

Computing lecture 1

Getting started

I. Your computing environment

operating systems

networks

windowing systems

II. Data analysis tools

PAW

III. Programming

languages

compiling, linking

IV. More utilities

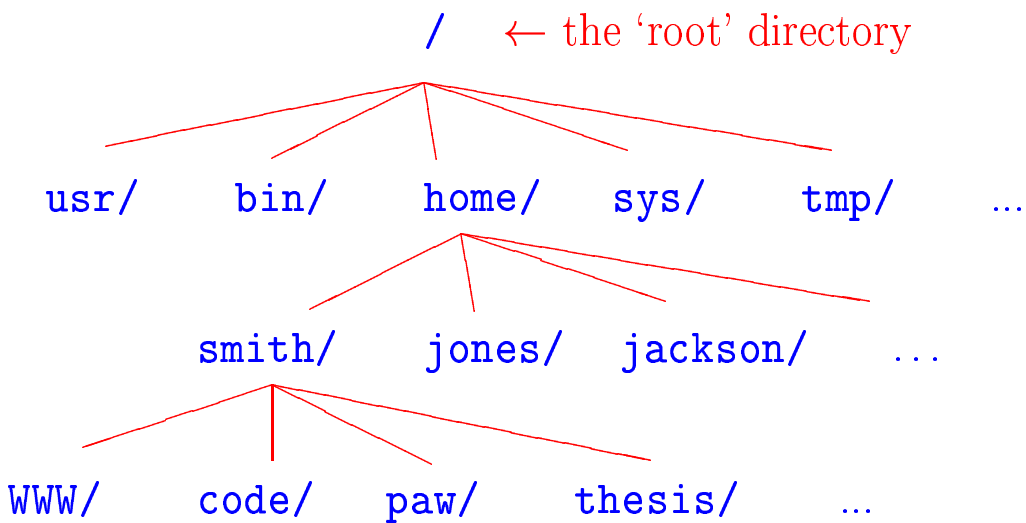
document preparation

manipulating plots

:

Operating systems

- Currently most widely used operating system in HEP is Unix:
many books, online tutorials (see course web site).
- Several shells (i.e. command sets) available: `sh`, `csh`, `tcsh`, `bash`, ...
- Shell/environment variables, shell scripts, redirection of i/o ...
- Tree-like structure for files and directories:



- A complete file name specifies the entire ‘path’:

`/home.smith.thesis.chapter1.tex`

- A tilde points to the home directory:

`~/thesis/chapter1.tex` ← the logged in user (e.g. smith)

`~jones/analysis/result.dat` ← a different user

- Single dot points to current directory, two dots for the one above:

`~smith/thesis` ← current directory

`../code` ← same as `~smith/code`

A few Unix commands (case sensitive!)

<code>pwd</code>	Show present working directory
<code>ls</code>	List files in present working directory
<code>ls -l</code>	List files of present working directory with details
<code>man ls</code>	Show manual page for <code>ls</code> . Works for all commands.
<code>cd</code>	Change present working directory to home directory
<code>mkdir foo</code>	Create subdirectory <i>foo</i>
<code>cd foo</code>	Change to subdirectory <i>foo</i> (go down in tree)
<code>cd ..</code>	Go up one directory in tree
<code>rmdir foo</code>	Remove subdirectory <i>foo</i> (must be empty)
<code>xemacs foo &</code>	Edit file <i>foo</i> with XEmacs (& to run in background)
<code>more foo</code>	Display file <i>foo</i> (space for next page)
<code>rm foo</code>	Delete file <i>foo</i>
<code>cp foo bar</code>	Copy file <i>foo</i> to file <i>bar</i> , e.g. <code>cp ~smith/foo ./</code> copies Smith's file <i>foo</i> to my current directory
<code>mv foo bar</code>	Rename file <i>foo</i> to <i>bar</i>
<code>lpr foo</code>	Print file <i>foo</i> . Use <code>-P</code> to specify print queue, e.g. <code>lpr -P<code>laser1</code> foo</code> (site dependent)
<code>ps</code>	Show existing processes
<code>kill 345</code>	Kill process number 345 (<code>kill -9</code> as last resort)
<code>./foo</code>	Run the executable program <i>foo</i> in current directory
<code>ctrl-c</code>	Terminate currently executing program

Better to read a book or online tutorial and use `man` pages

Networks

- Accounts usually at home lab and also SLAC, CERN, RAL, ...

<code>ftp shire.slac.stanford.edu</code>	file transfer
<code>telnet hpplus.cern.ch</code>	remote login
<code>ssh csfsun.rl.ac.uk</code>	'secure shell' login

- Many disks accessible world-wide via AFS (Andrew File System).

Typical (long) AFS file name:

`/afs/cern.ch/user/c/cowan/public/conference_plot.ps`



- Files (can be) visible world-wide; access determined by

Access Control Lists (ACL), and
whether you have a 'token' for the cell.

- You get a token (expires after 25 hours) by

`klog -cell cell name`

- Once you have a token, you can e.g. run editor locally and edit a file at a remote site (usually much faster).
- Useful commands:

<code>kpasswd</code>	Set AFS password
<code>tokens</code>	Show currently held tokens
<code>fs listquota</code>	Check AFS disk quota

N.B. Some sites use ARLA: same system, different command names.

X

- Windowing system, consisting of

X-server: process running on local machine (workstation or X-terminal); talks to monitor, keyboard, mouse, ...

X-clients: programs talking to X-server to request e.g. drawing windows, lines, etc. Can run on remote machines.

- Some important clients:

xterm terminal emulator

xman X-windows version of Unix manual

xemacs X-windows version of emacs text editor

paw CERN data analysis and display program

- Best with AFS or ssh. But if you need telnet e.g. to run on hpplus, specify hpplus as an allowed host to your local machine:

```
my_machine> xhost +hpplus.cern.ch
```

Log in to the remote machine:

```
my_machine> telnet hpplus.cern.ch
```

Set environment variable DISPLAY to tell X-client who to talk to; use internet address of your local machine plus :0

```
hpplus> setenv DISPLAY my_machine.ac.uk:0
```

Run X-client on hpplus; window appears on your local monitor:

```
hpplus> xterm &
```

Physics Analysis Workstation (PAW)

- PAW is an interactive program for plotting and manipulating
 - vectors
 - histograms
 - n -tuples
- To run, type **paw**. Enter workstation type, default probably OK.
- Many commands (hierarchical structure), e.g.
 - help** (type Q to return to command mode)
 - help plot** (help on command ‘plot’)
- Customize set-up, define new commands, etc. with file
.pawlogon.kumac
in home directory (Unix systems).
- Online tutorial and manual via PAW home page:
<http://wwwinfo.cern.ch/asd/paw/index.html>
- ‘Advanced’ features:
 - mathematical functions and operations (SIGMA)
 - FORTRAN interpreter (COMIS)
 - fitting with MINUIT

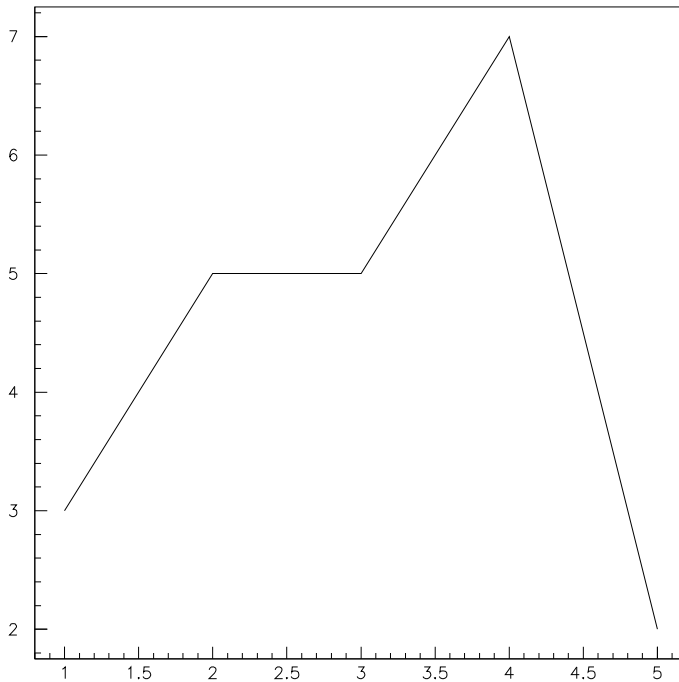
Using PAW

- Enter commands interactively or with command file (*.kumac)

Example – file **test.kumac** contains

```
vec/create x(5) R 1 2 3 4 5  
vec/create y(5) R 3 5 5 7 2  
graph 5 x y
```

Typing **exec test** produces



- To get an encapsulated PostScript file for printing, enter

```
pict/print filename.eps
```

(Needs **graphics/opt zfl1** e.g. in **.pawlogon.kumac.**)

Programming

- Many books on FORTRAN, C++ (see course web site).
- FORTRAN ('structured'), e.g. file `hello.f`,

```
program HELLO
    write (*, *) 'hello world'
END
```

- To compile, link and run,

```
f77 -o hello hello.f
```

 rename the executable program

./hello ← type this to run program

hello world ← output appears on screen

- List extra files separated by spaces; new line with backslash,

```
f77 -o greetings greetings.f bonjour.f \
hola.f gruezi.f yo_dude.f
```

- List CERN libraries at end in backwards single quotes, e.g.

‘cernlib’

‘cernlib graflib mathlib kernlib packlib’

- C++ ('object oriented'). `g++` runs the GNU C++ compiler `gcc`,

```
g++ -o hello hello.cc
```

A FORTRAN example

```
program MAKE_DATA

c Author: Glen Cowan
c Date: 21 August, 1999
c Test program to make a simple data set and write it to a file.

implicit NONE

c Constants

integer num_points
parameter (num_points = 11)

c Local variables

character*80 outfile

integer i

real x
real x_max
real x_min
real y

c Initialize some variables and open output file

x_min = 0.
x_max = 2.
outfile = 'test_data.dat'
open (unit = 20, file = outfile, form = 'formatted',
& status = 'unknown', carriagecontrol = 'list')

c Make the data and write to file

do i = 1, num_points
  x = (x_max-x_min)*FLOAT(i-1)/FLOAT(num_points-1) + x_min
  y = x**3 - 2.*x**2 + x
  write (20, *) x, y
end do

close (20)

stop
END
```

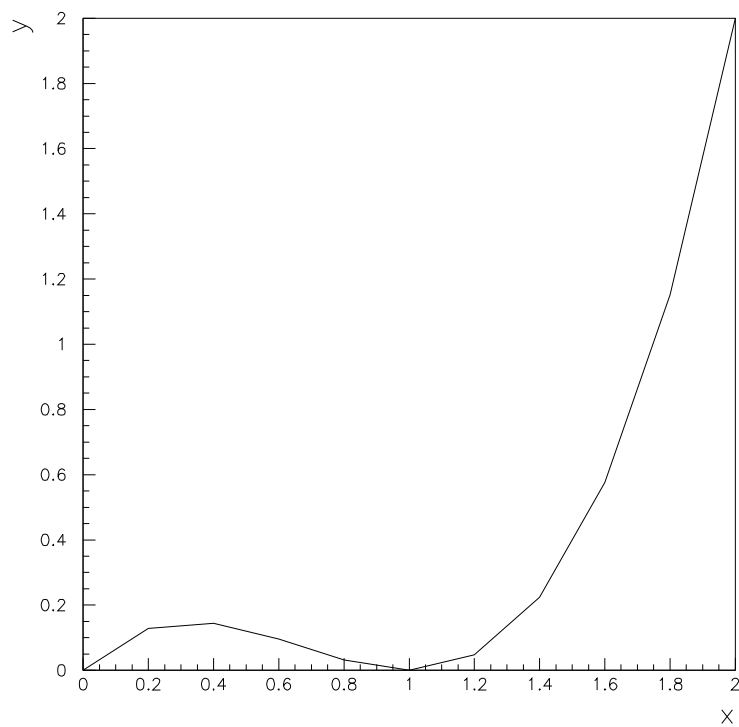
Looking at the output

- Running the program produces the file **test_data.dat**:

0.000000E + 00	0.000000E + 00
0.200000	0.1280000
0.400000	0.1440000
0.600000	9.6000016E - 02
0.800000	3.1999946E - 02
1.000000	0.0000000E + 00
1.200000	4.8000097E - 02
1.400000	0.2240001
1.600000	0.5759999
1.800000	1.152000
2.000000	2.000000

- With PAW, read in the file, create vectors and plot ...

```
vec/read x,y test_data.dat | create vectors x,y
null 0. 2. 0. 2.          | sets min/max values
graph $VLEN(x) x y 'L'   | $VLEN(x) = length of x
atit 'x' 'y'             | axis titles
```



Compiling and linking with gmake

- Often a program is in many files, e.g. `hello.cc` contains

```
#include <iostream.h>
#include "goodbye.h"      // contains prototype of function goodbye

void main(){
    cout << "Hello world" << endl;
    goodbye();
}
```

Function `goodbye` is in `goodbye.cc`, prototype in `goodbye.h`.

- We could compile and link with

```
g++ -o hello hello.cc goodbye.cc
```

which is really a short-cut for

```
g++ -c hello.cc      ← -c to compile (produces hello.o)
```

```
g++ -c goodbye.cc
```

```
g++ -o hello hello.o goodbye.o   ← link object files
```

- If e.g. `goodbye.cc` changed, we don't need to recompile `hello.cc`, but `g++ -o hello hello.cc goodbye.cc` does the whole lot; in large programs it's difficult to know what to recompile.
- With the Unix program `make` (GNU version `gmake`):
 - user supplies a *makefile* (called `GNUmakefile`, `makefile`, or `Makefile`), specifying how files depend on each other.
- Type `gmake` (plus optional argument); `gmake` looks at file dates and figures out what to do.

A simple makefile

- Makefiles have several types of statements, most importantly *rules*:

target : *dependencies* ...

command ← commands preceded by tab character

: ← new line (and tab) for each command

- A possible makefile for the previous example:

```
hello : hello.o goodbye.o
        g++ -o hello hello.o goodbye.o
hello.o : hello.cc goodbye.h
        g++ -c hello.cc
goodbye.o : goodbye.cc
        g++ -c goodbye.cc
```

- Type **gmake target** (if target name omitted, first one used), e.g.

gmake ← makes executable program **hello**

gmake goodbye.o ← compiles **goodbye.cc** only

- In practice, more complicated (see note on web):

```
# A simple makefile

objects = hello.o goodbye.o

hello : $(objects)
        g++ -o hello $(objects)
hello.o : hello.cc goodbye.h
goodbye.o : goodbye.cc

.PHONY : clean
clean :
        -rm hello $(objects)
```

Add comments, define variables, use implicit commands and dependencies, supply phony targets (e.g. **clean**), ...

Preparing documents with L^AT_EX

- Many online information sources and books; see e.g.
Kopka and Daly, *A Guide to L^AT_EX2e*, Addison-Wesley, 1995
- An almost minimal L^AT_EX source file (more samples on web):

```
%\documentstyle[12pt,epsfig]{article}      % use with version 2.09
\documentclass[a4paper,12pt]{article}        % use with version 2e
\usepackage{epsfig}                          % use with version 2e
\begin{document}

The text of your document goes here. \LaTeX is very good
at writing formulae as in equation (\ref{exp_dist}),

\begin{equation}
\label{exp_dist}
F(t) = \frac{1}{\tau} \int_0^t e^{-t'/\tau} dt' ,
\end{equation}

\end{document}
```

- To see output, type (default file extensions may be omitted)

<code>latex <i>file.tex</i></code>	← produces <i>file.dvi</i>
<code>dvips <i>file.dvi</i></code>	← produces <i>file.ps</i> , print with <code>lpr</code>
<code>gv <i>file.ps</i> &</code>	← GhostView (& to run in background)

- Also useful ...
- | | |
|---|---------------------------|
| <code>ps2pdf <i>file.ps</i></code> | ← creates <i>file.pdf</i> |
| <code>convert <i>file.gif</i> <i>file.ps</i></code> | (many formats possible) |

- See example files on web to see how to
 - include figures, create equations, lists, tables;
 - include sections, subsections, bibliography ...

More stuff

- Things we'll see later in the course:

Histogramming, n -tuples (HBOOK)

Structure of analysis programs in HEP

Monte Carlo (random numbers)

Function minimization (MINUIT)

- Things you'll have to pick up on your own:

A text editor: emacs, xemacs, vi, xedit, ...

E-mail: pine, outlook, Netscape

The web: HTML

Other operating systems: Windows NT, ...

and their utility programs: PowerPoint, Excel, Word, ...

Debugging programs: ddd, dbx, ...

Command languages: shell scripts, tcl, perl

Source code management: CVS, SRT, ...

Databases: Oracle, Objectivity, ...

Batch queues: LSF (CERN, SLAC, ...)

NQS (RAL, RH, UCL, ...)

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