Research Computing with C++

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Chapter 1

C++ for Research

Course Overview

Part 1

- Using C++ in research
- Better C++
  - Reliable
  - Reproducible
  - Good science

Part 2

- HPC concepts
- Shared memory parallelism - OpenMP
- Distributed memory parallelism - MPI
- Accelerators (GPU/Thrust)

Course Aims

- Teach how to do research with C++
- Optimise your research output
- A taster for various technologies
- Not just C++ syntax, Google/Compiler could tell you that!
Pre-requisites

- You are already doing some C++
- You are familiar with your compiler
- You are happy with the concept of classes
- You know C++ up to templates?
- You are familiar with development eg. version control
  - Git: https://git-scm.com/

Course Notes

- Revise some software Engineering: MPHYG001
- Register with Moodle: MPHYG002
  - key is “performance”
- Online notes: MPHYG002

Course Assessment

- 3 hour exam
- 2 pieces coursework - 40 hours each
  - 1 due 3rd week March
  - 1 due last week April
  - (roughly)

Course Community

- UCL Research Programming Hub: http://research-programming.ucl.ac.uk
- Slack: https://ucl-programming-hub.slack.com

Today’s Lesson

- Introduction, course overview, admin
- Using C++ in research
- Using Git
- Using CMake
- Using Catch unit testing framework
C++ In Research

Problems In Research

• Poor quality software
• Excuses
  – I’m not a software engineer
  – I don’t have time
  – It’s just a prototype
  – I’m unsure of my code (scared to share)

C++ Disadvantages

Some people say:

• Compiled language
  – (compiler versions, libraries, platform specific etc)
• Perceived as difficult, error prone, wordy, unfriendly syntax
• Result: It’s more trouble than its worth?

C++ Advantages

• Fast, code compiled to machine code
• Stable, evolving standard, powerful notation, improving
• Lots of libraries, Boost, Qt, VTK, ITK etc.
• Nice integration with CUDA, OpenACC, OpenCL, OpenMP, OpenMPI
• Result: Good reasons to use it, or you may have to use it

Research Programming

• Software is already expensive
  – Famous Book: Mythical Man Month
  – Famous People: Edsger W. Dijkstra
• Research programming is different
  – What is the end product?
Development Methodology?

- Will software engineering methods help?
  - Waterfall
  - Agile
- At the ‘concept discovery’ stage, probably too early to talk about product development

Approach

- What am I trying to achieve?
- How do I maximise my output?
- What is the best pathway to success?
- How do I de-risk (get results, meet deadlines) my research?

1. Types of Code

- What are you trying to achieve?
- Divide code:
  - Your algorithm: NiftyReg
  - Testing code
  - Data analysis
  - User Interface
  - Glue code
  - Deployment code
  - Scientific paper production

Examples: NiftyReg 300 citations in 5 years!

2. Maximise Your Value

- Developer time is expensive
- Your brain is your asset
- Write as little code as possible
- Focus tightly on your hypothesis
- Write the minimum code that produces a paper

Don’t fall into the trap “Hey, I’ll write a framework for that”
3. Ask Advice

- Before contemplating a new piece of software
  - Ask advice - Slack Channel
  - Review libraries and use them.
  - Check libraries are suitable, and sustainable.
  - Read Libraries section from Software Engineering course
  - Ask about best practices

Example - NiftyCal

- We should: Practice What We Preach
- Small, algorithms
- Unit tested
- Version controlled
- Small number of libraries
- Increased research output

Debunking The Excuses

- I’m not a software engineer
  - Learn effective, minimal tools
- I don’t have time
  - Unit testing to save time
  - Choose your battles/languages wisely
- I’m unsure of my code
  - Share, collaborate

What Isn’t This Course?

We are NOT suggesting that:

- C++ is the solution to all problems.
- You should write all parts of your code in C++. 
What Is This Course?

We aim to:

- Improve your C++ (and associated technologies).
- Do High Performance Computing (HPC).

So that:

- Apply it to research in a pragmatic fashion.
- You use the right tool for the job.

Git

Git Introduction

- This is a practical course
- We will use git for version control
- Submit git repository for coursework
- Here we provide a very minimal introduction

Git Resources

- Complete beginner - Try Git
- Git book by Scott Chacon
- Git section of MPHYG001
- MPHYG001 repo

Git Walk Through

(demo on command line)

- git init
- git add
- git commit
- git status
- git log
- git push
- git pull
- git clone
- forking
Homework

- Register Github
- Create new empty repository - CPPCW1
- Ensure it is a private repository - free
- Find project of interest - try cloning it
- Find project of interest - try forking it

CMake

CMake Introduction

- This is a practical course
- We will use CMake as a build tool
- CMake produces
  - Windows: Visual Studio project files
  - Linux: Make files
  - Mac: XCode projects, Make files
- This course will provide CMake code and boiler plate code

CMake Usage Linux/Mac

Demo an “out-of-source” build

cd ~/build
git clone https://github.com/MattClarkson/CMakeHelloWorld
mkdir CMakeHelloWorld-build
cd CMakeHelloWorld-build
ccmake ../CMakeHelloWorld
make

CMake Usage Windows

Demo an “out-of-source” build

- git clone https://github.com/MattClarkson/CMakeHelloWorld
- Run cmake-gui.exe
- Select source folder (CMakeHelloWorld downloaded above)
- Specify new build folder (CMakeHelloWorld-build next to, but not inside CMakeHelloWorld)
b • Hit configure
b • When asked, specify compiler
b • Set flags and repeatedly configure
b • When generate option is present, hit generate
b • Compile, normally using Visual Studio

Unit Testing

What is Unit Testing?

At a high level

b • Way of testing code.
b • Unit
  – Smallest ‘atomic’ chunk of code
  – i.e. Function, could be a Class
b • See also:
  – Integration/System Testing
  – Regression Testing
  – User Acceptance Testing

Benefits of Unit Testing?

b • Certainty of correctness
b • (Scientific Rigour)
b • Influences and improves design
b • Confidence to refactor, improve

Drawbacks for Unit Testing?

b • Don’t know how
  – This course will help
b • Takes too much time
  – Really?
  – IT SAVES TIME in the long run
Unit Testing Frameworks

Generally, all very similar

- JUnit (Java), NUnit (.net?), CppUnit, phpUnit,
- Basically
  - Macros (C++), methods (Java) to test conditions
  - Macros (C++), reflection (Java) to run/discover tests
  - Ways of looking at results.
    * Java/Eclipse: Integrated with IDE
    * Log file or standard output

Unit Testing Example

How To Start

We discuss

- Basic Example
- Some tips

Then its down to the developer/artist.

C++ Frameworks

To Consider:

- Catch
- GoogleTest
- QTestLib
- BoostTest
- CppTest
- CppUnit

Worked Example

- Borrowed from
  - Catch Tutorial
  - and Googletest Primer

- We use Catch, so notes are compilable
- But the concepts are the same
Code

To keep it simple for now we do this in one file:

```cpp
#define CATCH_CONFIG_MAIN // This tells Catch to provide a main() - only do this in one cpp file
#include "../catch/catch.hpp"

unsigned int Factorial( unsigned int number ) {
    return number <= 1 ? number : Factorial(number-1)*number;
}

TEST_CASE( "Factorials are computed", "[factorial]" ) {
    REQUIRE( Factorial(1) == 1 );
    REQUIRE( Factorial(2) == 2 );
    REQUIRE( Factorial(3) == 6 );
    REQUIRE( Factorial(10) == 3628800 );
}
```

Produces this output when run:

===============================================================================
All tests passed (4 assertions in 1 test case)

Principles

So, typically we have

- Some `#include` to get test framework
- Our code that we want to test
- Then make some assertions

Catch / GoogleTest

For example, in Catch:

```cpp
// TEST_CASE(<unique test name>, <test case name>)
TEST_CASE( "Factorials are computed", "[factorial]" ) {
    REQUIRE( Factorial(2) == 2 );
    REQUIRE( Factorial(3) == 6 );
}
```

In GoogleTest:
TEST(FactorialTest, HandlesPositiveInput) {
    EXPECT_EQ(2, Factorial(2));
    EXPECT_EQ(6, Factorial(3));
}

all done via C++ macros.

Tests That Fail

What about Factorial of zero? Adding

    REQUIRE( Factorial(0) == 1 );

 Produces something like:

    factorial2.cc:9: FAILED:
    REQUIRE( Factorial(0) == 1 )
    with expansion:
    0 == 1

Fix the Failing Test

Leading to:

    #define CATCH_CONFIG_MAIN    // This tells Catch to provide a main() - only do this in one cpp
    //include "../catch/catch.hpp"

    unsigned int Factorial( unsigned int number ) {
        //return number <= 1 ? number : Factorial(number-1)*number;
        return number > 1 ? Factorial(number-1)*number : 1;
    }

TEST_CASE( "Factorials are computed", "[factorial]" ) {
    REQUIRE( Factorial(0) == 1 );
    REQUIRE( Factorial(1) == 1 );
    REQUIRE( Factorial(2) == 2 );
    REQUIRE( Factorial(3) == 6 );
    REQUIRE( Factorial(10) == 3628800 );
}

which passes:
All tests passed (5 assertions in 1 test case)

Test Macros

Each framework has a variety of macros to test for failure. [Check][Check] has:

```c
REQUIRE(expression); // stop if fail
CHECK(expression);   // doesn't stop if fails
```

If an exception is thrown, it’s caught, reported and counts as a failure.

Examples:

```c
CHECK( str == "string value" );
CHECK( thisReturnsTrue() );
REQUIRE( i == 42 );
```

Others:

```c
REQUIRE_FALSE( expression )
CHECK_FALSE( expression )
REQUIRE_THROWS( expression ) # Must throw an exception
CHECK_THROWS( expression ) # Must throw an exception, and continue testing
REQUIRE_THROWS_AS( expression, exception_type )
CHECK_THROWS_AS( expression, exception_type )
REQUIRE_NOTHROW( expression )
CHECK_NOTHROW( expression )
```

Testing for Failure

To re-iterate:

- You should test failure cases
  - Force a failure
  - Check that exception is thrown
  - If exception is thrown, test passes
  - (Some people get confused, expecting test to fail)

- Examples
  - Saving to invalid file name
  - Negative numbers passed into double arguments
  - Invalid Physical quantities (e.g. -300 Kelvin)
Setup/Tear Down

- Some tests require objects to exist in memory
- These should be set up
  - for each test
  - for a group of tests
- Frameworks do differ in this regards

Setup/Tear Down in Catch

Referring to the Catch Tutorial:

```cpp
TEST_CASE( "vectors can be sized and resized", "[vector]" ) {

  std::vector<int> v( 5 );

  REQUIRE( v.size() == 5 );
  REQUIRE( v.capacity() >= 5 );

  SECTION( "resizing bigger changes size and capacity" ) {
    v.resize( 10 );

    REQUIRE( v.size() == 10 );
    REQUIRE( v.capacity() >= 10 );
  }

  SECTION( "resizing smaller changes size but not capacity" ) {
    v.resize( 0 );

    REQUIRE( v.size() == 0 );
    REQUIRE( v.capacity() >= 5 );
  }

  SECTION( "reserving bigger changes capacity but not size" ) {
    v.reserve( 10 );

    REQUIRE( v.size() == 5 );
    REQUIRE( v.capacity() >= 10 );
  }

  SECTION( "reserving smaller does not change size or capacity" ) {
    v.reserve( 0 );

    REQUIRE( v.size() == 5 );
    REQUIRE( v.capacity() >= 5 );
  }

}
```
So, Setup/Tear down is done before/after each section.

**Unit Testing Tips**

**C++ design**

- Stuff from above applies to Classes / Functions
- Think about arguments:
  - Code should be hard to use incorrectly.
  - Use `const, unsigned` etc.
  - Testing forces you to sort these out.

**Test Driven Development (TDD)**

- Methodology
  1. Write a test
  2. Run test, should fail
  3. Implement/Debug functionality
  4. Run test
     1. if succeed goto 5
     2. else goto 3
  5. Refactor to tidy up

**TDD in practice**

- Aim to get good coverage
- Some people quote 70% or more
- What are the downsides?
- Don’t write ‘brittle’ tests

**Behaviour Driven Development (BDD)**

- Behaviour Driven Development (BDD)
  - Refers to a whole area of software engineering
  - With associated tools and practices
  - Think about end-user perspective
  - Think about the desired behaviour not the implementation
  - See Jani Hartikainen article.
TDD Vs BDD

- **TDD**
  - Test/Design based on methods available
  - Often ties in implementation details
- **BDD**
  - Test/Design based on behaviour
  - Code to interfaces (later in course)
- Subtly different
- Aim for BDD

Anti-Pattern 1: Setters/Getters

Testing every Setter/Getter.
Consider:

```cpp
class Atom {
    public:
    void SetAtomicNumber(const int& number) { m_AtomicNumber = number; }
    int GetAtomicNumber() const { return m_AtomicNumber; }
    void SetName(const std::string& name) { m_Name = name; }
    std::string GetName() const { return m_Name; }
    
    private:
    int m_AtomicNumber;
    std::string m_Name;
};
```

and tests like:

```cpp```
TEST_CASE( "Testing Setters/Getters", "[Atom]" ) {
    Atom a;
    a.SetAtomicNumber(1);
    REQUIRE( a.GetAtomicNumber() == 1);
    a.SetName("Hydrogen");
    REQUIRE( a.GetName() == "Hydrogen");
}
```

- It feels tedious
- But you want good coverage
- This often puts people off testing
- It also produces “brittle”, where 1 change breaks many things

**Anti-Pattern 1: Suggestion.**

- Focus on behaviour.
  - What would end-user expect to be doing?
  - How would end-user be using this class?
  - Write tests that follow the use-case
  - Gives a more logical grouping
  - One test can cover > 1 function
  - i.e. move away from slavishly testing each function

- Minimise interface.
  - Provide the bare number of methods
  - Don’t provide setters if you don’t want them
  - Don’t provide getters unless the user needs something
  - Less to test. Use documentation to describe why.

**Anti-Pattern 2: Constructing Dependent Classes**

- Sometimes, by necessity we test groups of classes
- Or one class genuinely Has-A contained class
- But the contained class is expensive, or could be changed in future

**Anti-Pattern 2: Suggestion**

- Read up on Dependency Injection
- Enables you to create and inject dummy test classes
- So, testing again used to break down design, and increase flexibility

**Summary BDD Vs TDD**

Aim to write:

- Most concise description of requirements as unit tests
- Smallest amount of code to pass tests
- … i.e. based on behaviour
Any Questions?

End of Lecture?

• Example git repo, CMake, Catch template project:
  – https://github.com/MattClarkson/CMakeCatchTemplate
Chapter 2

Including Libraries

Why use Libraries?

Scientific Coding

- Explain your science
  - Should not just distribute binaries and papers
  - Source code is best description of your science
- Reproducible science
  - Collaborators must know how you build your project and libraries
- Time is short
  - Don’t re-invent the wheel
  - Use libraries if they provide bits you need

What are libraries?

- Libraries provide collections of useful classes and functions, ready to use
- C++ libraries can be somewhat harder to use than pure Python modules
- This lecture will get you started

Aim for this chapter

- How to choose a library
  - Licensing, longevity, developer community, technical implementation, feature list etc.
• Working with libraries
  – Including them
  – C++ concepts
  – Not an exhaustive product specific tutorial

Choosing libraries

Libraries are Awesome

A great set of libraries allows for a very powerful programming style:

• Write minimal code yourself
  – Choose the right libraries
  – Plug them together
  – Create impressive results

Libraries for Efficiency

Not only is this efficient with your programming time, it’s also more efficient with computer time.

The chances are any general algorithm you might want to use has already been programmed better by someone else.

Software Licenses

• Consider both:
  – License for 3rd party code / dependencies
  – License for your code when you distribute it

• So, even if you aren’t distributing code yet, you need to understand the licenses of your dependencies.

Third Party Licenses

CAVEAT: This is not legal advice. If in doubt, seek your own legal advice.

• When you distribute your code, the licenses of any libraries you use takes effect
• If library has:
- MIT and BSD are permissive. So you can do what you want, including sell it on.
- Apache handles multiple contributors and patent rights, but is basically permissive.

**Third Party Licenses**

CAVEAT: This is not legal advice. If in doubt, seek your own legal advice.

- When you distribute your code, the licenses of any libraries you use take effect
- If library has:
  - GPL requires you to open-source your code, including changes to the library you imported, and your work is considered a “derivative work”, so must be shared.
  - LGPL for libraries, but use dynamic not static linkage. If you use static linking its basically as per GPL.
- Still some debate on GPL/LGPL and derivative works - only true test is in court.

**Choosing a License**

CAVEAT: This is not legal advice. If in doubt, seek your own legal advice.

- When you plan to distribute code:
  - Read this book, and/or GitHub’s advice, and OSI for choosing your own license.
  - Don’t write your own license, unless you use legal advice.
  - Try to pick one of the standard ones for compatibility.
- Note: Once a 3rd party has your code under a license agreement, their restrictions are determined by that version of the code.

**Choose Stability**

- So, take care in your choice of 3rd party library
  - Don’t want to redo work later.
  - Don’t want to rely too heavily on non-distributable code.
  - But if you do, understand what that means.
Is Library Updated?

- Code that is not developed, rots.
  - When was the last commit?
  - How often are there commits?
  - Is the code on an open version control tool like GitHub?

Are Developers Reachable?

- Can you find the lead contributor on the internet?
  - Do they respond when approached?
- Are there contributors other than the lead contributor?

Is Code Tested?

- Are there many unit tests, do they run, do they pass?
- Does the library depend on other libraries?
- Are the build tools common?
- Is there a sensible versioning scheme (e.g. semantic versioning).
- Is there a suitable release schedule?

Is Library High Quality?

- Shouldn’t need to look excessively closely, but consider
  - Documentation
  - Number of ToDo’s
  - Dependencies (recursively)
  - Data Structures? How to import/export image
  - Can you write a convenient wrapper?

Library Features

- Then look at features
  - Manual
  - Easy to use
Summary

- In comparison with languages such as MATLAB/Python
  - In C++ prefer few well chosen libraries
  - Be aware of their licenses for future distribution
  - Keep a log of any changes, patches etc. that you make
  - Be able to compile all your own code, including libraries
    * so need common build environment. (eg. CMake, Make, Visual Studio).

Library Basics

Aim

You need your compiler to find:

- Headers: `#include`
- Library:
  - Dynamic: `.dylib, .so, .lib / .dll`
  - Static: `.a, .lib`

Linux/Mac

g++ -c -I/users/me/myproject/include main.cpp
g++ -o main main.o -L/users/me/myproject/lib -lmylib

- `-I` to specify include folder
- `-L` to specify library folder
- `-l` to specify the actual library

Compiler switches

So, that means

- Include directory
  - `-I /some/directory`
- Library directory
  - `-L /some/directory`
• Library
  – -l library

Similar concept on Windows, Linux and Mac.

Check Native Build Platform

• Look inside
  – Makefile
  – Visual Studio options
• Basically constructing -I, -L, -l switches to pass to command line compiler.

Windows Compiler Switches

• Visual Studio (check version)
• Project Properties
  – C/C++ -> Additional Include Directories.
  – Configuration Properties -> Linker -> Additional Library Directories
  – Linker -> Input -> Additional Dependencies.
• Check compile line - its equivalent to Linux/Mac, -I, -L, -l

Difficulties with Libraries

• Discuss
• (confession time)

Location Issues

When you use a library:

• Where is it?
• Header only?
• System version or your version?
• What about bugs? How do I upgrade?
Compilation Issues

When you use a library:

- Which library version?
- Which compiler version?
- Debug or Release?
- Static or Dynamic?
- 32 bit / 64 bit?
- Platform specific flags?
- Pre-installed, or did you compile it?

A Few Good Libraries

Due to all those issues shown above, again, the main advice for libraries:

- As few as possible
- As good as possible

Linking libraries

Linking

- From first lecture
  - Code is split into functions/classes
  - Related functions get grouped into libraries
  - Libraries get compiled / archived into one file
- End User needs
  - Header file = declarations (and implementation if header only)
  - Object code / library file = implementations

Static Linking

- Windows (.lib), Mac/Linux (.a)
- Compiled code from static library is copied into the current translation unit.
- Increases disk space compared with dynamic linking.
- Current translation unit then does not depend on that library.
Dynamic Linking

- Windows (.dll), Mac (.dylib), Linux (.so)
- Compiled code is left in the library.
- At runtime,
  - OS loads the executable
  - OS / Linker finds any unresolved libraries
  - Recursive process
- Saves disk space compared with static linking.
- Faster compilation/linking times?
- Current translation unit has a known dependency remaining.

Dynamic Loading

- System call to load a library (dlopen/LoadLibrary)
- Dynamically discover function names and variables
- Execute functions
- Normally used for plugins
- Not covered here

Space Comparison

- If you have many executables linking a common set of libraries
  - Static
    - Code gets copied - each executable bigger
    - Doesn’t require searching for libraries at run-time
  - Dynamic
    - Code gets referenced - smaller executable
    - Requires finding libraries at run-time
- Space - less of a concern these days

For Scientists

- Ease of use
- Ease of distribution to collaborators
- Prefer static if possible for ease of deployment
Packaging

- Packaging large apps takes effort
- hire Research Software Engineers

How to Check

- Windows - Dependency Walker
- Linux - ldd
- Mac - otool -L
  - (live demo on Mac)

Using libraries

Package Managers

- Package Manager (Linux/Mac)
  - Precompiled
  - Stable choice
  - Inter-dependencies work
- Linux
  - sudo apt-get install
  - sudo yum install
- Mac
  - sudo port install
  - brew install

Windows

- Libraries typically:
  - Randomly installed location
  - In system folders
  - In developer folders
  - In build folder
- Please clean your machine!!
- Try Chocolatey package manager
Package Managers

- So, if you can use standard versions of 3rd party libraries
- Package managers are a good way to go
- You just need to specify what versions so your collaborator can check

Problems

- As soon as you hit a bug in a library
  - How to update?
  - Knock on effects
    * Cascading updates
    * Inconsistent development environment

Build Your Own

- 2 basic approaches
  - External / Individual build
    * Build dependencies externally
    * Point your software at those packages
  - SuperBuild / Meta-Build
    * Write code to download all dependencies
    * The correct version numbers is stored in code

External / Individual Build

For example

C:\build\ITK-v1
C:\build\ITK-v1-build
C:\build\ITK-v1-install
C:\build\VTK-v2
C:\build\VTK-v2-build
C:\build\VTK-v2-install
C:\build\MyProject
C:\build\MyProject-build

We setup MyProject-build to know the location of ITK and VTK install folder.
Meta-Build / Super-Build

For example

C:\build\MyProject
C:\build\MyProject-SuperBuild\ITK\src
C:\build\MyProject-SuperBuild\ITK\build
C:\build\MyProject-SuperBuild\ITK\install
C:\build\MyProject-SuperBuild\VTK\src
C:\build\MyProject-SuperBuild\VTK\build
C:\build\MyProject-SuperBuild\VTK\install
C:\build\MyProject-SuperBuild\MyProject-build

We setup MyProject-build to know the location of ITK and VTK that it itself compiled.

Pro’s / Con’s

• External Build
  – Pro’s - build each dependency once
  – Con’s - collaborators will do this inconsistently
  – Con’s - how to manage multiple versions of all dependencies

• Meta Build
  – Pro’s - all documented, all self-contained, easier to share
  – Con’s - Slow build? Not a problem if you only run make in sub-folder MyProject-build

Examples

Reminder

• Fundamentally, we need:
  – Access to header files - declaration
  – Access to compiled code - definition

Header Only?

• After all this:
  – Static/Dynamic
– Package Managers / Build your own
– External build / Internal Build
– Release / Debug
– -I, -L, -l

• Header only libraries are very attractive.

**Use of CMake**

• Several ways depending on your setup
  – Specify paths and library names
  – Use `find_package`
    * Use 3rd party projects own config, eg. `VTKConfig.cmake`
    * Use a FindModule, some come with CMake
    * Write your own FindModule
    * Write your own FindModule with generated / substituted variables

**CMake - Header Only**

• `catch.hpp` Header files in project `CMakeCatchTemplate`
• From `CMakeCatchTemplate/CMakeLists.txt`

```cmake
include_directories(${CMAKE_SOURCE_DIR}/Code/)
add_subdirectory(Code)
if(BUILD_TESTING)
  include_directories(${CMAKE_SOURCE_DIR}/Testing/)
  add_subdirectory(Testing)
endif()
```

**CMake - Header Only**

• Options are:
  – Check small/medium size project into your project

• For example:

```
CMakeCatchTemplate/3rdParty/libraryA/version1/Class1.hpp
CMakeCatchTemplate/3rdParty/libraryA/version1/Class2.hpp
```

• Add to `CMakeLists.txt`

```
include_directories(${CMAKE_SOURCE_DIR}/3rdParty/libraryA/version1/)
```

• You’d have audit trail (via git repo) of when updates to library were made.
CMake - Header Only, External

- If larger, e.g. Eigen

C:\3rdParty\Eigen
C:\build\MyProject
C:\build\MyProject-build

- Add to CMakeLists.txt

  include_directories("C:\3rdParty\Eigen\install\include\eigen3")

- Hard-coded, but usable if you write detailed build instructions
- Not very platform independent
- Not very flexible
- Can be self contained if you have a Meta-build - read on.

CMake - find_package

- For example:

  find_package(OpenCV REQUIRED)
  include_directories(${OpenCV_INCLUDE_DIRS})
  list(APPEND ALL_THIRD_PARTY_LIBRARIES ${OpenCV_LIBS})
  add_definitions(-DBUILD_OpenCV)

- So a 3rd party package can provide information on how you should use it
- If its written in CMake code, even better!

find_package - Intro

- CMake comes with scripts to include various 3rd Party Libraries
- 3rd Parties can write these scripts aswell
- You can also write your own scripts
find_package - Search Locations

A Basic Example (for a full example - see docs)

- Given:

  ```cmake
  find_package(SomeLibrary [REQUIRED])
  ```

- CMake will search
  - all directories in CMAKE_MODULE_PATH
  - for SomeLibraryConfig.cmake - does config mode
  - for FindSomeLibrary.cmake - does module mode
  - case sensitive

find_package - Result

- `find_package(SomeLibrary)` should return `SomeLibrary_FOUND:BOOL=TRUE` if library was found
- Sets any other variables necessary to use the library
- Check CMakeCache.txt to see result

find_package - Usage

- So many 3rd party libraries are CMake ready.
- If things go wrong, you can debug it - CMake is all text based.

find_package - Fixing

- You can provide patched versions
- Add your source/build folder to the CMAKE_MODULE_PATH

```cmake
set(CMAKE_MODULE_PATH
  ${CMAKE_SOURCE_DIR}/CMake
  ${CMAKE_BINARY_DIR}
  ${CMAKE_MODULE_PATH}
)
```

- So CMake will find your version before the system versions
find_package - Tailoring

- You can write your own
  - e.g. FindEigen in CMakeCatchTemplate
- Use CMake to substitute variables
  - Force include/library dirs
  - Useful for vendors API that isn’t CMake compatible
  - Useful for meta-build. Force directories to match the package you just compiled.

Provide Build Flags

- If a package is found, you can add compiler flag.

  add_definitions(-DBUILD_OpenCV)

- So, you can optionally include things:

  #ifdef BUILD_OpenCV
  #include <cv.h>
  #endif

  Best not to do too much of this.

- Useful to provide build options, e.g. for running on clusters

Check Before Committing

- Before you commit code to git,

  #ifdef BUILD_OpenCV
  blah blah
  #include <cv.h>
  #endif

  should fail compilation
Summary

• Basic aim:
  – include_directories() generates -I
  – link_directories() generates -L
  – target_link_libraries(mylibrary PRIVATE ${libs}) generates -l for each library

• Might not need link_directories() if libraries fully qualified
• Try default CMake find_package
• Or write your own and add location to CMAKE_MODULE_PATH

CMakeCatchTemplate

Intro

• Demo project on GitHub: CMakeCatchTemplate
• No functional code, other than adding 2 numbers
• Basically shows how to use CMake

Features

• Full feature list in README.md
• SuperBuild or use without
• Downloads Boost, Eigen, glog, gflags, OpenCV, PCL, FLANN, VTK
• Example GUI apps (beyond scope of course)
• Unit testing
• Script rename.sh to create your own project

Main CMake flags

These were covered earlier - here’s how you set them.

• CMAKE_BUILD_TYPE:String=[Debug|Release]
• BUILD_SHARED_LIBS:BOOL=[OFF|ON]
• Compile flags
  – CMAKE_C_FLAGS, CMAKE_CXX_FLAGS, CMAKE_EXE_LINKER_FLAGS all String
32 or 64 bit

- Generally stick with default
- 32 or 64 bit?
  - Visual Studio - chose generator, check in Visual Studio (x86, Winn64)
  - CMAKE_C_FLAGS = -m32 etc. For example

SuperBuild

- See flag: BUILD_SUPERBUILD:BOOL=[ON|OFF]
- If OFF
  - Just compiles this project in current folder
- If ON
  - Dependencies in current folder
  - Compiles this project in sub-folder
- Try it
- A SuperBuild with no dependencies just does nothing

Home Work 1a - Basic Build

- Build CMakeCatchTemplate BUILD_SUPERBUILD=OFF
- Modify some C++, recompile
- Choose a project relevant to you - eg. ITK
- Compile it separately
- Or install via package manager

Home Work 1b - Basic Build

- Try to find_package(PackageName) to see what happens
- Set include_directories
- Add library to target_link_libraries
- Work out if its necessary to build it yourself

Home Work 2 - SuperBuild

- Not for the faint-hearted
  - 3rd party examples in CMake/ExternalProjects
  - Create a new one for your project
Add your project name to loop in SuperBuild.cmake
Add top-level control variables like BUILD_MyExternalProject
Pass variables from SuperBuild to main project build, as shown in SuperBuild.cmake

Summary

No Magic Answer

• C++ is more tricky than MATLAB / Python
• While package managers make it easier, you still need to understand what you’re building.

A Few Good Libraries

Main advice for libraries:

• As few as possible
• As high quality as possible

Ways to Use

• Easiest - header only
  – just include directly in your source code
  – use cmake include_directories()
  – compile it in
• use packages from package manager
  – use cmake to find_package
  – set variables to your installed version
• use build system to build everything
  – you control ALL flags
  – cmake does find_package on things you just compiled

Git Submodule Anyone?

• git submodule
  – If all dependencies are cmake’d
– Put each project in a git submodule
– Cmake can configure whole project
– Not really used on this course
Chapter 3

Better C++

Better C++

The Story So Far

- Git
- CMake
- Use libraries with CMake

Today’s Lesson

- Beginners C++ not sufficient
- Look at
  - C++ recap
  - Templates - generic programming
  - Exceptions - error handling
Chapter 4

C++ Libraries

C++ Libraries

The Story So Far

- Git
- CMake
- Use libraries
- More C++

Today’s Lesson

- C++ libraries
  - Language features
  - Using libraries

Using Eigen

Introduction

- Eigen is:
  - C++ template library,
  - Linear algebra, matrices, vectors, numerical solvers, etc.
Tutorials

Obviously, you can read:

- the existing manual pages
- tutorials (short, long).
- the Quick Reference

Getting started

- Header only, just need `#include`
- Uses CMake, but that’s just for
  - documentation
  - run unit tests
  - do installation.

C++ Principles

(i.e. why introduce Eigen on this course)

- Eigen uses
  - Templates
  - Loop unrolling, traits, template meta programming

Matrix Class

- This:

### Module | Header File | Contents
--- | --- | ---
Core | `#include <Eigen/Eigen>` | Matrix and Array classes, basic linear algebra (including triangular and symmetric products), array manipulation
Geometry | `#include <Eigen/Geometry>` | Transforms, Translation, Scaling, Rotation2D and 3D rotations (Quaternions, Angles)
LU | `#include <Eigen/LU>` | Inverse, determinant, LU decompositions with solver (FullPivLU, PartialPivLU)
Cholesky | `#include <Eigen/Cholesky>` | LLT and LDLT Cholesky factorization with solver
Householder | `#include <Eigen/Householder>` | Householder transformations, this module is used by several linear algebra modules
SVD | `#include <Eigen/SVD>` | SVD decompositions with least squares solver (JacobiSVD)
QR | `#include <Eigen/QR>` | QR decompositions with solver (HouseholderQR, CpuHouseholderQR, FullPivotHouseholderQR)
Eigenvalues | `#include <Eigen/Eigenvalues>` | Eigenvalue, eigenvector decompositions (EigenSolver, SelfAdjointEigenSolver, ComplexEigenSolver)
Sparse | `#include <Eigen/Sparse>` | Sparse matrix storage and related basic linear algebra (SparseMatrix, DynamicSparseMatrix, SparseVector)

columns | `#include <Eigen/Geometry>` | Includes Core, Geometry, LU, Cholesky, SVD, QR, and Eigenvalues header files

columns | `#include <Eigen/Eigen>` | Includes Dense and Sparse header files (the whole Eigen library)
```cpp
#include <iostream>
#include <Eigen/Dense>
using Eigen::MatrixXd;

int main()
{
    MatrixXd m(2,2);
    m(0,0) = 3;
    m(1,0) = 2.5;
    m(0,1) = -1;
    m(1,1) = m(1,0) + m(0,1);
    std::cout << m << std::endl;
}

• Produces:
    3  -1
    2.5 1.5

Matrix Class Declaration

Matrix Class

template<typename _Scalar, int _Rows, int _Cols, int _Options, int _MaxRows, int _MaxCols>
class Matrix
    : public PlainObjectBase<Matrix<_Scalar, _Rows, _Cols, _Options, _MaxRows, _MaxCols> >
{

So, it's templates, so review last week's lecture.

Matrix Class Construction

But in documentation:

All combinations are allowed: you can have a matrix with a fixed number of rows and a dynamic number of columns, e are all valid:

- Matrix<double, 6, Dynamic>  // Dynamic number of columns (heap allocation)
- Matrix<double, Dynamic, 2>  // Dynamic number of rows (heap allocation)
- Matrix<double, Dynamic, Dynamic, RowMajor>  // Fully dynamic, row major (heap allocation)
- Matrix<double, 13, 3>  // Fully fixed (usually allocated on stack)

Figure 4.1: Matrix construction

It took a while but I searched and found:

src/Core/util/Constants.h:const int Dynamic = -1;

67
and both fixed and dynamic Matrices come from same template class???
How do they do that?

**DenseStorage.h - 1**

In src/Core/DenseStorage.h:

```cpp
template <typename T, int Size, int MatrixOrArrayOptions,
          int Alignment = (MatrixOrArrayOptions&DontAlign) ? 0
           : (((Size*sizeof(T))%16)==0) ? 16
           : 0 >
struct plain_array
{
    T array[Size];
}
```

So, a `plain_array` structure containing a stack allocated array.

**DenseStorage.h - 2**

In src/Core/DenseStorage.h:

```cpp
// purely fixed-size matrix
template<typename T, int Size, int _Rows, int _Cols, int _Options>
class DenseStorage
{
    internal::plain_array<T,Size,_Options> m_data;
}
```

There is a default template class for DenseStorage, and specialisation for fixed arrays.

**DenseStorage.h - 3**

In src/Core/DenseStorage.h:

```cpp
// purely dynamic matrix.
template<typename T, int _Options> class DenseStorage<T, Dynamic, Dynamic, Dynamic, _Options>
{
    T *m_data;
    DenseIndex m_rows;
    DenseIndex m_cols;
}
```

There is a default template class for DenseStorage, and specialisation for Dynamic arrays.
Eigen Matrix Summary

- Templated type supports dynamic and fixed arrays seamlessly on stack or heap
- typedef's to make life easier: Matrix3d = 3 by 3 of double
- Uses TMP to generate generate code at compile time
- Benefit from optimisations such as loop unrolling when using fixed size constant arrays

Eigen Usage - CMake Include

- Need to set include path
- You could download and ‘install’ eigen into your project, and commit it.
  e.g.

```cpp
include_directories(${CMAKE_SOURCE_DIR}/session03/cpp/Eigen/eigen-3.2.3/include/eigen3)
```

Eigen Usage - CMake Module

- CMake (3.1) does not have a Find Module for eigen, but eigen provides one.
- So, in your source tree

```sh
mkdir CMake
cp <path_to_eigen>/cmake/FindEigen3.cmake ./CMake
```

- Then in your CMakeLists.txt

```cpp
set(CMAKE_MODULE_PATH "${CMAKE_SOURCE_DIR}/CMake;${CMAKE_MODULE_PATH}"
find_package(Eigen3)
include_directories(${EIGEN3_INCLUDE_DIR})
```

Eigen Usage - CMake External

- NiftySeg uses

```cpp
option(USE_SYSTEM_EIGEN "Use an already installed version of the Eigen library" OFF)
if(USE_SYSTEM_EIGEN)
  find_package(EIGEN REQUIRED)
else()
  set(${PROJECT_NAME}_VERSION_EIGEN "ffa86ff5570" CACHE STRING "Version of EIGEN" FORCE)
```
ExternalProject_Add(Eigen
  URL ${${PROJECT_NAME}_LOCATION_EIGEN}
  URL_MD5 ${${PROJECT_NAME}_MD5_SUM_EIGEN}
  PREFIX ${PROJECT_BINARY_DIR}/Eigen
  DOWNLOAD_DIR ${PROJECT_BINARY_DIR}/Eigen/download
  SOURCE_DIR ${PROJECT_BINARY_DIR}/Eigen/source
  STAMP_DIR ${PROJECT_BINARY_DIR}/Eigen/stamps
  TMP_DIR ${PROJECT_BINARY_DIR}/Eigen/tmp
  BINARY_DIR ${PROJECT_BINARY_DIR}/Eigen/build
  CMAKE_ARGS
    ${CMAKE_PROPAGATED_VARIABLES}
    -DCMAKE_INSTALL_PREFIX:PATH=${PROJECT_BINARY_DIR}/Eigen/install
    -DBUILD_TESTING=1
  )
set(Eigen_INCLUDE_DIR ${PROJECT_BINARY_DIR}/Eigen/install/include/eigen3)
endif()
include_directories(${Eigen_INCLUDE_DIR})

Eigen Example - in PCL

Point Cloud Library uses Eigen. Lets look at point based registration of two, same length, point sets.

PCL - Manual Registration

- Class to hold point lists
- Callbacks (not shown here) to add points to list

```cpp
class ManualRegistration : public QMainWindow {
  protected:
    pcl::PointCloud<pcl::PointXYZ> src_pc_;  
    pcl::PointCloud<pcl::PointXYZ> dst_pc_;  
    Eigen::Matrix4f transform_;  
}
```

PCL - SVD class

- In apps/src/manual_registration/manual_registration.cpp
- Create a class to estimate SVD of two point sets
• See paper Arun et. al. 1987
• It's an example of the orthogonal procrustes problem

```cpp
tfe.estimateRigidTransformation(src_pc_, dst_pc_, transform_);
```

### PCL - Correlation

After subtracting each point from the mean point we have in registration/include/pcl/registration/impl/

```cpp
template <typename PointSource, typename PointTarget, typename Scalar> void pcl::registration::TransformationEstimationSVD<PointSource, PointTarget, Scalar>::getTransformationFromCorrelation (const Eigen::Matrix<Scalar, Eigen::Dynamic, Eigen::Dynamic> &cloud_src_demean, const Eigen::Matrix<Scalar, 4, 1> &centroid_src, const Eigen::Matrix<Scalar, Eigen::Dynamic, Eigen::Dynamic> &cloud_tgt_demean, const Eigen::Matrix<Scalar, 4, 1> &centroid_tgt, Matrix4 &transformation_matrix) const
{
  transformation_matrix.setIdentity ();

  // Assemble the correlation matrix H = source * target'
  Eigen::Matrix<Scalar, 3, 3> H = (cloud_src_demean * cloud_tgt_demean.transpose ()).topLeftCorner (3, 3);

  // Compute the Singular Value Decomposition
  Eigen::JacobiSVD<Eigen::Matrix<Scalar, 3, 3> > svd (H, Eigen::ComputeFullU | Eigen::ComputeFullV);
  Eigen::Matrix<Scalar, 3, 3> u = svd.matrixU ();
  Eigen::Matrix<Scalar, 3, 3> v = svd.matrixV ();

  // Compute R = V * U'
  if (u.determinant () * v.determinant () < 0)
  {
    for (int x = 0; x < 3; ++x)
      v (x, 2) *= -1;
  }

  Eigen::Matrix<Scalar, 3, 3> R = v * u.transpose ();

  // Return the correct transformation
  transformation_matrix.topLeftCorner (3, 3) = R;
  const Eigen::Matrix<Scalar, 3, 1> Rc (R * centroid_src.head (3));
  transformation_matrix.block (0, 3, 3, 1) = centroid_tgt.head (3) - Rc;
}
```
PCL - Summary

- Use of Eigen implements Arun et. al. 1987 in one main function.
- Relatively close matching of algorithm to code.
- Template notation a bit awkward, but now, not so insurmountable.
- So, using library, we gain power and benefit of very experienced library programmers.

Eigen Summary

- Header only
- Use CMake to set the include path
- Templated, so its compiled in, no link or run-time dependencies
- Simple to use linear algebra library
- Advise not to mix with GUI code
- Consider static linking as using templates anyway - ease of distribution

Using Boost

Introduction

- Boost is “…one of the most highly regarded and expertly designed C++ library projects in the world.”
- A large (121+) collection of C++ libraries
- Aim to establish standards, contribute to C++11, C++17 etc.
- Hard to use C++ without bumping into Boost at some point
- It’s heavily templated
- Many libraries header only, some require compiling.

Libraries included

- Log, FileSystem, Asio, Serialization, Pool (memory) …
- Regexp, String Algo, DateTime, …
- Math, Odeint, Graph, Polygon, Rational, …
- Each has good documentation, and tutorial, and unit tests, and is widely compiled.

Getting started

- Default build system: bjam
• Also CMake version of boost project, possible deprecated.
• Once installed, many header only libraries, so similar to Eigen.

Installing pre-compiled

• Linux:
  – sudo apt-get install boost
  – sudo apt-get install libboost1.53-dev
• Mac
  – Homebrew (brew) or Macports (port)
• Windows
  – Precompiled binaries? Probably you need to build from source.

Compiling from source

• Follow build instructions here
• Or use bigger project with it as part of build system
  – NifTK, MITK, Slicer, Gimias (medical imaging)

C++ Principles

(i.e. why introduce Boost on this course)

• Boost uses
  – Templates
  – Widespread use of:
    * Generic Programming
    * Template Meta-Programming
  – Functors

C Function Pointers - 1

• Useful if using Numerical Recipes in C
• See Wikipedia article and tutorials online

This:
```c
#include <stdio.h> /* for printf */

double cm_to_inches(double cm) {
    return cm / 2.54;
}

int main(void) {
    double (*func1)(double) = cm_to_inches;
    printf("Converting %f cm to %f inches by calling function.\n", 5.0, cm_to_inches(5.0));
    printf("Converting %f cm to %f inches by deref pointer.\n", 15.0, func1(15.0));
    return 0;
}

Produces:

Converting 5.000000 cm to 1.968504 inches by calling function.
Converting 15.000000 cm to 5.905512 inches by deref pointer.

C Function Pointers - 2

- Function pointers can be passed to functions

This:

```c
#include <stdio.h>
#include <math.h>

double integrate(double (*funcp)(double), double lo, double hi) {
    double sum = 0.0;
    for (int i = 0; i <= 100; i++)
    {
        sum += (*funcp)(i / 100.0 * (hi - lo) + lo);
    }
    return sum / 100.0;
}

int main(void) {
    double (*fp)(double) = sin;
    printf("sum(sin): %f\n", integrate(fp, 0.0, 1.0));
    return 0;
}

Produces:

sum(sin): 0.463901
```
C Function Pointers - 3
• Function pointers
– often used for callbacks, cost functions in optimisation etc.
– called by name, or dereference pointer
– are generally stateless

C++ Function Objects - 1
• We can define an object to represent a function
– Called Function Object or Functor
This:
#include <vector>
#include <algorithm>
#include <iostream>
struct IntComparator
{
bool operator()(const int &a, const int &b) const
{
return a < b;
}
};
/*
template <class RandomIt, class Compare>
void sort(RandomIt first, RandomIt last, Compare comp);
*/
int main()
{
std::vector<int> items;
items.push_back(1);
items.push_back(3);
items.push_back(2);
std::sort(items.begin(), items.end(), IntComparator());
std::cout << items[0] << "," << items[1] << "," << items[2] << std::endl;
return 0;
}
Produces:
1,2,3

75


C++ Function Objects - 2

- But function objects can
  - Have state
  - Have member variables
  - Be complex objects, created by any means
  - e.g. Cost function, similarity between two images

CMake for Boost

- If installed correctly, should be something like:

```cmake
set(Boost_ADDITIONAL_VERSIONS 1.53.0 1.54.0 1.53 1.54)
find_package(Boost 1.53.0)
if(Boost_FOUND)
  include_directories(${Boost_INCLUDE_DIRS})
endif()
```

Boost Example

- With 121+ libraries, can’t give tutorial on each one!
- Pick one small numerical example
- Illustrate the use of functors in ODE integration

Using Boost odeint

- Its a numerical example, as we are doing scientific computing!
- As with many libraries, just include right header

```cpp
#include <boost/numeric/odeint.hpp> // Include ODE solver library
  // just to check our build system found it

// See this tutorial
```

Boost odeint - 1

Given these global definitions:

```cpp
const double gam = 0.15;
typedef std::vector< double > state_type;

// This is functor class
```
Boost odeint - 2

First define a functor for the function to integrate:

class harm_osc {
    double m_gam; // class can have member variables, state etc.
public:
    harm_osc( double gam ) : m_gam(gam) { }

    // odeint integrators normally call f(x, dxdt, t)
    void operator()( const state_type &x, state_type &dxdt, const double /* t */ )
    {
        dxdt[0] = x[1];
        dxdt[1] = -x[0] - m_gam*x[1];
    }
};

// This is observer to record output, and is also a functor class

Boost odeint - 3

Define an observer to collect graph-points:

struct push_back_state_and_time
{
    std::vector< state_type >& m_states;
    std::vector< double >& m_times;

    push_back_state_and_time( std::vector< state_type > &states, std::vector< double > &times ) { }

    void operator()( const state_type &x, double t )
    {
        m_states.push_back( x );
        m_times.push_back( t );
    }
};

Boost odeint - 4

The run it:
int main(void) {

  state_type x(2);
  x[0] = 1.0; // start at x=1.0, p=0.0
  x[1] = 0.0;

  std::vector<state_type> x_vec; // vector of vectors
  std::vector<double> times; // stores each time point

  harm_osc harmonic_oscillator(0.15);
  size_t steps = boost::numeric::odeint::integrate(
    harmonic_oscillator,
    x, 0.0 , 10.0 , 0.1 ,
    push_back_state_and_time( x_vec , times ) );

  for( size_t i=0; i<=steps; i++ )
  {
    std::cout << times[i] << 't' << x_vec[i][0] << 't' << x_vec[i][1] << '
';
  }
}

Boost odeint - 4

Produces:

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>0.1</td>
<td>0.995029</td>
<td>-0.0990884</td>
</tr>
<tr>
<td>0.342911</td>
<td>0.942763</td>
<td>-0.32773</td>
</tr>
<tr>
<td>0.59639</td>
<td>0.832373</td>
<td>-0.537274</td>
</tr>
<tr>
<td>0.865439</td>
<td>0.662854</td>
<td>-0.714058</td>
</tr>
<tr>
<td>1.15445</td>
<td>0.436595</td>
<td>-0.839871</td>
</tr>
<tr>
<td>1.44346</td>
<td>0.184494</td>
<td>-0.892171</td>
</tr>
<tr>
<td>1.70128</td>
<td>-0.044646</td>
<td>-0.875731</td>
</tr>
<tr>
<td>1.9591</td>
<td>-0.262234</td>
<td>-0.80313</td>
</tr>
<tr>
<td>2.21692</td>
<td>-0.454545</td>
<td>-0.681207</td>
</tr>
<tr>
<td>2.48815</td>
<td>-0.616993</td>
<td>-0.510476</td>
</tr>
<tr>
<td>2.77613</td>
<td>-0.733844</td>
<td>-0.296963</td>
</tr>
<tr>
<td>3.06411</td>
<td>-0.7866</td>
<td>-0.0685387</td>
</tr>
<tr>
<td>3.35209</td>
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<td>0.155753</td>
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<td>3.64008</td>
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<tr>
<td>3.92806</td>
<td>-0.571123</td>
<td>0.522852</td>
</tr>
<tr>
<td>4.21604</td>
<td>-0.402601</td>
<td>0.638582</td>
</tr>
<tr>
<td>4.50402</td>
<td>-0.20875</td>
<td>0.697943</td>
</tr>
</tbody>
</table>
Why Boost for Numerics

- Broader question is
  - Why someone else’s library? Boost or some other.
- Advanced use of Template Meta Programming, Traits
  - Performance optimisations
  - Alternative implementations
    * CUDA via Thrust
    * MPI
    * etc
- You just focus on your bit

Using ITK

Introduction

- Insight Segmentation and Registration Toolkit
- Insight Journal for library additions
- Large community in medical image processing
- Deliberately no visualisation, see VTK.
C++ Principles

- Heavy use of Generic Programming
- Use of Template Meta-Programming
- Often perceived by “scientific programmers” (Matlab) as difficult
- Aim: demonstrate here, that we can now use it!
- Of particular interest
  - typedefs - make life easier
  - SmartPointers - reduce leaking memory
  - Iterators - fast image access
  - Object Factories - extensibility

Architecture Concept

- Use of pipeline of filters
- Simple to plug image processing filters together
- Sometimes difficult to manage memory for huge images

Filter Usage - 1

We work through a simple filter program. First, typedefs are aliases.

```cpp
int main(int argc, char** argv)
{
    const unsigned int Dimension = 2;
    typedef int PixelType;
    typedef itk::Image<PixelType, Dimension> ImageType;
    typedef itk::AddImageFilter<ImageType, ImageType> AddFilterType;
    typedef itk::ImageFileReader<ImageType> ImageReaderType;
    typedef itk::ImageFileWriter<ImageType> ImageWriterType;
```

Filter Usage - 2

Objects are constructed:

```cpp
ImageReaderType::Pointer reader1 = ImageReaderType::New();
ImageReaderType::Pointer reader2 = ImageReaderType::New();
AddFilterType::Pointer addFilter = AddFilterType::New();
```
ImageWriterType::Pointer writer = ImageWriterType::New();

// eg. if not using typedefs
//itk::ImageFileWriter< itk::Image<int, 2> >::Pointer writer
// = itk::ImageFileWriter< itk::Image<int, 2> >::New();

Filter Usage - 3

Pipeline is executed:

reader1->SetFileName("inputFileName1.nii");
reader2->SetFileName("inputFileName2.nii");
addFilter->SetInput(0, reader1->GetOutput());
addFilter->SetInput(1, reader2->GetOutput());
writer->SetInput(addFilter->GetOutput());
writer->SetFileName("outputFileName1.nii");
//writer->Update(); // commented out, as filenames are fake.
// and build system for lecture notes
// tries to run the program.

return 0;
}

More information on ITK Pipeline can be found in the ITK Software Guide.

Smart Pointer Intro

Lets look at some interesting features.

- Smart Pointer
  - Class, like a pointer, but ‘Smarter’ (clever)
  - Typically, once allocated will automatically destroy the pointed to object
  - Implementations vary, STL, ITK, VTK, Qt, so read the docs
- So, in each class e.g. itkAddImageFilter

```cpp
typedef AddImageFilter Self
typedef SmartPointer<Self> Pointer
```

and so, its used like

```cpp
ClassName::Pointer variableName = ClassName::New();
```
Smart Pointer Class

In theSmartPointer itself

```cpp
/** Constructor to pointer p */
SmartPointer (ObjectType *p):
    m_Pointer(p)
    { this->Register(); }

/** Destructor */
~SmartPointer ()
{    
    this->UnRegister();
    m_Pointer = ITK_SP_NULLPTR;
    }
```

and

```cpp
private:
    /** The pointer to the object referred to by this smart pointer. */
    ObjectType *m_Pointer;

    void Register()
    {
    if ( m_Pointer ) { m_Pointer->Register(); }
    }
```

General Smart Pointer Usage

- Avoid use of explicit pairs of new/delete
- Immediately assign object toSmartPointer
- Consistently (i.e. always) useSmartPointer
  - Pass (reference to)SmartPointer to function.
  - Can (but should you?) returnSmartPointer from function.
  - Don’t use raw pointer, and don’t store raw pointers to objects.
  - You can’t test raw pointer to check if object still exists.
- Object is deleted when lastSmartPointer reference goes out of scope

ITKSmartPointer

- ITK keeps reference count in itk::LightObject base class
- So, it can only be used by sub-classes of itk::LightObject
• Reference is held in the object
• Same method used in MITK, as MITK uses ITK concepts
• VTK has a SmartPointer that requires calling Delete explicitly (!!)
• STL has much clearer definition of different types of smart pointer
• Read THIS tutorial

Implementing a Filter

• ITK provides many image processing filters.
• But you can write your own easily
  – Single Threaded - override GenerateData()
  – Multi-Threaded - override ThreadedGenerateData()
• Now we see an example - thresholding, as we want to study the C++ not the image processing.

Filter Impl - 1

Basic filter:

```cpp
namespace itk
{
  template< class TInputImage, class TOutputImage = TInputImage>
  class MyThresholdFilter:
  public ImageToImageFilter< TInputImage, TOutputImage >
  {
    public:
```

Filter Impl - 2

Boilerplate nested typedefs:

```cpp
/** Standard class typedefs. */
typedef MyThresholdFilter Self;
typedef ImageToImageFilter< TInputImage, TOutputImage > Superclass;
typedefSmartPointer< Self > Pointer;
typedef typename TInputImage::PixelType InputPixelType;
typedef typename TOutputImage::PixelType OutputPixelType;

/** Method for creation through the object factory. */
itkNewMacro(Self);
```
/** Run-time type information (and related methods). */

 itkTypeMacro(ImageFilter, ImageToImageFilter);

Filter Impl - 3

Look at ITK Macros:

  itkSetMacro(Low, InputPixelType);
  itkGetMacro(Low, InputPixelType);
  itkSetMacro(High, InputPixelType);
  itkGetMacro(High, InputPixelType);

protected:
  MyThresholdFilter()
  ~MyThresholdFilter()

Filter Impl - 4

The main method:

/** Does the real work. */
virtual void GenerateData()
{
  TInputImage *inputImage = static_cast<TInputImage *>(this->ProcessObject::GetInput(0));
  TOutputImage *outputImage = static_cast<TOutputImage *>(this->ProcessObject::GetOutput(0));
  ImageRegionConstIterator<TInputImage> inputIterator = ImageRegionConstIterator<TInputImage>(inputImage, inputImage->GetLargestPossibleRegion());
  ImageRegionIterator<TOutputImage> outputIterator = ImageRegionIterator<TOutputImage>(outputImage, outputImage->GetLargestPossibleRegion());

  for (inputIterator.GoToBegin(), outputIterator.GoToBegin();
       !inputIterator.IsAtEnd() && !outputIterator.IsAtEnd();
       ++inputIterator,
       ++outputIterator)
  {
    if (*inputIterator >= m_Low && *inputIterator <= m_High)
    {
      *outputIterator = 1;
    }
    else

84
{  *outputIterator = 0;  }
}
}

private:
MyThresholdFilter(const Self &); // purposely not implemented
void operator=(const Self &); // purposely not implemented
InputPixelType m_Low;
InputPixelType m_High;
};
} // end namespace

int main(int argc, char** argv)
{
  // Not providing a real example,
  // as I don't know how to read/write images within the
  // dezy framework.
  return 0;
}

Iterators

- ITK provides many iterators
- Generic Programming means:
  - Suitable for n-dimensions
  - Suitable for all types of data
- Also, different image access concepts
  - Region of Interest
  - Random subsampling
  - No change in code
- So iterators enable you to traverse image and encapsulate the traversal
  mechanism in an iterator
- Similar concept to STL .begin(), .end()
- See ITK Software Guide

Private Constructors?

If you look at an ITK filter, you may notice for example
protected:
    AddImageFilter() {}
    virtual ~AddImageFilter() {}

private:
    AddImageFilter(const Self &);
    void operator=(const Self &);

- Copy constructor and copy assignment are private and not implemented
- Constructor and Destructor private. So how do you use?

Static New Method
You will then see

    /** Method for creation through the object factory. */
    itkNewMacro(Self);

which if you hunt for long enough, you find this snippet

#define itkSimpleNewMacro(x)
static Pointer New(void)
{
    Pointer smartPtr = ::itk::ObjectFactory< x >::Create();
    if ( smartPtr.GetPointer() == ITK_NULLPTR )
    {
        smartPtr = new x;
    }
    return smartPtr;
}

So, either this ObjectFactory creates it, or a standard new call.

ObjectFactory::Create
In itk::ObjectFactory we ask factory to CreateInstance using a char*

    static typename T::Pointer Create()
    {
        LightObject::Pointer ret = CreateInstance( typeid( T ).name() );
        return dynamic_cast< T * >( ret.GetPointer() );
    }

CreateInstance works with either a base class name, or a class name to return either a specific class, or a family of classes derived from a common base class.
Why Object Factories?

- Rather than create objects directly
- Ask a class (ObjectFactory) to do it
- This class contain complex logic, not just a new operator
- So, we can
  - dynamically load libraries from ITK_AUTOLOAD_PATH at runtime
  - Have a list/map of current classes, and provide overrides
  - i.e swap in a GPU version instead of CPU
- More dynamic variant of FactoryMethod, AbstractFactory (See GoF)

File IO Example

In itkImageFileReader.hxx

```cpp
std::list< LightObject::Pointer > allObjects =
  ObjectFactoryBase::CreateAllInstance("itkImageIOBase");
```

- We ask the factory for every class that is a sub-class of itkImageIOBase.
- Then we can ask each ImageIOBase sub-class if it can read a specific format.
- First one to reply true reads the image.
- In general case, ask ObjectFactoryBase for any class.

Object Factory List of Factories

In class ObjectFactoryBase

```cpp
class ObjectFactoryBase: public Object
{
  public:
    static std::list< ObjectFactoryBase * > GetRegisteredFactories();

This class maintains a static vector of ObjectFactoryBase. These are added programatically, via static initialisation or dynamically via the ITK_AUTO_LOAD_PATH.
PNG IO Factory

- Given `itkPNGImageIO.h/cxx` can read PNG images
- We see in `itkPNGImageIOFactory.cxx`

```cpp
PNGImageIOFactory::PNGImageIOFactory()
{
    this->RegisterOverride( "itkImageIOBase",
                           "itkPNGImageIO",
                           "PNG Image IO",
                           1,
                           CreateObjectFunction< PNGImageIO >::New() );
}
```

So, PNG factory says it implements a type of `itkImageIOBase`, will return an `itkPNGImageIO`, and instantiates a function object that calls the right constructor.

ObjectFactory Summary

- ObjectFactory defines a static vector of ObjectFactory
- ObjectFactory objects loaded:
  - Directly named in code at compile time
  - Via static initialisers when a dynamic library is loaded
  - Or from `ITK_AUTOLOAD_PATH`
- ObjectFactory returns one/all classes that implement a given class
- Static New method now asks factory for a class.
- So, you can override any ITK class.
- Why is above example not an infinite loop?

ITK Summary

- Pipeline architecture for most filters
- Also includes a registration framework (see ITK Software Guide)
- Smart Pointers - reference counting, automatic deletion
- Static New method with ObjectFactory to enable overriding any class at runtime
- Dynamic loading via `ITK_AUTOLOAD_PATH`
- Pipeline architecture - easy to prototype, once you know C++
- Write your own filter, unit test, generalise to n-dimension, of n-vectors.
- Easy to extend to multi-threading
Summary

Learning Objectives

• Use well written libraries
• Understand enough C++ for some common C++ libraries

Further Reading

Please refer to the following:

• Short Eigen Tutorial
• Longer Eigen Tutorial
• Boost Tutorials for each library
• ITK Software Guide
• Simple ITK: Template free, wrapper, python, paper
Chapter 5

Patterns

The Story So Far

- Git
- CMake
- Use libraries
- C++ features
- Seeing those C++ in libraries

Today’s Lesson

- How to assemble classes
- Beginner mistakes
  - 1 class - all functionality
  - Deep inheritance trees
- You need
  - How to assemble / organise classes

RAII Pattern

What is it?

- Resource Allocation Is Initialisation (RAII)
• Obtain all resources in constructor
• Release them all in destructor

Why is it?

• Guaranteed fully initialised object once constructor is complete
• Objects on stack are guaranteed to be destroyed when an exception is thrown and stack is unwound
  – Including smart pointers to objects

Example

• You may already be using it: STL example
• Another example

Program To Interfaces

Why?

• In research code we often “just start hacking”
• You tend to mix interface and implementation
• Results in client of a class having implicit dependency on the implementation
• So, define a pure virtual class, not for inheritance, but for clean API

Example

```cpp
#include <memory>
#include <vector>
#include <string>

class DataPlayerI {
public:
    virtual void StartPlaying() = 0;
    virtual void StopPlaying() = 0;
};

class FileDataPlayer : public DataPlayerI {
public:
    FileDataPlayer(const std::string& fileName){}; // opens file (RAII)
```
-FileDataPlayer(); // releases file (RAII)
public:
  virtual void StartPlaying() {};
  virtual void StopPlaying() {};
};

class Experiment {
public:
  Experiment(DataPlayerI *d) { m_Player.reset(d); } // takes ownership
  void Run() {};
  std::vector<std::string> GetResults() const {};
private:
  std::unique_ptr<DataPlayerI> m_Player;
};

int main(int argc, char** argv)
{
  FileDataPlayer fdp(argv[1]); // Or some class WebDataPlayer derived from DataPlayerI
  Experiment e(&fdp);
  e.Run();

  // etc.
}

Comments

- Useful between sub-components of a system
  - GUI front end, Web back end
  - Logic and Database
- Is useful in general to force loose connections between areas of code
  - e.g. different libraries that have different dependencies
  - e.g. camera calibration depends on OpenCV
  - define an interface that just exports standard types
  - stops the spread of dependencies
  - Lookup Pimpl idiom

Inheritance

Don’t overuse Inheritance

- Inheritance is not just for saving duplication
• It MUST represent derived/related types
• Derived class must truly represent ‘is-a’ relationship
• eg ‘Square’ is-a ‘Shape’
• Deep inheritance hierarchies are almost always wrong
• If something ‘doesn’t quite fit’ check your inheritance

Surely Its Simple?

• Common example: Square/Rectangle problem, here

```cpp
#include <iostream>

class Rectangle {
public:
    Rectangle() : m_Width(0), m_Height(0) {};
virtual ~Rectangle(){};
    int GetArea() const { return m_Width*m_Height; }
virtual void SetWidth(int w) { m_Width=w; }
virtual void SetHeight(int h) { m_Height=h; }
protected:
    int m_Width;
    int m_Height;
};

class Square : public Rectangle {
public:
    Square(){};
virtual ~Square(){};
    virtual void SetWidth(int w) { m_Width=w; m_Height=w; }
    virtual void SetHeight(int h) { m_Width=h; m_Height=h; }
};

int main()
{
    Rectangle *r = new Square();
r->SetWidth(5);
r->SetHeight(6);
    std::cout << "Area = " << r->GetArea() << std::endl;
}
```

Liskov Substitution Principal

• Wikipedia
• “if S is a subtype of T, then objects of type T may be replaced with objects of type S without altering any of the desirable properties of that program”
• Can something truly be substituted?
• If someone else filled a vector of type T, would I care what type I have?
• Look for:
  – Preconditions cannot be strengthened
  – Postconditions cannot be weakened
  – Invariants preserved

What to Look For

• If you have:
  – Methods you don’t want to implement in derived class
  – Methods that don’t make sense in derived class
  – Methods that are unnecessary in derived class
  – If you have a list of something, and someone else swapping a derived type would cause problems
• Then you have probably got your inheritance wrong

Composition Vs Inheritance

• Lots of Info online eg. wikipedia
• In basic OO Principals
  – ‘Has-a’ means ‘pointer or reference to’
  – eg.Car has-a Engine
• But there is also:
  – Composition: Strong ‘has-a’. Component parts are owned by thing pointing to them.
  – Aggregation: Weak ‘has-a’. Component part has its own lifecycle.
  – Association: General term, referring to either composition or aggregation, just a ‘pointer-to’

Examples

• House ‘has-a’ set of Rooms. Destroying House, you destroy all Room objects. No point having a House on its own.
• Department ‘has-a’ Professor. If a department is shutdown (deleted), Professor should not be deleted. Needs assigning to another department.
But Why?

- Good article: Choosing Composition or Inheritance
- Inheritance has much tighter definition than you realise
- Composition is more flexible

Dependency Injection

Construction

- What could be wrong with this:

```cpp
#include <memory>

class Bar {
};

class Foo {
public:
    Foo()
    {
        m_Bar = new Bar();
    }

private:
    Bar* m_Bar;
};

int main()
{
    Foo a;
}
```

Unwanted Dependencies

- If constructor instantiates class directly:
  - Hard-coded class name
  - Duplication of initialisation code
Dependency Injection

- Read Martin Fowler’s Inversion of Control Containers and the Dependency Injection Pattern
- Type 2 - Constructor Injection
- Type 3 - Setter Injection

Constructor Injection Example

```cpp
#include <memory>

class Bar {
};

class Foo {
public:
    Foo(Bar* b)
        : m_Bar(b)
    {
    }

private:
    Bar* m_Bar;
};

int main()
{
    Bar b;
    Foo a(&b);
}
```

Setter Injection Example

```cpp
#include <memory>

class Bar {
};

class Foo {
public:
    Foo()
    {
    }
```
void SetBar(Bar *b) { m_Bar = b; }

private:
    Bar* m_Bar;
};

int main()
{
    Bar b;
    Foo a();
    a.SetBar(&b);
}

Question: Which is better?

Advantages of Dependency Injection

• Using Dependency Injection
  – Removes hard coding of new ClassName
  – Creation is done outside class, so class only uses public API
  – Leads towards fewer assumptions in the code

Construction Patterns

Constructional Patterns

• Other methods include
  – See Gang of Four book
  – Strategy Pattern
  – Factory Pattern
  – Abstract Factory Pattern
  – Builder Pattern

• Also look up Service Locator Pattern

Managing Complexity

• Rather than monolithic code (bad)
• We end up with many smaller classes (good)
• So, it's more flexible (good)
• It does look more complex at first (don't panic)
• You get used to thinking in small objects

Summary

Putting It Together - Subsystems

• Program to interfaces
• Inject dependencies
• Always make fully initialised object
• Always use smart pointers
• Use exceptions for errors
• RAII very useful
• Results in a more flexible, extensible, robust design

Putting It Together - OOP

• Prefer composition over inheritance
• Use constructional patterns to assemble code

Object Oriented Review

C-style Programming

• Procedural programming
• Pass data to functions

Function-style Example

```c
double compute_similarity(double *imag1, double* image2, double *params)
{
    // stuff
}
```

```c
double calculate_derivative(double* params)
{
    // stuff
}
```
double convert_to_decimal(double numerator, double denominator)
{
    // stuff
}

Disadvantages

- Can get out of hand as program size increases
- Can’t easily describe relationships between bits of data
- Relies on method documentation, function and variable names
- Can’t easily control/(enforce control of) access to data

C Struct

- So, in C, the struct was invented
- Basically a class without methods
- This at least provides a logical grouping

Struct Example

struct Fraction {
    int numerator;
    int denominator;
};

double convertToDecimal(const Fraction& f)
{
    return f.numerator/static_cast<double>(f.denominator);
}

C++ Class

- C++ provides the class to enhance the language with user defined types
- Once defined, use types as if native to the language

Abstraction

- C++ class mechanism enables you to define a type
  - independent of its data
– independent of its implementation
– class defines concept or blueprint
– instantiation creates object

Abstraction - Grady Booch

“An abstraction denotes the essential characteristics of an object that distinguish it from all other kinds of objects and thus provide crisply defined conceptual boundaries, relative to the perspective of the viewer.”

Class Example

class Atom {
public:
    Atom();
    ~Atom();
    int GetAtomicNumber();
    double GetAtomicWeight();
};

int main(int argc, char** argv)
{
    Atom a;
}

Encapsulation

• Encapsulation is:
  – Bundling together methods and data
  – Restricting access, defining public interface

• Describes how you correctly use something

Public/Private/Protected

• For class methods/variables:
  – private: only available in this class
  – protected: available in this class and derived classes
  – public: available to anyone with access to the object

• (public, protected, private inheritance comes later)
Class Example

```cpp
// Defining a User Defined Type.
class Fraction {

public: // access control
    // How to create
    Fraction();
    Fraction(const int &num, const int &denom);

    // How to destroy
    ~Fraction();

    // How to access
    int numerator() const;
    int denominator() const;

    // What you can do
    const Fraction operator+(const Fraction &another);

private: // access control
    // The data
    int m_Numerator;
    int m_Denominator;
};
```

Inheritance

- Used for:
  - Defining new types based on a common type
- Careful:
  - Beware - “Reduce code duplication, less maintenance”
  - Types in a hierarchy MUST be related
  - Don’t over-use inheritance
  - We will cover other ways of object re-use

Class Example

```cpp
class Shape {
public:
    Shape();
    void setVisible(const bool &isVisible) { m_IsVisible = isVisible; }
};
```
virtual void rotate(const double &degrees) = 0;
virtual void scale(const double &factor) = 0;
// + other methods
private:
bool m_IsVisible;
unsigned char m_Colour[3]; // RGB
double m_CentreOfMass[2];
};

class Rectangle : public Shape {
public:
Rectangle();
virtual void rotate(const double &degrees);
virtual void scale(const double &factor);
// + other methods
private:
double m_Corner1[2];
double m_Corner2[2];
};

class Circle : public Shape {
public:
Circle();
virtual void rotate(const double &degrees);
virtual void scale(const double &factor);
// + other methods
private:
float radius;
};

Polymorphism

- Several types:
  - (normally) “subtype”: via inheritance
  - “parametric”: via templates
  - “ad hoc”: via function overloading
- Common interface to entities of different types
- Same method, different behaviour

Class Example

#include "shape.h"
int main(int argc, char** argv)
{  
  Circle c1;  
  Rectangle r1;  
  Shape *s1 = &c1;  
  Shape *s2 = &r1;  

  // Calls method in Shape (as not virtual)
  bool isVisible = true;
  s1->setVisible(isVisible);
  s2->setVisible(isVisible);

  // Calls method in derived (as declared virtual)
  s1->rotate(10);
  s2->rotate(10);
}

Essential Reading

General Advice

- Do a version control course such as Software Carpentry, or MPHYG001.
- Contribute to online open-source project
- For your repository - pick a coding style, and stick to it

Daily Reading

- Every C++ developer should keep repeatedly reading at least:
  - Effective C++, Meyers  
  - More Effective C++, Meyers  
  - Effective STL, Meyers

Additional Reading

- Recommended
  - Accelerated C++, Koenig, Moo.
  - Design Patterns (1994), Gamma, Help, Johnson and Vlassides
  - Modern C++ Design, Andrei Alexandrescu
C++ tips

Numbers in brackets refer to Scott Meyers “Effective C++” book.

- Top C++ tips such as:
  - Declare data members private (22)
  - Use `const` whenever possible (3)
  - Make interfaces easy to use correctly and hard to use incorrectly (18)
  - Avoid returning “handles” to object internals (28)
  - Initialise objects properly. Throw exceptions from constructors. Fail early. (4)

OO tips

- More general OO tips such as:
  - Never throw exceptions from destructors
  - Prefer non-member non-friend functions to member functions (better encapsulation) (23)
  - Make sure public inheritance really models “is-a” (32)
  - Learn alternatives to polymorphism (Template Method, Strategy) (35)
  - Model “has-a” through composition (38)

Using Templates

What Are Templates?

- C++ templates allow functions/classes to operate on generic types.
- See: Generic Programming.
- Write code, where ‘type’ is provided later
- Types instantiated at compile time, as they are needed
- (Remember, C++ is strongly typed)

You May Already Use Them!

You probably use them already. Example type (class):

```cpp
std::vector<int> myVectorInts;
```

Example algorithm: C++ sort
std::sort(myVectorInts.begin(), myVectorInts.end());

Aim: Write functions, classes, in terms of future/other/generic types, type provided as parameter.

Why Are Templates Useful?

- Generic programming:
  - not pre-processor macros
  - so maintain type safety
  - separate algorithm from implementation
  - extensible, optimisable via Template Meta-Programming (TMP)

Book

- You should read “Modern C++ Design”
- 2001, but still excellent text on templates, meta-programming, policy based design etc.
- This section of course, gives basic introduction for research programmers

Are Templates Difficult?

- Some say: notation is ugly
  - Does take getting used to
  - Use typedef to simplify
- Some say: verbose, confusing error messages
  - Nothing intrinsically difficult
  - Take small steps, compile regularly
  - Learn to think like a compiler
- Code infiltration
  - Use sparingly
  - Hide usage behind clean interface

Why Templates in Research?

- Generalise 2D, 3D, n-dimensions, (e.g. ITK)
- Test numerical code with simple types, apply to complex/other types
- Several useful libraries for research
Why Teach Templates?

- Standard Template Library uses them
- More common in research code, than business code
- In research, more likely to ‘code for the unknown’
- Boost, Qt, EIGEN uses them

Function Templates

Function Templates Example

• Credit to www.cplusplus.com

// function template
#include <iostream>
using namespace std;

template <class T> // class/typename
T sum (T a, T b)
{
    T result;
    result = a + b;
    return result;
}

int main () {
    int i=5, j=6;
    double f=2.0, g=0.5;
    cout << sum<int>(i,j) << ' 
';
    cout << sum<double>(f,g) << ' 
';
    return 0;
}

• And produces this output when run

11
2.5

Why Use Function Templates?

• Instead of function overloading
  – Reduce your code duplication
– Reduce your maintenance
– Reduce your effort
– Also see this Additional tutorial.

Language Definition 1

• From the language reference

```
template < parameter-list > function-declaration

• so

template < class T > // note 'class'
void MyFunction(T a, T b)
{
  // do something
}

• or

template < typename T1, typename T2 > // note 'typename'
T1 MyFunctionTwoArgs(T1 a, T2 b)
{
  // do something
}
```

Language Definition 2

• Also
  – Can use class or typename.
  – I prefer typename.
  – Template parameter can apply to references, pointers, return types, arrays etc.

Default Argument Resolution

• Given:

```
double GetAverage<typeName T>(const std::vector<T>& someNumbers);
```

• then:
std::vector<double> myNumbers;
double result = GetAverage(myNumbers);

• will call:

double GetAverage<double>(const std::vector<double>& someNumbers);

• So, if function parameters can inform the compiler uniquely as to which
  function to instantiate, its automatically compiled.

Explicit Argument Resolution - 1

• However, given:

double GetAverage<typename T>(const T& a, const T& b);

• and:

int a, b;
int result = GetAverage(a, b);

• But you don’t want the int version called (due to integer division perhaps),
you can:

double result = GetAverage<double>(a, b);

Explicit Argument Resolution - 2

• equivalent to

GetAverage<double>(static_cast<double>(a), static_cast<double>(b));

• i.e. name the template function parameter explicitly.
• Cases for Explicit Template Argument Specification
  – Force compilation of a specific version (eg. int as above)
  – Also if method parameters do not allow compiler to deduce anything
    eg. PrintSize() method.
Beware of Code Bloat

- Given:

```cpp
global GetMax<typename T1, typename T2>(const &T1, const &T2);
```

- and:

```cpp
double r1 = GetMax(1, 2);
double r2 = GetMax(1, 2.0);
double r3 = GetMax(1.0, 2.0);
```

- The compiler will generate 3 different max functions.
- Be Careful
  - Executables/libraries get larger
  - Compilation time will increase
  - Error messages get more verbose

Two Stage Compilation

- Basic syntax checking (e.g. brackets, semi-colon, etc), when `#include’d`
- But only compiled when instantiated (e.g. check existence of + operator).

Instantiation

- Object Code is only really generated if code is used
- Template functions can be
  - .h file only
  - .h file that includes separate .cxx/.txx/.hxx file (e.g. ITK)
  - .h file and separate .cxx/.txx file (sometimes by convention a .hpp file)
- In general
  - Most libraries/people prefer header only implementations

Explicit Instantiation - 1

- Language Reference [here](#)
- Microsoft Example
• Given (library) header:

```cpp
#ifndef explicitInstantiation_h
#define explicitInstantiation_h
template <typename T> void f(T s);
#endif
```

• Given (library) implementation:

```cpp
#include <iostream>
#include <typeinfo>
#include "explicitInstantiation.h"
template<typename T>
void f(T s)
{
    std::cout << typeid(T).name() << " " << s << \\

    template void f<double>(double);  // instantiates f<double>(double)
template void f<char>(char);      // instantiates f<char>(char), template argument deduced
template void f<int>();          // instantiates f<int>(int), template argument deduced
```

Explicit Instantiation - 2

• Given client code:

```cpp
#include <iostream>
#include "explicitInstantiation.h"

int main(int argc, char** argv)
{
    std::cout << "Matt, double 1.0=" << std::endl;
    f(1.0);
    std::cout << "Matt, char a=" << std::endl;
    f('a');
    std::cout << "Matt, int 2=" << std::endl;
    f(2);
    // std::cout << "Matt, float 3.0=" << std::endl;
    // f<float>(static_cast<float>(3.0));  // compile error
}
```

• We get:

Matt, double 1.0=

Matt, char a=

Matt, int 2=

Matt, float 3.0=

110
Explicit Instantiation - 3

- Explicit Instantiation:
  - Forces instantiation of the function
  - Must appear after the definition
  - Must appear only once for given argument list
  - Stops implicit instantiation
- So, mainly used by compiled library providers
- Clients then \#include header and link to library

Linking CXX executable explicitInstantiationMain.x
Undefined symbols for architecture x86_64:
  "void f<float>(float)", referenced from:

Implicit Instantiation - 1

- Instantiated as they are used
- Normally via \#include header files.
- Given (library) header, that contains implementation:

```cpp
#ifndef explicitInstantiation_h
#define explicitInstantiation_h
#include <iostream>
#include <typeinfo>
template <typename T> void f(T s) { std::cout << typeid(T).name() << " " << s << '\n'; }
#endif
```

Implicit Instantiation - 2

- Given client code:

```cpp
#include <iostream>
#include "implicitInstantiation.h"
```
```cpp
int main(int argc, char** argv)
{
    std::cout << "Matt, double 1.0=" << std::endl;
    f(1.0);
    std::cout << "Matt, char a=" << std::endl;
    f('a');
    std::cout << "Matt, int 2=" << std::endl;
    f(2);
    std::cout << "Matt, float 3.0=" << std::endl;
    f<float>(static_cast<float>(3.0)); // no compile error
}
```

- We get:

```
Matt, double 1.0=
d 1
Matt, char a=
c a
Matt, int 2=
i 2
Matt, float 3.0=
f 3
```

## Class Templates

### Class Templates Example - 1

- If you understand template functions, then template classes are easy!
- Refering to this tutorial, an example:

Header:

```cpp
template <typename T> class MyPair {
    T m_Values[2];

public:
    MyPair(const T &first, const T &second);
    T getMax() const;
};

#include "pairClassExample.cc"
```
Class Templates Example - 2

Implementation:

```cpp
template <typename T>
MyPair<T>::MyPair(const T& first, const T& second)  
{
    m_Values[0] = first;
    m_Values[1] = second;
}

template <typename T>
T
MyPair<T>::getMax() const
{
    if (m_Values[0] > m_Values[1])
        return m_Values[0];
    else
        return m_Values[1];
}
```

Class Templates Example - 3

Usage:

```cpp
#include "pairClassExample.h"
#include <iostream>

int main(int argc, char** argv)
{
    MyPair<int> a(1,2);
    std::cout << "Max is:" << a.getMax() << std::endl;
}
```

Quick Comments

- Implementation, 3 uses of parameter T
- Same Implicit/Explicit instantiation rules
- Note implicit requirements, eg. operator >
  - Remember the 2 stage compilation
  - Remember code not instantiated until its used
  - Take Unit Testing Seriously!
Template Specialisation

- If template defined for type T
- Full specialisation - special case for a specific type eg. char
- Partial specialisation - special case for a type that still templates, e.g. T*

```cpp
template <typename T> class MyVector {
    template <> class MyVector<char> { // full specialisation
        template <typename T> MyVector<T*> { // partial specialisation
```

Nested Types

In libraries such as ITK, we see:

```cpp
template< typename T, unsigned int NVectorDimension = 3 >
class Vector:public FixedArray< T, NVectorDimension >
{
    public:
    // various stuff
    typedef T ValueType;
    // various stuff
    T someMemberVariable;

```

- typedef is just an alias
- using nested typedef, must be qualified by class name
- can also refer to a real variable

Smart Pointers

Use of Raw Pointers

- Given a pointer passed to a function

```cpp
void DoSomethingClever(int *a)
{
    // write some code
}
```

- How do we use the pointer?
- What problems are there?
Problems with Raw Pointers

- From “Effective Modern C++”, Meyers, p117.
  - If you are done, do you destroy it?
  - How to destroy it? Call `delete` or some method first: `a->Shutdown();`
  - Single object or array?
  - `delete` or `delete[]`?
  - How to ensure the whole system only deletes it once?
  - Is it dangling, if I don’t delete it?

Use Smart Pointers

- `new/delete` on raw pointers not good enough
- So, use Smart Pointers
  - automatically delete pointed to object
  - explicit control over sharing
  - i.e. smarter
- Smart Pointers model the “ownership”

Further Reading

- Notes here are based on these:
  - David Kieras online paper
  - “Effective Modern C++”, Meyers, ch4

Standard Library Smart Pointers

- Here we teach Standard Library
  - `std::unique_ptr` - models `has-a` but also unique ownership
  - `std::shared_ptr` - models `has-a` but shared ownership
  - `std::weak_ptr` - temporary reference, breaks circular references

Stack Allocated - No Leak.

- To recap:
```cpp
#include "Fraction.h"
int main() {
    Fraction f(1,4);
}
```

- Gives:

I'm being deleted
- So stack allocated objects are deleted, when stack unwinds.

**Heap Allocated - Leak.**
- To recap:

```cpp
#include "Fraction.h"
int main() {
    Fraction *f = new Fraction(1,4);
}
```

- Gives:

- So heap allocated objects are not deleted.
- Its the pointer (stack allocated) that’s deleted.

**Unique Ptr - Unique Ownership**
- So:

```cpp
#include "Fraction.h"
#include <memory>
int main() {
    std::unique_ptr<Fraction> f(new Fraction(1,4));
}
```

- Gives:

I'm being deleted
- And object is deleted.
- Is that it?
Unique Ptr - Move?

- Does move work?

```cpp
#include "Fraction.h"
#include <memory>
#include <iostream>

int main() {
    std::unique_ptr<Fraction> f(new Fraction(1,4));
    // std::unique_ptr<Fraction> f2(f); // compile error
    std::cerr << "f=" << f.get() << std::endl;

    std::unique_ptr<Fraction> f2;
    // f2 = f; // compile error
    // f2.reset(f.get()); // bad idea

    f2.reset(f.release());
    std::cout << "f=" << f.get() << ", f2=" << f2.get() << std::endl;

    f = std::move(f2);
    std::cout << "f=" << f.get() << ", f2=" << f2.get() << std::endl;
}
```

- Gives:

```
f=0, f2=0x1aad010
f=0x1aad010, f2=0
I'm being deleted
```

- We see that API makes difficult to use incorrectly.

Unique Ptr - Usage 1

- Forces you to think about ownership
  - No copy constructor
  - No assignment
- Consequently
  - Can’t pass pointer by value
  - Use move semantics for placing in containers
Unique Ptr - Usage 2

- Put raw pointer STRAIGHT into unique_ptr
- see std::make_unique in C++14.

```cpp
#include "Fraction.h"
#include <memory>
int main() {
    std::unique_ptr<Fraction> f(new Fraction(1, 4));
}
```

Shared Ptr - Shared Ownership

- Many pointers pointing to same object
- Object only deleted if no pointers refer to it
- Achieved via reference counting

Shared Ptr Control Block

- Won’t go to too many details:
- From “Effective Modern C++”, Meyers, p140

Shared Ptr - Usage 1

- Place raw pointer straight into shared_ptr
- Pass to functions, reference or by value
- Copy/Move constructors and assignment all implemented

Shared Ptr - Usage 2

```cpp
#include "Fraction.h"
#include <memory>
#include <iostream>
void divideBy2(const std::shared_ptr<Fraction>& f) {
    f->denominator *= 2;
}
void multiplyBy2(const std::shared_ptr<Fraction> f) {
    f->numerator *= 2;
}
```
int main() {
    std::shared_ptr<Fraction> f1(new Fraction(1,4));
    std::shared_ptr<Fraction> f2 = f1;
    divideBy2(f1);
    multiplyBy2(f2);
    std::cout << "Value= " << f1->numerator << "/" << f1->denominator << std::endl;
    std::cout << "f1= " << f1.get() << ", f2= " << f2.get() << std::endl;
}

Shared Ptr - Usage 3

- Watch out for exceptions.
- “Effective Modern C++”, Meyers, p140

#include "Fraction.h"
#include <memory>
#include <stdexcept>
#include <vector>
int checkSomething(const std::shared_ptr<Fraction>& f, const int& i) {
    // whatever.
}
int computeSomethingFirst()
{
    // what if this throws?
}
int main()
{
    std::vector<std::shared_ptr<Fraction>> spaceForLotsOfFractions;
    int result = checkSomething(std::shared_ptr<Fraction>(new Fraction(1,4)),
                                  computeSomethingFirst());
}

Shared Ptr - Usage 4

- Prefer std::make_shared
- Exception safe

#include "Fraction.h"
#include <memory>
```cpp
#include <stdexcept>
#include <vector>
int checkSomething(const std::shared_ptr<Fraction>& f, const int& i)
{
    // whatever.
}
int computeSomethingFirst()
{
    // what if this throws?
}
int main()
{
    std::vector<std::shared_ptr<Fraction> > spaceForLotsOfFractions;
    int result = checkSomething(std::make_shared<Fraction>(1,4),
                                computeSomethingFirst());
}

Weak Ptr - Why?

- Like a shared pointer, but doesn’t actually own anything
- Use for example:
  - Caches
  - Break circular pointers
- Limited API
- Not terribly common as most code ends up as hierarchies

Weak Ptr - Example

- See David Kiers online paper

```
// Needs converting to shared, and checking
std::shared_ptr<Fraction> s2 = w1.lock();
if (s2)
{
    std::cout << "Object w1 exists=\" " << s2->numerator << "/" << s2->denominator << std::endl;
}

// Or, create shared, check for exception
std::shared_ptr<Fraction> s3(w2);
std::cout << "Object must exists=\" " << s3->numerator << "/" << s3->denominator << std::endl;

Final Advice

- Benefits of immediate, fine-grained, garbage collection
- Just ask Scott Meyers!
  - Use unique_ptr for unique ownership
  - Easy to convert unique_ptr to shared_ptr
  - But not the reverse
  - Use shared_ptr for shared resource management
  - Avoid raw new - use make_shared, make_unique
  - Use weak_ptr for pointers that can dangle (cache etc)

Comment on Boost

- Boost has become a sandbox for standard C++
- Boost features become part of standard C++, (different name space)
- So if you are forced to use old compiler
  - You could use boost - lecture 5.

Intrusive Vs Non-Intrusive

- Intrusive - Base class maintains a reference count eg. ITK
- Non-intrusive
  - std::unique_ptr
  - std::shared_ptr
  - std::weak_ptr
  - works for any class
ITK (intrusive) Smart Pointers

class MyFilter : // Other stuff
{
public:
    typedef MyFilter Self
    typedef SmartPointer<Self> Pointer;
    itkNewMacro(Self);
protected:
    MyFilter();
    virtual ~MyFilter();
};

double someFunction(MyFilter::Pointer p)
{
    // stuff
}

int main()
{
    MyFilter::Pointer p = itk::MyFilter::New();
}

Conclusion for Smart Pointers

• Default to standard library, check compiler
• Lots of other Smart Pointers
  – Boost (use STL).
  – ITK
  – VTK
  – Qt Smart Pointers
• Don’t be tempted to write your own
• Always read the manual
• Always consistently use it

Error Handling

Exceptions

• Exceptions are the C++ or Object Oriented way of Error Handling
• Read this example
Exception Handling Example

```cpp
#include <stdexcept>
#include <iostream>
bool someFunction() { return false; }

int main()
{
    try
    {
        bool isOK = false;
        isOK = someFunction();
        if (!isOK)
        {
            throw std::runtime_error("Something is wrong");
        }
    }
    catch (std::exception& e)
    {
        std::cerr << "Caught Exception:" << e.what() << std::endl;
    }
}
```

What’s the Point?

- Have separated error handling logic from application logic
- First, lets look at C-style return codes

Error Handling C-Style

```cpp
int foo(int a, int b)
{
    // stuff

    if(some error condition)
    {
        return 1;
    } else if (another error condition) {
        return 2;
    } else {
        return 0;
    }
}
```
void caller(int a, int b)
{
    int result = foo(a, b);
    if (result == 1) // do something
    else if (result == 2) // do something difference
    else
    {
        // All ok, continue as you wish
    }
}

Outcome

- Can be perfectly usable
- Depends on depth of function call stack
- Depends on complexity of program
- If deep/large, then can become unwieldy

Error Handling C++ Style

#include <stdexcept>
#include <iostream>

int ReadNumberFromFile(const std::string& fileName)
{
    if (fileName.length() == 0)
    {
        throw std::runtime_error("Empty fileName provided");
    }

    // Check for file existence etc. throw io errors.
    // do stuff
    return 2; // returning dummy number to force error
}

void ValidateNumber(int number)
{
    if (number < 3)
    {
        throw std::logic_error("Number is < 3");
    }
if (number > 10)
{
    throw std::logic_error("Number is > 10");
}

int main(int argc, char** argv)
{
    try
    {
        if (argc < 2)
        {
            std::cerr << "Usage: " << argv[0] << " fileName" << std::endl;
            return EXIT_FAILURE;
        }

        int myNumber = ReadNumberFromFile(argv[1]);
        ValidateNumber(myNumber);

        // Compute stuff.

        return EXIT_SUCCESS;
    }
    catch (std::exception& e)
    {
        std::cerr << "Caught Exception:" << e.what() << std::endl;
    }
}

Outcome

- Code that throws does not worry about the catcher
- Exceptions are classes, can carry data
- Exceptions can form class hierarchy

Consequences

- More suited to larger libraries of re-usable functions
- Many different catchers, all implementing different error handling
- Lends itself to layered software (draw diagram)
- Generally scales better, more flexible
Practical Tips For Exception Handling

- Decide on error handling strategy at start
- Use it consistently
- Create your own base class exception
- Derive all your exceptions from that base class
- Stick to a few obvious classes, not one class for every single error

More Practical Tips For Exception Handling

- Look at C++ standard classes and tutorial
- An exception macro may be useful, e.g. `mitk::Exception` and `mth::Throw()`
- Beware side-effects
  - Perform validation before updating any member variables
Chapter 6

HPC Concepts

High Performance Computing Overview

Faster Code

How can we make a computer go faster?

- Do instructions faster
- Do more instructions at once

Doing Instructions Faster

- How?
  - Reduce wait time. I.E cache misses, data loading, etc.
    * Can be complex and hardware dependant
    * Often automated
  - Write machine code rather than interpreted code
    * JIT does this for you
    * Allows for cool tracing optimisations
  - Increase CPU speed
    * CPU instructions happen faster
    * Data might not move faster

Doing More Instructions at Once

Image variant, original by Benjamin D. Esham / Wikimedia Commons, CC BY-SA 3.0
Figure 6.1: Dining Philosophers
Concurrency Problems

- Deadlock
  - All philosophers have one fork each and are waiting for another to appear
- Starvation
  - A philosopher never gets to eat
- Race conditions
  - A bug where the order of operations matters. This is usually either an algorithm design bug, or two philosophers both using the same fork at the same time.

Some Better Names

- Philosopher: thread, process, task, etc.
- Fork: resource, lock, etc.
- Spaghetti: Critical section.

Flynn’s Taxonomy

- SISD: You doing something on your own
- SIMD: You and your clones doing the same things on different inputs
- MISD: You and your friends doing different things with the same inputs
- MIMD: You and your friends doing unrelated things
- SIMT: You and your clones doing the same things on different inputs, but you can choose to skip steps

Amdahl’s Law

You cannot make your code go faster than the non-parallel bit.

Background Reading

- This section is based on background reading outside the classroom
- This will save time in the classroom for practical work
- But you do need to know this content
- Read:
  - Some background history on Wikipedia
  - Blaise Barney’s overview of parallel computing
Essential Reading

- For the exam you will need:
  - Amdahl’s law
  - Flynn’s Taxonomy

- For tutorial’s and practical work you will need:
  - Use of unix shell
  - Submitting jobs on a cluster
  - See RITS HPC Training
  - We’ll recap this now

Aim

For the remainder of the course, we need to develop

- Skills to run jobs on a cluster, e.g. Legion.
- A locally installed development environment, so you can develop
- Familiarity with new technologies, OpenMP, MPI, Accelerators, Cloud, so you can make a reasoned choice
Chapter 7

Shared memory parallelism

Shared Memory Parallelism

OpenMP

- Shared memory only parallelization
- Useful for parallelization on a single cluster node
- MPI next week for inter-node parallelization
- Can write hybrid code with both OpenMP and MPI

About OpenMP

- Extensions of existing programming languages
- Standardized by international committee
- Support for Fortran, C and C++
- C/C++ uses the same syntax
- Fortran is slightly different

How it works

- Thread based parallelization
- A master thread starts executing the code.
- Sections of the code is marked as parallel
  - A set of threads are forked and used together with the master thread
  - When the parallel block ends the threads are killed or put to sleep
Typical use cases

- A loop with independent iterations
- Hopefully a significant part of the execution time
- More complicated if an iterations have dependencies

OpenMP

OpenMP basic syntax

- Annotate code with #pragma omp ...
  - This instruct the compiler in how to parallize the code
  - #pragmas are a instructions to the compiler
  - Not part of the language
  - i.e. #pragma once alternative to include guards
  - Compiler will usually ignore pragmas that it doesn’t understand
  - All OpenMP pragmas start with #pragma omp

- OpenMP must typically be activated when compiling code

OpenMP library

- OpenMP library:
  - It provides utility functions.
  - omp_get_num_threads() ...
  - Use with #include <omp.h>

Compiler support

OpenMP is supported by most compilers.

- OpenMP must typically be activated with a command line flags at compile time. Different for different compilers. Examples:
  - Intel, Linux, Mac -openmp
  - Intel, Windows /Qopenmp
  - GCC/G++, -fopenmp

Clang OpenMP is slightly different from other OpenMP code, and as such may not be able to work together (as clang does not support 128-bit floats).
CMake Support

CMake knows how to deal with OpenMP, mostly:

```cmake
find_package(OpenMP)

add_program(my_threaded_monster main.cc)
if(OPENMP_FOUND)
  target_compile_options(my_threaded_monster PUBLIC "${OpenMP_CXX_FLAGS}"
  target_link_libraries(my_threaded_monster PUBLIC "${OpenMP_CXX_FLAGS}")
endif()
```

Hello world

```c
#include <iostream>
#include <omp.h>

int main(int argc, char ** argv)
{
  #pragma omp parallel
  {
    int threadnum = 0;
    int numthreads = 0;
    threadnum = omp_get_thread_num();
    numthreads = omp_get_num_threads();
    std::cout << "Hello World, I am " << threadnum << " of " << numthreads << std::endl;
  }
}
```

- #pragma omp parallel marks a block is to be run in parallel
- In this case all threads do the same
- No real work sharing

Issues with this example

- `std::cout` is not thread safe. Output from different threads may be mixed
  - Try running the code
  - Mixed output?
- All threads call `omp_get_num_threads()` with the same result
  - Might be wasteful if this was a slow function
  - Everybody stores a copy of numthreads
  - Waste of memory
Slightly improved hello world

```cpp
#include <iostream>
#if defined _OPENMP
#include <omp.h>
#endif

int main(int argc, char ** argv)
{
    int threadnum = 0;
    int numthreads = 0;
    #pragma omp parallel shared(numthreads), private(threadnum)
    {
        #ifdef _OPENMP
            threadnum = omp_get_thread_num();
            #pragma omp single
            {
                numthreads = omp_get_num_threads();
            }
        #endif
        #pragma omp critical
        {
            std::cout << "Hello World, I am " << threadnum << " of " << numthreads << std::endl;
        }
    }
}
```

Improvements:

- Use `#pragma omp critical` to only allow one thread to write at a time
  - Comes with a performance penalty since only one thread is running this code at a time
- Use Preprocessor `#ifdef _OPENMP` to only include code if OpenMP is enabled
  - Code works both with and without OpenMP
- Variables defined outside parallel regions
  - Must be careful to tell OpenMP how to handle them
    - `shared, private, first private`
  - More about this later
- `#pragma omp single`
  - Only one thread calls `get_num_threads()`
Running OpenMP code for the course

If you have a multicore computer with GCC or other suitable compiler you can run it locally.

Otherwise you can use GCC on aristotle

- ssh username@aristotle.rc.ucl.ac.uk
- g++ -fopenmp -O3 mycode.cc

References

- OpenMP homepage
- OpenMP cheat sheet
- OpenMP specifications

Parallelizing loops with OpenMP

Simple example

Integrate:

\[ \int_0^1 \frac{4}{1+x^2} \, dx = \pi \]

```cpp
#include <iostream>

int main(int argc, char ** argv)
{

    double pi,sum,x;
    const int N = 1000000;
    const double w = 1.0/N;

    pi = 0.0;
    sum = 0.0;
    #pragma omp parallel private(x), firstprivate(sum), shared(pi)
    {
        #pragma omp for
        for (int i = 0; i < N; ++i)
        {
            x = w*(i-0.5);
            sum = sum + 4.0/(1.0 + x*x);
        }
        #pragma omp critical
        { pi = pi + sum; }
    }

    return 0;
}
```
\{ 
    pi = pi + w*sum;
\}

\}
std::cout << "Result is " << pi << std::endl;
}

Variable scope

- Private: Each thread has its own copy. Any values from before the \texttt{parallel} block are ignored, and are not affected after.
- Shared: Only one shared variable. Shared both between threads and with the non parallel code.
- Firstprivate: Private variable but initialized with value from before the \texttt{parallel} block.

Details of example

- Important that \texttt{x} and \texttt{sum} are private
  - Try making them shared and see what happens
- Note that the default is shared
  - Can be controlled with the default clause
  - \texttt{default(none)} is safer
  - “Explicit is better that implicit”
- We use a critical region to add safety without a race condition

Reduction

- Aggregating a result from multiple threads with a single mathematical operation
- Is a very common pattern
- OpenMP has build in support for doing this
- Simplifies the code and avoids the explicit critical region
- Easier to write and may perform better

Reduction example

\#include <iostream>

\texttt{int main(int argc, char ** argv)}
{ 
    double pi,sum,x;
    const int N = 1000000;
    const double w = 1.0/N;

    pi = 0.0;
    sum = 0.0;

    #pragma omp parallel private(x), reduction(+:sum)
    {
        #pragma omp for
        for (int i = 0; i < N; ++i)
        {
            x = w*(i-0.5);
            sum = sum + 4.0/(1.0 + x*x);
        }
    }
    pi = w*sum;
    std::cout << "Result is " << pi << std::endl;
}

Races, locks and critical regions

Introduction

In the best of worlds our calculations can be done independently. However, even in our simplest examples we saw issues.

- std::cout is not thread safe. Garbage mixed output
- Needs to use critical to merge output
- Real world examples may be more complicated
- Incorrectly shared variable leads to random and typically wrong results

Race conditions

When the result of a calculation depends on the timing between threads.

- Example: threads writing to same variable
- Can be hard to detect
- May only happen in rare cases
- May only happen on specific platforms
- Or depend on system load from other applications
Barriers and synchronisation

Typically it is necessary to synchronize threads. Make sure that all threads are done with a piece of work before moving on. Barriers synchronizes threads.

- Parallel regions such as `omp for` have an implicit barrier at the end
  - Threads wait for the last to finish before moving on
  - May waste significant amount of time
  - We will return to look at load balancing later
  - Sometimes there is no need to wait
  - Disable implicit barrier with `nowait`

- Sometimes you need a barrier where there is no implicit barrier
  - `#pragma omp barrier` inserts a barrier
  - Overusing this may lead to a performance drop

Protecting code and variables

- `#pragma omp critical`
  - Only one task can execute at a time
  - Protect non thread-safe code

- `#pragma omp single`
  - Only one thread executes this block
  - The first thread that arrives will execute the code

- `#pragma omp master`
  - Similar to `single` but uses the master thread

- `#pragma omp atomic`
  - Protect a variable by changing it in one step.

Mutex locks

Sometimes the critical regions are not flexible enough to implement your algorithm.

Examples:

- Need to prevent two different pieces of code from running at the same time.
- Need to lock only a fraction of a large array.
OpenMP locks

OpenMP locks is a general way to manage resources in threads.

- A thread tries to set the lock.
- If the lock is not held by any other thread it is successful and free to carry on.
- If not it will wait until the lock becomes unset.
- Important to remember to unset the lock when done.
- Might otherwise result in a deadlock. Program hangs.

Example

Replace the critical region with a lock. In this case there is no real gain from using a lock.

```cpp
#include <iostream>
#include <omp.h>

int main(int argc, char ** argv)
{
    double pi,sum,x;
    const int N = 10000000;
    const double w = 1.0/N;
    omp_lock_t writelock;
    pi = 0.0;
    sum = 0.0;
    #pragma omp parallel private(x), firstprivate(sum), shared(pi)
    {
        #pragma omp for
        for (int i = 0; i < N; ++i)
        {
            x = w*(i-0.5);
            sum = sum + 4.0/(1.0 + x*x);
        }
        omp_set_lock(&writelock);
        pi = pi + w*sum;
        omp_unset_lock(&writelock);
    }
    omp_destroy_lock(&writelock);
    std::cout << "Result is " << pi << std::endl;
}
```
Multiple locks

Sometimes it is useful to lock multiple resources with different locks.

- Use multiple locks protecting different resources
- Can result in deadlocks if two threads needs both needs the same locks
- One thread holds one lock and the other one holds the other
- Both are waiting for a lock to be free

Notes

OpenMP implements two types of locks. We have only considered simple locks. Consult the OpenMP specifications for nested locks.

OpenMP Tasks

Introduction

- Not all problems are easily expressed as for loops.
- The task construction creates a number of tasks
- The tasks are added to a queue
- Threads take a task from the queue

Example

Calculate Fibonacci numbers by recursion.

- Only as an example:
  - Hard to get any performance improvement. Usually slower than serial code
  - Inefficient algorithm in any case. Why?
  - Consider limiting the number of tasks. Why?
- Use taskwait to ensure results are done before adding their results together

Code

```c
#include <iostream>
#ifdef _OPENMP
```
```c
#include <omp.h>
#endif

int fib(int n)
{
    if (n < 2)
        return n;
    int x;
    int y;
    const int tune = 40;
    #pragma omp task firstprivate(n) shared(x)
    {
        x = fib(n-1);
    }
    #pragma omp task firstprivate(n) shared(y)
    {
        y = fib(n-2);
    }
    #pragma omp taskwait

    return x + y;
}

Main function

int main(int argc, char ** argv)
{
    #ifdef _OPENMP
    omp_set_dynamic(0);
    #endif
    const int num = 20;
    int a;
    #pragma omp parallel shared(a)
    {
        #pragma omp single nowait
        {
            a = fib(num);
        }
    }
    std::cout << "fib " << num << " is " << a << std::endl;
}
```

Note only one thread initially creates tasks. Tasks are still running in parallel.
Advanced usage

- Task dependency:
  - Depends on child tasks. `#taskwait`
  - Real cases may be more complicated
  - May need to explicitly set dependency
  - `#pragma omp task depends(in/out/inout:variable)`
  - See OpenMP docs for details

- `taskyield` Allows a task to be suspended in favour of a different task:
  - Could be useful together with locks

Controlling task generation

- `if(expr) expr==false` create an undeferred task
  - Suspend the present task and execute the new task immediately on the same thread

- `final(expr) expr==true` This is the final task
  - All child tasks are included in the present task

- `mergeable`
  - Included and undeferred tasks may be merged into the parent task

May useful to avoid creating too many small tasks. I.e. in our Fibonacci example.

Scheduling and Load Balancing

Number of threads

The number of threads executing an OpenMP code is determined by the environmental variable `OMP_NUM_THREADS`.

Normally `OMP_NUM_THREADS` is equal to the number of CPU cores

Load balancing

Consider our earlier example of a for loop. 10,000,000 iterations split on 4 cores

Two obvious strategies:

- Split in 4 chunks:
- 2,500,000 iterations for each core
- Minimal overhead for managing threads
- Probably a good solution if the cost is independent of thread
- But what if the cost depends on the thread.
- One thread might be slower than the rest

- Give each thread one iteration at a time
  - No idling thread
  - But huge overhead

The best solution is probably somewhere in between.

**OpenMP strategies**

OpenMP offers a number of different strategies for load balancing set by the following key words. The default is static with one chunk per thread.

- **static**: Iterations are divided into chunks of size `chunk_size` and assigned to threads in round-robin order
- **dynamic**: Each thread executes a chunk of iterations then requests another chunk until none remain
- **guided**: Like dynamic but the chunk size depends on the number of remaining iterations
- **auto**: The decision regarding scheduling is delegated to the compiler and/or runtime system
- **runtime**: The schedule and chunk size are controlled by runtime variables

**Which strategy to use**

It is hard to give general advice on the strategy to use. Depends on the problem and platform. Typically needs benchmarking and experimentation.

- If there is little variation in the runtime of iteration static with a large chunk size minimizes overhead
- In other cases it might make sense to reduce the chunk size or use dynamic or guided
- Note that both dynamic and guided comes with additional overhead to schedule the distribution of work
Alternatives to OpenMP

Overview

- OpenACC:
  - Similar to OpenMP but intended for accelerators. Lecture 9
- MPI:
  - For distributed memory systems. Next lecture
- POSIX and Windows threads. Not portable across operation systems
- C++ 11 threads
- Intel Threading Building Blocks C++ only template library
  - Somewhat more complicated. Requires good understanding of templated code
- Intel Cilk Plus C and C++, Intel and GCC >= 4.9 only

C++11

Simple example:

```cpp
#include <thread>
#include <iostream>

void f()
{
    std::cout << "Hello" << std::endl;
}

void g()
{
    std::cout << "world" << std::endl;
}

int main(int argc, char ** argv)
{
    std::thread t1 {f};
    std::thread t2 {g};

    t1.join();
    t2.join();
}
Details

- Same problem as first OpenMP example. `std::cout` is not thread safe
  - Use mutex and locks
- Queues
- Futures and promise
- `packed_task`

Likely more suited for multi threaded desktop applications than scientific software.

**Cilk Plus**

```cpp
#include <iostream>
#include <time.h>
#include <cilk/cilk.h>
#include <cilk/cilk_api.h>

int fib(int n)
{
    if (n < 2)
        return n;
    int x = cilk_spawn fib(n-1);
    int y = fib(n-2);
    cilk_sync;
    return x + y;
}

int main(int argc, char ** argv)
{
    const int n = 35;
    if (argc > 1)
    {
        // Set the number of workers to be used
        __cilkrts_set_param("nworkers", argv[1]);
    }
    int a = fib(n);
    std::cout << "fib(" << n << ") is " << a << std::endl;
}
```
Further Reading

Tutorials

Any additional tutorials can be added here:

- Blaise Barney, Lawrence Livermore National Laboratory

Wavelet decomposition

Wavelet transforms

- Wavelet transforms decompose a signal into details of increasingly large scales
- Given a signal \( s[j] \), with \( j \in [1, N] \), we want to split
  - an approximation \( A_n[i] \) with \( i \in [1, N/2] \)
  - the details \( D_n[i] \) with \( i \in [1, N/2] \)
- Repeat the process on the approximaton

Pseudo-code

- loop over levels in \([0, L^{\text{max}}]\)

For simplicity, the signal is periodic: \( s[j] = s[j + N] \).

- set \( N' = N \) if \( N \) is even, \( N' = N + 1 \) otherwise
- with \( j \in [0, N' - 1] \), \( i \in [0, n - 1] \)
- set \( D^s[j] = \sum_{i=0}^{n-1} s[2 \ast j + i] h[i] \)
- set \( A^s[j] = \sum_{i=0}^{n-1} s[2 \ast j + i] l[i] \)
- set \( s = A^s \)

Instructions

Given a header and the unit-tests:

1. reconstruct the serial version using the jigsaw implementation file
2. Parallelize with openmp
How many parallelization schemes did you come up with?

If you are not having enough fun, figure out the inverse operation. Wavelets transforms are bijective.

Header

#ifndef CPP_COURSE_WAVELETS_H

#include <cassert>
#include <iostream>
#include <vector>

namespace wavelets {

//! \brief Underlying type of all real numbers in the wavelets
//! \details Defining a "type hierarchy" makes it easy to modify all the basic
//! types underlying a coding project
typedef double Scalar;

//! \brief Represents 1-D signal of any size
typedef std::vector<Scalar> Signal;

//! \brief High and low-pass filter data
//! \details Note
//! ----
//! This structure declares it's attribute public. And later we declare global
//! variables as instances of this type. Both of these aspects are frowned
//! upon, *except* in this one case. The wavelet filters are constants for a
//! given Daubechey wavelet. For constants of nature and constants of math,
//! whether scalar or vectorial (and small), it's okay to use global variable
//! and public attributes. Otherwise, beware!
//!
//! Premature Optimization?
//! -----------------------
//! We could have \texttt{std::array<Scalar, N> high_pass} and the same for
//! \texttt{low_pass}. However, \texttt{N}, the number of coefficients, depends on the actual
//! wavelet. And in \texttt{std::array<Scalar, N>}, \texttt{N} needs to be known at compile
//! time. We could make this change and the compiler might be able to make use
//! of this extra information to produce faster code. However, it would mean
//! that \texttt{DaubecheyData} needs to be declared as \texttt{template<int N> struct}
//! \texttt{DaubecheyData}, and all the functions taking \texttt{DaubecheyData<N>} as input
would also need to be templates.

Templating is cool, fun, but viral. So we might as well use the simpler implementation and wait for benchmarks and profiling results to tell us whether we really need the extra complexity.

```cpp
struct DaubechyData {
    std::vector<Scalar> low_pass;
    std::vector<Scalar> high_pass;
};
```

- **brief** Data for the Daubechy wavelets of type 1
- **details** This data does not change from code to code. In fact, I got them from wikipedia. Constants of nature and constants of math are the only variable that should be declared global (and `const`).

```cpp
extern DaubechyData const daubechy1;
```

- **brief** Data for the Daubechy wavelets of type 2
- **details**

```cpp
extern DaubechyData const daubechy2;
```

- **brief** Applies a filter starting from a given location of the signal
- **details** The signal is defined by the range `start`, and `end`. `location` must be a position inside that range. The filter is applied starting from that location of the signal.

```cpp
Scalar apply_cyclical_filter(Signal::const_iterator const &start, Signal::const_iterator location, 
                           Signal::const_iterator const &end, Signal const &filter);
```

- **brief** Applies a filter starting from a given location of the signal
- **details** Accumulates the result of `signal[location + i] * filter[i]` for `i` in `[0, filter.size())`. If `location + i` goes out of range, then it cycles back to the beginning of the signal. In practice, this means the signal is periodic.

```cpp
Scalar apply_cyclical_filter(Signal const &signal, Signal::size_type location, Signal const &filter);
```

- **brief** Applies the wavelet transform once to the signal
- **details** The output iterator should point to a valid range of the same size as the input signal *if the size of the signal is even*, and one larger than the signal *if the size of the signal is odd*. The result is undefined when the signal and output arrays overlap.

```cpp
void single_direct_transform(Signal::const_iterator const &start, Signal::const_iterator const &end, 
                            Signal::iterator const &out, DaubechyData const &wavelet);
```

- **brief** Applies high and low pass once to the signal range
- **details** This wrapper is also to make testing somewhat simpler.
Signal single_direct_transform(Signal const &signal, DaubechyData const &wavelet);

//! 
//   Applies the wavelet transform `levels` time to the signal
// !/ \details The number of coefficients is given by the function
// !/ `number_of_coefficients`.
void direct_transform(Signal::const_iterator start, Signal::const_iterator end,
                       Signal::iterator out, DaubechyData const &wavelet, unsigned int levels = 1);

//! 
//   Figures out number of coeffs
unsigned int number_of_coefficients(unsigned int signal_size, unsigned int levels);

//! 
//   Transforms the input signal
Signal direct_transform(Signal const &signal, DaubechyData const &wavelet, unsigned int levels = 1);
}
#endif

Tests

define CATCH_CONFIG_MAIN // This tells Catch to provide a main() - only do this in one cpp
#include "wavelets.h"
#include <algorithm>
#include <catch/catch.hpp>
#ifdef _OPENMP
#include <omp.h>
#endif
#endif

TEST_CASE("Application of Cyclical Filters")
{
    wavelets::Signal const signal{ 1, 2, 3, 5, 5, 6 };  
    wavelets::Signal const filter{ 3, -3 };

    SECTION("No cyclical overlap problem")
    {
        CHECK(wavelets::apply_cyclical_filter(signal, 0, filter) == Approx(-3));
        CHECK(wavelets::apply_cyclical_filter(signal, 1, filter) == Approx(-3));
        CHECK(wavelets::apply_cyclical_filter(signal, 2, filter) == Approx(-6));
    }
    SECTION("Filter reaches exactly end of signal")
    {
        CHECK(wavelets::apply_cyclical_filter(signal, 4, filter) == Approx(-3));
    }
    SECTION("Filter reaches past end of signal")
    {
        CHECK(wavelets::apply_cyclical_filter(signal, 5, filter) == Approx(15));
    }
}
SECTION("Empty filter")
{
    CHECK(wavelets::apply_cyclical_filter({ 5, 2 }, 0, {}) == Approx(0));
}
SECTION("Empty signal")
{
    CHECK(wavelets::apply_cyclical_filter({}, 0, { 1, 2 }) == Approx(0));
}
SECTION("Empty signal and filter")
{
    CHECK(wavelets::apply_cyclical_filter({}, 0, {}) == Approx(0));
}
SECTION("Filter much much bigger than signal")
{
    CHECK(wavelets::apply_cyclical_filter({ 1, 2 }, 0, { 1, 2, 3, 4, 5 }) == Approx(21));
}

TEST_CASE("Single pass wavelet transform")
{
    // Use unnormalized haar wavelets, because simple
    // Normalize by multiplying the coeffs by std::sqrt(2)
    wavelets::DaubechyData const haar{ { 1, 1 }, { 1, -1 } };
    SECTION("Haar")
    {
        // Odd number of elements to test periodicity
        auto const actual = single_direct_transform({ 1, 2, 3, 5, 5, 6, 8 }, haar);
        // high pass  low pass
        // details  approximation
        wavelets::Signal const expected{ -1, -2, -1, 7, /* */ 3, 8, 11, 9 }; //
        REQUIRE(actual.size() == expected.size());
        CHECK(std::equal(actual.begin(), actual.end(), expected.begin()));
    }
    SECTION("Empty signal")
    {
        CHECK(wavelets::single_direct_transform({}, haar).size() == 0);
    }
}

TEST_CASE("Multi-pass wavelet transform")
{
    wavelets::DaubechyData const haar{ { 1, 1 }, { 1, 1 } };
    SECTION("Empty signal")
    {
    }
}
SECTION("Level 0 is a copy")
{
    wavelets::Signal const signal{ 1, 2, 3, 4 };  
    auto const actual = wavelets::direct_transform(signal, haar, 0);
    REQUIRE(actual.size() == signal.size());
    CHECK(std::equal(actual.begin(), actual.end(), signal.begin()));
}

SECTION("Haar")
{
    wavelets::Signal const signal{ 1, 2, 3, 5, 5, 6, 8 };
    std::vector<wavelets::Signal> const expecteds{
        signal,  // level 0
        {-1, -2, -1, 7, 3, 8, 11, 9 },  // level 1
        {-1, -2, -1, 7, -5, 2, 11, 20 }, // level 2
        {-1, -2, -1, 7, -5, 2, -9, 31 }, // level 3
        {-1, -2, -1, 7, -5, 2, -9, 0, 62 }, // level 4
        {-1, -2, -1, 7, -5, 2, -9, 0, 0, 124 }, // level 5
        {-1, -2, -1, 7, -5, 2, -9, 0, 0, 0, 248 }, // level 6
    };
    for (decltype(expecteds.size()) levels(0); levels < expecteds.size(); ++levels) {
        auto const actual = direct_transform(signal, haar, levels);
        REQUIRE(actual.size() == expecteds[levels].size());
        CHECK(std::equal(actual.begin(), actual.end(), expecteds[levels].begin()));
    }
}

#if defined(_OPENMP)
    // Catch offers to print out the duration of a test, so we can use it as a
    // make-do benchmarking framework
    // Run the executable with --duration yes
    TEST_CASE("OpenMP")
    {
        wavelets::Signal input(1000000);
        wavelets::Signal output(1000000);
        std::generate(input.begin(), input.end(), std::rand);
        std::fill(output.begin(), output.end(), 0);
        SECTION("Serial")
        {
            auto const nthreads = omp_get_max_threads();
            omp_set_num_threads(1);
            for (int i(0); i < 1000; ++i)
                wavelets::direct_transform(}
SECTION("Parallel")
{
for (int i(0); i < 1000; ++i)
    wavelets::direct_transform(
        input.begin(), input.end(), output.begin(), wavelets::daubechy2, 8);
}

Implementation
#include "wavelets.h"
#include <cassert>
#include <numeric>
#ifdef _OPENMP
#include <omp.h>
#endif
namespace wavelets {

    DaubechyData const daubechy1{
        { 7.071067811865475244008443621048490392848359376884740365883398e-01,
           7.071067811865475244008443621048490392848359376884740365883398e-01 },
        { -7.071067811865475244008443621048490392848359376884740365883398e-01,
           7.071067811865475244008443621048490392848359376884740365883398e-01 }
    };

    DaubechyData const daubechy2{
        { 4.8296291314453414337487159988644486838169524195042022752011715e-01,
           8.365163037378079055752937809168732034593703883484392934953414e-01,
           2.241438680420133810259727622404003554678835181824717613871683e-01,
           -1.294095225512603811744494188120241641745344506599652569070016e-01 },
        { 1.294095225512603811744494188120241641745344506599652569070016e-01,
           2.241438680420133810259727622404003554678835181824717613871683e-01,
           -8.365163037378079055752937809168732034593703883484392934953414e-01,
           4.8296291314453414337487159988644486838169524195042022752011715e-01 }
    };

    Scalar apply_cyclical_filter(
        Signal const& signal, Signal::size_type location, Signal const& filter)
    {
        return apply_cyclical_filter(}
signal.begin(), signal.begin() + location, signal.end(), filter);
}

Signal single_direct_transform(Signal const& signal, DaubechyData const& wavelet)
{
    Signal coefficients(number_of_coefficients(signal.size(), 1));
    single_direct_transform(signal.begin(), signal.end(), coefficients.begin(), wavelet);
    return coefficients;
}

Signal direct_transform(Signal const& signal, DaubechyData const& wavelet, unsigned int levels)
{
    // shifting the bit left is the same as multiplying by two
    Signal coefficients(number_of_coefficients(signal.size(), levels));
    direct_transform(signal.begin(), signal.end(), coefficients.begin(), wavelet, levels);
    return coefficients;
}

unsigned int number_of_coefficients(unsigned int signal_size, unsigned int levels)
{
    if (levels == 0)
        return signal_size;
    auto const even_size = signal_size + signal_size % 2;
    if (levels == 1)
        return even_size;
    return even_size / 2 + number_of_coefficients(even_size / 2, levels - 1);
}

Scalar apply_cyclical_filter(
    Signal::const_iterator const& start,
    Signal::const_iterator location,
    Signal::const_iterator const& end,
    Signal const& filter)
{
    if (start == end)
        return 0;

    assert(location >= start);
    assert(location < end);

    Signal::value_type result(0);
    auto i_filter = filter.begin();
    while (i_filter != filter.end()) {
        for (; i_filter != filter.end() and location != end; ++i_filter, ++location)
            result += (*i_filter) * (*location);
location = start;
return result;
}

void single_direct_transform(
    Signal::const_iterator const& start, Signal::const_iterator const& end,
    Signal::iterator const &out, DaubechyData const& wavelet)
{
    assert(start <= end);
    int const half = ((end - start) + (end - start) % 2) / 2;
#pragma omp parallel
    {
#pragma omp for
    for (int i=0; i < half; ++i)
        *(out + i) = apply_cyclical_filter(start, start + 2 * i, end, wavelet.high_pass);
#pragma omp for
    for (int i=0; i < half; ++i)
        *(out + i + half) = apply_cyclical_filter(start, start + 2 * i, end, wavelet.low_pass);
    }
}

void direct_transform(
    Signal::const_iterator start, Signal::const_iterator end,
    Signal::iterator out, DaubechyData const& wavelet,
    unsigned int levels)
{
    if (start == end)
        return
    assert(start <= end);
    if (levels == 0) {
        std::copy(start, end, out);
        return;
    }
    single_direct_transform(start, end, out, wavelet); // first iteration
    auto const half = [](Signal::size_type n) { return (n + n % 2) / 2; };
    auto approx_size = half(end - start);
    Signal work_array(approx_size);
    auto const i_work = work_array.begin();
    for (unsigned int i(1); i < levels; ++i, approx_size = half(approx_size)) {
        out += approx_size;
        std::copy(out, out + approx_size, i_work);
        single_direct_transform(i_work, i_work + approx_size, out, wavelet);
Chapter 8

Distributed memory parallelism

Distributed Memory Parallelism

The basic idea

• many processes, each with their own data

• each process is independent

• processes can send messages to one another

MPI in practice

Specification and implementation

• in practice, we use MPI, the Message Passing Interface
• MPI is a specification for a library
• It is implemented by separate vendors/open-source projects
  – OpenMPI
  – mpich
• It is a C library with many many bindings:
  – Fortran (part of official MPI specification)
  – Python: boost, mpi4py
  – R: Rmpi
  – c++: boost

Programming and running

• an MPI program is executed with mpiexec -n N [options] nameOfProgram [args]

• MPI programs call methods from the mpi library

```c
int MPI_Bcast(void *buffer, int count, MPI_Datatype datatype, int root, MPI_Comm comm)
```

• vendors provide wrappers (mpiCC, mpic++) around compilers. Wrappers point to header file location and link to right libraries. MPI program can be (easily) compiled by substituting (g++|icc) -> mpiCC

Hello, world!: hello.cc

```c
#include <mpi.h>
#include <iostream>

int main(int argc, char * argv[]) {
  /// Must be first call
  MPI_Init (&argc, &argv);
  /// Now MPI calls possible

  /// Size of communicator and process rank
  int rank, size;
  MPI_Comm_rank (MPI_COMM_WORLD, &rank);
  MPI_Comm_size (MPI_COMM_WORLD, &size);

  std::cout << "Processor " << rank << " of " << size << " says hello\n";

  /// Must be last MPI call
  MPI_Finalize();
  /// No more MPI calls from here
  return 0;
}
```
Hello, world!: CMakeLists.txt

```cmake
find_package(MPI REQUIRED)
add_executable(hello hello.cc)
target_include_directories(hello SYSTEM PUBLIC ${MPI_INCLUDE_DIRS})
target_link_libraries(hello PUBLIC ${MPI_LIBRARIES})
```

Hello, world!: compiling and running

On aristotle.rc.ucl.ac.uk:

- load modules: `module load GCC/4.7.2 OpenMPI/1.6.4-GCC-4.7.2`
- module load cmake/2.8.10.2
- create files “hello.cc” and “CMakeLists.txt” in some directory
- create build directory `mkdir build && cd build`
- run cmake and make `cmake .. && make`
- run the code `mpiexec -n 4 hello`

Hello, world! dissected

- MPI calls *must* appear between `MPI_Init` and `MPI_Finalize`
- Groups of processes are handled by a communicator. `MPI_COMM_WORLD` handles the group of all processes.
- Size of group and rank (order) of process in group
- By *convention*, process of rank 0 is *special* and called *root*

MPI with CATCH

Running MPI unit-tests requires `MPI_Init` and `MPI_Finalize` before and after the test framework (*not* inside the tests).
#include <mpi.h>
// Next line tells CATCH we will use our own main function
#define CATCH_CONFIG_RUNNER
#include "catch.hpp"

TEST_CASE("Just test I exist") {
    int rank, size;
    MPI_Comm_rank (MPI_COMM_WORLD, &rank);
    MPI_Comm_size (MPI_COMM_WORLD, &size);
    CHECK(size > 0); CHECK(rank >= 0);
}

int main(int argc, char * argv[]) {
    MPI_Init (&argc, &argv);
    int result = Catch::Session().run(argc, argv);
    MPI_Finalize();
    return result;
}

Point to point communication

Many point-2-point communication schemes

Can you think of two behaviours for message passing?

- Process 0 can (i) give message and then either (ii) leave or (iii) wait for acknowledgements
- Process 1 can (i) receive message
- MPI can (i) receive message, (ii) deliver message, (iii) deliver acknowledgments

Blocking synchronous send
### Stage Figure

<table>
<thead>
<tr>
<th>Stage</th>
<th>Figure</th>
</tr>
</thead>
<tbody>
<tr>
<td>a. 0, 1, and MPI stand ready:</td>
<td><img src="image1.png" alt="Image" /></td>
</tr>
<tr>
<td>b. message dropped off by 0:</td>
<td><img src="image2.png" alt="Image" /></td>
</tr>
<tr>
<td>c. transit:</td>
<td><img src="image3.png" alt="Image" /></td>
</tr>
<tr>
<td>d. message received by 1</td>
<td><img src="image4.png" alt="Image" /></td>
</tr>
<tr>
<td>e. receipt received by 0</td>
<td><img src="image5.png" alt="Image" /></td>
</tr>
</tbody>
</table>

**Blocking send**

<table>
<thead>
<tr>
<th>Stage</th>
<th>Figure</th>
</tr>
</thead>
<tbody>
<tr>
<td>a. 0, 1, and MPI stand ready:</td>
<td><img src="image6.png" alt="Image" /></td>
</tr>
<tr>
<td>b. message dropped off by 0:</td>
<td><img src="image7.png" alt="Image" /></td>
</tr>
<tr>
<td>c. transit, 0 leaves</td>
<td><img src="image8.png" alt="Image" /></td>
</tr>
</tbody>
</table>
Non-blocking send

Stage | Figure
-----|------
da. message received by 1 | ![Figure](image)

a. 0, 1, and MPI stand ready:

b. 0 leaves message in safebox

c. transit

d. message received by 1

e. receipt placed in safebox

Blocking synchronous send

```c
int MPI_Ssend(const void *buf, int count, MPI_Datatype datatype, int dest, int tag, MPI_Comm comm)
```

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>buf</td>
<td>Pointer to buffer. Always <code>void</code> because practical C is not type safe.</td>
</tr>
<tr>
<td>Parameter</td>
<td>Meaning</td>
</tr>
<tr>
<td>-----------</td>
<td>---------</td>
</tr>
<tr>
<td>count</td>
<td>Size of the buffer. I.e. length of the message to send, in units of the specified datatype (not bytes)</td>
</tr>
<tr>
<td>datatype</td>
<td>Encodes type of the buffer. MPI_INT for integers, MPI_CHAR for characters. Lots of others.</td>
</tr>
<tr>
<td>dest</td>
<td>Rank of the receiving process</td>
</tr>
<tr>
<td>tag</td>
<td>A tag for message book-keeping</td>
</tr>
<tr>
<td>comm</td>
<td>The communicator – usually just MPI_COMM_WORLD</td>
</tr>
<tr>
<td>return</td>
<td>An error tag. Equals MPI_SUCCESS on success.</td>
</tr>
</tbody>
</table>

**Blocking receive**

```c
int MPI_Recv(void *buf, int count, MPI_Datatype datatype, int source, int tag, MPI_Comm comm, MPI_Status *status)
```

Good for both synchronous and asynchronous communication

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>buf</td>
<td>Pointer to receiving pre-allocated buffer</td>
</tr>
<tr>
<td>count</td>
<td>Size of the buffer. I.e. maximum length of the message to receive. See MPI_Get_count</td>
</tr>
<tr>
<td>datatype</td>
<td>Informs on the type of the buffer</td>
</tr>
<tr>
<td>source</td>
<td>Rank of the sending process</td>
</tr>
<tr>
<td>tag</td>
<td>A tag for message book-keeping</td>
</tr>
<tr>
<td>status</td>
<td>‘MPI_STATUS_IGNORE’ for now. See MPI_Get_count.</td>
</tr>
<tr>
<td>comm</td>
<td>The communicator</td>
</tr>
<tr>
<td>return</td>
<td>Error tag</td>
</tr>
</tbody>
</table>

**Example: Blocking synchronous example**

Inside a new section in the test framework:

```c
std::string const peace = "I come in peace!";
if(rank == 0) {
    int const error = MPI_Ssend(
        (void*) peace.c_str(), peace.size() + 1, MPI_CHAR, 1, 42, MPI_COMM_WORLD);
```
// Here, we guarantee that Rank 1 has received the message.
REQUIRE(error == MPI_SUCCESS);
}
if(rank == 1) {
    char buffer[256];
    int const error = MPI_Recv(
        buffer, 256, MPI_CHAR, 0, 42, MPI_COMM_WORLD, MPI_STATUS_IGNORE);
    REQUIRE(error == MPI_SUCCESS);
    CHECK(std::string(buffer) == peace);
}

Common bug: Set both sender and receiver to 0. What happens?

**Example: Do you know your C vs C++ strings?**

Why the +1?

```cpp
int const error = MPI_Ssend(
    (void*) peace.c_str(), peace.size() + 1, MPI_CHAR, 1, 42, MPI_COMM_WORLD);
```

Because C and C++ `char const*` strings are null-terminated to indicate the string is finished, which adds an extra character. However, `std::string` abstracts it away. And so its length does not include the null-termination.

**Example: Causing a dead-lock**

Watch out for order of send and receive!

Bad:

```cpp
if(rank == 0) {
    MPI_Ssend (sendbuf, count, MPI_INT, 1, tag, comm);
    MPI_Recv (recvbuf, count, MPI_INT, 1, tag, comm, &status);
} else {
    MPI_Ssend (sendbuf, count, MPI_INT, 0, tag, comm);
    MPI_Recv (recvbuf, count, MPI_INT, 0, tag, comm, &status);
}
```

Good:
if(rank == 0) {
    MPI_Ssend (sendbuf, count, MPI_INT, 1, tag, comm);
    MPI_Recv (recvbuf, count, MPI_INT, 1, tag, comm, &status);
} else {
    MPI_Recv (recvbuf, count, MPI_INT, 0, tag, comm, &status);
    MPI_Ssend (sendbuf, count, MPI_INT, 0, tag, comm);
}

Send vs SSend

Why would we use Send instead of SSend?

```
std::string peace = "I come in peace!";
if(rank == 0) {
    int const error = MPI_Send(
        (void*) peace.c_str(), peace.size() + 1, MPI_CHAR, 1, 42, MPI_COMM_WORLD);
    // We do not guarantee that Rank 1 has received the message yet
    // But nor do we necessarily know it hasn't.
    // But we are definitely allowed to change the string, as MPI promises
    // it has been buffered
    peace = "Shoot to kill!"; // Safe to reuse the send buffer.
    REQUIRE(error == MPI_SUCCESS);
}
if(rank == 1) {
    char buffer[256];
    int const error = MPI_Recv(
        buffer, 256, MPI_CHAR, 0, 42, MPI_COMM_WORLD, MPI_STATUS_IGNORE);
    REQUIRE(error == MPI_SUCCESS);
    CHECK(std::string(buffer) == peace);
}
```

Both guarantee the buffer is safe to reuse. Send makes no guarantee as to whether it returns early or not. But SSend forces a synchronisation point: the codes reach the matching places, with all processes waiting until all reach that point.

It may come out slightly faster to use Send, since having a synchronisation point when you don’t need one can slow things down: Suppose (A) runs slightly faster, then (B) does; at the end, they’ve both been running fully efficiently.

With a synchronisation point in between, you’ll have wasted time:

This is only important when there is noise or variability in the execution time on different processes, but this is often the case.
So unnecessary synchronisation points are bad. The MPI Implementation may choose to buffer, or synchronise in Send; you’re letting MPI guess.

However, if you want to fine tune this to get the best performance, you should use ISend.

**Non-blocking: ISend/IRecv**

With ISend, we indicate when we want the message to set off.

We receive a handle to the message, of type `MPI_Request*` which we can use to require it has been received, or check.

This produces more complicated code, but you can write code which **overlaps calculation with communication**: the message is travelling, while you get on with something else. We’ll see a practical example of using this next lecture.

```c
std::string peace = "I come in peace!";
if(rank == 0) {
    MPI_Request request;
    int error = MPI_Isend(
        (void*) peace.c_str(), peace.size() + 1, MPI_CHAR, 1, 42,
        MPI_COMM_WORLD, &request);
    // We do not guarantee that Rank 1 has received the message yet
    // We can carry on, and ANY WORK WE DO NOW WILL OVERLAP WITH THE
    // COMMUNICATION
    // BUT, we can't safely change the string.
    REQUIRE(error == MPI_SUCCESS);
    // Do some expensive work here
    for (int i=0; i<1000; i++) {}
    // BUSINESS FOR EXAMPLE
    MPI_Status status;
    error = MPI_Wait(&request, &status);
    REQUIRE(error == MPI_SUCCESS);
    // Here, we run code that requires the message to have been
    // successfully sent.
}
if(rank == 1) {
    char buffer[256];
    int const error = MPI_Recv(
        buffer, 256, MPI_CHAR, 0, 42, MPI_COMM_WORLD, MPI_STATUS_IGNORE);
    REQUIRE(error == MPI_SUCCESS);
    CHECK(std::string(buffer) == peace);
}

int MPI_Isend(const void *buf, int count, MPI_Datatype datatype, int dest, int tag,
              MPI_Comm comm, MPI_Request *request)
```
Pass the parcel: SendRecv

Consider a group of N processes in a ring: each has a value, and wants to “pass the parcel” to the left. How would you achieve this with SSend and Receive?

```c
int message = rank*rank;
int received = -7;

// Define the ring
int left = rank-1;
int right = rank+1;
if (rank==0) {
    left = size-1;
}
if (rank == size-1){
    right = 0;
}

With synchronous calls each process can only either be sending or receiving. So the even processes need to send, while the odd ones receive, then vice-versa. This is clearly inefficient.

```c
if (rank%2 == 0) {
    int error = MPI_Ssend(
        &message, 1, MPI_INT, left, rank, MPI_COMM_WORLD);
    error = MPI_Recv(
        &received, 1, MPI_INT, right, right, MPI_COMM_WORLD, MPI_STATUS_IGNORE);
}
if (rank%2 == 1) {
    int error = MPI_Recv(
        &received, 1, MPI_INT, right, right, MPI_COMM_WORLD, MPI_STATUS_IGNORE);
    error = MPI_Ssend(
        &message, 1, MPI_INT, left, rank, MPI_COMM_WORLD);
    REQUIRE( received == right*right );
}

With ISend/IRrecv, this can be achieved in one go: each process posts its send, then posts its receive, then waits for completion.
MPI_Request request;
// Everyone sets up their messages to send
int error = MPI_Isend(
  &message, 1, MPI_INT, left, rank, MPI_COMM_WORLD, &request);

// Recv acts as our sync-barrier
error = MPI_Recv(
  &received, 1, MPI_INT, right, right, MPI_COMM_WORLD, MPI_STATUS_IGNORE);

// But let's check our send completed:
error = MPI_Wait(&request, MPI_STATUS_IGNORE);
REQUIRE(error == MPI_SUCCESS);
REQUIRE(received == right*right);

However, this is such a common pattern, that there is a separate MPI call to make this easier:

int MPI_Sendrecv(void *sendbuf, int scount, MPI_Datatype stype, int dest, int stag,
                  void *recvbuf, int rcount, MPI_Datatype rtype, int source, int rtag,
                  MPI_Comm comm, MPI_Status *status)

Each argument is duplicated for the send and receive payloads.
Classroom exercise: implement ring-send using Sendrecv.

Al(most all) point to point

Sending messages:

<table>
<thead>
<tr>
<th>name</th>
<th>Blocking</th>
<th>forces synchronisation point</th>
<th>Buffer-safe</th>
</tr>
</thead>
<tbody>
<tr>
<td>MPI_Send</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
</tr>
<tr>
<td>MPI_Send</td>
<td>maybe</td>
<td>no</td>
<td>yes</td>
</tr>
<tr>
<td>MPI_Isend</td>
<td>no</td>
<td>no</td>
<td>no</td>
</tr>
</tbody>
</table>

Receiving messages:

<table>
<thead>
<tr>
<th>name</th>
<th>blocking</th>
</tr>
</thead>
<tbody>
<tr>
<td>MPI_Recv</td>
<td>yes</td>
</tr>
</tbody>
</table>
Collective Communication

Many possible communication schemes

Think of two possible forms of collective communications:

- give a beginning state
- give an end state

Broadcast: one to many

<table>
<thead>
<tr>
<th>State</th>
<th>Figure</th>
</tr>
</thead>
<tbody>
<tr>
<td>data in 0, no data in 1, 2</td>
<td></td>
</tr>
<tr>
<td>data from 0 sent to 0, 1</td>
<td></td>
</tr>
</tbody>
</table>

Gather: many to one
State Figure

data in 0, 1, 2

data from 1, 2 sent to 0

Scatter: one to many

State Figure

data in 0

data from 0 in 0, 1, 2

All to All: many to many

State Figure

data in 0, 1, 2

from each to each

Reduce operation
Wherefrom the baby bunny?

... 

Sum, difference, or any other binary operator:

\[
\frac{(1 + 0)}{2} + 1 = \boxed{1} \quad \quad \frac{(1 - 0)}{2} - 1 = \boxed{0}
\]

**Collective operation API**

Group synchronisation:

```c
int MPI_Barrier(MPI_Comm comm);
```

Broadcasting:

```c
int MPI_Bcast(void *buffer, int count, MPI_Datatype datatype, int root, MPI_Comm comm)
```

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Content</th>
</tr>
</thead>
<tbody>
<tr>
<td>buf</td>
<td>Pointer to sending/receiving buffer</td>
</tr>
<tr>
<td>count</td>
<td>Size of the buffer/message</td>
</tr>
<tr>
<td>datatype</td>
<td>Informs on the type of the buffer</td>
</tr>
<tr>
<td>root</td>
<td>Sending processor</td>
</tr>
<tr>
<td>comm</td>
<td>The communicator!</td>
</tr>
<tr>
<td>return</td>
<td>Error tag</td>
</tr>
</tbody>
</table>
Example of collective operation (1)

Insert into a new CATCH section the following commands

```cpp
std::string const peace = "I come in peace!";
std::string message = "";
int error;
if(rank == 0) {
    message = peace;
    error = MPI_Bcast((void*) peace.c_str(), peace.size() + 1, MPI_CHAR, 0, MPI_COMM_WORLD);
} else {
    char buffer[256];
    error = MPI_Bcast(buffer, 256, MPI_CHAR, 0, MPI_COMM_WORLD);
    message = std::string(buffer);
}
```

Example of collective operations (2)

And then insert the following right after it

```cpp
for(int i(0); i < size; ++i) {
    if(rank == i) {
        INFO("Current rank is " << rank);
        REQUIRE(error == MPI_SUCCESS);
        CHECK(message == peace);
    }
    MPI_Barrier(MPI_COMM_WORLD);
}
```

Causing deadlocks

Explain why the following two codes fail.

1. Replace the loop in the last fragment with:

   ```cpp
   for(int i(1); i < size; ++i) ...
   ```

2. Refactor and put everything inside the loop
std::string const peace = "I come in peace!";
std::string message;
for(int i(0); i < size; ++i) {
    if(i == 0 and rank == 0) { /* broadcast */ }
    else if(rank == i) { /* broadcast */ }
    if(rank == i) { /* testing bit */ }
    MPI_Barrier(MPI_COMM_WORLD);
}

NOTE: a loop with a condition for i == 0 is a common anti-pattern (eg bad)

All to all operation

int MPI_Scatter(const void *sendbuf, int sendcount, MPI_Datatype sendtype,
                    void *recvbuf, int recvcount, MPI_Datatype recvtype, int root,
                    MPI_Comm comm)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Content</th>
</tr>
</thead>
<tbody>
<tr>
<td>sendbuf</td>
<td>Pointer to sending buffer (only significant at root)</td>
</tr>
<tr>
<td>sendcount</td>
<td>Size of a single message</td>
</tr>
<tr>
<td>datatype</td>
<td>Type of the buffer</td>
</tr>
<tr>
<td>recvbuf</td>
<td>Pointer to receiving buffers (also at root)</td>
</tr>
<tr>
<td>recvcount</td>
<td>Size of the receiving buffer</td>
</tr>
<tr>
<td>recvtype</td>
<td>Informs on the type of the receiving buffer</td>
</tr>
</tbody>
</table>

Exercise: Have the root scatter "This....." "message." "is.split." to 3 processors (including it self).

Splitting the communicators

Groups of processes can be split according to color:

int MPI_Comm_split(MPI_Comm comm, int color, int key, MPI_Comm *newcomm)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Content</th>
</tr>
</thead>
<tbody>
<tr>
<td>comm</td>
<td>Communicator that contains all the processes to be split</td>
</tr>
<tr>
<td>Parameter</td>
<td>Content</td>
</tr>
<tr>
<td>-----------</td>
<td>---------</td>
</tr>
<tr>
<td>color</td>
<td>All processes with same color end up in same group</td>
</tr>
<tr>
<td>int key</td>
<td>Controls rank in final group</td>
</tr>
<tr>
<td>newcomm</td>
<td>Output communicator</td>
</tr>
</tbody>
</table>

### Splitting communicators: example

The following splits processes into two groups with ratio 1:2.

```c
bool const is_apple = rank % 3 == 0;
SECTION("split 1:3 and keep same process order") {
    MPI_Comm apple_orange;
    MPI_Comm_split(MPI_COMM_WORLD, is_apple ? 0: 1, rank, &apple_orange);

    int nrank, nsize;
    MPI_Comm_rank(apple_orange, &nrank);
    MPI_Comm_size(apple_orange, &nsize);

    int const div = (size - 1) / 3, napples = 1 + div;
    if(is_apple) {
        CHECK(nsize == napples);
        CHECK(nrank == rank / 3);
    } else {
        CHECK(nsize == size - napples);
        CHECK(nrank == rank - 1 - (rank / 3));
    }
}
```

### Splitting communicators: Exercise

Exercise:

- use “-rank” as the key: what happens?
- split into three groups with ratios 1:1:2
- use one of the collective operation on a single group

### Scatter operation solution

```c
std::string const message = "This message is going to come out in separate channels";
```
```c
int N = message.size() / size;
if (message.size() < size) return;

char buffer[256];
if (rank == 0) {
  int const error = MPI_Scatter(
    (void*) message.c_str(), N, MPI_CHAR,
    buffer, 256, MPI_CHAR, 0, MPI_COMM_WORLD
  );
  REQUIRE(error == MPI_SUCCESS);
  CHECK(message.substr(rank*N, N) == std::string(buffer, N));
} else {
  int const error = MPI_Scatter(
    NULL, -1, MPI_CHAR, // not significant outside root
    buffer, 256, MPI_CHAR, 0, MPI_COMM_WORLD
  );
  REQUIRE(error == MPI_SUCCESS);
  CHECK(message.substr(rank*N, N) == std::string(buffer, N));
}
```

More advanced MPI

Architecture and usage

- depending on library, MPI processes can be placed on specific node...
- ... and even chained to specific cores
- Fewer processes than core means we can do MPI + openmp:
  - some data is distributed (MPI)
  - some data is shared (openMP)
• MPI-3 allows for creating/destroying processes dynamically

Splitting communicators

MPI_Group_* specify operations to create sets of processes. In practice, it defines operations on sets:

• union
• intersection
• difference

And allows the creation of a communicator for the resulting group.

More MPI data-types

It is possible to declare complex data types

• strided vectors, e.g. only one of every element (MPI_Type_vector)
• sub-matrices (strided in n axes, n >= 2) (MPI_Type_Create_struct)
• irregular strides (MPI_Type_indexed)

One sided communication

Prior to MPI-3, both sending and receiving processes must be aware of the communication.
One-sided communication allow processes to define a buffer that other processes can access without their explicit knowledge.

And also

• Cartesian grid topology where process (1, 1) is neighbor of (0, 1), (1, 0), (2, 1), (1, 2). With simplified operations to send data EAST, WEST, UP, DOWN. . . .
• More complex graph topologies
• non-blocking collective operations
Chapter 9

MPI Design Example

Objectives for this chapter

In this chapter, we give an extended introduction to a particular parallel programming example, introducing some common programming patterns which occur in working with MPI.

Smooth Life

Conway’s Game of Life

The Game of Life is a cellular automaton devised by John Conway in 1970. A cell, on a square grid, is either alive or dead. On the next iteration:

- An alive cell:
  - remains alive if it has 2 or 3 alive neighbours out of 8
  - dies (of isolation or overcrowding) otherwise
- A dead cell:
  - becomes alive if it has exactly 3 alive neighbours
  - stays dead otherwise

Conway’s Game of Life

This simple set of rules produces beautiful, complex behaviours:
Smooth Life

Smooth Life, proposed by Stephan Rafler, extends this to continuous variables:

- The neighbourhood is an integral over a ring centered on a point.
- The local state is an integral over the disk inside the ring.

(The ring has outer radius 3*inner radius, so that the area ratio of 1:8 matches the grid version.)

Smooth Life

- A point has some degree of aliveness.
- Next timestep, a point’s aliveness depends on these two integrals \( D(r) \) and \( R(r) \)
- The new aliveness \( S(D(r), R(d)) \) is a smoothly varying function such that:
  - If \( D(d) \) is 1, S will be 1 if \( d^{(1)} \leq D(r) \leq d^{(2)} \)
  - If \( R(d) \) is 0, S will be 1 if \( b^{(1)} \leq R(r) \leq b^{(2)} \)

A “Sigmoid” function is constructed that smoothly blends between these limits.

Smooth Life on a computer

We discretise Smooth Life using a grid, so that the integrals become sums. The aliveness variable becomes a floating point number.
To avoid the hard-edges of a “ring” and “disk” defined on a grid, we weight the sum by the fraction of a cell that would fall inside the ring or disk:

If the distance $d$ from the edge of the ring is within 0.5 units, we weight the integral by $2d - 1$, so that it smoothly varies from 1 just inside to 0 just outside.

**Smooth Life**

Smooth Life shows even more interesting behaviour:

**SmoothLifeVideo**

- Gliders moving any direction
- “Tension tubes”

**Exercise**

We will create the following two functions:

- a function to compute the two integrals $D(r)$ and $R(r)$ as $D(x_0, y_0) = \sum_{i,j \in \text{grid}} F(i, j) \text{Disk}(r = ||(i - x_0, j - y_0)||)$ with $F(i, j)$ the current value at point $(i, j)$
- the main update function:
  - loop over all points $(x, y)$ in field:
  - compute integrals centered at $(x, y)$
  - update the field using the transition function

All other functions, including the transition function, Disk, etc are given.

**Square domain into a 1-d vector**

We wrap the 2d-grid into a one d vector in row-major format: $F(i, j) \leftrightarrow F(I)$ with $i = I/N_x$ and $j = I \% N_x$, with $N_x$ the number of points in direction $x$.

**Distances wrap around a torus**

We use periodic boundary conditions: the field is a torus.

```cpp
int Smooth::TorusDistance(int x1, int x2, int size) const {
    auto const remainder = std::abs(x1 - x2) % size;
    return std::min(remainder, std::abs(remainder - size));
}
```
Smoothed edge of ring and disk.

Disk($r$):

double Smooth::Disk(distance radius) const {
    if(radius > inner + smoothing / 2)
        return 0.0;
    if(radius < inner - smoothing / 2)
        return 1.0;
    return (inner + smoothing / 2 - radius) / smoothing;
}

Automated tests for mathematics

SECTION("Sigmoid function is correct") {
    double e = std::exp(1.0);
    REQUIRE(Smooth::Sigmoid(1.0, 1.0, 4.0) == 0.5);
    REQUIRE(std::abs(Smooth::Sigmoid(1.0, 0.0, 4.0) - e / (1 + e)) < 0.0001);
    REQUIRE(Smooth::Sigmoid(10000, 1.0, 4.0) == 1.0);
    REQUIRE(std::abs(Smooth::Sigmoid(0.0, 1.0, 0.1)) < 0.001);
}

Comments

We can see that this is pretty slow:

If the overall grid is $M$ by $N$, and the range of interaction ($3\times$ the inner radius), is $r$, then each time step takes $MNr^2$ calculations: if we take all of these proportional as we “fine grain” our discretisation (a square domain, and a constant interaction distance in absolute units), the problem grows like $N^4$!

So let’s parallelize!

“Ideal” Domain decomposition

Domain decomposition

One of the most important problems in designing a parallel code is “Domain Decomposition”:

- How will I divide up the calculation between processes?
Design objectives in decomposition are:

- Minimise communication
- Optimise load balance (Share out work evenly)

**Decomposing Smooth Life**

We’ll go for a 1-d spatial decomposition for smooth life, dividing the domain into “Stripes” along the x-axis.

If we have an $N \times M$ domain, and $p$, processes, each process will be responsible for $NM/p$ cells, and $NMr^2/p$ calculations.

**Static and dynamic balance**

This will achieve perfect static load balance: the average work done by each process is the same. If we were not solving in a rectangular grid, this would have been harder.

However, since the calculation can (perhaps) be done quicker when the domain is empty, and our field will vary as time passes, we will not achieve perfect dynamic load balance.

**Communication in Smooth Life**

Given that each cell needs to know the state of cells within a range $r = 3r_d$, where $r_d$ is the inner radius of the neighbourhood ring, and $r$ the outer radius, we need to get this information to the neighbouring sites.

This is great: we only need to transfer information to neighbours, not all the other processes. Such “local” communication results in fast code.

The amount of communication to take place each time step is proportional to $rMp$, but assuming an appropriate network topology exists, each pair of neighbours can look after their communication at the same time, so communication will take time proportional to $rMp/p = rM$.

**Strong scaling**

We therefore expect the time taken for a simulation to vary like: $Mr(k + Nr/p)$.
(Where $k$ is a parameter describing the relative time to communicate one cell’s state compared to the time for calculating one cell)

Thus, we see that for a FIXED problem size, the benefit of parallelism will disappear and communication will dominate: this is Amdahl’s law again.
Weak scaling

However, if we consider larger and larger problems, growing $N$ as $p$ grows, then we can stop communication overtaking us. This is a common outcome: local problems provide perfect weak scaling (until network congestion or IO problems dominate).

Any NONLOCAL communication, where the total amount of time for communication to take place grows as the number of processes does (such as a gather, which takes $p$, a reduction, like $ln(p)$, or an all-to-all, like $p^2$, means that perfect weak scaling can’t be achieved.)

Exercise: Blocking Collective

We will parallelize the update and integral functions by having each process work on a contiguous strip of the whole field.

For simplicity, each process owns a full replica of the field in memory. This is inefficient since each process owns memory describing a part of the field it will never use. Improving this is fairly easy, but require some more bookkeeping. Do try it at home!

Exercise: parallelize using a blocking collective

Exercise: Halo update

Parallelize using a non-blocking collective and layer computation and calculation:

1. send data (part of the field) other processes need
2. update part of owned field that does not need data from other processes
3. receive data from other processes
4. update part of owned field that needs data from other processes

This is called a halo update and quite common to domain decomposition problems. We can layer communication and computation even more by splitting over data on the left and on the right boundaries.

Header

```cpp
#include <string>
#include <vector>
#include <mpi.h>

typedef double density;
```
typedef double distance;
typedef double filling;

class Smooth {
public:
    Smooth(int sizex = 100, int sizey = 100, distance inner = 21.0, filling birth_1 = 0.278,
           filling birth_2 = 0.365, filling death_1 = 0.267, filling death_2 = 0.445,
           filling smoothing_disk = 0.147, filling smoothing_ring = 0.028);
    int Size() const;
    int Sizex() const;
    int Sizey() const;
    int Range() const;
    int Frame() const;
    const std::vector<density> &Field() const;
    void Field(std::vector<density> const &input);

    //! Piecewise linear function defining the disk
    //! - 1 inside the disk
    //! - 0 outside the disk
    //! - 0 < x < 1 in the smoothing region
    double Disk(distance radius) const;

    double Ring(distance radius) const;

    //! Smooth step function: 0 at -infty, 1 at +infty
    static double Sigmoid(double x, double width);

    static double Sigmoid(double variable, double center, double width);

    density Transition(filling disk, filling ring) const;
    int TorusDistance(int x1, int y1, int x2, int y2) const;
    double Radius(int x1, int y1, int x2, int y2) const;
    double NormalisationRing() const;
    double NormalisationDisk() const;

    //! Sets the playing field to random values
    void SeedRandom();

    //! Sets the playing field to constant values
    void SeedConstant(density constant = 0);

    //! Adds a disk to the playing field
    void AddDisk(int x0 = 0, int y0 = 0);

    //! Adds a ring to the playing field
void AddRing(int x0 = 0, int y0 = 0);
//! Sets a single pixel in the field
void AddPixel(int x0, int y0, density value);
//! Moves to next step
void Update();
//! Prints current field to standard output
void Write(std::ostream &out);

//! Returns {disk, ring} integrals at point (x, y)
std::pair<density, density> Integrals(int x, int y) const;

//! Linear index from cartesian index
int Index(int i, int j) const;
//! Cartesian index from linear index
std::pair<int, int> Index(int i) const;

private:
int sizex, sizey;
std::vector<density> field, work_field;
filling birth_1, death_1;
filling birth_2, death_2;
filling smoothing_disk, smoothing_ring;
distance inner, outer, smoothing;
int frame;
double normalisation_disk, normalisation_ring;

#ifdef HAS_MPI
public:
MPI_Comm const &Communicator() const { return communicator; }
void Communicator(MPI_Comm const &comm) { communicator = comm; }

//! Update which layers computation and communication
void LayeredUpdate();

//! Figure start owned sites for given rank
static int OwnedStart(int nsites, int ncomms, int rank);

//! \brief Syncs fields between processes
//! \details Assumes that each rank owns the sites given by OwnedRange.
static void WholeFieldBlockingSync(std::vector<density> &field, MPI_Comm const &comm);

//! \brief Syncs fields between processes without blocking
//! \details Assumes that each rank owns the sites given by OwnedRange.
static MPI_Request WholeFieldNonBlockingSync(std::vector<density> &field, MPI_Comm const &comm);

private:
MPI_Comm communicator;
# ifndef HAS_MPI
# define CATCH_CONFIG_MAIN
# else
# define CATCH_CONFIG_RUNNER
#include <catch.hpp>
#endif

#include <cmath>
#include <random>
#include "catch.hpp"
#include "smooth.h"

TEST_CASE("Compute Integrals") {
  Smooth smooth(300, 300);
  smooth.SeedConstant(0);

  // check for different positions in the torus
  for(auto const x : {150, 298, 0})
    for(auto const y : {150, 298, 0}) {
      SECTION("At position (" + std::to_string(x) + ", " + std::to_string(y) + ")") {
        SECTION("Ring only") {
          smooth.AddRing(150, 150);
          auto const result = smooth.Integrals(150, 150);
          // 0.1 accuracy because of smoothing
          CHECK(std::get<0>(result) == Approx(0).epsilon(0.1));
          CHECK(std::get<1>(result) == Approx(1).epsilon(0.1));
        }
        SECTION("Disk only") {
          smooth.AddDisk(150, 150);
          auto const result = smooth.Integrals(150, 150);
          CHECK(std::get<0>(result) == Approx(1).epsilon(0.1));
          CHECK(std::get<1>(result) == Approx(0).epsilon(0.1));
        }
        SECTION("Disk and ring") {
          smooth.AddRing(150, 150);
          smooth.AddDisk(150, 150);
          auto const result = smooth.Integrals(150, 150);
        }
      }
    }

  // check for different positions in the torus
  for(auto const x : {150, 298, 0})
    for(auto const y : {150, 298, 0}) {
      SECTION("At position (" + std::to_string(x) + ", " + std::to_string(y) + ")") {
        SECTION("Ring only") {
          smooth.AddRing(150, 150);
          auto const result = smooth.Integrals(150, 150);
          // 0.1 accuracy because of smoothing
          CHECK(std::get<0>(result) == Approx(0).epsilon(0.1));
          CHECK(std::get<1>(result) == Approx(1).epsilon(0.1));
        }
        SECTION("Disk only") {
          smooth.AddDisk(150, 150);
          auto const result = smooth.Integrals(150, 150);
          CHECK(std::get<0>(result) == Approx(1).epsilon(0.1));
          CHECK(std::get<1>(result) == Approx(0).epsilon(0.1));
        }
        SECTION("Disk and ring") {
          smooth.AddRing(150, 150);
          smooth.AddDisk(150, 150);
          auto const result = smooth.Integrals(150, 150);
        }
      }
    }
TEST_CASE("Update") {
  // just test playing with a single pixel lit up sufficiently that the
  // transition is non-zero in the ring.
  auto const radius = 5;
  Smooth smooth(100, 100, radius);
  smooth.AddPixel(50, 50, 0.3 * smooth.NormalisationRing());
  CHECK(std::get<0>(smooth.Integrals(50, 50))
       == Approx(0.3 * smooth.NormalisationRing() / smooth.NormalisationDisk()));

  // check the integrals are numbers for which Transition gives non-zero result
  // in the ring
  CHECK(std::get<1>(smooth.Integrals(50, 50)) == Approx(0));
  CHECK(std::get<0>(smooth.Integrals(40, 40)) == Approx(0));
  CHECK(std::get<1>(smooth.Integrals(40, 40)) == Approx(0.3));
  CHECK(std::get<0>(smooth.Integrals(42, 39)) == Approx(0));
  CHECK(std::get<1>(smooth.Integrals(42, 39)) == Approx(0.3));

  // Now call update
  smooth.Update();
  auto const field = smooth.Field();

  // And check death in the disk
  CHECK(field[smooth.Index(50, 50)] == Approx(0));
  CHECK(field[smooth.Index(51, 52)] == Approx(0));
  // And check life in the ring
  CHECK(field[smooth.Index(45, 45)] == Approx(smooth.Transition(0, 0.3)));
  CHECK(field[smooth.Index(42, 39)] == Approx(smooth.Transition(0, 0.3)));
  // And check death outside
  CHECK(field[smooth.Index(15, 15)] == Approx(0));
}

#ifdef HAS_MPI
TEST_CASE("Arithmetics for splitting a field on different nodes") {
  CHECK(Smooth::OwnedStart(5, 2, 0) == 0);
  CHECK(Smooth::OwnedStart(5, 2, 1) == 3);

  for(int i(0); i < 5; ++i)
    CHECK(Smooth::OwnedStart(5, 5, i) == i);

  // with too many procs, some procs have empty ranges

  #endif
for(int i(5); i < 10; ++i)
    CHECK(Smooth::OwnedStart(5, 10, i) == 5);
}

TEST_CASE("Sync whole field") {
    int rank, ncomms;
    MPI_Comm_rank(MPI_COMM_WORLD, &rank);
    MPI_Comm_size(MPI_COMM_WORLD, &ncomms);

    // Create known field: -1 outside owned range, equal to rank inside
    // Different on each process!
    // Also, we make sure the size does not split evenly with the number of procs,
    // because that is a harder test.
    std::vector<density> field(5 * ncomms + ncomms / 3, -1);
    std::fill(field.begin() + Smooth::OwnedStart(field.size(), ncomms, rank),
               field.begin() + Smooth::OwnedStart(field.size(), ncomms, rank + 1), rank);

    SECTION("Blocking synchronisation") {
        Smooth::WholeFieldBlockingSync(field, MPI_COMM_WORLD);

        for(int r(0); r < ncomms; ++r)
            CHECK(std::all_of(field.begin() + Smooth::OwnedStart(field.size(), ncomms, r),
                             field.begin() + Smooth::OwnedStart(field.size(), ncomms, r + 1),
                             [r](density d) { return std::abs(d - r) < 1e-8; }));
    }

    SECTION("Non blocking synchronisation") {
        auto request = Smooth::WholeFieldNonBlockingSync(field, MPI_COMM_WORLD);
        MPI_Wait(&request, MPI_STATUS_IGNORE);

        for(int r(0); r < ncomms; ++r)
            CHECK(std::all_of(field.begin() + Smooth::OwnedStart(field.size(), ncomms, r),
                             field.begin() + Smooth::OwnedStart(field.size(), ncomms, r + 1),
                             [r](density d) { return std::abs(d - r) < 1e-8; }));
    }
}

TEST_CASE("Serial vs parallel") {
    Smooth serial(100, 100, 5);
    Smooth parallel(100, 100, 5);
    parallel.Communicator(MPI_COMM_WORLD);

    // generate one field for all Smooth instances
    std::vector<density> field(100 * 100);
    std::random_device rd; // Will be used to obtain a seed for the random number engine
    std::mt19937 gen(rd());
std::uniform_real_distribution<> randdist(0, 1);
std::generate(field.begin(), field.end(), [randdist, &gen]() { return randdist(gen); });
MPI_Bcast(field.data(), field.size(), MPI_DOUBLE, 0, MPI_COMM_WORLD);

// set the fields for both Smooth instances
serial.Field(field);
parallel.Field(field);

int rank, ncomms;
MPI_Comm_rank(MPI_COMM_WORLD, &rank);
MPI_Comm_size(MPI_COMM_WORLD, &ncomms);

auto const start = Smooth::OwnedStart(field.size(), ncomms, rank);
auto const end = Smooth::OwnedStart(field.size(), ncomms, rank);
// if this is false, then the test itself is wrong
CHECK(std::equal(serial.Field().begin() + start, serial.Field().begin() + end,
                 parallel.Field().begin() + start));

SECTION("Blocking synchronization") {
  // check fields are the same in parallel and in serial for a few iterations
  for(int i(0); i < 3; ++i) {
    serial.Update();
    parallel.Update();
    CHECK(std::equal(serial.Field().begin() + start, serial.Field().begin() + end,
                     parallel.Field().begin() + start));
  }
}

SECTION("Layered communication-computation") {
  for(int i(0); i < 3; ++i) {
    serial.Update();
    parallel.LayeredUpdate();
    CHECK(std::equal(serial.Field().begin() + start, serial.Field().begin() + end,
                     parallel.Field().begin() + start));
  }
}

TEST_CASE("Smooth model can be instantiated and configured", "[Smooth]") {

SECTION("Smooth can be constructed") {
  Smooth smooth;
  REQUIRE(smooth.Size() == 10000);
  REQUIRE(smooth.Field().size() == smooth.Size());
}
TEST_CASE("Smooth mathematical functions are correct", "[Smooth]") {
    Smooth smooth;
    SECTION("Disk support function is correct") {
        REQUIRE(smooth.Disk(500) == Approx(0));
        REQUIRE(smooth.Disk(21.6) == 0.0);
        REQUIRE(smooth.Disk(21.4) > 0.0);
        REQUIRE(smooth.Disk(20.6) > 0.0);
        REQUIRE(smooth.Disk(20.6) < 1.0);
        REQUIRE(smooth.Disk(20.4) == Approx(1.0));
        REQUIRE(smooth.Disk(21.0) == Approx(0.5));
    }
    SECTION("Ring support function is correct") {
        REQUIRE(smooth.Ring(22) == 1.0);
        REQUIRE(smooth.Ring(21.6) == 1.0);
        REQUIRE(smooth.Ring(21.4) > 0.0);
        REQUIRE(smooth.Ring(20.6) > 0.0);
        REQUIRE(smooth.Ring(20.4) == 0.0);
        REQUIRE(smooth.Ring(21.0) == 0.5);
        REQUIRE(smooth.Ring(64.0) == 0.0);
        REQUIRE(smooth.Ring(63.6) == 0.0);
        REQUIRE(smooth.Ring(63.4) > 0.0);
        REQUIRE(smooth.Ring(62.6) > 0.0);
        REQUIRE(smooth.Ring(62.4) == 1.0);
        REQUIRE(smooth.Ring(63.0) == 0.5);
    }
    /// Sigmoid_Test
    SECTION("Sigmoid function is correct") {
        double e = std::exp(1.0);
        REQUIRE(Smooth::Sigmoid(1.0, 1.0, 4.0) == 0.5);
        REQUIRE(std::abs(Smooth::Sigmoid(1.0, 0.0, 4.0) - e / (1 + e)) < 0.0001);
        REQUIRE(Smooth::Sigmoid(10000, 1.0, 4.0) == 1.0);
        REQUIRE(std::abs(Smooth::Sigmoid(0.0, 1.0, 0.1)) < 0.001);
    }
    /// end
    SECTION("Transition function is correct") {
        REQUIRE(std::abs(smooth.Transition(1.0, 0.3) - 1.0) < 0.1);
        REQUIRE(smooth.Transition(1.0, 1.0) == Approx(0));
        REQUIRE(std::abs(smooth.Transition(0.0, 0.3) - 1.0) < 0.1);
    }
}
TEST_CASE("NormalisationsAreCorrect") {
  Smooth smooth(100, 100, 10);
  SECTION("Disk Normalisation is correct") {
    // Should be roughly \pi*radius*radius,
    REQUIRE(std::abs(smooth.NormalisationDisk() - 314.15) < 1.0);
  }
  SECTION("Ring Normalisation is correct") {
    // Should be roughly \pi*outer*outer-\pi*inner*inner, \pi*100*(9-1) = 2513.27
    REQUIRE(std::abs(smooth.NormalisationRing() - 2513.27) < 2.0);
  }
}

TEST_CASE("FillingsAreUnityWhenSeeded") {
  Smooth smooth;
  smooth.SeedConstant(0);
  SECTION("DiskFillingUnityWithDiskSeed") {
    smooth.AddDisk();
    REQUIRE(std::get<0>(smooth.Integrals(0, 0)) == Approx(1).epsilon(0.1));
  }
  SECTION("Disk Filling Zero With Ring Seed") {
    smooth.AddRing();
    REQUIRE(std::get<0>(smooth.Integrals(0, 0)) == Approx(0).epsilon(0.1));
  }
  SECTION("RingFillingUnityWithRingSeed") {
    smooth.AddRing();
    REQUIRE(std::get<1>(smooth.Integrals(0, 0)) == Approx(1).epsilon(0.1));
  }
}

TEST_CASE("FillingFieldHasRangeofValues") {
  Smooth smooth(300, 300);
  smooth.SeedConstant(0);
  smooth.AddRing();
  double min = 1.0;
  double max = 0.0;
for(int x = 0; x < 300; x++) {
    double filling = std::get<1>(smooth.Integrals(x, 0));
    min = std::min(min, filling);  
    max = std::max(max, filling);
}
REQUIRE(min < 0.2);
REQUIRE(max > 0.4);
}

int main(int argc, const char **argv) {

    // There must be exactly one instance
    Catch::Session session;
    MPI_Init(&argc, const_cast<char ***>(&argv));
    auto const result = session.run();
    MPI_Finalize();
    return result;
} #endif

Implementation
#include <cassert>
#include <cmath>
#include <cstdlib>
#include <iostream>
#include "smooth.h"

Smooth::Smooth(int sizex, int sizey, distance inner, filling birth_1, filling birth_2,
                filling death_1, filling death_2, filling smoothing_disk, filling smoothing_ring
: sizex(sizex), sizey(sizey), field(sizex * sizey), work_field(sizex * sizey), inner(inner),
    birth_1(birth_1), birth_2(birth_2), death_1(death_1), death_2(death_2),
    smoothing_disk(smoothing_disk), smoothing_ring(smoothing_ring), outer(inner * 3),
    smoothing(1.0)
#ifndef HAS_MPI
    communicator(MPI_COMM_SELF)
#endif
{
    normalisation_disk = NormalisationDisk();
    normalisation_ring = NormalisationRing();
}
const std::vector<density> &Smooth::Field() const { return field; }
void Smooth::Field(std::vector<density> const &input) {
    assert(field.size() == input.size());
    field = input;
}

int Smooth::Range() const { return outer + smoothing / 2; }
int Smooth::Sizex() const { return sizex; }
int Smooth::Sizey() const { return sizey; }
int Smooth::Size() const { return sizex * sizey; }

/// "Disk_Smoothing"
double Smooth::Disk(distance radius) const {
    if(radius > inner + smoothing / 2)
        return 0.0;
    if(radius < inner - smoothing / 2)
        return 1.0;
    return (inner + smoothing / 2 - radius) / smoothing;
}

/// end

double Smooth::Ring(distance radius) const {
    if(radius < inner - smoothing / 2)
        return 0.0;
    if(radius < inner + smoothing / 2)
        return (radius + smoothing / 2 - inner) / smoothing;
    if(radius < outer - smoothing / 2)
        return 1.0;
    if(radius < outer + smoothing / 2)
        return (outer + smoothing / 2 - radius) / smoothing;
    return 0.0;
}

double Smooth::Sigmoid(double variable, double center, double width) {
    return Sigmoid(variable - center, width);
}
double Smooth::Sigmoid(double x, double width) {
    return 1.0 / (1.0 + std::exp(-4.0 * x / width));
}
density Smooth::Transition(filling disk, filling ring) const {
    auto const sdisk = Sigmoid(disk - 0.5, smoothing_disk);
    auto const t1 = birth_1 * (1.0 - sdisk) + death_1 * sdisk;
    auto const t2 = birth_2 * (1.0 - sdisk) + death_2 * sdisk;
    return Sigmoid(ring - t1, smoothing_ring) * Sigmoid(t2 - ring, smoothing_ring);
}
int Smooth::Index(int i, int j) const { return i * Sizex() + j; }
std::pair<int, int> Smooth::Index(int i) const { return {i / Sizex(), i % Sizex()}; }

/// "Torus_Difference"
int Smooth::TorusDistance(int x1, int x2, int size) const {
    auto const remainder = std::abs(x1 - x2) % size;
    return std::min(remainder, std::abs(remainder - size));
}

/// end

double Smooth::Radius(int x1, int y1, int x2, int y2) const {
    int xdiff = TorusDistance(x1, x2, sizex);
    int ydiff = TorusDistance(y1, y2, sizey);
    return std::sqrt(xdiff * xdiff + ydiff * ydiff);
}

double Smooth::NormalisationDisk() const {
    double total = 0.0;
    for(int x = 0; x < sizex; x++)
        for(int y = 0; y < sizey; y++)
            total += Disk(Radius(0, 0, x, y));
    return total;
}

double Smooth::NormalisationRing() const {
    double total = 0.0;
    for(int x = 0; x < sizex; x++)
        for(int y = 0; y < sizey; y++)
            total += Ring(Radius(0, 0, x, y));
    return total;
}

void Smooth::Update() {
#ifdef HAS_MPI
    int rank, ncomms;
    MPI_Comm_rank(Communicator(), &rank);
    MPI_Comm_size(Communicator(), &ncomms);

    WholeFieldBlockingSync(field, communicator);
    auto const start = OwnedStart(Size(), ncomms, rank);
    auto const end = OwnedStart(Size(), ncomms, rank + 1);
#else
    auto const start = 0;
    auto const end = field.size();
#endif
}
for(int i(start); i < end; ++i) {
    auto const xy = Index(i);
    auto const integrals = Integrals(xy.first, xy.second);
    work_field[i] = Transition(integrals.first, integrals.second);
}

std::swap(field, work_field);
frame++;
}

#elif HAS_MPI

void Smooth::LayeredUpdate() {
    int rank, ncomms;
    MPI_Comm_rank(Communicator(), &rank);
    MPI_Comm_size(Communicator(), &ncomms);

    auto request = WholeFieldNonBlockingSync(field, communicator);
    auto const start = OwnedStart(Size(), ncomms, rank);
    auto const end = OwnedStart(Size(), ncomms, rank + 1);
    auto const interaction = Sizex() * static_cast<int>(std::floor(outer + smoothing / 2 + 1));

    auto const set_work_field_at_index = [this](int i) {
        auto const xy = Index(i);
        auto const integrals = Integrals(xy.first, xy.second);
        work_field[i] = Transition(integrals.first, integrals.second);
    };

    for(int i(start + interaction); i < end - interaction; ++i)
        set_work_field_at_index(i);

    MPI_Wait(&request, MPI_STATUS_IGNORE);

    for(int i(start); i < std::min(end, start + interaction); ++i)
        set_work_field_at_index(i);

    for(int i(std::min(end, end - interaction)); i < end; ++i)
        set_work_field_at_index(i);

    std::swap(field, work_field);
    frame++;
}
#endif

std::pair<density, density> Smooth::Integrals(int x, int y) const {
    density ring_total(0), disk_total(0);
    for(std::vector<density>::size_type i(0); i < field.size(); ++i) {
        ...
auto const cartesian = Index(i);
int deltax = TorusDistance(x, cartesian.first, sizex);
if(deltax > outer + smoothing / 2)
{
    continue;
}

int deltay = TorusDistance(y, cartesian.second, sizey);
if(deltay > outer + smoothing / 2)
{
    continue;
}

double radius = std::sqrt(deltax * deltax + deltay * deltay);

double fieldv = field[i];
ring_total += fieldv * Ring(radius);
disk_total += fieldv * Disk(radius);
}
return {disk_total / NormalisationDisk(), ring_total / NormalisationRing();}
}

void Smooth::SeedRandom()
{
    for(int x = 0; x < sizex; x++)
    for(int y = 0; y < sizey; y++)
        field[Index(x, y)] += (static_cast<double>(rand()) / static_cast<double>(RAND_MAX));
}

void Smooth::SeedConstant(density constant) { std::fill(field.begin(), field.end(), constant); }

void Smooth::AddDisk(int x0, int y0) {
    for(int x = 0; x < sizex; x++)
    for(int y = 0; y < sizey; y++)
        field[Index(x, y)] += Disk(Radius(x0, y0, x, y));
}

void Smooth::AddRing(int x0, int y0) {
    for(int x = 0; x < sizex; x++)
    for(int y = 0; y < sizey; y++)
        field[Index(x, y)] += Ring(Radius(x0, y0, x, y));
}

void Smooth::AddPixel(int x0, int y0, density value) { field[Index(x0, y0)] = value; }

void Smooth::Write(std::ostream &out) {
    for(int x = 0; x < sizex; x++)
    for(int y = 0; y < sizey; y++)
        out << field[Index(x, y)] << " , ";
    out << std::endl;
}
out << std::endl;

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int Smooth::Frame() const { return frame; }

#ifdef HAS_MPI
int Smooth::OwnedStart(int nsites, int ncomms, int rank) {
        assert(nsites >= 0);
        assert(ncomms > 0);
        assert(rank >= 0 and rank <= ncomms);
        return rank * (nsites / ncomms) + std::min(nsites % ncomms, rank);
    }

void Smooth::WholeFieldBlockingSync(std::vector<density> &field, MPI_Comm const &comm) {
    int rank, ncomms;
    MPI_Comm_rank(comm, &rank);
    MPI_Comm_size(comm, &ncomms);

    if(ncomms == 1)
        return;

    std::vector<int> displacements{0}, sizes;

    for(int i(0); i < ncomms; ++i) {
        displacements.push_back(Smooth::OwnedStart(field.size(), ncomms, i + 1));
        sizes.push_back(displacements.back() - displacements[i]);
    }

    MPI_Allgatherv(MPI_IN_PLACE, sizes[rank], MPI_DOUBLE, field.data(), sizes.data(),
                   displacements.data(), MPI_DOUBLE, comm);
}

MPI_Request Smooth::WholeFieldNonBlockingSync(std::vector<density> &field, MPI_Comm const &comm) {
    int rank, ncomms;
    MPI_Comm_rank(comm, &rank);
    MPI_Comm_size(comm, &ncomms);

    std::vector<int> displacements{0}, sizes;

    for(int i(0); i < ncomms; ++i) {
        displacements.push_back(Smooth::OwnedStart(field.size(), ncomms, i + 1));
        sizes.push_back(displacements.back() - displacements[i]);
    }

    MPI_Request request;
    MPI_Iallgatherv(MPI_IN_PLACE, sizes[rank], MPI_DOUBLE, field.data(), sizes.data(),
                    displacements.data(), MPI_DOUBLE, comm, &request);
    return request;
}
Halo swap

Where do we put the communicated data?

The data we need to receive from our neighbour needs to be put in a place where it can be conveniently used to calculate the new state of cells within distance \( r \) of the boundary.

The standard design pattern for this is to use a **Halo Swap**: we extend the memory buffer used for our state, adding new space either side of our main domain to hold a **halo**.

Domains with a halo

Thus, each process now holds \( N + 2r \) cells:

- \( 0 \leq x < r \), the left halo, holding data calculated by the left neighbour
- \( r \leq x < 2r \), data which we calculate, which will form our left neighbour’s right halo
- \( 2r \leq x < N \), data which we calculate, unneeded by neighbours
- \( N \leq x < N + r \), data which we calculate, which will form our right neighbour’s left halo
- \( N + r \leq x < N + 2r \), the right halo.

Domains with a halo

\[
\begin{align*}
\text{range(outer + smoothing / 2)}, \ & \text{local}_x\_size(sizex / mpi\_size), \\
\text{local}_x\_size\_with\_halo(local\_x\_size + 2 \times \text{range}), \ & \text{total}_x\_size(sizex), \\
\text{x_coordinate_offset(rank \times local\_x\_size - \text{range})}, \ & \text{local}_x\_min\_calculate(\text{range}), \\
\text{local}_x\_max\_needed\_left(2 \times \text{range}), \ & \text{local}_x\_max\_calculate(\text{range} + local\_x\_size), \\
\text{local}_x\_min\_needed\_right(local\_x\_size),
\end{align*}
\]

Coding with a halo

We will thus **update** the field only from \( r \) to \( N + r \), but we will **access** the field from 0 to \( N + 2r \):
int from_x=0; int to_x=local_x_size_with_halo

void Smooth::QuickUpdateStripe(int from_x, int to_x) {
    for(int x = from_x; x < to_x; x++) {
        for(int y = 0; y < sizey; y++) {
            double ring_total = 0.0;
            double disk_total = 0.0;
            for(int x1 = 0; x1 < local_x_size_with_halo; x1++) {
                int deltax = TorusDifference(x + x_coordinate_offset, x1 + x_coordinate_offset, total_x_size);
                if(deltax > outer + smoothing / 2)
                    continue;
                for(int y1 = 0; y1 < sizey; y1++) {
                    int deltay = TorusDifference(y, y1, sizey);
                    if(deltay > outer + smoothing / 2)
                        continue;
                    double radius = std::sqrt(deltax * deltax + deltay * deltay);
                    double fieldv = Field(x1, y1);
                    ring_total += fieldv * Ring(radius);
                    disk_total += fieldv * Disk(radius);
                }
            }
            SetNewField(x, y, transition(disk_total / normalisation_disk, ring_total / normalisation_ring));
        }
    }
}

Transferring the halo

We need to pass data left at the same time as we receive data from the right.
If we use separate Send and Recv calls, we’ll have a deadlock: process 2 will be
sending to process 1, while process 3 is sending to process 2. No process will be
executing a Recv.
Fortunately, MPI provides Sendrecv: expressing that we want to do a blocking
Send to one process while we simultaneously do a blocking Recv from another.
Exactly what we need for pass-the-parcel.
Transferring the halo

```cpp
void Smooth::CommunicateMPI() {
    BufferLeftHaloForSend();
    MPI_Sendrecv(send_transport_buffer, range * sizey, MPI_DOUBLE, left, rank,
                 receive_transport_buffer, range * sizey, MPI_DOUBLE, right, right, MPI_COMM_WORLD,
                 MPI_STATUS_IGNORE);
    UnpackRightHaloFromReceive();
    BufferRightHaloForSend();
    MPI_Sendrecv(send_transport_buffer, range * sizey, MPI_DOUBLE, right, mpi_size + rank,
                 receive_transport_buffer, range * sizey, MPI_DOUBLE, left, mpi_size + left,
                 MPI_COMM_WORLD, MPI_STATUS_IGNORE);
    UnpackLeftHaloFromReceive();
}
```

Noncontiguous memory

Our serial solution uses a C++ `vector<vector<double>>` to store our field. That means each column of data starts at a different location in memory; our data is not contiguous in memory, the outer vector holds a series of pointers to the start of each inner vector.

This means we can’t just transmit $Mr$ doubles in one go by sending from the address of `&field1[0][0]`.

Buffering

We’ll get around this by copying all the data into a send buffer, and unpacking from a receive buffer. This adds to our communication overhead, but doesn’t change its scaling behaviour, both overheads are $O(Mr)$.

We’ll look in a later section how this can be avoided.

Buffering

```cpp
void Smooth::BufferLeftHaloForSend() {
    for(int x = local_x_min_calculate; x < local_x_min_calculate + range; x++) {
        for(int y = 0; y < sizey; y++) {
            send_transport_buffer[y * range + x - local_x_min_calculate] = Field(x, y);
        }
    }
}
```
void Smooth::UnpackRightHaloFromReceive() {
    for(int x = local_x_max_calculate; x < local_x_size_with_halo; x++) {
        for(int y = 0; y < sizey; y++) {
            SetField(x, y, receive_transport_buffer[y * range + x - local_x_max_calculate]);
        }
    }
}

Testing communications

We implement a copy of our buffers which doesn’t use MPI:

void Smooth::CommunicateLocal(Smooth &left, Smooth &right) {
    BufferLeftHaloForSend();
    std::memcpy(left.receive_transport_buffer, send_transport_buffer,
                sizeof(density) * range * sizey);
    left.UnpackRightHaloFromReceive();
    BufferRightHaloForSend();
    std::memcpy(right.receive_transport_buffer, send_transport_buffer,
                sizeof(density) * range * sizey);
    right.UnpackLeftHaloFromReceive();
}

which allows us to check our halos are all set up correctly.

Testing communications

We test that our copy works as expected:

TEST_CASE("CommunicationBufferingFunctionsCorrectly") {
    Smooth smooth(200, 100, 5, 0, 2);
    Smooth smooth2(200, 100, 5, 1, 2);
    REQUIRE(smooth.LocalXSize() == 100);
    REQUIRE(smooth.LocalXSizeWithHalo() == 130);
    REQUIRE(smooth.Radius(0, 0, 0, 0) == 0);
    REQUIRE(smooth2.Radius(0, 0, 0, 0) == 0);
    smooth.SeedDisk(); // Half the Seeded Disk falls in smooth2's domain, so total filling will
                      // half a disk.
    REQUIRE(smooth.Field(15, 0) == 1.0);
    REQUIRE(std::abs(smooth.FillingDisk(15, 0) - 0.5) < 0.1);
}
REQUIRE(smooth2.FillingDisk(85, 0) == 0.0);
smooth.CommunicateLocal(smooth2, smooth2); // Transport the data
REQUIRE(std::abs(smooth.FillingDisk(15, 0) - 0.5) < 0.1);
REQUIRE(smooth2.Field(115, 0) == 1.0);
REQUIRE(std::abs(smooth2.FillingDisk(115, 0) - 0.5) < 0.1);
}

Testing communications

We test that the MPI comms results are the same as serial

smooth.QuickUpdate();
paraSmooth.CommunicateMPI();
paraSmooth.QuickUpdate();
paraSmooth.CommunicateMPI();
for(unsigned int x = 0; x < 50; x++) {
    for(unsigned int y = 0; y < 100; y++) {
        REQUIRE(std::abs(smooth.Field(x + 15 + rank * 50, y) - paraSmooth.Field(x + 15, y)) < 0.00001);
    }
}

Deployment

Getting our code onto Legion

We now have to

- Clone our code base onto our cluster
- Load appropriate modules to get the compilers we need
- Build our code
- Construct an appropriate submission script
- Submit the submission script
- Wait for the job to queue and run
- Copy the data back from the cluster
- Analyse it to display our results.

This is a pain in the neck
Scripting deployment

There are various tools that can be used to automate this process. You should use one.

Since I like Python, I use Fabric to do this.

You create `fabfile.py` in your top level folder, and at the shell you can write:

```bash
fab legion.cold
fab legion.sub
fab legion.stat
fab legion.fetch
```

to build and run code on Legion, without ever using ssh. You can be in any folder below the `fabfile.py` level, such as inside a build folder; the command line tool recurses upward to look for this file, (Just like git looks for the .git folder.)

Writing fabric tasks

If you know Python, writing fabric tasks is easy:

`fabfile.py`:

```python
@task
def build():
    with cd('/home/ucgajhe/smooth/build'):
        with prefix('module load cmake'):
            run('make')

fab build
```

Templating jobscripts

Editing a jobscript every time you want to change the number of cores you want to run on is tedious. I use a templating tool Mako to generate the jobscript:

```
#$ -pe openmpi ${processes}
```

the templating tool fills in anything in `{}` from a variable in the fabric code.
Configuration files

Avoid using lots of command line arguments to configure your program. It’s easier to just have one argument, a configuration file path.

- The configuration file can be kept with results for your records.
- Configuration files can be shipped back and forth to the cluster with fabric.

Even if your code is using simple C++ I/O rather than a nice formatting library, it’s best to design your config file in a format which other frameworks can easily read. My favourite is Yaml.

```
width: 200
height: 100
range: 5
```

Results

It works

Derived Datatypes

Avoiding buffering

We’ve still got our ugly re-buffering of the halo data into contiguous memory, arising from our use of `vector<vector<double>>`

We can get around that by changing to a flat 1-d array of memory, and storing the \((x, y)\) element at `Field[Mx + y]`. This works fine, especially if we refactor all access to get and update from the field into accessors, so we only need make the change in one place.

Wrap Access to the Field

```cpp
density Smooth::Field(int x, int y) const { return (*field)[sizey * x + y]; }
```
void Smooth::SetNewField(int x, int y, density value) { (*fieldNew)[sizey * x + y] = value; }

void Smooth::SetField(int x, int y, density value) { (*field)[sizey * x + y] = value; }

void Smooth::SeedField(int x, int y, density value) {
    SetField(x, y, value + Field(x, y));
    if(Field(x, y) > 1.0) {
        SetField(x, y, 1.0);
    }
}

Copy Directly without Buffers

void Smooth::CommunicateMPIUnbuffered() {
    MPI_Sendrecv((*field) + sizey * range, range * sizey, MPI_DOUBLE, left, rank,
                 (*field) + sizey * local_x_max_calculate, range * sizey, MPI_DOUBLE, right, MPI_COMM_WORLD, MPI_STATUS_IGNORE);
    MPI_Sendrecv((*field) + sizey * local_x_min_needed_right, range * sizey, MPI_DOUBLE, right, mpi_size + rank, (*field), range * sizey, MPI_DOUBLE, left, mpi_size + left, MPI_COMM_WORLD, MPI_STATUS_IGNORE);
}

Defining a Halo Datatype

It’s usually the case in programming that thinking of a whole body of data as a single entity produces cleaner, faster code than programming at the level of the individual datum.

We want to be able to think of the Halo as a single object to be transferred, rather than as a series of doubles.

We can do this using an MPI Derived Datatype:

Declare Datatype

void Smooth::DefineHaloDatatype() {
    MPI_Type_contiguous(sizey * range, MPI_DOUBLE, &halo_type);
    MPI_Type_commit(&halo_type);
}
Use Datatype

```cpp
void Smooth::CommunicateMPIDerivedDatatype() {
    MPI_Sendrecv(*field + sizey * local_x_min_calculate, 1, halo_type, left, rank,
                 *field + sizey * local_x_max_calculate, 1, halo_type, right, right, MPI_COMM_WORLD,
                 MPI_STATUS_IGNORE);
    MPI_Sendrecv(*field + sizey * local_x_min_needed_right, 1, halo_type, right, mpi_size + rank,
                 *field, 1, halo_type, left, mpi_size + left, MPI_COMM_WORLD, MPI_STATUS_IGNORE);
}
```

Strided datatypes

Supposing we wanted to use a 2-d decomposition. Our y-direction halo’s data would not be contiguous in memory.

Let’s imagine we used $N_y + x$ to index into the field instead of $M_x + y$: we can define a derived datatype which specifies data as a series of stretches, with gaps. 

```
MPI_Type_Vector(M, r, N)
```

would define the relevant type for this: $M$ chunks, each $r$ doubles long, each $N$ doubles apart in memory.

Overlapping computation and communication

Wasteful blocking

We are calculating our whole field, then sharing all the halo.

This is wasteful: we don’t need our neighbour’s data to calculate the data in the middle of our field.

We can start transmitting our halo and receiving our neighbour’s, while calculating our middle section. Thus, our communication will not take up extra time, as it is overlapped with calculation.

To do this, we need to use asynchronous communication.

Asynchronous communication strategy

- Start sending/receiving data
- Do the part of calculation independent of needed data
- Wait until the communication is complete (hopefully instant, if it’s already finished)
- Do the part of the calculation that needs communicated data

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Asynchronous communication for 2-d halo:

- Start clockwise send/receive
- Calculate middle
- Finish clockwise send
- Start anticlockwise send
- Calculate right
- Finish anticlockwise send
- Calculate left

Asynchronous MPI

```cpp
template <typename density, int rank, int halo_type, int sizey>
void Smooth::InitiateLeftComms() {
    MPI_Isend(*field + sizey * local_x_min_calculate, 1, halo_type, left, rank, MPI_COMM_WORLD,
              &request_left);
    MPI_Irecv(*field + sizey * local_x_max_calculate, 1, halo_type, right, right, MPI_COMM_WORLD,
              &request_left);
}
```

```cpp
template <typename density, int rank, int halo_type, int sizey>
void Smooth::ResolveLeftComms() { MPI_Wait(&request_left, MPI_STATUS_IGNORE); }
```

Implementation of Asynchronous Communication

```cpp
template <typename density, int rank, int halo_type, int sizey>
void Smooth::UpdateAndCommunicateAsynchronously() {
    InitiateLeftComms();
    // Calculate Middle Stripe
    QuickUpdateStripe(local_x_max_needed_left, local_x_min_needed_right);
    ResolveLeftComms(); // So we now have the right halo
    InitiateRightComms();
    QuickUpdateStripe(local_x_min_needed_right, local_x_max_calculate);
    ResolveRightComms();
    QuickUpdateStripe(local_x_min_calculate, local_x_max_needed_left);
    SwapFields();
}
```

```cpp
template <typename density, int rank, int halo_type, int sizey>
density *Smooth::StartOfWritingBlock() { return &(*field)[local_x_min_calculate * sizey]; }
```
Chapter 10

MPI File I/O

Parallel Input and Output

Objectives for the Chapter

In this chapter, we discuss how to get information out of our parallel programs, and how to configure them.

Because we need to reconcile the outputs from different processes, and distribute configuration amongst them, this can be quite complex.

We will look at a few different choices of how to do this.

Visualisation

Visualisation is important

We’ve put a lot of emphasis onto using Unit Tests to verify your code. However, it’s just as important to visualise your results.

“The graph looks OK” is not a good enough standard of truth.

But “the graph looks wrong” is a strong indication of problems!

Visualisation is hard

Three-D slices through your data, contour surfaces, animations…

These are all really important for understanding scientific output.

But doing the visualisation in C++ can be very very difficult.
Simulate remotely, analyse locally

Visualise locally if you can.

Use a tool like fabric to organise your data and make it easy to ship it back from the cluster.

Use dynamic languages like Python or Matlab to visualise your data, or tools like Paraview or VisIt.

In-Situ visualisation

When working at the highest scales of parallel, saving out raw simulation data becomes impossible: compute power scales faster than storage.

Under these circumstances, it is necessary to do data reduction on the cluster.

In-situ visualisation, where animations or graphs are constructed as part of the analysis, is one approach.

Visualising with NumPy and Matplotlib

The Python toolchain for visualising quantitative data is powerful, fast and easy.

Here’s how we turn a NumPy Matrix view of our data, with axes (frames, width, height), into an animation and save it to disk as an mp4:

```python
import matplotlib.animation
import matplotlib.pyplot

def plot(frames, outpath):
    figure = matplotlib.pyplot.figure()
    def _animate(frame_id):
        print "Processing frame", frame_id
        matplotlib.pyplot.imshow(frames[frame_id], vmin=0, vmax=1)
    anim = matplotlib.animation.FuncAnimation(figure, _animate, len(frames), interval=100)
    anim.save(outpath)

try doing that with C++ libraries!
```

Formatted Text IO

Formatted Text IO

- Is easy
When and when not to use text IO

Use text IO:

- Metadata
- Configuration
- Logging

Don’t use it:

- For output of large numerical datasets
- For initial conditions
- For checkpoint/restart

Use libraries to generate formatted text

For a templating library like Mako in C++ try CTemplate

This is a great way to create XML and YAML files.

Raw CSV file generation with built-in C++ `<iostream>` is not very robust.

Libraries will automatically quote strings which contain commas.

Boost’s Spirit library is good for this too.

Low-level IO exemplar

Nevertheless, in simple cases where text is of a known format, C++’s built in formatted IO is quick and easy:

```cpp
void TextWriter::Header(int frames) {
    *outfile << smooth.LocalXSize() << "", " << smooth.Sizey() << ", " << rank << ", " << size
       << std::endl;
}
```
void TextWriter::Write() {
    for(int x = smooth.Range(); x < smooth.LocalXSize() + smooth.Range(); x++) {
        for(int y = 0; y < smooth.Sizey(); y++) {
            *outfile << smooth.Field(x, y) << " , "
        }
        *outfile << std::endl;
    }
    *outfile << std::endl;
}

Text Data and NumPy

Parsing text data in NumPy is also very easy:

```python
buffer=numpy.genfromtxt(data,delimiter="",)[::,-1]
```

```python
frames_data=buffer.reshape([frame_count, width/size, height])
```

this is how the animations you saw last lecture were created.

Text File Bloat

However, representing $1.3455188104 \cdot 10^{-10}$ in text:

```
1.3455188104e-10,
```

uses many bytes per number, (one per character in ASCII, more in unicode) whereas recording in a binary representation, even at double precision, typically uses 8.

It’s also harder do parallel IO in such files, as the distance of a certain quantity from the start of the file can’t be predicted.

Binary IO in C++

Binary IO in C++

To do basic binary file IO in C++, we set `std::ios::binary` as we open the file.

However, we can’t use the nice `<<` operator, as these still generate formatted `chars`. We have to use `ostream::write()`, which is a low-level, C-style function.
Binary IO in C++

```cpp
void BinaryWriter::Header(int frames) {
    outfile->write(reinterpret_cast<char *>(&sizex), sizeof(int));
    outfile->write(reinterpret_cast<char *>(&sizey), sizeof(int));
    outfile->write(reinterpret_cast<char *>(&rank), sizeof(int));
    outfile->write(reinterpret_cast<char *>(&size), sizeof(int));
    outfile->write(reinterpret_cast<char *>(&frames), sizeof(int));
}

void BinaryWriter::Write() {
    outfile->write(reinterpret_cast<char *>(smooth.StartOfWritingBlock()),
                   local_element_count * sizeof(double));
}
```

Binary IO in NumPy

```python
buffer = numpy.fromfile(data, bulk_type, frame_count*height*width/size)
```

... more on that ‘bulk_type’ parameter later.

Endianness and Portability

Portability

There's a big problem with binary files: the bytes that get written for a `double` on one platform are different from those on others.

This is particularly problematic in HPC: the architecture of your laptop is unlikely to be the same as on ARCHER, so if when ship your output files back, your nice visualisation code will not work.

Length of datatypes

The C++ standard does not specify how many bytes are used to represent a `double`: just that it is more than a `float`! (And similarly for other datatypes.)

This means that your data will be represented differently on different platforms.

Therefore, don’t forget to use `sizeof(type)` whenever working with byte-level routines like MPI, if you want your code to work on both your laptop, and your local supercomputer.
Endianness

Another problem is Endianness: a double is actually almost always 8 bytes, with a 1 bit sign, 11 bit exponent, and 52 bit mantissa. (The IEEE standard).

But there’s still ambiguity in how these bytes are ordered.

The 4-byte standard signed integer “5” can be represented as:

```
00000000 00000000 00001001 00000000 (little endian)
```

or as

```
00000101 00000000 00000000 00000000 (big endian)
```

NumPy dtypes

As long as you’re aware of these problems, you can usually make your visualiser compatible with the endianness and byte counts of the data you’re getting off your system.

If you’re visualising with python you can set your datatype to be e.g. `<f8` for a little endian 8-byte floating point, or `>i4` for a big-endian 4-byte signed integer.

XDR

XDR, or ‘extensible data representation’ is a portability standard defined for binary IO. If you write through XDR, data will be converted to the XDR standard representation.

XDR data is big endian, has everything in multiples of 4 bytes, and supports a rich library of appropriate types.

Writing with XDR

```c
#include <cstdio>
#include <rpc/types.h>
#include <rpc/xdr.h>

void XDRWriter::Write() {
    char *start_to_write = reinterpret_cast<char *>(smooth.StartOfWritingBlock());
```
Writing from a single process

One process, one file

So far, we’ve been writing one process from each file, calling them frames.dat.0, frames.dat.1 etc.

```python
def append_process_suffix(prefix, process):
    return prefix + '.' + str(process)

def process_many_files(folder, prefix, size, header_type, bulk_type):
    process_frames=[]
    for process in range(size):
        path=os.path.join(folder, append_process_suffix(prefix, process))
        # code snippet...
    return numpy.concatenate(process_frames, 1)
```

One process, one file

This is necessary, because we cannot simply have multiple processes writing to the same file, without considerable care, as they will:

- Block while waiting for access to the file
- Overwrite each others’ content

However, the one-process-one-file approach does not scale to large numbers of files: the file system can be overwhelmed once we’re running at the thousands-of-processors level.
Writing from a single process

An alternative approach is to share all the data to a master process. (Which can be done in $O(lnp)$ time), and then write from that file. This avoids the complexity of reconciling datafiles locally, and avoids overwhealming the cluster filesystem with many small files.

void SingleWriter::Write() {
    double *receive_buffer;
    if(rank == 0) {
        receive_buffer = new double[total_element_count];
    }
    MPI_Gather(smooth.StartOfWritingBlock(), local_element_count, MPI_DOUBLE, receive_buffer, local_element_count, MPI_DOUBLE, 0, MPI_COMM_WORLD);
    if(rank == 0) {
        xdr_vector(&xdrfile, reinterpret_cast<char *>(receive_buffer), total_element_count, sizeof(double), reinterpret_cast<xdrproc_t>(xdr_double));
    }
}

Writing from a single process

It’s important to create the file and write the header only on the master process:

SingleWriter::SingleWriter(Smooth &smooth, int rank, int size) : SmoothWriter(smooth, rank, size)
if(rank != 0) {
    return;
}

Parallel IO

Serialising on a single process

We know that *any task which is not $O(1/p)$ will eventually dominate the cost as number of processes increases, preventing scaling.
This is Amdahl’s law again.

The master-process-writes approach introduces this problem; IO quickly becomes the dominant part of the task, preventing weak scaling as problem sizes and processor counts increase.

**Parallel file systems**

Supercomputers provide **parallel file systems**. These store each file in multiple “stripes”: one can obtain as much parallelism in IO as there are stripes in files. To make use of this, it is necessary to use MPI’s parallel IO library, MPI-IO.

**Introduction to MPI-IO**

MPI-IO works by accessing files as data buffers like core MPI_Send and so on:

```c
MPI_File_write(outfile, smooth.StartOfWritingBlock(), local_element_count, MPI_DOUBLE, MPI_STATUS_IGNORE);
```

**Opening parallel file**

All processes work together to open and create the file:

```c
MPI_File_open(MPI_COMM_WORLD, const_cast<char *>(fname.str().c_str()), MPI_MODE_CREATE | MPI_MODE_RDWR, MPI_INFO_NULL, &outfile);
```

**Finding your place**

The hard part is synchronising things so that each process writes it’s section of the file:

```c
int offset = 4 * sizeof(int) + 
            rank * local_element_count * sizeof(double) + 
            smooth.Frame() * total_element_count * sizeof(double); // Frame offset in the file

MPI_File_seek(outfile, offset, MPI_SEEK_SET);
```
High level research IO libraries

Standard libraries for scientific data formats such as HDF5 support parallel IO. You should use these if you can, as you’ll get the benefits of both endianness-portability and parallel IO, together with built-in metadata support and compatibility with other tools.

Introduction to HDF5 or NetCDF is beyond the scope of this course, but familiarising yourself with these is strongly encouraged!
Chapter 11

Accelerators

Introduction to Accelerators

Gamers vs Researchers

- Global entertainment industry (2016): $1800bn
- Global video-game industry (PC and console) (2016): $100bn
- Global video-game industry – software expenditure (2016): $50bn
- US government research efforts (pre-Trump), ~$100bn
- Global HPC industry (2011): $10bn

What is an accelerator?

- An accelerator is a piece of computing hardware that performs some function faster than is possible in software running on a general purpose CPU
  - an example is the Floating Point Unit inside a CPU that performs calculations on real numbers faster than the Arithmetic and Logic Unit
  - better performance is achieved through concurrency - the ability to perform several operations at once, in parallel
- When an accelerator is separate from the CPU it is referred to as a “hardware accelerator”

Why would I want to use one?

- Available on many laptop, desktops, phones, and supercomputers
• Can speed-up calculations 2-20x by off-loading from CPU to GPU
• Especially good at streaming
  – 3D graphics
  – MPEG decoding
  – cryptography

CPU Vectorisation

Using an accelerator within the CPU

• Consider the following code which performs \( y = a \cdot x + y \) on vectors \( x \) and \( y \):

```c
int saxpy(int n, float a, const float * restrict x, int incx,
          float * restrict y, int incy) {
    if (n < 0)
        return 1;
    for (int i = 0; i < n; i++)
        y[i * incy] += a * x[i * incx];
    return 0;
}
```

• Assuming single precision floating point multiply-add is implemented as one CPU instruction the for loop executes \( n \) instructions (plus integer arithmetic for the loop counter, etc.).

CPUs as multicore vector processors

• In the mid-1990s, Intel were investigating ways of increasing the multimedia performance of their CPUs without being able to increase the clock speed
• Their solution was to implement a set of registers capable of executing the same operation on multiple elements of an array in a single instruction
  – known as SIMD (Single Instruction, Multiple Data)
  – branded as MMX (64-bit, integer only), SSE (128-bit, integer + floating point) and AVX (256-bit)
Compiler Autovectorisation

- Modern compilers are able to automatically recognise when SIMD instructions can be applied to certain loops
  - elements are known to be contiguous
  - arrays are not aliased

```c
int saxpy_fast(int n, float a, const float * restrict x, int incx,
               float * restrict y, int incy) {
    if (n < 0)
        return 1;

    if (incx == 1 && incy == 1) {
        for (int i = 0; i < n; i++)
            y[i] += a * x[i];
    } else {
        for (int i = 0; i < n; i++)
            y[i * incy] += a * x[i * incx];
    }

    return 0;
}
```

Autovectorisation results

- Running this and timing the invocations produces the following output:

```
n = 10000, incx = 1, incy = 1
saxpy: 0.008837ms
saxpy_fast: 0.002802ms
```

- Since Intel’s SIMD registers operate on 4 single precision floats at once the first loop executes approximately \(n/4\) instructions
  - resulting in a (near) 4x speedup (Amdahl’s Law)
- Can be combined with OpenMP directives (lecture 5) to use SIMD units on multiple cores of the CPU

Support for Autovectorisation

- GCC
- use `-ftree-loop-vectorize` to activate
- use `-fopt-info-vec` to check whether loops are vectorised
- automatically performed with `-O3`

- **ICC**
  - use `-vec` (Linux/OSX) or `/Qvec` (Windows) to activate
  - use `-vec-report=n`/`/Qvec-report:n` to check whether loops are vectorised (`n > 0`)
  - automatically performed with `-O2`/`/O2` and higher

### Support for Autovectorisation

- **Clang**
  - use `-fvectorize` to activate
  - automatically performed at `-O2` and higher
  - no way to see compiler auto-detection

- **MSVC**
  - activated by default
  - use `/Qvec-report:n` to check whether loops are vectorised (`n > 0`)

### Using a GPU as an Accelerator

#### GPUs for 3D graphics

- 3D graphics rendering involves lots of operations on 3 and 4 dimensional vectors
  - positions of vertices (x, y, z)
  - colour and transparency of pixels (RGBA)
- Realtime graphics rendering needs to be fast so accuracy is often sacrificed
  - GPUs are good at 32-bit integer and single precision floating point arithmetic
  - not as good at 64-bit integer and double precision floating point
- 3D graphics operations are independent of one another
  - and can be performed in parallel
GPUs as multicore vector processors

- Multicore vector processors.
- Much faster memory access to their own memory
- Slow transfer between host and device (~5GB/s)
- Constrained and complex access to different types of device memory
- SIMD: Single Instruction, Multiple Data

<table>
<thead>
<tr>
<th>Property</th>
<th>CPU</th>
<th>GPU</th>
</tr>
</thead>
<tbody>
<tr>
<td>No of threads</td>
<td>&lt;10</td>
<td>1000s</td>
</tr>
<tr>
<td>SIMD width</td>
<td>256 bits</td>
<td>1024 bits</td>
</tr>
<tr>
<td>Memory bandwidth</td>
<td>&lt;25 GB/s</td>
<td>&lt;500 GB/s</td>
</tr>
</tbody>
</table>

GPU Hardware

- Each core (multiprocessor - often abbreviated as “SM" or “MP") has:
  - 128 Single Precision Floating Point Units
  - 32 Double Precision FPUs (that also perform single precision transcendental functions)
  - 16384 32-bit registers (very fast access)
  - 64KB of “shared” cache memory (fast access)
  - 10KB of “constant” cache memory (writable by the CPU, fast, read-only access by the GPU)
- GPUs also have “global” memory which is the figure quoted by graphics card manufacturers (2GB, etc.)
  - accessed by the CPU across the PCI-Express bus
  - high latency, slow access by the GPU (but still up to 20x faster than CPU RAM access)

General Purpose Programming for GPUs

- Threads are organised in groups of 32 called “warps”
- Threads also organised “Thread blocks” of several warps
- “Thread blocks” are organised in 1-d, 2-d, or 4-d grids
- “Thread blocks” are assigned to Streaming Multiprocessors
- A GPU cards is contains 1-50 Streaming Multiprocessor
GPU Architecture

SIMT

- Threads within a block have access to the same “shared” memory
- Threads within a block can synchronise
- No communication or synchronisation primitives across blocks/SMs
  - can use atomic operations on variables in global memory (slow)
- This type of programming is a hybrid between threaded programming and SIMD and hence is called SIMT by Nvidia

Executing code on the GPU

- Functions executed on the GPU and called from the CPU are called “kernels”
  - kernel execution is asynchronous to the CPU
  - CPU must call a function which blocks until the GPU has finished executing the current kernel before attempting to download results
  - there is an implicit synchronisation barrier on the GPU at the start of each kernel
- Common programming pattern:
  - upload data from CPU RAM to GPU global memory (slow)
– execute kernel(s) on GPU (fast)
– synchronise
– download results (slow)

GPU-accelerating your code

• Four options to GPU accelerate code:
  – replace existing libraries with GPU-accelerated ones (ArrayFire, cuFFT, cuBLAS, Magma, …)
  – use compiler directives to automatically generate GPU-accelerated portions of code (OpenMP 4.5, OpenACC)
  – use CUDA::Thrust C++ template library to build your own kernels
  – write your own kernels in CUDA-C

Streaming computations

*Single Instruction*, Multiple Data

All threads in a warp perform the exact same hardware instruction, but on different data.

On thread 0:

```cpp
a[0] = alpha * x[0] + 1;
__syncthreads();
y[0] -= a[31];
```

On thread 31:

```cpp
a[31] = alpha * x[31] + 1;
__syncthreads();
y[31] -= a[0];
```

Is this single instruction?

```cpp
a[thread_id] = ((thread_id < 16) ? 1: -1) + alpha * x[thread_id];
if(thread_id < 16)
  a[thread_id] = alpha * x[thread_id] + 1;
else
  a[thread_id] = alpha * x[thread_id] - 1;
__syncthreads();
y[thread_id] -= a[31 - thread_id];
```
Single Instruction, *Multiple Contiguous Data*

Each thread in a warp should access a consecutive address in memory. Lets create a kernel for copying a vector using a single block consisting of a single warp (of 32 threads). Assume that the size of the vector is a multiple of 32. threadIdx.x is the thread id in Cuda, and automagically passed to each kernel.

The following is a jumble of too many expressions: put it back in order.

```c
__global__ void copy(float *odata, const float *idata, int n)
{
    for (int j = 0; j < n; ++j)
        odata[j + threadIdx.x] = idata[j + threadIdx.x];
    for (int j = 0; j < n; j += 32)
        odata[j + threadIdx.x] = idata[j + threadIdx.x];
    for (int j = 0; j < n; ++j)
        odata[j + threadIdx.x * 32] = idata[j + threadIdx.x * 32];

    odata[j + threadIdx.x] = idata[j + threadIdx.x];
    odata[j + threadIdx.x * 32] = idata[j + threadIdx.x * 32];
}
```

The joys of indexing

Now imagine rewriting the same code with n by m by p blocks of threads, where each block is u by v by w threads. You have access to the size of the block blockDim.x, the index and size of the grid (of blocks) blockIdx.(x, y, z) and gridDim.(x, y, z).

To limit memory transfers, each warp should read and write to contiguous arrays in memory.

See also: shared memory, bank conflicts

Structure of Arrays or Arrays of structures

```c
struct Atom { int x, y, z; }; 
std::vector<Atom> molecule;

or

struct Molecule {
    std::vector<int> x;
    std::vector<int> y;
    std::vector<int> z;
};
```
Using Somebody Else’s Software is a good way to avoid becoming a GPU expert. But which? Check for usability and sustainability:

- Does it do what you need?
- What license is it under?
- Is it simply available on Github/BitBucker/Gitlab?
- Are there automatic tests?
- Is it an active development repo? (number of commits, date of last commit)
- How many people/labs/companies are committing to it? (check pull-requests)
- Is there a community of users? (check issues, wiki)
- Is there documentation?

Broadcasting (Vectorization)

GPU require running the same operations over multiple data (SIMD). Broadcasting transforms nested loops into a set of matrix or array operations amenable to SIMD, and likely to be already GPU-ized by external libraries.

Also useful for MATLAB, Python/numpy, etc...

Example:

\[ G = \max(||A - R_i||) \]

for all \( i \), where \( A \) and \( R_i \) are vectors

Naive solution with explicit loops and STL:

```cpp
double naive(std::vector<std::array<double, 3>> const &Rs,
             std::array<double, 3> const &A) {
    double result(0);
    for (auto const &R : Rs) {
        if (result < norm)
            result = norm;
    }
    return std::sqrt(result);
}
```
Broadcasting (Vectorization)

1. transform A from a vector to a matrix A_{3xn}
2. compute pow2 = (R - A_{3xn})^2 elementwise
3. sum pow2 over columns
4. reduce final vector using max

```cpp
double broadcasting(std::vector<std::array<double, 3>> const &Rs,
std::array<double, 3> const &A) {
    af::array A_3x1(A.size(), A.data());
    af::array gpuRs(A.size(), Rs.size(), Rs[0].data());

    auto const A_3xn = af::tile(A_3x1, 1, Rs.size());
    auto const norms = af::sum(af::pow(gpuRs - A_3xn, 2), 0);
    auto const result = af::max(norms);
    std::shared_ptr<double> const host_array(result.host<double>(),
        [](double *ptr) { delete[] ptr; });

    return std::sqrt(*host_array);
}
```

A word on transfer rate

- Transfer to GPU takes time $T_0$

```cpp
af::array gpuRs(A.size(), Rs.size(), Rs[0].data());
af::array gpuA(A.size(), A.data());
```

- Compute takes $C/n$, $n$ the number of GPU threads

```cpp
auto const result
    = af::max(af::sum(af::pow(gpuRs - af::tile(gpuA, 1, Rs.size()), 2), 0));
```

- Transfer to CPU takes time $T_1$

```cpp
af::array gpuRs(A.size(), Rs.size(), Rs[0].data());
af::array gpuA(A.size(), A.data());
```

- Then, possibly $T_0 + T_1 + C/n > C$
Exercise: Broadcasting

\[ G = \sum_{i, j, k} \cos(K_k \cdot (R_i - R_j)) \]

Given

```cpp
std::vector<std::array<double, 3>> const Rs = {...};
std::vector<std::array<double, 3>> const Ks = {...};
```

Write code to compute \( G \) (pseudo, or real code):

1. Create an \( R_s \) array of \((3, 1, n)\) using `af::moddims` \((n = R_s\text{.size()}\))
2. Create an \( R_s \) array of \((3, n, 1)\) using `af::moddims`
3. Tile appropriately
4. Compute the dot product using `af::moddims, af::matmul, af::transpose`
5. Sum over the cosine of the result

Exercise: Line of Sight

Given an \( n \) by \( m \) matrix of altitudes, with \( n \) orientations and \( m \) distances:

```cpp
std::vector<double> altitudes(nOrientations * mDistances) = { ... };
```

Compute the whether any point \( i, j \) is in sight. A point is in sight if, for that orientation, all previous angles between X-horizon and X-point are smaller

\[
\text{atan}(z_i / (\text{stepsize} \times i))
\]

`af::scanf` might come in handy: - given \([a0, a1, a2, ..., aN]\) - it computes \([0, a0, a0 + a1, a0 + a1 + a2, ..., a0 + ... + aN]\) - leading zero is removed in exclusive scans

This function looks intrinsically difficult to parallelize, but there are well known solutions. It is useful in many fields.
The end of brace wars: brainless auto formatting

Code is read more often than written:

```cpp
int GCD(int a, int b) {int r; while(b) {r = a % b; a = b; b = r;} return a;}
```

`clang-format` is one possible code formatter. Add it (or any equivalent) to your editor for automatic formatting.

Linting

Software to check code for correctness:

- the compilers themselves: `--Wall`
- clang-tidy
- cppcheck

Example:

```cpp
if (FFTW_plan_flag != FFTW_ESTIMATE | FFTW_PRESERVE_INPUT) {
...
}
```

Refactoring

Once tests exist, it is easy and safe to modify the code:

Refactoring means rewriting:

- to simplify existing code
- to simplify future development
- for legibility
- to decrease tech-debt
- to consolidate similar code (avoid copy-pasta)

Checking memory allocation intrusively

There are a variety of compiler flags to check standard memory errors:

- g++: `-fsanitize=address`
- clang: `address sanitizer`

Good when debugging/testing, but may impact performance. May not detect all memory errors (e.g. read before initialization).
Checking memory allocation non-intrusively

Valgrind is an instrumentation framework for Linux and (older) Mac.
It detects memory errors and leaks by intercepting every memory access, allocation, and deallocation.
Unfortunately, Valgrind currently does not work with Mac OS/X > 10.11.
So let’s use docker (on Linux) and docker-machine (Windows, Mac OS/X)!

Exercise: Traveling salesman solved by Simulated Annealing

Traveling Salesman Problem:
A salesman living in an n-dimensional world must visit N cities. What is the shortest path?
Simulated Annealing:

• Start from a candidate A
• Create a neighbor B of A
• if path(A) > path(B), then swap A and B
• else if \( \exp(\beta \cdot (\text{path}(B) - \text{path}(A))) > \text{random()} \), then swap A and B
• loop until satisfied

Setting up a docker VM and docker

First download/update to the latest course:

\[ \text{git clone https://github.com/UCL-RITS/research-computing-with-cpp} \]

Creating a virtual machine is optional on Linux, and necessary on Windows and Mac OS/X

\[ \text{> docker-machine create cpp_course} \]
\[ \quad \text{--driver virtualbox} \]
\[ \quad \text{--virtualbox-memory 4000} \]
\[ \quad \text{--virtualbox-cpu-count 2} \]
\[ \text{> eval $(docker-machine env cpp_course)} \]
The last lines lets docker know on which VM it should create containers.

Then ssh into the machine and look for your home directory:

```bash
> docker-machine ssh cpp_course
> pwd
# Mac users
> ls /Users/
# Linux users
> ls /home
# Windows users
# uhm, no idea :(, look around and let me know!
```

Then, create a Dockerfile specifying the container we want:

```bash
> mkdir docker_dir
> cat > docker_dir/Dockerfile <<EOF
FROM ubuntu:latest
RUN apt-get update && apt-get install -y cmake g++ valgrind
EOF
```

Build an image of the container

```bash
> docker build -t course_container /path/to/docker_dir
```

Now build the code in `11Performance/cpp` using an instance (a.k.a container) of the image.

First, check you can see the directory with the source:

```bash
> docker run --rm \
-\v /path/to/source/on/vm:/path/to/source/on/container \
-w /path/to/source/on/container \
course_container \
ls
```

This should print the content of the directory on your machine, if:

- the virtual box VM was set-up to mount your home directory (automatic)
- the container was set up to mount the VM directory (`-v path/VM:/path/container`)

Finally, replace the `ls` command to:

1. create a build directory in the source code directory
2. run cmake from the build directory
3. run make in the build directory

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Running valgrind on program called awful

Assuming everything went well, there should be a compiled program called awful.
It can only run inside the container!
It has memory leaks and bugs. Investigate and correct using valgrind:

> docker run --rm
   
   -v /path/to/source/on/vm:/path/to/source/on/container
   -w /path/to/source/on/container
   course_container
   valgrind -v --leak-check=full --show-leak-kinds=all
   --track-origin=yes ./awful

Running valgrind on program called less_bad

Even programs written without explicit memory allocations can have memory
bugs.
The next version uses Eigen to solve the same problem.

1. add libeigen3-dev to the Dockerfile
2. rebuild the image
3. re-build the code
4. run valgrind on less_bad
5. investigate and correct the code

Amdahl’s law

Maximum theoretical speedup for parallelizing a given task.

- Given a program that takes a time $T$ to execute
- When parallelizing a task $A$ taking time $P$ over $n$ threads
- Then the maximum speedup is:
  $S = \frac{T}{T + \frac{P}{n}}$

Some simple cases:

- $P \to 0$, $n \to \infty$, then $S \to 1$
\[ P = 0.5, \ n \to \infty, \text{ then } S \to 2 \]

In practice, it means programmers/researchers should measure performance before jumping to “optimize” code (a.k.a. apply the scientific method?).

**Profiling**

Refers to measuring how much time the program spends in each function:

- Valgrind, via callgrind provides an accurate, non-intrusive solution
- gprof require “instrumenting” the program, e.g. recompiling with the gprof library. It polls the program every so often, a.k.a. “sampling”
- gperftools, intrusive, thread-capable, can select parts of code to profile, info can be visualized by kcachegrind and/or pprof
- XCode instruments provides “sampling” without requiring instrumentation

Other considerations to take into account: threads, GPU-specifics, MPI...

**Exercise: Profiling the correct less_bad**

1. install kcachegrind or qcachegrind (latter on Mac + Homebrew, or Windows)
2. recompile in release mode + debug info cmake -DCMAKE_BUILD_TYPE=RelWithDebInfo
3. Run the following command:

   ```
   > docker run --rm \
   -v /path/to/source/on/vm:/path/to/source/on/container \
   -w /path/to/source/on/container \
   course_container \
   valgrind -v --tool=callgrind ./awful
   ```

   1. Then run qcachegrind on the artifact file:

   ```
   > /usr/local/Cellar/qcachegrind/16.12.0/bin/qcachegrind callgrind.out.1
   ```

Notice the difference between include and self, e.g. the time spent in the function as whole, vs the time spent strictly in the function excluding sub-function calls.

Questions:
1. How much time is spent on `neighbor`
2. How much time is spent on `TravelDistance::operator()`
3. What happens when you increase the number of cities to 100, or 10000
4. What should we optimize or parallelize?

**Micro-benchmarking**

meaningful bit of code, and systematically run performance tests for each
Scientific method applied to performance: measure the time taken by each commit.
Profiling can tell us what part to include in a benchmark.
Possible micro-benchmarking frameworks:

- timers in unit test framework (not really accurate)
- google/benchmark
- hayai, a “google-test”-like framework
- others

Questions micro-benchmarking can answer:

1. How long does it take on average
2. Standard-deviation from average
3. Worst case

Questions it doesn’t always answer:

1. Performance of large subsets or whole application
2. Parallelization: communication vs computation

**Exercise: build and run micro_benchmark**

`micro_benchmark` reproduces the evaluation function from the travelling salesman problem.
It reproduces how a micro-benchmark framework works:

1. run code N times for warm-up
2. run code N’ times for actual measurement
Make sure the code is built in \texttt{Release} mode.

Questions:

1. Why is the float implementation faster/slower? Does the speed-up change with the number of dimensions or the number of cities? What happens for \texttt{Nrow} = 3?
2. Write a function that computes the distance manually (without Eigen syntactic sugar). Is it faster for \texttt{Nrows}=2? What about \texttt{Nrows}=8?

Remark:
None of these have tests (this is an exercise in bad code, after all). Do you trust we are solving the travelling salesman problem? I know that \texttt{awful} does not, beyond the memory bugs... Because I added a bug. But maybe there are further bugs still.

How much easier would it be to test the \texttt{manual} code above if we had tests for the evaluation function?
Chapter 12

Post-coding medley, memory leaks, and performance measurements

Please install docker and docker-machine

Mac OS/X:

> brew install Caskroom/cask/virtualbox
> brew install docker-machine
> brew install docker

or

> brew install Caskroom/cask/docker-toolbox


Windows or Mac OS/X: - https://www.docker.com/products/docker-toolbox

Side-note: Flynn’s Taxonomy of Parallelization

- SISD: Single instruction single data

prototypical serial code
• SIMD: Single instruction multiple data

Same instruction is performed in parallel over different inputs. Necessary in GPU (at the level of a warp of 32 threads). Likely in OpenMP (for loop parallelization) and MPI.

• MIMD: Multiple instruction multiple data

Basically, different threads or different nodes doing different things, e.g. computing different terms in an equation, dealing one with the GUI, the other with a database, etc...

• MISD: Multiple instructions single data

Weird... Used for fault tolerance (different algorithm that should lead to same output).

All tests pass, the code works: are we done yet?

Lots of changes can be made to a code, from cosmetic to crucial:

• formatting, linting, and refactoring
• checking for memory leaks
• profiling and performance
• benchmarking
Chapter 13

Appendix - STL

Appendix - STL

Standard Template Library

This lecture got compressed into lecture 3.

- Notes provided here in their entirety, for historical reasons.

The Standard Template Library

Contents

STL reference

- Containers
  - Sequences
  - Associative
- Iterators
- I/O streams
- Functors/functions
- Algorithms
Motivation

Motivation - why STL?

Because you have better things to do than rediscovering the wheel.

Task: Read in two files with an unknown number of integers, and sort them altogether.

The WRONG way to do it!

```c
#include <stdio.h>
#include <stdlib.h>

int compare(const void* a, const void* b)
{
    return *(int*)a - *(int*)b;
}

int main()
{
    FILE* if1 = fopen("95stl/cpp/randomNumbers1.txt","r");
    FILE* if2 = fopen("95stl/cpp/randomNumbers2.txt","r");

    // First determine size of array needed
    int size1=0, size2=0;
    while (!feof(if1)) {
        fscanf(if1, "%*d");
        if (!feof(if1)) size1++;
    }
    rewind(if1);
    while (!feof(if2)) {
        fscanf(if2, "%*d");
        if (!feof(if2)) size2++;
    }
    rewind(if2);

    // Read in the data
    int theArray[size1+size2];
    for (int i=0;i<size1;++i) {
        fscanf(if1, "%d", &theArray[i]);
    }
    for (int i=size1;i<size1+size2;++i) {
        fscanf(if2, "%d", &theArray[i]);
    }
```
fclose(if1);
fclose(if2);

// Sort and output
qsort(theArray, size1+size2, sizeof(int), compare);
for (int i=0; i<size1+size2; ++i) {
    printf("%d ", theArray[i]);
}

STL solution

#include <vector>
#include <algorithm>
#include <iostream>
#include <fstream>

int main()
{
    std::ifstream if1("95stl/cpp/randomNumbers1.txt", std::ifstream::in);
    std::ifstream if2("95stl/cpp/randomNumbers2.txt", std::ifstream::in);

    // Read in the data.
    int number;
    std::vector<int> theArray;
    while (!if1.eof()) {
        if1 >> number;
        if (!if1.eof()) theArray.push_back(number);
    }
    while (!if2.eof()) {
        if2 >> number;
        if (!if2.eof()) theArray.push_back(number);
    }
    if1.close();
    if2.close();

    // Sort and output
    std::sort(theArray.begin(), theArray.end());
    for (int i=0; i<theArray.size(); ++i) {
        std::cout << theArray[i] << " ";
    }
}
More conveniences

- Similar API between different STL containers. eg. myContainer->begin(),
  end(), at(), erase(), clear(), size()....
- Algorithms and data structures optimised for speed and memory.

STL Containers

Two general types:

- **Sequences**: Elements are ordered in a linear sequence. Individual elements
  are accessed by their position in this sequence/index. eg. myVector[3],
  myList.front(), etc.

- **Associative**: Elements are referenced by their key and not by their position
  in the container. eg. myMap['key1']

Sequences

Properties:

- **size**: fixed/dynamic
- **access**: random/sequential
- underlying memory structure: contiguous/not
  - random access in non-contiguous memory is tricky
  - affects how efficiently inserting/removing of elements can be done
  - affects if pointer arithmetic can be done
- optimised insert/remove operations

Sequences

<table>
<thead>
<tr>
<th>Name</th>
<th>size</th>
<th>access</th>
<th>memory</th>
<th>efficient insert/remove</th>
</tr>
</thead>
<tbody>
<tr>
<td>array</td>
<td>fixed</td>
<td>random</td>
<td>contiguous</td>
<td>-</td>
</tr>
<tr>
<td>vector</td>
<td>dynamic</td>
<td>random</td>
<td>contiguous</td>
<td>at end only</td>
</tr>
<tr>
<td>deque</td>
<td>dynamic</td>
<td>random</td>
<td>non-contiguous</td>
<td>both ends</td>
</tr>
<tr>
<td>list</td>
<td>dynamic</td>
<td>sequential</td>
<td>non-contiguous</td>
<td>anywhere</td>
</tr>
<tr>
<td>forward_list</td>
<td>dynamic</td>
<td>sequential, only forward</td>
<td>non-contiguous</td>
<td>anywhere</td>
</tr>
</tbody>
</table>
More on vector

It’s dynamic, so you can add/erase elements:

```cpp
std::vector<int> myVec;
for (int i=0;i<10;++i) {
    myVec.push_back(i);
}
myVec.insert(myVec.begin(),-1);
myVec.erase(1);

myVec = -1 1 2 3 4 5 6 7 8 9
... or manipulate its size:

std::vector<int> myVec(10);
std::cout << "Vector size before = " << myVec.size();
myVec.resize(5);
std::cout << " after = " << myVec.size() << "\n";

Vector size before = 10 after = 5

Note on C++11

- For all containers, `emplace(const_iterator position, Args&&... args)` is preferable to `insert(const_iterator position, const value_type& val)`, as it doesn’t create any copies of the object you add to the container.
- Cases you might prefer something other than `emplace`:
  - backward compatibility
  - `insert` has more constructors
  - at least `emplace_back` might not work as expected in some implementations

Exercise

Think of cases where you’d use a specific container

Associative containers

Properties:
• key: is it separate from value?
  – maps: key-value pair
  – sets: value is the key (and thus it’s const!)
• ordering. Affects performance:
  – unordered containers fastest to fill and access by key
  – ordered containers fastest to iterate through, and they’re already ordered :o)
• unique values?

**Associative containers**

<table>
<thead>
<tr>
<th></th>
<th>ordered</th>
<th>unordered</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>unique</strong></td>
<td>map</td>
<td>unordered_map</td>
</tr>
<tr>
<td><strong>non-unique</strong></td>
<td>multimap</td>
<td>unordered_multimap</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>ordered</th>
<th>unordered</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>unique</strong></td>
<td>set</td>
<td>unordered_set</td>
</tr>
<tr>
<td><strong>non-unique</strong></td>
<td>multiset</td>
<td>unordered_multiset</td>
</tr>
</tbody>
</table>

**More on maps**

Fill them with

```cpp
std::map<std::string, int> myMap;
myMap["broccoli"] = 2;
myMap["garlic"] = 1;
myMap["broccoli"] = 1;  // returns reference to element => element is updated
myMap.insert( std::pair<std::string, int>("bread",4) );
myMap.insert( std::pair<std::string, int>("broccoli",3) );  // returns iterator
  // to existing element => element is not updated

typedef std::multimap<std::string, int> MMMapType;
MMMapType myMMMap;  // there’s no [] for multimap
myMMMap.insert( std::pair<std::string, int>("broccoli",2) );
myMMMap.insert( std::pair<std::string, int>("bread",4) );
myMMMap.insert( std::pair<std::string, int>("broccoli",3) );
```

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More on maps

Access elements with

```cpp
std::cout << "myMap[brocolli] = " << myMap["brocolli"] << "\n";
std::cout << "myMap.find(bread) = " << myMap.find("bread")->second << "\n";
for (std::map<std::string, int>::iterator it=myMap.begin(); it!=myMap.end(); ++it)
    std::cout << it->first << " " << it->second << "\n";
```

```cpp
std::cout << "myMMap.count(brocolli) = " << myMMap.count("brocolli") << "\n";
std::pair < MMapType::iterator, MMapType::iterator> range = myMMap.equal_range("brocolli");
for (MMapType::iterator it=range.first; it!=range.second; ++it)
    std::cout << it->first << " " << it->second << "\n";
for (MMapType::iterator it=myMMap.begin(); it!=myMMap.end(); ++it)
    std::cout << it->first << " " << it->second << "\n";
```

myMap[brocolli] = 1
myMap.find(bread) = 4
bread 4
broocolli 1
garlic 1
myMMap.count(brocolli) = 2
broocolli 2
broocolli 3
bread 4
broocolli 2
broocolli 3

Example

Read in file with unknown number of particle-momentum pairs.

proton 4.5
electron 17.8
pion 4.6
muon 12.0
pion 3.2
neutrino 8.2
pion 23.7
proton 9.4
neutrino 6.7

• Then print out
1. list of all particle-momentum pairs in alphabetical order
   — how about in ascending momentum order?
2. list of types of particles in the file in alphabetical order
   — how about in mass order?
3. list of particle-max momentum pairs

Task 1

```cpp
#include <map>
#include <string>
#include <iostream>
#include <fstream>

int main()
{
    std::ifstream ifs("95stl/cpp/particleList.txt",std::ifstream::in);
    // Read in the data
    std::multimap<std::string,double> theParticles;
    std::string name;
    double momentum;
    while (!ifs.eof()) {
        ifs >> name >> momentum;
        if (!ifs.eof())
            theParticles.insert( std::pair<std::string,double>(name,momentum) );
    }
    ifs.close();
    // Output - it's already sorted!
    std::multimap<std::string,double>::iterator iter = theParticles.begin();
    for ( ; iter!=theParticles.end(); ++iter) {
        std::cout << iter->first << " " << iter->second << std::endl;
    }
}
```

Task 2

- Hint1: use a set
- Hint2: write a custom comparator

Task 2

- Hint1: use a set
- Hint2: write a custom comparator
class compMass {
public:
    compMass() :
        m_particlesOrdered({"neutrino", "electron", "muon", "pion", "kaon", "proton"})
    {};
    bool operator() (const std::string& s1, const std::string& s2) const
    {
        int index1=-1, index2=-1;
        for (int i=0;i<m_particlesOrdered.size();++i) {
            if (m_particlesOrdered[i]==s1) index1 = i;
            if (m_particlesOrdered[i]==s2) index2 = i;
        }
        return index1<index2; // unknown particles appear first (index=-1)
    }
private:
    std::vector<std::string> m_particlesOrdered;
};

then use it in the set constructor:
std::set<std::string,compMass> theParticles;

Task 3

Hint: extend the map class. - How?

- inherit from std::map - NO!
  - STL containers are not designed to be polymorphic, as they don’t have virtual destructors (Meyers 1:7)
- composition - write a class that contains either an std::map object or smart pointer to it.
- free functions with STL containers/iterators as arguments

Task 3 - with function

void keepMax(std::map<std::string,double>& theMap, const std::string key, const double value) {
    if (theMap.find(key)==theMap.end()) //element doesn’t already exist in map
        theMap[key] = value;
    else
        // any logic can go in here, eg a counter, an average, etc...
theMap[key] = std::max(theMap[key],value);
}

Then use it in the loop when filling the container:
if (!feof(ifp)) keepMax( theParticles,name,momentum );

**Task 3 - with composition**

class maxMap
{
 public:
 typedef std::map<std::string,double>::iterator iterator;

 void insert(const std::pair<std::string,double> theElement)
 {
   std::string key = theElement.first;
   double value = theElement.second;
   if (m_map.find(key)==m_map.end()) //element doesn't already exist in map
     m_map[key] = value;
   else // any logic can go in here, eg a counter, an average, etc...
     m_map[key] = std::max(m_map[key],value);
 }

 iterator begin() { return m_map.begin(); }
 iterator end() { return m_map.end(); }
 private:
   std::map<std::string,double> m_map;
};

**Task 3 - with composition**

Then use it in main:

int main()
{
  std::ifstream ifs("95stl/cpp/particleList.txt",std::ifstream::in);
  // Read in the data
  maxMap theParticles;
  std::string name;
  double momentum;
  while (!ifs.eof()) {

ifs >> name >> momentum;
if (!ifs.eof())
    theParticles.insert( std::pair<std::string,double>(name,momentum) );
}
ifs.close();
// Output - it's already sorted!
maxMap::iterator iter = theParticles.begin();
for ( ; iter!=theParticles.end(); ++iter) {
    std::cout << iter->first << " " << iter->second << std::endl;
}
}

Accessing containers: iterators

For random-access containers you can do

std::vector<int> myVector = {1,2,3,4};
for (int i=0; i<myVector.size(); ++i) {
    std::cout << myVector[i] << std::endl;
}

But for sequential-access ones, you can only do

#include <iterator>

std::list<int> mylist = {1,2,3,4};
std::list<int>::iterator it=mylist.begin()
for ( ; it!=mylist.end(); ++it) {
    std::cout << *it << std::endl;
}

Iterators

An iterator is any object that, pointing to some element in a container, has the ability to iterate through the elements of that range using a set of operators.

- Minimum operators needed: increment (++) and dereference (*).
- A pointer is the simplest iterator.
- Brings some container-independence.
  - especially when using typedef

http://www.cplusplus.com/reference/iterator/
Pairs

```cpp
#include <utility> // std::pair

int main () { 
    std::pair<std::string,double> product1; // default constructor
    std::pair<std::string,double> product2("tomatoes",2.30); // value init

    auto foo = std::make_pair (10,20); // created a std::pair<int,int>
    product1 = std::make_pair(std::string("lightbulbs"),0.99); // using make_pair

    product2.first = "shoes"; // the type of first is string
    product2.second = 39.90; // the type of second is double

    std::cout << "foo: " << foo.first << " , " << foo.second << '\n';

    return 0;
}
```

Tuples

```cpp
#include <tuple> // std::tuple, std::make_tuple, std::get

int main() {
    std::tuple<int,char> one; // default
    std::tuple<int,char> two(10,'a'); // initialization

    auto first = std::make_tuple (10,'a'); // tuple < int, char >
    const int a = 0; int b[3]; // decayed types:
    auto second = std::make_tuple (a,b); // tuple < int, int* >

    std::cout << "two contains: " << std::get<0>(two);
    std::cout << " and " << std::get<1>(two);
    std::cout << std::endl;

    return 0;
}
```
Assignment

- Download a human genome file from ftp://ftp.ensembl.org/pub/release-87/fasta/homo_sapiens/dna/ . This is a sequence of characters from the dictionary \{A,C,G,T,N\}.
- List all the possible 3-letter combinations (k-mers for k=3) that appear in this file together with the number of appearances of each in the file, in order of number of appearances.

  - Output should be something like

\[
\begin{align*}
\text{NNN} & \quad 3345028 \\
\text{AGT} & \quad 2348 \\
\text{CTT} & \quad 1578 \\
\ldots
\end{align*}
\]

- Hint: use only associative containers. You’ll need to extend their functionality to some kind of counter.

Other STL stuff

Contents

- I/O with streams
- Function objects
- Algorithms

Streams

- `iosstream` for `std::cout`, `std::cerr` and `std::cin`
- `fstream` for file I/O
- `sstream` for string manipulation

```cpp
std::ifstream myfile(filename,std::ifstream::in);
if (!myfile.good()) {
    stringstream mess;
    mess << "Cannot open file " << filename << " . It probably doesn't exist." << endl;
    throw runtime_error(mess.str());
}

myfile.seekg(0, std::ios::end);
m_size = myfile.tellg();
```
m_buffer = new char[m_size];
myfile.seekg(0, std::ios::beg);
myfile.read(m_buffer, m_size);
myfile.close();

Function objects

• Remember std::set<std::string, compMass> theParticles;?
• There are several ways to define function-type stuff in c++, and
  std::function wraps them all
  – functions and function pointers
  – functors (i.e. an object with an operator() member function)
  – lambdas (i.e. nameless inline functions)
• Such functions can be used as comparators in STL containers or algorithms,
among other things.

# include <functional> // std::function, std::negate

// a function:
int half(int x) {return x/2;}

// a function object class:
struct third_t {
  int operator()(int x) {return x/3;}
};

// a class with data members:
struct MyValue {
  int value;
  int fifth() {return value/5;}
};

int main () {
  std::function<int(int)> fn1 = half; // function
  std::function<int(int)> fn2 = &half; // function pointer
  std::function<int(int)> fn3 = third_t(); // function object
  std::function<int(int)> fn4 = [](int x){return x/4;}; // lambda expression
  std::function<int(int)> fn5 = std::negate<int>(); // standard function object

  std::cout << "fn1(60): " << fn1(60) << \n';
  std::cout << "fn2(60): " << fn2(60) << \n';
std::cout << "fn3(60): " << fn3(60) << 'n';
std::cout << "fn4(60): " << fn4(60) << 'n';
std::cout << "fn5(60): " << fn5(60) << 'n';

// stuff with members:
std::function<int(MyValue&)> value = &MyValue::value; // pointer to data member
std::function<int(MyValue&)> fifth = &MyValue::fifth; // pointer to member function

MyValue sixty {60};
std::cout << "value(sixty): " << value(sixty) << 'n';
std::cout << "fifth(sixty): " << fifth(sixty) << 'n';
}

Function objects

Output:

fn1(60): 30
fn2(60): 30
fn3(60): 20
fn4(60): 15
fn5(60): -60
value(sixty): 60
fifth(sixty): 12

Algorithms

http://www.cplusplus.com/reference/algorithm/

- A collection of functions especially designed to be used on ranges of elements.
- Don’t start coding any task, before checking if it’s already there!

std::sort(RandomAccessIter first, RandomAccessIter last, Compare comp)
std::transform(InIter first, InIter last, OutIter res, UnaryOp op)
std::transform(InIter1 first1, InIter1 last1, InIter2 first2, OutIter res, BinaryOp op)
std::for_each (InIter first, InIter last, Function fn)
std::max_element(ForwardIter first, ForwardIter last, Compare comp)
std::min_element(ForwardIter first, ForwardIter last, Compare comp)
std::minmax_element(ForwardIter first, ForwardIter last, Compare comp)
Chapter 14

Appendix - TMP

Appendix - Template Meta-Programming

No Longer In Course

- Notes provided here for historical reasons.
- Also, read “Modern C++ Design”

Template Meta-Programming (TMP)

What Is It?

- See Wikipedia, Wikibooks, Keith Schwarz
- C++ Template
  - Type or function, parameterised over, set of types, constants or functions
  - Instantiated at compile time
- Meta Programme
  - Program that produces or manipulates constructs of target language
  - Typically, it generates code
- Template Meta-Programme
  - C++ programme, uses Templates, generate C++ code at compile time
TMP is Turing Complete

- Given: A Turing Machine
  - Tape, head, states, program, etc.
- A language is “Turing Complete” if it can simulate a Turing Machine
  - e.g. Conditional branching, infinite looping
- Turing’s work underpins much of “what can be computed” on a modern computer
  - C, C++ no templates, C++ with templates, C++ TMP
  - All Turing Complete
- Interesting that compiler can generate such theoretically powerful code.
- But when, where, why, how to use TMP?
- (side-note: Its not just a C++ pre-processor macro)

Why Use It?

- Use sparingly as code difficult to follow
- Use for
  - Optimisations
  - Represent Behaviour as a Type
  - Traits classes
- But when you see it, you need to understand it!

Factorial Example

See Wikipedia Factorial Example

- This:

```cpp
#include <iostream>
using namespace std;

template <int n>
struct factorial {
    enum { value = n * factorial<n - 1>::value }
};
```
template <>
struct factorial<0> {
    enum { value = 1 }; 
};

int main () {
    std::cout << factorial<0>::value << std::endl;
    std::cout << factorial<8>::value << std::endl;
}

• Produces:

1
40320

Factorial Notes:

• Compiler must know values at compile time
  – i.e. constant literal or constant expression
  – See also constexpr
• Generates/Instantiates all functions recursively
• Factorial 16 = 2004189184
• Factorial 17 overflows
• This simple example to illustrate “computation”
• But when is TMP actually useful?
• Notice that parameter was an integer value … not just “int” type

Loop Example

• This:

#include <iostream>
#include <vector>
using namespace std;

template<typename T>
T Sum(const std::vector<T>& data)
{
    T total = 0;
    for (size_t i = 0; i < data.size(); i++)
    {
        total += data[i];
    }
    return total;
}
int main () {
    size_t numberOfInts = 3;
    size_t numberOfLoops = 1000000000;
    vector<int> a(numberOfInts);
    int total = 0;

    std::cout << "Started" << std::endl;
    for (size_t j = 0; j < numberOfLoops; j++)
    {
        for (size_t i = 0; i < numberOfInts; i++)
        {
            total = Sum(a);
        }
    }
    std::cout << "Finished:" << total << std::endl;
}

• Time: numberOfInts=3 took 40 seconds

Loop Unrolled

• This:

#include <iostream>
#include <vector>
using namespace std;

template <typename T, int length>
class FixedVector {
    T data[length];
public:
    FixedVector()
    {
        // Initialise
        for (size_t i = 0; i < length; i++)
        {
            data[i] = 0;
        }
    }
    T Sum()
```cpp
T sum = 0;
for (size_t i = 0; i < length; i++)
{
    sum += data[i];
}
return sum;
}

int main () {
    const size_t numberOfInts = 3;
    const size_t numberOfLoops = 1000000000;
    FixedVector<int, numberOfInts> a;
    int total = 0;

    std::cout << "Started" << std::endl;
    for (size_t j = 0; j < numberOfLoops; j++)
    {
        for (size_t i = 0; i < numberOfInts; i++)
        {
            total = a.Sum();
        }
    }
    std::cout << "Finished:" << total << std::endl;
}
```

- Time: numberOfInts=3 took 32 seconds when switch to fixed vector, and 23 when a raw array.

**Policy Checking**

- Templates parameterised by type not by behaviour
- But you can make a class to represent the behaviour
- See Keith Schwarz for longer example.

**Simple Policy Checking Example**

- This:

```cpp
#include <iostream>
#include <vector>
#include <stdexcept>
```
class NoRangeCheckingPolicy {
    public:
        static void CheckRange(size_t pos, size_t n) { return; } // no checking
    }

class ThrowErrorRangeCheckingPolicy {
    public:
        static void CheckRange(size_t pos, size_t n)
        {
            if (pos >= n) { throw std::runtime_error("Out of range!"); }
        }
    }

    template <typename T,
        typename RangeCheckingPolicy = NoRangeCheckingPolicy
    >
    class Vector : public RangeCheckingPolicy
    {
        private:
            std::vector<T> data;
        public:
            // other methods etc.
            const T& operator[](size_t pos) const
            {
                RangeCheckingPolicy::CheckRange(pos, data.size());
                return data[pos];
            }
    }

    int main () {
        Vector<int, ThrowErrorRangeCheckingPolicy> a;
        // a.push_back(1); or similar
        // a.push_back(2); or similar
        try {
            std::cout << a[3] << std::endl;
        } catch (const std::runtime_error& e) {
            std::cerr << e.what();
        }
        return 0;
    }

    • Produces:

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Summary of Policy Checking Example

- Define interface for behaviour
- Parameterize over all behaviours
- Use multiple-inheritance to import policies
- e.g. logging / asserts

Traits

- From C++ standard 17.1.18
  - “a class that encapsulates a set of types and functions necessary for
    template classes and template functions to manipulate objects of
    types for which they are instantiated.”
- Basically: Traits represent details about a type
- You may be using them already!
- Start with a simple example

Simple Traits Example

- This:

```cpp
#include <iostream>

template <typename T>
struct is_void {
    static const bool value = false;
};

template <>
struct is_void<void> {
    static const bool value = true;
};

int main () {
    std::cout << "is_void(void)=" << is_void<void>::value << std::endl;
    std::cout << "is_void(int)=" << is_void<int>::value << std::endl;
    return 0;
}
```

- Produces:
is_void(void)=1
is_void(int)=0

Traits Principles

- Small, simple, normally public, eg. struct
- else/if
  - Else template
  - partial specialisations
  - full specialisations
- Probably using them already
  - std::numeric_limits<double>::max()
  - ITK has similar itk::NumericTrait< PixelType>
- Applies to primitives as well as types

Traits Examples

- Simple Tutorial from Aaron Ballman
- Boost meta-programming support
- Boost type_traits tutorial
- C++11 has many traits

Wait, Inheritance Vs Traits?

- We said inheritance is often overused in OO
- We say that too frequent if/switch statements based on type are bad in OO
- C++11 providing many is_X type traits returning bool, leading to if/else
- So, when to use it?

When to use Traits

- Some advice
  - Sparingly
  - To add information to templated types
  - Get algorithm to work for 1 data type
  - If you extend to multiple data types and consider templates
    - When you need type specific behaviour
· traits probably better than template specialisation
· traits better than inheritance based template hierarchies

- Remember
  - Scientist = few use-cases
  - Library designer = coding for the unknown, and potentially limitless use-cases
    * More likely of interest to library designers

**TMP Use in Medical Imaging - 1**

Declare an ITK image

template< typename TPixel, unsigned int VImageDimension = 2 >
class Image: public ImageBase< VImageDimension >
{
    public:
    // etc

    - TPixel, int, float etc.
    - VImageDimension = number of dimensions

**TMP Use in Medical Imaging - 2**

But what type is origin/spacing/dimensions?

template< unsigned int VImageDimension = 2 >
class ImageBase: public DataObject
{
    typedef SpacePrecisionType SpacingValueType;
    typedef Vector< SpacingValueType, VImageDimension > SpacingType;

**TMP Use in Medical Imaging - 3**

So now look at Vector

template< typename T, unsigned int NVectorDimension = 3 >
class Vector: public FixedArray< T, NVectorDimension >
{
    public:
Now we can see how fixed length arrays are used

template< typename T, unsigned int TVectorDimension >
const typename Vector< T, TVectorDimension >::Self &
Vector< T, TVectorDimension >::operator+=(const Self & vec)
{
    for ( unsigned int i = 0; i < TVectorDimension; i++ )
    {
        ( *this )[i] += vec[i];
    }
    return *this;
}

which may be unrolled by compiler.

TMP Use in Medical Imaging - 5

- ITK
  - uses `itk::NumericTraits<>` adding mathematical operators like multiplicative identity, additive identity
  - uses traits to describe features of meshes, like `numeric_limits`, but more generalised
- MITK (requires coffee and a quiet room)
  - uses `mitkPixelTypeList.h` for multiplexing acrosstemplated imageto non-templated image type
  - uses `mitkGetClassHierarchy.h` to extract a list of class names in the inheritance hierarchy
- TMP in B-spline based registration:

Further Reading For Traits

- Keith Schwarz
- Nathan Meyers
- Todd Veldhuizen, traits scientific computing
- Thaddaeus Frogley, ACCU, traits tutorial
- Aaron Ballman
- Andrei Alexandrescu
- Andrei Alexandrescu traits with state
• Boost meta-programming support
• Boost type_traits tutorial
• C++11 has many traits

Further Reading In General

• Andrei Alexandrescu’s Book
• Herb Sutter’s Guru of The Week, especially 71 and this article
• And of course, keep reading Meyers

Summary

• Learnt
  – Notation for template function/class/meta-programming
  – Uses and limitations of template function/class
  – Template Meta-Programming
    * Optimisation, loop unrolling
    * Policy classes
    * Traits
Chapter 15

Appendix - Cloud

Appendix - Cloud computing

No Longer In Course

- Notes provided here for historical reasons.

Big data

Some data requires special treatment for collection, storage, and analysis.

The ‘three vs’:

- volume
- velocity
- variety

Cloud computing

Cloud computing is an approach built around the concept of shared resources.

- dynamically add and remove computing resources as you need them.
- bring up large clusters of computing power, then shut it own just as quickly
X* as a service

Emergence of service oriented approaches:

- Platform as a service (PaaS)
- Software as a service (SaaS)
- Infrastructure as a service (SaaS)

Exercise 1: Working in the cloud

Spinning up an instance

This example briefly run through the process of:

- creating an account with a provider of cloud services
- spinning up a single cloud instance using a web interface

Create an account

A growing number of companies are offering cloud computing services, for example:

- Amazon Web Services: http://aws.amazon.com
- Cloudera: http://www.rackspace.co.uk/cloud
- Google Cloud Platform: https://cloud.google.com
- Microsoft Azure: http://azure.microsoft.com

Create a key pair

In this exercise we will be working with Amazon Web Services.

Amazon uses public key cryptography to authenticate users, so we’ll need to create a private/public key pair:

```bash
# Create a key pair
$ ssh-keygen -t rsa -f ~/.ssh/ec2 -b 4096
```

We then need to register our public key with Amazon Web Services.
Create a single instance using the web interface

Navigate to the EC2 Dashboard and create a ‘micro’ instance (1 CPU, 2.5GHZ, 1GB RAM):

Connect to the instance with SSH

For Linux instances, the username is ‘ec2-user’:

```
# <key_file>: ~/.ssh/ec2
# <public_ID>: 52.16.106.209
$ ssh ec2-user@<public_ID> -i <key_file>
```

![SSH Connection](image)

```
# Connected
     _-|__|__|__|__|
     |            |
     |           /|
     |          / |
     |         /  |
     |        /   |

[ec2-user@ip-xxx ~]$ 266
```
Terminate the server

When finished with the instance, remember to terminate it!

Exercise 2: Working in the cloud

Again, from the command line . . .

Install Amazon Web Services Command Line Interface:
http://docs.aws.amazon.com/cli/latest/userguide/installing.html

# install the tools with pip
sudo pip install awscli
aws ec2 help

Configure the tools

To use the command line tools, you’ll need to configure your AWS Access Keys, region, and output format:

$ aws configure

# This configuration is stored locally in a home folder named .aws
# On Unix systems: ~/.aws/config;
# On Windows: %UserProfile%\.aws\config
AWS Access Key ID [****************VDLA]: EXXXXXAMPLE
AWS Secret Access Key [****************pa8o]: EXXXXXAMPLE
Default region name [eu-west-1]: eu-west-1
Default output format [json]: json

Test our connection to Amazon Web Services

If our connection has been set up correctly, ‘describe-regions’ will return a list of Amazon Web Service regions:

# Successful connection will return list of AWS regions
# HTTPSConnectionPool error? Try changing region to eu-west-1

$ aws ec2 describe-regions

Create a key pair

To connect to the instance, we will need a key pair. If you haven’t already done so, create one now:

# Create an SSH key pair
$ ssh-keygen -t rsa -f ~/.ssh/ec2 -b 4096

Transfer the public key to AWS:

# <key_name> is a unique name for the pair (e.g. my-key)
# <key_blob> is the public key: "$(cat ~/.ssh/ec2.pub)"
$ aws ec2 import-key-pair --key-name <key_name> --public-key-material <key_blob>

Create a security group

We’ll also need to create a security group...

# creates security group named my-security-group
$ aws ec2 create-security-group --group-name "My security group" --description "SSH access from my local IP address"

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Configure the security group

...and allow inbound connections from our local IP address:

```bash
$ aws ec2 authorize-security-group-ingress \
  --group-name "My security group" \
  --cidr <local_IP_address>/32 \
  --port 22 \
  --protocol tcp
```

Note: the /32 at the end of the IP address is the bit number of the CIDR netmask. The /32 mask is equivalent to 255.255.255.255, so defines a single host. Lower values broaden the range of allowed addresses. An IP of 0.0.0.0/0 would allow all inbound connections.

Locate an appropriate Machine Image

An Amazon Machine Image contains the software configuration (operating system, software...) needed to launch an instance. AMIs are provided by:

- Amazon Web Services
- the user community
- AWS Marketplace

We will search for an Amazon Machine Image ID (AMI-ID) using the command line tools:

```bash
$ aws ec2 describe-images --owners amazon \n  --filters "Name=name,Values=amzn-ami-hvm-2016.09.1.20170119-x86_64-gp2"
```

Launch an instance

Launch an instance using the Amazon Machine Image ID:

```bash
$ aws ec2 run-instances --image-id <AMI-ID> \
  --key-name <key_name> \
  --instance-type t2.micro \
  --security-groups <group_name>
```

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View the instance

We can check the instance and find the public IP:

```bash
# View information about the EC2 instances
# e.g. state, root volume, IP address, public DNS name
$ aws ec2 describe-instances
```

Connect to the instance

Use the public IP to connect:

```bash
# for Linux instances, the username is ec2-user
# <public_IP>: 52.16.106.209
# <key_file>: ~/.ssh/ec2
$ ssh -i <key_file> ec2-user@<public_IP>
```

You should now be connected!!:

```bash
__| __|_ )
_| \( /  Amazon Linux AMI
___\\___\___|
[ec2-user@ip-172-31-5-39 -]$
```

Terminate the instance

Don’t forget to terminate the instance when you have finished:

```bash
# terminate the instance
# <InstanceId>: i-87086760
$ aws ec2 terminate-instances --instance-ids <InstanceId>
```

```
TERMINATINGINSTANCES  i-87086760
CURRENTSTATE  32 shutting-down
PREVIOUSSTATE  16 running
```
Virtualisation

Reproducible research

The ability to reproduce the analyses of research studies is increasingly recognised as important.

Several approaches have developed that help researchers to package up code so that their code and dependencies can be distributed and run by others.

Virtual machines

A popular method for creating a shareable environment is with the use of virtual machines.

The isolated system created by virtual machines can be beneficial, but criticisms include:

• size: virtual machines can be bulky
• performance: virtual machines may use significant system resources

Tools such as Vagrant have helped to simplify the process of creating and using virtual machines:
https://www.vagrantup.com

Virtual environments

Virtual environments offer an alternative to virtual machines. Rather than constructing an entirely new system, virtual environments in general seek to provide reproducible ‘containers’ which are layered on top of an existing environment.

A popular tool for creating virtual environments is Docker:
https://www.docker.com

Exercise 3: Virtualisation

Reproducible research

In this example, we spin up a single EC2 instance and reproduce the analysis from a study in a virtual environment.

The simple analysis uses countwords, a shell script, to count the occurrences of words in a book (in our case dorian.txt, The Picture of Dorian Gray).

We will be using Docker to manage our virtual environment: https://docs.docker.com/userguide/dockerimages/
Create a new security group

For this exercise we will be setting up a web connection to the EC2 instance, allowing us to connect to IPython Notebook in a browser. To enable this connection, we will create a new security group:

```
$ aws ec2 create-security-group --group-name "ipython_notebook" \
   --description "web access to ipython notebook"
```

We then need to configure this group to allow inbound connections:

```
# SSH
$ aws ec2 authorize-security-group-ingress \
   --group-name "ipython_notebook" \
   --cidr 0.0.0.0/0 \
   --port 22 \
   --protocol tcp
# port 8888
aws ec2 authorize-security-group-ingress \
   --group-name "ipython_notebook" \
   --cidr 0.0.0.0/0 \
   --port 8888 \
   --protocol tcp
```

Spin up a cloud instance

As before, we will now spin up a new cloud instance:

```
# <AMI-ID>: e.g. ami-9d23aeea
# <key_name>: e.g. "my_key"
# <group_name>: e.g. "ipython_notebook"
$ aws ec2 run-instances --image-id <AMI-ID> \
   --key-name <key_name> \
   --instance-type t2.micro \
   --security-groups <group_name>
```

Connect with SSH

When the instance is running, a public IP will be available:

```
$ aws ec2 describe-instances
"NetworkInterfaces": [
```
Use the public IP to connect over SSH:

```
# <key_file>: e.g. ~/.ssh/ec2
$ ssh ec2-user@<public_IP> -i <key_file>
```

### Install and run Docker on the remote system

Docker needs to be installed on the Cloud instance:

```
[ec2-user@ip-xxx ~]$ sudo yum install -y docker
[ec2-user@ip-xxx ~]$ sudo service docker start
```

### Docker environments are created by a set of commands

Docker environments are created by running a set of commands held within a Dockerfile. A snippet of the Dockerfile used to create our environment is shown below:

```bash
# Set the source image
FROM ipython/scipystack:master

# Specify commands to run inside the image
RUN apt-get update
RUN apt-get install -y wget

# Create directory for container
RUN mkdir /analysis

# Get the code
ADD https://raw.githubusercontent.com/tompollard/dorian/master/mapper.py ~/analysis/mapper.py

# Run notebook
RUN echo "ipython notebook --ip=0.0.0.0 --port=8888 --no-browser" > /usr/bin/notebook.sh; 
chmod 755 /usr/bin/notebook.sh
```

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Pull, build, and run the Docker image

Build and run the docker environment:

```
# NB: sudo is not needed when running inside boot2docker
[ec2-user@ip-xxx ~]$ sudo docker pull tompollard/dorian:master
[ec2-user@ip-xxx ~]$ sudo docker run -t --rm=true -p 8888:8888 -i \
tompollard/dorian:master
```

IPython Notebook is now running in the virtual environment and available to
the outside environment on port 8888:

```
[NotebookApp] Using existing profile dir: '/root/.ipython/profile_default'
[NotebookApp] Serving notebooks from local directory: /analysis
[NotebookApp] The IPython Notebook is running at: http://0.0.0.0:8888/
[NotebookApp] Use Control-C to stop this server and shut down all kernels (twice to skip confirm)
```

Connect to the IPython Notebook

We can now connect to the IPython Notebook in a browser at the public IP of
the Amazon Instance on port 8888 (i.e. http://{publicIP}:8888):

Now use the IPython Notebook to complete the exercise

Using the IPython Notebook, we can reproduce the analysis on both existing
and new datasets.

Distributed computing

Distributed computing

Dividing a problem into many tasks means analysis can be shared across multiple
computers, in a parallel fashion.
Figure 15.1: IPython Notebook

Figure 15.2: Common words in Dorian Gray
Cloud computing systems have made distributed computing increasingly accessible to individual users.

Introduced by Google in 2004, MapReduce is a popular model that supports distributed computing on large data sets across clusters of computers.

**MapReduce**

MapReduce systems are built around the concepts of:

- a mapper, in which the master node takes an input, separates it into sub-problems, and distributes those to worker notes
- a ‘shuffle’ step to distribute data from the mapper
- a reducer which collates the answers to the sub-problems and combines them to solve the initial question.

**Distributed file systems**

Distributed file systems

- developed to provide reliable data storage across distributed systems
• replicates data across multiple hosts to achieve reliability

Hadoop

Apache Hadoop:

• framework for distributed processing of large data sets
• scales from single servers to many machines
• includes Hadoop MapReduce and the Hadoop Distributed File System

Exercise 4: MapReduce

The ‘Hello World’ of MapReduce

In this example, we will demonstrate how the mapper and reducer can be applied on our local machines to count the number of times each word appears in a book.

Choose a book

Find a good book on Project Gutenberg and download it: http://www.gutenberg.org/browse/scores/top

# The Picture of Dorian Gray
$ wget http://www.gutenberg.org/cache/epub/174/pg174.txt -O dorian.txt

$ head dorian.txt

Title: The Picture of Dorian Gray

The artist is the creator of beautiful things. To reveal art and conceal the artist is art’s aim. The critic is he who can translate into another manner or a new material his impression of beautiful things.

Hadoop streaming

Hadoop streaming enables MapReduce jobs to be run using any executable as the mapper and reducer: http://hadoop.apache.org/docs/r1.2.1/streaming.html

The mapper and reducer are written to take line-by-line inputs from stdin and emit the output to stdout.
Mapper

Our mapper:

- takes lines from our book
- extracts words from the line with a regular expression
- outputs a tab-separated string, value pair for each word (e.g. theword 1)

```
#!/usr/bin/env python
import sys
import re

def mapper(stream):
    pattern = re.compile('^[a-zA-Z][a-zA-Z0-9]*$')
    for line in stream:
        for word in pattern.findall(line):
            print word.lower() + '	' + 1

mapper(sys.stdin)
```

Reducer

Our reducer:

- takes the word, value pairs generated by the mapper
- sums the count for each word
- outputs a word, count pair

```
#!/usr/bin/env python
import sys

def reducer(stream):
    mydict = {}
    line = stream.readline()
    while line:

        # Get the key/value pair
        line = line.strip()
        word, count = line.split('	')
        count = int(count)

        # Add to the dict
        if word in mydict:
```
mydict[word] += 1
else:
    mydict[word] = 1

# Get the next line
line = stream.readline()

# Print the aggregated key/value pairs
# for word in sorted(mydict.keys()): # order by word
for word in sorted(mydict, key=mydict.get, reverse=True): # or by count
    print word, '	', mydict[word]

reducer(sys.stdin)

Demonstrate locally

Download the mapper and reducer:

$ git clone https://github.com/tompollard/dorian

Create a pipeline to process the book:

# sort represents the Hadoop shuffle
$ cat dorian.txt | ./mapper.py | ./reducer.py | sort

the   3948
of    2298
and   2279
to    2181
a     1730
i     1694
he    1544
...

Exercise 5: Distributed computing

Again, with Hadoop

In the previous exercise, we demonstrated MapReduce on our local computer. In this exercise we will spin up multiple cloud instances, making use of Hadoop to carry out a distributed MapReduce operation.

We could set up multiple instances in the cloud (for example with EC2), then configure Hadoop to run across the instances. This is not trivial and takes time.
Hadoop in the cloud

Fortunately, several cloud providers offer configurable Hadoop services. One such service is Amazon Elastic MapReduce (EMR).

Elastic MapReduce provides a framework for:

- uploading data and code to the Simple Storage Service (S3)
- analysing with a multi-instance cluster on the Elastic Compute Cloud (EC2)

Create an S3 bucket

Create an S3 bucket to hold the input data and our map/reduce functions:

1. Open the Amazon Web Services Console: http://aws.amazon.com
2. Select “Create Bucket” and enter a globally unique name
3. Ensure the S3 Bucket shares the same region as other instances in your cluster

Or, through the command line interface:

$ aws s3 mb s3://ucl-jh-books-example

Copy data and code to S3

Sample map and reduce functions are available on GitHub:

# clone the code from a remote repository
$ git clone https://github.com/tompollard/dorian

Copy the data and code to S3:

# Copy input code and data to S3
# No support for unix-style wildcards
$ aws s3 cp dorian.txt s3://my-bucket-ucl123/input/
$ aws s3 cp mapper.py s3://my-bucket-ucl123/code/
$ aws s3 cp reducer.py s3://my-bucket-ucl123/code/
Launch the compute cluster

Create a compute cluster with one master instance and two core instances:

```bash
# Start an EMR cluster
# <ami-version>: version of the machine image to use
# <instance-type>: number and type of Amazon EC2 instances
# <key_name>: "mykeypair"
$ aws emr create-cluster --ami-version 3.11.0
   --instance-groups InstanceGroupType=MASTER,InstanceCount=1,InstanceType=m3.xlarge
   InstanceGroupType=CORE,InstanceCount=2,InstanceType=m3.xlarge
   --use-default-roles

"ClusterId": "j-3HGKJHENDDX8"
```

Get the cluster ID

Get the cluster-id:

```bash
# Get the cluster-id
$ aws emr list-clusters
...
"Id": "j-3HGKJHENDDX8",
"Name": "Development Cluster"
```

Get the public DNS name of the cluster

When the cluster is up and running, get the public DNS name:

```bash
# Get the DNS
$ aws emr describe-cluster --cluster-id j-3HGKJHENDDX8
   | grep MasterPublicDnsName
...
"MasterPublicDnsName": "ec2-52-16-235-144.eu-west-1.compute.amazonaws.com"
```

Connect to the cluster

SSH into the cluster using the username ‘hadoop’:

```bash
# SSH into the master node
# <key_file>: ~/.ssh/ec2
# <MasterPublicDnsName>: ec2-52-16-235-144.eu-west-1.compute.amazonaws.com
$ ssh hadoop@<MasterPublicDnsName> -i <key_file>
```
Run the analysis

To process multiple input files, use a wildcard

```
[hadoop@ip-xxx ~]$ hadoop \
   jar contrib/streaming/hadoop-*streaming*.jar \
   -files s3://my-bucket-ucl123/code/mapper.py,s3://my-bucket-ucl123/code/reducer.py \
   -input s3://my-bucket-ucl123/input/* \
   -output s3://my-bucket-ucl123/output/ \
   -mapper mapper.py \
   -reducer reducer.py
```

View the results

Results are saved to the output folder. Each reduce task writes its output to a separate file:

```
# List files in the output folder
$ aws s3 ls s3://my-bucket-ucl123/output/
```

Download the output:

```
# Copy the output files to our local folder
# No support for unix-style wildcards, so use --recursive
$ aws s3 cp s3://my-bucket-ucl123/output . --recursive
```

```
# View the file
$ head part-00001

the 3948
and 2279
in 1266
his 996
lord 248
...
Terminate the cluster

Once our analysis is complete, terminate the cluster:

```
# get cluster id: aws emr list-clusters
# <cluster_ID>: j-3HGKJHENDODX8
$ aws emr terminate-clusters --cluster-id <cluster_ID>
```

Delete the bucket

```
$ aws s3 rb s3://my-bucket-ucl123 --force
```
Chapter 16

Linux Install

Git

If git is not already available on your machine you can try to install it via your distribution package manager (e.g. \texttt{apt-get} or \texttt{yum}).

On ubuntu or other Debian systems:

\texttt{sudo apt-get install git}

On RedHat based systems:

\texttt{sudo yum install git}

CMake

Again, install the appropriate package with apt-get or yum (\texttt{cmake}). Minimum version 3.5.

Editor and shell

Many different text editors suitable for programming are available. If you don’t already have a favourite, you could look at \texttt{Kate}.

Regardless of which editor you have chosen you should configure git to use it. Executing something like this in a terminal should work:
git config --global core.editor NameofYourEditorHere

The default shell is usually bash but if not you can get to bash by opening a terminal and typing bash.
Chapter 17

Windows Install

Editor

Unless you already use a specific editor which you are comfortable with we recommend using Notepad++ on windows.

Using Notepad++ to edit text files including code should be straight forward but in addition you should configure git to use Notepad++ when writing commit messages (see below).

Git

Install Git for Windows.

During the installation, you can select “Use Notepad++ as Git’s default editor” if you installed Notepad++ above. Make sure you tick “Use Git from the Windows Command Prompt” so other Unix tools can find git. The defaults should be suitable for other options.

Then install the GitHub for Windows client.

CMake

Install cmake. Minimum version 3.5.

And choose to add it to the path for all users if so prompted. (You may need to log out and log back in again before this takes effect!)
Unix tools

Install MinGW by following the download link. It should install MinGW’s package manager. On the left, select Basic Setup, and on the right select mingw32-base, mingw-developer-toolkit, mingw-gcc-g++ and msys-base. On some systems these package might be selected from start. Finally, click the installation menu and Apply Changes.

Locating your install

Now, we need to find out where Git and Notepad++ have been installed, this will be either in C:\Program Files (x86) or in C:\Program Files. The former is the norm on more modern versions of windows. If you have the older version, replace Program\ Files\ \(x86\) with Program\ Files in the instructions below.

Telling Shell where to find the tools

We need to tell the new shell installed by MinGW where Notepad++ is.
To do this, use NotePad++ to edit the file at C:\MinGW\msys\1.0\etc\profile and toward the end, above the line alias clear=clsb add the following:

```
# Path settings from SoftwareCarpentry
export PATH=$PATH:/c/Program Files\ \(x86\)/Notepad++
# End of Software carpentry settings
```

Finding your terminal

Check this works by opening MinGW shell, with the start menu (Start->All programs->MinGW->MinGW Shell). This should open a terminal window, where commands can be typed in directly.

On windows 8 and 10, there may be no app for MinGW. In that case, open the run app and type in

C:\MinGW\msys\1.0\msys.bat

You can also create a shortcut to this file on your Desktop for quicker access.
Checking which tools you have

Once you have a terminal open, type

`which notepad++`

which should produce readout similar to

`/c/Program Files (x86)/Notepad++/notepad++.exe`

Also try:

`which git`

which should produce

`/c/Program Files/Git/cmd/git.exe`

The `which` command is used to figure out where a given program is located on disk.

Tell Git about your editor

If you didn’t do this as part of the Git install, you now need to update the default editor used by Git.

`git config --global core.editor "'C:/Program Files (x86)/Notepad++/notepad++.exe' -multiInst -nosession -noPlugin"`

Note that it is not obvious how to copy and paste text in a Windows terminal including Git Bash. Copy and paste can be found by right clicking on the top bar of the window and selecting the commands from the drop down menu (in a sub menu).

You should now have a working version of git and notepad++, accessible from your shell.
Chapter 18

Mac Install

XCode and command line tools

Install XCode using the Mac app store.
Then, go to Xcode...Preferences...Downloads... and install the command line tools option

A note on Homebrew Versus MacPorts

If you already have either Homebrew or MacPorts installed, do NOT install both. This usually creates a confusing mess for students. If you have either installed already for other courses, then as long as the minimum compiler versions, cmake versions and git versions seem to work, there is no reason to swap. If you have neither installed, we prefer Homebrew, and the notes below are for Homebrew.

Installing Homebrew

Install Homebrew via typing this at a terminal:

```
ruby -e "$(curl -fsSL https://raw.githubusercontent.com/mxcl/homebrew/go)"
```

Git

Once Homebrew is installed:
brew install git

Then install the GitHub for Mac client. (If you have problems with older versions of OSX, it’s safe to skip this.)

CMake

Just do

brew install cmake

Minimum version 3.5.

Editor and shell

The default text editor on OS X textedit should be sufficient for our use. Alternatively choose from a list of other good editors.

To setup git to use textedit executing the following in a terminal should do.

```
git config --global core.editor
   /Applications/TextEdit.app/Contents/MacOS/TextEdit
```

The default terminal on OSX should also be sufficient. If you want a more advanced terminal iTerm2 is an alternative.
Chapter 19

Installing Boost

Linux Users

Install the appropriate package with apt-get or yum, for example:

```
sudo apt-get install boost
```

On Ubuntu 13.04 boost is spilt into multiple packages. The package needed here is “libboost1.53-dev” and to install it you should do:

```
sudo apt-get install libboost1.53-dev
```

Check that your package manager is delivering at least version 1.53, if you have an earlier version, you will need to download from source, following the windows instructions below.

Mac Users

```
brew install boost
```

Windows Users

You will need to download and install boost manually.

Create an appropriate folder, somewhere near your working code for the course.

For example, if you are working in `/home/myusername/devel/cppcourse/rsd-cppcourse/reactor` you could do:
mkdir -p ~/devel/libraries/boost

cd ~/devel/libraries/boost

Now, download and unzip boost, which will take quite a while:

curl -L http://sourceforge.net/projects/boost/files/... 
   ...boost/1.54.0/boost_1_54_0.zip/download > boost.zip
unzip boost.zip

(All one line, not literally two lines with dots!)

CMake and Boost

If we download from source, before we next build, we will need to tell our shell
where the boost library can be found

cd build
export CMAKE_INCLUDE_PATH=/home/myusername/devel/libraries/boost_1_54_0

cmake ..
make
cetest

... or your equivalent.
Chapter 20

Installation

Installation Instructions

Introduction

This document contains instructions for installation of the packages we’ll be using during the course. You will be following the training on your own machines, so please complete these instructions. You don’t need everything before lecture one: you can refer to these instructions as the course proceeds.

What you’ll need by the end of the course

- CMake build tool.
- Catch unit testing framework.
- Git and the GitHub website.
- Some C++ libraries: The Boost C++ library package, and Eigen linear algebra library.
- A code editor of your choice.

Eduroam

We will be using UCL’s eduroam service to connect to the internet for this work. So you should ensure you have eduroam installed and working.
Assignment 2

Serial Solution

Build a C++ implementation of Conway’s Game of Life. See for example https://en.wikipedia.org/wiki/Conway%27s_Game_of_Life [5 Marks]

Marks Scheme:

- Valid code: 1 mark.
- Readable code: 1 marks.
- Appropriate unit tests: 1 marks.
- Well-structured project layout and build system: 1 mark.
- Use of version control: 1 mark.

Organising remote computation

Define an appropriate output file format, and use a script in a dynamic language (Python, MATLAB, R, Ruby, Perl etc) to create an mpeg video of your results. Use a script to deploy your serial code on a remote cluster with qsub. [5 marks]

Marks scheme:

- Output file works: 1 mark.
- Visualiser works: 1 mark.
- Automated deployment script with fabric or bash: 1 mark.
- Output file and script organisation support reproducible research: 1 mark.
- Valid job submission script: 1 mark.

Shared memory parallelism

Parallelise your code using OpenMP. Create a graph of performance versus core count on a machine with at least 12 cores. Comment on your findings. [5 Marks]

Marks scheme:

- Valid OpenMP parallelisation: 1 mark.
- Preprocessor usage so that code remains valid without OpenMP: 1 mark
- Script to organise job runs and results for performance measurement: 1 mark.
- Clear and meaningful scaling graph: 1 mark.
- Discussion: 1 mark
Distributed memory parallelism

Parallelise your code using MPI, with a 2-dimensional spatial decomposition scheme. Create a performance graph with at least 12 cores. Comment on your findings. [5 marks]

Marks scheme:

- Valid MPI parallelisation: 1 mark.
- Valid 2-dimensional decomposition scheme: 1 mark.
- Unit tests to exercise decomposition scheme: 1 mark.
- Performance graph, with script to organise measurement runs and clear graph: 1 mark
- Discussion: 1 mark

Accelerators

Accelerate your serial solution using at least one of CUDA Thrust, CUDA C or OpenCL. Experiment with different thread counts and performance optimisations. Measure speedup compared to the serial code, either on a cluster or using a graphics card on your own computer. Comment on your findings. (There is one mark available for a second accelerator solution.) [5 marks]

Marks scheme:

- Valid accelerator parallelisation: 1 mark
- Clean, readable, tested code: 1 mark
- Optimisation by exploring different thread counts and other configurable parameters: 1 mark
- Speedup analysis and discussion: 1 mark
- Second accelerator parallelisation: 1 mark.

Submission of the assignment

You should submit your solution via Moodle, including the entire code repository with version control history. Your solution should include a text report with a discussion of your project and a clear explanation of how to build and invoke your code – the marker will deduct marks if the code cannot easily be built and run as indicated. Your report should be broken down into sections that reflect those of the assignment, and should include your graphs, performance measurements, and discussion of any problems or interesting design decisions you encounter along the way. You should separate your solutions for each section with version control revision numbers, tags or branches and describe these in the report.
Chapter 21

Assessment

Assessment Structure

- 3 hour exam
  - Past papers from 2015 and later are available, but syllabus does evolve
  - Past marks schemes and standard solutions are not available

- 2 pieces coursework - 40 hours each
  - 1 due 3rd week March
  - 1 due just after Easter