / 2 \) to \( N \)

Scale like \( N*^N \)
Fast Fourier Transform (FFT)

The DFT can be calculated very efficiently using the algorithm known as the FFT, which uses symmetry properties of the DFT's

\[ F_k = \sum_{j=0}^{N-1} e^{2\pi i j k/N} f_j \]

\[ = \sum_{j=0}^{N/2-1} e^{2\pi i k (2j)} f_{2j} + \sum_{j=0}^{N/2-1} e^{2\pi i k (2j+1)} f_{2j+1} \]

\[ = \sum_{j=0}^{N/2-1} e^{2\pi i k j/(N/2)} f_{2j} + W^k \sum_{j=0}^{N/2-1} e^{2\pi i k j/(N/2)} f_{2j+1} \]

\[ = F_k^e + W^k F_k^o \]
Fast Fourier Transform (FFT)

\[ F_k = F_k^e + W^k F_k^o \]

- \( F_k \): DFT of even terms
- \( F_k^e \): DFT of even terms
- \( F_k^o \): DFT of odd terms
- \( W^k = \exp(2\pi i/N) \)
Now Iterate:

\[ F_e = F_{ee} + W^{k/2} F_{eo} \]

\[ F_o = F_{oe} + W^{k/2} F_{oo} \]

You obtain a series for each value of \( f_n \)

\[ F_{oeoeoeeoeoeoe} = f_n \]

Scale like \( N \cdot \log N \) (binary tree)
Parallel Domain Decomposition

How to compute a FFT on a distributed memory system
On a 1D array:
  ◦ Algorithm limits:
    • All the tasks must know the whole initial array
    • No advantages in using distributed memory systems
  ◦ Solutions:
    • Using OpenMP it is possible to increase the performance on shared memory systems

On a Multi–Dimensional array:
  ◦ It is possible to use distributed memory systems
Multi-dimensional FFT (an example)

\[ H(n_1, n_2) = \text{FFT-on-index-1} \left( \text{FFT-on-index-2} \left[ h(k_1, k_2) \right] \right) \]

\[ = \text{FFT-on-index-2} \left( \text{FFT-on-index-1} \left[ h(k_1, k_2) \right] \right) \]

1) For each value of \( j \) and \( k \)
Apply FFT to \( h(1...N,j,k) \)

2) For each value of \( i \) and \( k \)
Apply FFT to \( h(i,1...N,k) \)

3) For each value of \( i \) and \( j \)
Apply FFT to \( h(i,j,1...N) \)
Distribute data along one coordinate (e.g. \( Z \))

This is known as "Slab Decomposition" or 1D Decomposition
Transform along x and y

Each processor transforms its own sub-grid along the x and y independently of the other.
Data redistribution involving x and z

The data are now distributed along x
FFT along $z$

each processor transform its own sub-grid

along the $z$ dimension independently of the other
Data are re-distributed, back from x to z

The 3D array now has the original layout, but each element
Has been substituted with its FFT.
Limit of Slab Decomposition

- **Pro:**
  - Simply to implement
  - Moderate communications

- **Con:**
  - Parallelization only along one direction
  - Maximum number of MPI tasks bounded by the size of the larger array index

- **Possible Solutions:**
  - 2D (Pencil) Decomposition
2D Domain Decomposition
Slab vs Pencil Decomposition

- Slab (1D) decomposition:
  - Faster on a limited number of cores
  - Parallelization is limited by the length of the largest axis of the 3D data array used

- Pencil (2D) decomposition:
  - Faster on massively parallel supercomputers
  - Slower using large size arrays on a moderate number of cores (more MPI communications)
Some useful papers

- **Auto-tuning of the FFTW Library for Massively Parallel Supercomputers.**
  - M. Guarrasi, G. Erbacci, A. Emerson;
  - 2012, PRACE white paper;
  - Available at [this link](#);

- **Scalability Improvements for DFT Codes due to the Implementation of the 2D Domain Decomposition Algorithm.**
  - M. Guarrasi, S. Frigio, A. Emerson, G. Erbacci
  - 2013, PRACE white paper;
  - Available at [this link](#);

- **Testing and Implementing Some New Algorithms Using the FFTW Library on Massively Parallel Supercomputers.**
  - M. Guarrasi, N. Li, S. Frigio, A. Emerson, G. Erbacci;
  - Accepted for ParCo 2013 conference proceedings.

- **2DECOMP&FFT – A highly scalable 2D decomposition library and FFT interface.**
  - N. Li, S. Laizet;
  - 2010, Cray User Group 2010 conference;
  - Available at [this link](#);

- **P3DFFT: a framework for parallel computations of Fourier transforms in three dimensions.**
  - D. Pekurovsky;

- **The Design and Implementation of FFTW3.**
  - M. Frigio, S. G. Johnson;
  - 2005, Proceedings of the IEEE.
FFT Numerical Libraries

The simplest way to compute a FFT on a modern HPC system
Introduction

FFTW is a C subroutine library for computing the Discrete Fourier Transform (DFT) in one or more dimensions, of both real and complex data, and of arbitrary input size. We believe that FFTW, which is free software, should become the FFT library of choice for most applications. Our benchmarks, performed on a variety of platforms, show that FFTW's performance is typically superior to that of other publicly available FFT software. Moreover, FFTW's performance is portable: the program will perform well on most architectures without modification.

It is difficult to summarize in a few words all the complexities that arise when testing many programs, and there is no "best" or "fastest" program. However, FFTW appears to be the fastest program most of the time for in-order transforms, especially in the multi-dimensional and real-complex cases (Kasparov is the best chess player in the world even though he loses some games). Hence the name, "FFTW," which stands for the somewhat whimsical title of "Fastest Fourier Transform in the West." Please visit the benchFFT home page for a more extensive survey of the results.

The FFTW package was developed at MIT by Matteo Frigo and Steven G. Johnson.
• Written in C

• Fortran wrapper is also provided

• FFTW adapt itself to your machines, your cache, the size of your memory, the number of register, etc...

• FFTW doesn't use a fixed algorithm to make DFT
  • FFTW chose the best algorithm for your machines

• Computation is split in 2 phases:
  • PLAN creation
  • Execution

• FFTW support transforms of data with arbitrary length, rank, multiplicity, and memory layout, and more....
Many different versions:

**FFTW 2:**
- Released in 2003
- Well tested and used in many codes
- Includes serial and parallel transforms for both shared and distributed memory system

**FFTW 3:**
- Released in February 2012
- Includes serial and parallel transforms for both shared and distributed memory system
- Hybrid implementation MPI-OpenMP
- Last version is FFTW 3.3.3
Some Useful Instructions
How can I compile a code that uses FFTW on PLX?

• Module Loading:

\[
\text{module load autoload fftw/3.3.3--openmpi--1.6.3--intel--cs-xe-2013--binary}
\]

• Including header:

• \(-\$\text{FFTW}\_\text{INC}\)

• Linking:

\[
\begin{align*}
\text{-L}\$\text{FFTW}\_\text{LIB} & -l\text{fftw3_mpi} & -l\text{fftw3_omp} & -l\text{fftw3f} & -l\text{m} & \text{ (single precision)} \\
\text{-L}\$\text{FFTW}\_\text{LIB} & -l\text{fftw3_mpi} & -l\text{fftw3_omp} & -l\text{fftw3} & -l\text{m} & \text{ (double precision)}
\end{align*}
\]

• An example:

\[$ \text{mpif90 -O3 -I}\$\text{FFTW}\_\text{INC} \text{ example.F90 -L}\$\text{FFTW}\_\text{LIB} \text{.fftw3_mpi -lfftw3_omp .fftw3 -lm} \]$
How can I compile a code that uses FFTW on FERMI?

• Module Loading:

```
module load autoload fftw/3.3.2--bgq-gnu--4.4.6
```

• Including header:

```
-I$FFTW3_INC
```

• Linking:

```
-L$FFTW3_LIB -lfftw3f_mpi -lfftw3f_omp -lfftw3f -lm (single precision)
-L$FFTW3_LIB -lfftw3_mpi -lfftw3_omp -lfftw3 -lm (double precision)
```

• An example:

```
$ mpif90 -O3 -I$FFTW3_INC example.F90 -L$FFTW3_LIB .lfftw3_mpi .lfftw3_omp .lfftw3 -lm
```
Some important Remarks for FORTRAN users

- Function in C became function in FORTRAN if they have a return value, and subroutines otherwise.
- All C types are mapped via the iso_c_binning standard.
- FFTW plans are type(C_PTR) in FORTRAN.
- The ordering of FORTRAN array dimensions must be reversed when they are passed to the FFTW plan creation.
Initialize FFTW

Including FFTW Lib:
• C:
  • Serial:
    
    \#include <fftw.h>
  
  • MPI:
    
    \#include <fftw-mpi.h>

• FORTRAN:
  • Serial:
    
    include 'fftw3.f03'
  
  • MPI:
    
    include 'fftw3-mpi.f03'

MPI initialization:
• C:
  
  void fftw_mpi_init(void)

• FORTRAN:
  
  fftw_mpi_init()
Array creation

C:

• Fixed size array:
  
  ```c
  fftx_complex data[n0][n1][n2]
  ```

• Dynamic array:
  
  ```c
  data = fftw_alloc_complex(n0*n1*n2)
  ```

• MPI dynamic arrays:
  
  ```c
  fftw_complex *data
  ptrdiff_t alloc_local, local_no, local_no_start
  alloc_local = fftw_mpi_local_size_3d(n0, n1, n2, MPI_COMM_WORLD, &local_n0,&local_n0_start)
  data = fftw_alloc_complex(alloc_local)
  ```

FORTRAN:

• Fixed size array (simplest way):
  
  ```fortran
  complex(C_DOUBLE_COMPLEX), dimension(n0,n1,n2) :: data
  ```

• Dynamic array (simplest way):
  
  ```fortran
  complex(C_DOUBLE_COMPLEX), allocatable, dimension(:,:,:) :: data
  allocate (data(n0, n1, n2))
  ```

• Dynamic array (fastest method):
  
  ```fortran
  complex(C_DOUBLE_COMPLEX), pointer :: data(:,:,:)
  type(C_PTR) :: cdata
  cdata = fftw_alloc_complex(n0*n1*n2)
  call c_f_pointer(cdata, data, [n0,n1,n2])
  ```

• MPI dynamic arrays:
  
  ```fortran
  complex(C_DOUBLE_COMPLEX), pointer :: data(:,:,:)
  type(C_PTR) :: cdata
  integer(C_INTPTR_T) :: alloc_local, local_n2, local_n2_offset
  alloc_local = fftw_mpi_local_size_3d(n2, n1, n0, MPI_COMM_WORLD, local_n2, local_n2_offset)
  cdata = fftw_alloc_complex(alloc_local)
  call c_f_pointer(cdata, data, [n0,n1,local_n2])
  ```
Plan Creation (C2C)

1D Complex to complex DFT:
• C:
  ```c
  fftw_plan = fftw_plan_dft_1d(int nx, fftw_complex *in, fftw_complex *out, fftw_direction dir, unsigned flags)
  ```
• FORTRAN:
  ```fortran
  plan = ftw_plan_dft_1d(nx, in, out, dir, flags)
  ```

2D Complex to complex DFT:
• C:
  ```c
  fftw_plan = fftw_plan_dft_2d(int nx, int ny, fftw_complex *in, fftw_complex *out, fftw_direction dir, unsigned flags)
  ```
  ```c
  fftw_plan = fftw_mpi_plan_dft_2d(int nx, int ny, fftw_complex *in, fftw_complex *out, MPI_COMM_WORLD, fftw_direction dir, int flags)
  ```
• FORTRAN:
  ```fortran
  plan = ftw_plan_dft_2d(nx, ny, in, out, dir, flags)
  ```
  ```fortran
  plan = ftw_mpi_plan_dft_2d(ny, nx, in, out, MPI_COMM_WORLD, dir, flags)
  ```

3D Complex to complex DFT:
• C:
  ```c
  fftw_plan = fftw_plan_dft_3d(int nx, int ny, int nz, fftw_complex *in, fftw_complex *out, fftw_direction dir, unsigned flags)
  ```
  ```c
  fftw_plan = fftw_mpi_plan_dft_3d(int nx, int ny, int nz, fftw_complex *in, fftw_complex *out, MPI_COMM_WORLD, fftw_direction dir, int flags)
  ```
• FORTRAN:
  ```fortran
  plan = ftw_plan_dft_3d(nx, ny, in, out, dir, flags)
  ```
  ```fortran
  plan = ftw_mpi_plan_dft_3d(nz, ny, nx, in, out, MPI_COMM_WORLD, dir, flags)
  ```
Plan Creation (R2C)

1D Real to complex DFT:
• C:
  
  ```c
  fftw_plan = fftw_plan_dft_r2c_1d(int nx, double *in, fftw_complex *out, fftw_direction dir, unsigned flags)
  ```

  ```fortran
  ftw_plan_dft_r2c_1d(nx, in, out, dir, flags)
  ```

2D Real to complex DFT:
• C:
  
  ```c
  fftw_plan = fftw_plan_dft_r2c_2d(int nx, int ny, double *in, fftw_complex *out, fftw_direction dir, unsigned flags)
  ```

  ```c
  fftw_plan = fftw_mpi_plan_dft_r2c_2d(int nx, int ny, double *in, fftw_complex *out, MPI_COMM_WORLD, fftw_direction dir, int flags)
  ```

  ```fortran
  ftw_plan_dft_r2c_2d(ny, nx, in, out, dir, flags)
  ```

  ```fortran
  ftw_mpi_plan_dft_r2c_2d(ny, nx, in, out, MPI_COMM_WORLD, dir, flags)
  ```

3D Real to complex DFT:
• C:
  
  ```c
  fftw_plan = fftw_plan_dft_r2c_3d(int nx, int ny, int nz, fftw_complex *in, fftw_complex *out, fftw_direction dir, unsigned flags)
  ```

  ```c
  fftw_plan = fftw_mpi_plan_dft_r2c_3d(int nx, int ny, int nz, fftw_complex *in, fftw_complex *out, MPI_COMM_WORLD, fftw_direction dir, int flags)
  ```

  ```fortran
  ftw_plan_dft_r2c_3d(nz, ny, nx, in, out, dir, flags)
  ```

  ```fortran
  ftw_mpi_plan_dft_r2c_3d(nz, ny, nx, in, out, MPI_COMM_WORLD, dir, flags)
  ```
Plan Execution

**Complex to complex DFT:**

- **C:**
  
  ```c
  void fftw_execute_dft(fftw_plan plan, fftw_complex *in, fftw_complex *out)
  void fftw_mpi_execute_dft (fftw_plan plan, fftw_complex *in, fftw_complex *out)
  ```

- **FORTRAN:**
  
  ```fortran
  fftw_execute_dft (plan, in, out)
  fftw_mpi_execute_dft (plan, in, out)
  ```

**Real to complex DFT:**

- **C:**
  
  ```c
  void fftw_execute_dft (fftw_plan plan, double *in, fftw_complex *out)
  void fftw_mpi_execute_dft (fftw_plan plan, double *in, fftw_complex *out)
  ```

- **FORTRAN:**
  
  ```fortran
  fftw_execute_dft (plan, in, out)
  Fftw_mpi_execute_dft (plan, in, out)
  ```
Finalizing FFTW

Destroying PLAN:
• C:
  void fftw_destroy_plan(fftw_plan plan)
• FORTRAN:
  fftw_destroy_plan(plan)

FFTW MPI cleanup:
• C:
  void fftw_mpi_cleanup ()
• FORTRAN:
  fftw_mpi_cleanup ()

Deallocate data:
• C:
  void fftw_free (fftw_complex data)
• FORTRAN:
  fftw_free (data)
Some Useful Examples
program FFTW1D
    use, intrinsic :: iso_c_binding
    implicit none
    include 'fftw3.f03'
    integer(C_INTPTR_T) :: L = 1024
    integer(C_INT) :: LL
    type(C_PTR) :: plan1
    complex(C_DOUBLE_COMPLEX), dimension(1024) :: idata, odata
    integer :: i
    character(len=41), parameter :: filename='serial_data.txt'
    LL = int(L,C_INT)

    !! create MPI plan for in-place forward DF
    plan1 = fftw_plan_dft_1d(LL, idata, odata, FFTW_FORWARD, FFTW_ESTIMATE)

    !! initialize data
    do i = 1, L
        if (i .le. (L/2)) then
            idata(i) = (1.,0.)
        else
            idata(i) = (0.,0.)
        endif
    end do

    !! compute transform (as many times as desired)
    call fftw_execute_dft(plan1, idata, odata)

    !! deallocate and destroy plans
    call fftw_destroy_plan(plan1)
end
program FFTW1D
use, intrinsic :: iso_c_binding
implicit none
include 'fftw3.f03'
integer(C_INTPTR_T):: L = 1024
integer(C_INT) :: LL
type(C_PTR) :: plan1
type(C_PTR) :: p_idata, p_odata
complex(C_DOUBLE_COMPLEX), dimension(:), pointer :: idata, odata
integer :: i
!! Allocate
LL = int(L,C_INT)
p_idata = fftw_alloc_complex(L)
p_odata = fftw_alloc_complex(L)
call c_f_pointer(p_idata,idata,(/L/))
call c_f_pointer(p_odata,odata,(/L/))
!! create MPI plan for in-place forward DF
plan1 = fftw_plan_dft_1d(LL, idata, odata, FFTW_FORWARD, FFTW_ESTIMATE)
!! initialize data
do i = 1, L
  if (i .le. (L/2)) then
    idata(i) = (1.,0.)
  else
    idata(i) = (0.,0.)
  endif
end do
!! compute transform (as many times as desired)
call fftw_execute_dft(plan1, idata, odata)
!! deallocate and destroy plans
call fftw_destroy_plan(plan1)
call fftw_free(p_idata)
call fftw_free(p_odata)
end
# include <stdlib.h>
# include <stdio.h>
# include <math.h>
# include <fftw3.h>

int main ( void )
{
    ptrdiff_t i;
    const ptrdiff_t n = 1024;
    fftw_complex *in;
    fftw_complex *out;
    fftw_plan plan_forward;
    /* Create arrays. */
    in = fftw_malloc ( sizeof ( fftw_complex ) * n );
    out = fftw_malloc ( sizeof ( fftw_complex ) * n );
    /* Initialize data */
    for ( i = 0; i < n; i++ ) {
        if ( i <= (n/2-1)) {
            in[i][0] = 1.;
            in[i][1] = 0.;
        } else {
            in[i][0] = 0.;
            in[i][1] = 0.;
        }
    }
    /* Create plans. */
    plan_forward = fftw_plan_dft_1d ( n, in, out, FFTW_FORWARD, FFTW_ESTIMATE );
    /* Compute transform (as many times as desired) */
    fftw_execute ( plan_forward );
    /* deallocate and destroy plans */
    fftw_destroy_plan ( plan_forward );
    fftw_free ( in );
    fftw_free ( out );
    return 0;
}
program FFT_MPI_3D
    use, intrinsic :: iso_c_binding
    implicit none
    include 'mpif.h'
    include 'fftw3-mpi.f03'
    integer(C_INTPTR_T), parameter :: L = 1024
    integer(C_INTPTR_T), parameter :: M = 1024
    type(C_PTR) :: plan, cdata
    complex(C_DOUBLE_COMPLEX), pointer :: fdata(:,:)
    integer(C_INTPTR_T) :: alloc_local, local_M, local_j_offset
    integer(C_INTPTR_T) :: i, j
    complex(C_DOUBLE_COMPLEX) :: fout
    integer :: ierr, myid, nproc

! Initialize
    call mpi_init(ierr)
    call MPI_COMM_SIZE(MPI_COMM_WORLD, nproc, ierr)
    call MPI_COMM_RANK(MPI_COMM_WORLD, myid, ierr)
    call fftw_mpi_init()

! get local data size and allocate (note dimension reversal)
    alloc_local = fftw_mpi_local_size_2d(M, L, MPI_COMM_WORLD, local_M, local_j_offset)
    cdata = fftw_alloc_complex(alloc_local)
    call c_f_pointer(cdata, fdata(:,), [L,local_M])

! create MPI plan for in-place forward DFT (note dimension reversal)
    plan = fftw_mpi_plan_dft_2d(M, L, fdata, fdata, MPI_COMM_WORLD, FFTW_FORWARD, FFTW_MEASURE)
! initialize data to some function my_function(i,j)
    do j = 1, local_M
        do i = 1, L
            call initial(i, (j + local_j_offset), L, M, fout)
            fdata(i, j) = fout
        end do
    end do

! compute transform (as many times as desired)
    call fftw_mpi_execute_dft(plan, fdata, fdata)

! deallocate and destroy plans
    call fftw_destroy_plan(plan)
    call fftw_mpi_cleanup()
    call fftw_free(cdata)
    call mpi_finalize(ierr)
end
# include <stdlib.h>
# include <stdio.h>
# include <math.h>
# include <mpi.h>
# include <fftw3-mpi.h>

int main(int argc, char **argv)
{
    const ptrdiff_t L = 1024, M = 1024;
    fftw_plan plan;
    fftw_complex *data;
    ptrdiff_t alloc_local, local_L, local_L_start, i, j, ii;
    double xx, yy, rr, r2, t0, t1, t2, t3, tplan, texec;
    const double amp = 0.25;
    /* Initialize */
    MPI_Init(&argc, &argv);
    fftw_mpi_init();

    /* get local data size and allocate */
    alloc_local = fftw_mpi_local_size_2d(L, M, MPI_COMM_WORLD, &local_L, &local_L_start);
    data = fftw_alloc_complex(alloc_local);
    /* create plan for in-place forward DFT */
    plan = fftw_mpi_plan_dft_2d(L, M, data, data, MPI_COMM_WORLD, FFTW_FORWARD, FFTW_ESTIMATE);
}
/* initialize data to some function my_function(x,y) */

/* compute transforms, in-place, as many times as desired */
    fftw_execute(plan);
/* deallocate and destroy plans */
    fftw_destroy_plan(plan);
    fftw_mpi_cleanup();
    fftw_free(data);
    MPI_Finalize();
}
The most important FFT Fortran Library that use 2D (Pencil) Domain Decomposition
• General-purpose 2D pencil decomposition module to support building large-scale parallel applications on distributed memory systems.

• Highly scalable and efficient distributed Fast Fourier Transform module, supporting three dimensional FFTs (both complex-to-complex and real-to-complex/complex-to-real).

• Halo-cell support allowing explicit message passing between neighbouring blocks.

• Parallel I/O module to support the handling of large data sets.

• Shared-memory optimisation on the communication code for multi-code systems.

• Written in Fortran

• Best performance using Fortran 2003 standard

• No C wrapper is already provided

• Structure: Plan Creation – Execution – Plan Destruction

• Uses FFTW lib (or ESSL) to compute 1D transforms

• More efficient on massively parallel supercomputers.

• Well tested

• Additional features
How can I compile a code that uses 2Decomp&FFT on PLX?

- **Module Loading:**
  
  module load autoload profile/advanced

  Module load 2Decomp_FFT/1.5.847--openmpi--1.6.3--intel--cs-xe-2013—binary

- **Including header:**
  
  -I$FFTW_INC -I$DECOMP_2D_FFT_INC

- **Linking (double precision):**
  
  -L$DECOMP_2D_FFT_LIB -L$FFTW_LIB -l2decomp_fft -lfftw3_mpi -lfftw3

- **Example:**

  mpif90 -I. -I$FFTW_INC -I$DECOMP_2D_FFT_INCLUDE prova.F90 -L$DECOMP_2D_FFT_LIB -L$FFTW_LIB -l2decomp_fft -lfftw3
How can I compile a code that uses 2Decomp&FFT on FERMI?

• Module Loading:

```bash
module load autoload profile/advanced
module load 2Decomp_fft/1.5.847--bgq-gnu--4.4.6
```

• Including header:

```bash
-I$FFTW3_INC -I$DECOMP_2D_FFT_INC
```

• Linking (double precision):

```bash
-L$DECOMP_2D_FFT_LIB -L$FFTW3_LIB -l2decomp_fft -lfftw3_mpi -lfftw3
```

• Example:

```bash
mpif90 -I. -I$FFTW3_INC -I$DECOMP_2D_FFT_INC example.F90 -L$FFTW3_LIB -L$DECOMP_2D_FFT_LIB -l2decomp_fft -lfftw3 -lm
```
Including 2Decomp&FFT Lib:
  use decomp_2d
  use decomp_2d_fft

Initial declarations:
  integer, parameter :: p_row = ..
  integer, parameter :: p_col = ...
  complex(mytype), allocatable, dimension(:,,:,:,:) :: in, out

Please note that: \( p_{row} \times p_{col} = N_{core} \)

Initialization:
  call decomp_2d_init(n0,n1,n2,p_row,p_col)
  call decomp_2d_fft_init

Allocation:
  call decomp_2d_fft_get_size(fft_start,fft_end,fft_size)
  allocate (in(xstart(1):xend(1),xstart(2):xend(2),xstart(3):xend(3)))
  allocate (out(fft_start(1):fft_end(1), fft_start(2):fft_end(2), fft_start(3):fft_end(3))) ! R2C FFT
  allocate (out(zstart(1):zend(1),zstart(2):zend(2),zstart(3):zend(3))) ! C2C FFT

Execution:
  call decomp_2d_fft_3d(in, out, DECOMP_2D_FFT_FORWARD)

Finalizing:
  call decomp_2d_fft_finalize
  call decomp_2d_finalize
An example
PROGRAM FFT_3D_2Decomp_MPI
   use mpi
   use, intrinsic :: iso_c_binding
   use decomp_2d
   use decomp_2d_fft
   implicit none
   integer, parameter :: L = 128
   integer, parameter :: M = 128
   integer, parameter :: N = 128
   integer, parameter :: p_row = 16
   integer, parameter :: p_col = 16
   integer :: nx, ny, nz
   complex(mytype), allocatable, dimension(:,:, :) :: in, out
   complex(mytype) :: fout
   integer :: ierror, i,j,k, numproc, mype
   integer, dimension(3) :: sizex, sizez

! ===== Initialize
   call MPI_INIT(ierr)
   call decomp_2d_init(L,M,N,p_row,p_col)
   call decomp_2d_fft_init
allocate (in(xstart(1):xend(1),xstart(2):xend(2),xstart(3):xend(3)))
allocate (out(zstart(1):zend(1),zstart(2):zend(2),zstart(3):zend(3)))

! ===== each processor gets its local portion of global data =====

do k=xstart(3),xend(3)
do j=xstart(2),xend(2)
do i=xstart(1),xend(1)
  call initial(i, j, k, L, M, N, fout)
in(i,j,k) = fout
end do
end do
end do

! ===== 3D forward FFT =====

call decomp_2d_fft_3d(in, out, DECOMP_2D_FFT_FORWARD)

! ==========================
call decomp_2d_fft_finalize
call decomp_2d_finalize
deallocate(in,out)
call MPI_FINALIZE(ierr)
end
Parallel Three-Dimensional Fast Fourier Transforms (P3DFFT)
• General-purpose 2D pencil decomposition module to support building large-scale parallel applications on distributed memory systems.

• Highly scalable and efficient distributed Fast Fourier Transform module, supporting three dimensional FFTs (both complex-to-complex and real-to-complex/complex-to-real).

• Sine/cosine/Chebyshev/empty transform

• Shared-memory optimisation on the communication code for multi-code systems.

• Written in Fortran 90

• C wrapper is already provided

• Structure: Plan Creation – Execution – Plan Destruction

• Uses FFTW lib (or ESSL) to compute 1D transforms

• More efficient on massively parallel supercomputers.

• Well tested but not stable as 2Decomp&FFT

• Additional features
How can I compile a code that uses P3DFFT on PLX?

• Module Loading:

```plaintext
module load autoload profile/advanced
module load p3dfft/2.5.1--openmpi--1.6.3--intel--cs-xe-2013--binary
```

• Including header:

```plaintext
-I$FFTW_INC  -I$P3DFFT_INC
```

• Linking (double precision):

```plaintext
-L$P3DFFT_LIB  -L$FFTW_LIB  -lp3dfft -lfftw3
```

• Example:

```plaintext
mpif90 -openmp -I. -I$P3DFFT_INC -I$FFTW_INC example.F90 -L$P3DFFT_LIB -L$FFTW_LIB -lp3dfft -lfftw3 -lmpi_f90 -lmpi_f77
```
How can I compile a code that uses P3DFFT on FERMI?

• Module Loading:

   module load autoload profile/advanced

   module load p3dfft/2.6.1.beta--bgq-gnu--4.4.6

• Including header:

   -I$FFTW_INC -I$P3DFFT_INC

• Linking (double precision):

   -L$P3DFFT_INC -L$FFTW_LIB -l2decomp_fft -lfftw3_mpi -lfftw3

• Example:
Including P3DFFT Lib:
#include "p3dfft.h"

Initial declarations:
MPI_Dims_create(nproc,2,dims);
if(dims[0] > dims[1]) {
    dims[0] = dims[1];
    dims[1] = nproc/dims[0]; }

Initialization:
p3dfft_setup(dims,nx,ny,nz,1,memsize);
p3dfft_get_dims(istart,iend,isize,1);
p3dfft_get_dims(fstart,fend,isize,2);

Allocation:
A = (double *) malloc(sizeof(double) * isize[0]*isize[1]*isize[2]);
B = (double *) malloc(sizeof(double) * fsize[0]*fsize[1]*fsize[2]*2);
C = (double *) malloc(sizeof(double) * isize[0]*isize[1]*isize[2]);

Execution:
p3dfft_ftran_r2c(A,B,op_f);
p3dfft_btran_c2r(B,C,op_b);

Finalizing:
p3dfft_clean();
MPI_Finalize();
An example
```c
#include "p3dfft.h"
#include <mpi.h>
#include <stdio.h>
#include <stdlib.h>
#include <math.h>

int main(int argc,char **argv)
{
    double *A,*B,*p,*C;
    int memsize[3];
    int istart[3], isize[3], iend[3];
    int fstart[3], fsize[3], fend[3];
    /* .... ... ...*/
    MPI_Init(&argc,&argv);
    MPI_Comm_size(MPI_COMM_WORLD,&nproc);
    MPI_Comm_rank(MPI_COMM_WORLD,&proc_id);
    dims[0]=dims[1]=0;
    MPI_Dims_create(nproc,2,dims);
    if(dims[0] > dims[1]) {
        dims[0] = dims[1];
        dims[1] = nproc/dims[0];
    }
    /* ... ... ...*/
```
p3dfft_setup(dims, nx, ny, nz, 1, memsize);
MPI_Barrier(MPI_COMM_WORLD);
p3dfft_get_dims(istart, iend, isize, 1);
p3dfft_get_dims(fstart, fend, fsize, 2);
/* ... ... ... */
A = (double *) malloc(sizeof(double) * isize[0] * isize[1] * isize[2]);
B = (double **) malloc(sizeof(double) * fsize[0] * fsize[1] * fsize[2] * 2);
C = (double **) malloc(sizeof(double) * isize[0] * isize[1] * isize[2]);
MPI_Barrier(MPI_COMM_WORLD);
/* ... Initialization of A ... */
p3dfft_ftran_r2c(A, B, op_f);
/* ... ... ... */
for (i = 0; i < Ntot; i++) {
    B[i] *= nx * ny * nz;
}
MPI_Barrier(MPI_COMM_WORLD);
/* ... ... ... */
p3dfft_btran_c2r(B, C, op_b);
/* ... ... ... */
MPI_Barrier(MPI_COMM_WORLD);
/* ... ... ... */
p3dfft_clean();
MPI_Finalize();
Performance results

- Simple c2c FFT code based on 2Decomp&FFT library vs standard FFTW library
- Tested on FERMI up to 4096 cores
- Execution times
Performance Results

- FFT Kernel of BlowupNS CFD code based on 2Decomp&FFT library
- Tested on FERMI up to 32768 cores
- Two resolution tested:
  - 4096x256x256
  - 4096x512x512
Links:

FFTW Homepage: http://www.fftw.org/

Download FFTW-3: http://www.fftw.org/fftw-3.3.3.tar.gz


2Decomp&FFT homepage: http://www.2decomp.org/

Download 2Decomp&FFT: http://www.2decomp.org/download/2decomp_fft-1.5.847.tar.gz

Online Manual 2Decomp&FFT: http://www.2decomp.org/decomp_api.html

P3DFFT homepage: http://code.google.com/p/p3dfft/

Download P3DFFT: http://p3dfft.googlecode.com/files/p3dfft-dist.2.6.1.tar.gz

Thank You

For any other info,
Send an email to
m.guarrasi@cineca.it