Advanced UNIX

Scripting, data exploration, text parsing

What you should know

- Basic shell navigation
- Making file and directories
- Moving files
- Basic file handling

- cd, ls, pwd
- touch, mkdir
- mv, cp
- head, tail,cat, less
- Some experience in a script editor

Advanced Unix Materials

Log onto the server using isu username and password. Fetch materials from github:

- \$ ssh isuusername@bcbio.gdcb.iastate.edu
- \$ git clone https://github.com/j23414/adv-unix-workshop.git

Slides: https://github.com/j23414/adv-unix-workshop

Workshop Outline

- 1. Creating shell scripts
- 2. Exploring data with grep, sort, uniq and wc
- 3. sed search/replace and line deletion
- 4. awk advanced data processing language

Extra: All Unix Commands

```
$ echo $PATH
                     # prints path to installed Unix commands
                     # go to folder containing Unix commands
$ cd <path there>
                     # list available commands
$ 1s *
$ man <command>
                     # show manual page for command
$ which <command>
                     # tells which executable is used
                     # who is also logged in?
$ who
                     # what commands are currently running?
$ top
```

Part One

Creating shell scripts

What are shellscripts

 Anything you type into your terminal, can be pasted into a file and executed

 The code in the shellscript is read line-byline by the bash interpreter, just like the lines you type into your terminal

hello.sh

Another 'Hello World' example...

```
#! /bin/bash
echo 'Hello world'

touch file.txt # creates a file
ls -ltr # lists files
```

Hashbang (#!)

You need to tell the system what program should interpret your script

Syntax: #! <path to program>

Examples (first line of file):

#! /bin/bash
#! /usr/bin/python

Comments

Anything following a '#' is ignored by bash

list doc files in the current directory ls *.doc

Comments are notes to human readers

Your turn...

Create hello.sh

```
$ cd # go to home directory
$ touch hello.sh # create a script file
$ vi hello.sh # open editor
```

```
#! /bin/bash echo 'Your message here....'
```

Run the script

```
$ bash hello.sh # Method 1

$ ls -l hello.sh # Change to executable

$ ls -l hello.sh # Change to executable

$ ls -l hello.sh # Method 2
```

'./' is used to run an executable file

Part Two

grep, sort, uniq, wc

Piping and Redirection

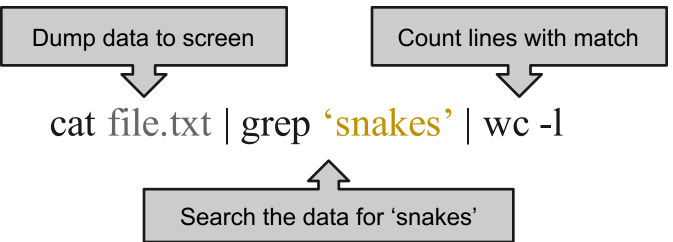
Review:

Redirection

Given A and B are programs and f is a file A | B | Pipe output of A to input of B | A > f | Overwrite f with A's output | A >> f | Append A's output to end of f

Pipelines

Many UNIX tools can be linked into piplines



Warnings

```
# a.txt will be empty after both commands
```

- **\$ head** a.txt > a.txt
- \$ head a.txt | A | B > a.txt

Never open a file for both reading and writing in one pipeline

Four powerful tools

- 1. wc count lines, words, or characters
- 2. grep search tool
- 3. sort flexible sort tool
- 4. uniq find unique lines

Sample data and exercises

Move to section-2/ which contains the files:

- 1. h[12345].txt (5 files)
- 2. script.sh
- 3. solutions.sh
- 4. unsorted.tab

WC

word count - count lines, words and characters

Options:

```
-l, --lines line count
```

-w, --words word count

-m, --chars character count

-L, --max-line-length

wc examples

```
# prints count of lines, words, and bytes
$ wc h*.txt
# Word count, like in MS Word
$ man bash | wc -w
# Count files in the working directory
$ ls | wc -l
```

grep

- prints lines matching the search pattern
- · for multiple files, tells which files matched
- has lots of very powerful options

Syntax:

```
$ grep [options] <pattern> <files>
$ <in> | grep [options] <pattern>
```

Examples 2.1

```
$ grep 'primrose' h*.txt
$ grep 'not to be' h*.txt
```

If your shell is not coloring the matches, run the following command:

```
$ alias grep='grep --color=auto'
```

Exercise 2.1

Practice we and basic grep

Navigate to section-2/

Make script.sh executable (chmod 755)

Open file script.sh

Follow the instructions for Exercise 2.1

some grep options

--help list of options and brief explanations

-c, --count

-v, --invert-match

-i, --ignore-case

-w,--word-regexp

-1, --files-with-match

-A, --after-context

-B, --before-context

-C, --context

-h, --no-filename

-L, --files-without-match

-o, --only-matching

-E, --extended-regexp

Examples 2.3

```
# Lines containing 'Fred'
$ grep 'Fred' m.tab
                             # Count 'Fred' matches
$ grep -c 'Fred' m.tab
$ grep -v 'Fred' m.tab
                             # Lines except 'Fred'
                             # Case insensitive match of 'g'
$ grep -i 'g' m.tab
$ grep -C1 'rose' h*.tab
                             # Shows the context, 1 line
$ grep -wC1 'rose' h*.tab
                             # Only match rose word
```

Two more options

-E, --extended-regexp

These commands require regular expressions to be really useful

Regular Expressions (1)

```
matches any character except a newline
       matches 0 or more of previous character
       matches 1 or more of previous character
      matches characters x, y and z
XYZ
[^xyz] matches characters OTHER than x, y and z
       anchors match at the BEGINNING of the line
       anchors match at the END of the line
       escapes the following special character
```

Example 2.4

```
$ grep -E '[aeiou]' m.tab
                            # Match vowels
$ grep -E '[^aeiou]' m.tab
                            # Match consonants
$ grep -E '^[1-5]' m.tab
                            # Begins with numbers 1-5
$ grep -E '[0-9]*G$' m.tab
                           # Ends in number and 'G'
$ grep -E '[a-z]+able' h*.txt # Match words that have 'able'
$ grep -oE '[a-z]+able' h*.txt # Only print out match
$ grep -E '^\[.*\]$' h*.txt # Prints stage directions
```

sort - reorder a file

--help list of options and brief explanations

```
-g, --general-numeric-sort
```

- -n, --numeric-sort
- -r, --reverse
- -u, --unique
- -k, --key=POS Sort by column

uniq

```
--help list of options and brief explanations
-c, --count count occurences of each line
-d, --repeated print only duplicated lines
-u, --unique print only uniq lines
```

Input must be sorted!
Only compares two adjacent lines!

Example 2.7

```
# The following two are identical
$ sort m.tab | uniq
$ sort -u m.tab
# Try these
$ sort m.tab | uniq -c
$ sort m.tab | uniq -d
$ sort m.tab | uniq -u
$ sort -k2 m.tab
```

Pipeline strategies

```
grep | sort | uniq
grep | sort | uniq | wc
<input> | sort | uniq -c | sort -n
```

Strategy: Build the pipelines up incrementally, checking output at each step

Exercise 2.2

Practice building pipelines

Navigate to section-2/

Follow the instructions for Exercise 2.2

Part Three

Substitution with sed

Sample Data

Move into section-3/, find the following:

- m.tab
- similar to unsorted tab in Part 2
- ids.txt a file of info on imaginary people
- s.fa

- a protein sequence file

The power of sed

\$ sed 's/This/That/g' yourfile.txt # text replacement

- search and replace (with style)
- extract specific patterns from files
- delete specific lines or ranges of lines

sed will not hurt your data

sed reads your data and writes to output.

The output will pour into your terminal unless redirected to a pipe or file.

Your original file is perfectly safe

sed won't, but YOU can

NEVER REDIRECT TO ORIGIN

--- Pipelines should not be circular ---

The following will destroy z.txt:

```
prog1 z.txt | prog2 > z.txt #BAD!!!
```

Test drive sed...

```
$ sed '' m.tab
                               # prints everything
$ sed -n '' m.tab
                               # prints nothing
$ sed -n '/Fred/p' m.tab
                               # same as "grep 'Fred' m.txt"
                               # duplicate 'Fred' lines
$ sed '/Fred/p' m.tab
$ sed 's/Fred/George/' m.tab
                               # 1st time text replacement
                               # global text replacement
$ sed 's/Fred/George/g' m.tab
```

sed syntax

sed [OPTIONS] <command>

sed [OPTIONS] '[LINE_ADDRESS] PROCEDURE'

sed workflow

for each line of input remove trailing newline character if line matches the address perform user's procedure if -n option is NOT set append newline and print

Addresses - by number

- 1 Matches line number 1
- 12 Matches line number 12
- 2,5 Matches lines 2 to 5
- 5,\$ Matches lines 5 and on

```
$ sed -n '1p' m.tab # prints 1st line
$ sed -n '5,$p' m.tab # prints lines 5 and on
```

Addresses - by expression

```
/ham/ Matches lines with pattern 'ham' /a/,/b/ Matches from lines matching a to b 1,/ham/ Matches lines 1 to matching 'ham' /ham/,$ Matches from 'ham' to the end
```

```
$ sed -n '/ham/p' m.tab # prints lines matching ham
$ sed -n '/start/,/stop/p' m.tab # print between two patterns
```

Procedure: deletion (d)

When the line matches the address, sed does not print, rather it moves onto the next line

Examples 3.1: deletion (d)

The address can be a number or a regular expression:

```
$ sed 'd' m.tab # delete everything
$ sed '1d' m.tab # delete 1st line
$ sed '/Fred/d' m.tab # delete lines containing 'Fred'
$ sed '5,10d' m.tab # delete lines 5 to 10
$ sed '10,$d' m.tab # delete lines from 10 on
$ sed '/R/,/T/d' m.tab # delete lines between R and T
$ sed '/Fred/,/Duffy/d' m.tab
```

! operator, invert selection

Addresses can be negated with !

```
1! Matches lines NOT equal to 1
```

2,5! Matches lines NOT between 2 and 5

```
$ sed -n '1!p' m.tab # prints all except 1st line
$ sed -n '2,5!p' m.tab # prints all except lines 5 and on
$ sed -n '/Bob/!p' m.tab# prints all except Bob lines
```

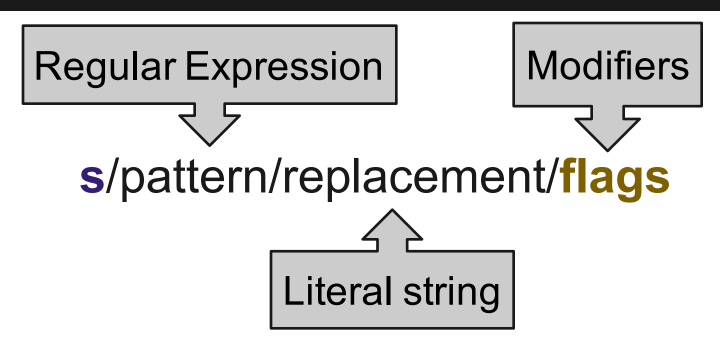
Regular Expressions (1)

```
matches any character except a newline
matches 0 or more of the previous char
[...] matches any of the enclosed
[^...] matches everything EXCEPT the enclosed
anchors match at the BEGINNING of the line
anchors match at the END of the line
escapes the following special character
```

Examples 3.2: regex

```
$ sed '/[TA]$/d' m.tab # Remove if ends in T or A
$ sed '/[^TA]$/d' m.tab # Remove if not ends in T or A
$ sed '/^Scene/d' h1.txt # Remove if starts with 'Scene'
$ sed '/^\[/d' h1.txt # Remove if starts with '['
```

substitution (s)



Examples

```
# replace each line's 1st 'this' with 'that' cat "this this" | sed 's/this/that/'
```

replace EVERY 'this' with 'that' (global flag) cat "this this" | sed 's/this/that/g'

Examples 3.3

```
$ sed 's/Fred/George/' m.tab
$ sed '/Feb/,/Sep/ s/Fred/George/' m.tab
$ sed 's/\[.*\]//' h*.txt
\$ sed 's/\n/' s.fa
                        # Instead use: tr ' '\n' s.fa
\$ sed 's/\|/\t/' s.fa
                        # Instead use: tr '|' '\t' s.fa
```

Exercise 3.1 (ids.txt)

- 1. cat ids.txt, check format (anything weird?)
- 2. delete ONLY those who are absent
- 3. delete ONLY those who are present
- 4. delete all entries after Mark
- 5. delete entries with an '*' after the name

Optional Challenge

Pull out gi, ref number, description and species from a protein sequence.

More advanced sed using a combination of pattern matching and substitution.

Extended Expressions (2)

```
    (...) captures the enclosed sequence
    n recalls nth captured sequence
    matches 1 or more of the previous characters
    OR
```

All of these require the -r argument (-E on mac)

Exercise 3.2

s.fa is formatted as so:

```
>gi|<gi>|ref|<ref>| <description> [<species>]
<sequence line 1>
...
<sequence line N>
```

- 1. Extract the 4 header regions individually
- 2. Write the gi and ref to a comma-delimited file

Print only if substituted

Problem:

You want a list of integers, but all the lines in the input still print

Solution:

Extraction strategy

To one of more words from a line:

>Start with the term to be extracted

sed -rn 's/.*([0-9]+).*
$$\wedge$$
1/p'

➤ Make pattern unambiguous by adding context

sed -r 's/.*id=(
$$[0-9]+$$
).*/\1/'

➤If no context is necessary, just use grep -o

Part Four

AWK: columnar data

Sample Data

Navigate to section-4

- diamonds.tab
- d.tab

- Borrowed from Hadley
- 25 lines of diamonds.tab

What is AWK?

- AWK is a full programing language
 - variables and arrays (like Perl hashes)
 - loops and conditional statements
 - o math, string processing, and user defined functions
- Sed-like addressing and regular expressions
- Automatically splits lines into words

Terms

- record
- field

- usually a line of input
- records are split into fields
- command a condition/procedure pair
- condition a logical test
 - procedure code block that is run if the condition is TRUE

AWK Pseudocode

```
BEGIN { do initial stuff }
for each record in input
     split record into fields
     for each command
           if condition is TRUE
                 do procedure
END { do final stuff }
```

AWK Fields

AWK breaks lines into fields

By default, fields are separated by whitespaces, e.g.

A field can be accessed by prefixing '\$' to the field number, e.g. \$2 is 'Fair', \$3 is 'G'

```
awk '$3 == "G" d.tab # print if 3rd field is G
```

Comparison Operators (1)

```
Regular expression match
       Regular expression non-match
       Equals (don't use '=')
       Not equals
       Less than
       Greater than
       Greater than or equal to
       Less than or equal to
/a/,/b/ TRUE between matches (like in sed)
```

Examples 4.1

Now we can test against a single column

```
$ awk '$2 == "Ideal"' d.tab
$ awk '$2 != "Ideal"' d.tab
$ awk '$3 ~ /[GFI]/' d.tab # reg exp
$ awk '$5 > 60' d.tab
```

print command

```
awk '{print $2, $1}'
```

Prints 2nd and 1st fields
Commas are special, they are field separators
Procedures can be used alone
'{' and '}' are **NOT** optional

Comparison to sed

Problem: Print 2nd and 1st fields of input

```
# solution in awk

$ awk '{print $2, $1}'

# solution in sed

$ sed -r 's/([^]+) ([^]+).*/\2 \1/'
```

Resetting Field Separator

You may reset the separator with option (-F)

```
# set field separator to comma $ awk -F', '$2,$1,"' d.csv
```

Logical Operators

```
|| Logical OR&& Logical AND! Logical NOT
```

These are used to string conditions together (<condition1> || <condition2>) &&! <condition3>

Conditional examples

```
$ awk '$1 > 1 && $7 < 5000' d.tab
$ awk '$2 == "Premium" || $3 == "E"' d.tab
$ awk '!/^#/ && ($1 > 1 || $2 == "Premium")' d.tab
```

Try a few other combinations
You can also use the full dataset, diamonds.tab

Condition only calls

AWK Rule 1: If the *command* consists only of a *condition*, the *procedure* defaults to print *record*.

Examples 4.2

```
awk '/Ideal/' d.tab # sed -n '/Ideal/p' d.tab
awk '/Fair/,/Ideal/' a.tab
awk '1,5' a.tab # fyi doesn't work
awk '/[GH]/' a.tab
```

Warning about quotes

awk "\$1 > 1" d.tab # WRONG

Here AWK gets the *shell variable* \$1 instead of a literal string '\$1'

This shell variable, will usually be undefined

Procedures

Syntax

condition { procedure }

When condition is TRUE, do procedure (implicit IF statements)

\$2 == "Fred" { print **\$3** }

Mathematical Operators

AWK will interpret variables as numbers if you perform mathematical operations on them.

```
+ - * / normal plus, minus, times, div

^ ** exponentiation

% modulo operator
```

Math examples

```
$ echo '1.1 _ 4' | awk 'print $1, $2, $1 + $2' 1.1 _ 4 _ 5.1
$ echo '2 _ 8' | awk 'print $1 ** $2'
$ echo '1 _ 2 _ 5' | awk 'print ($1 + $2) ** $3'
```

String concatenation

- Adjacent strings are concatenated
- Spaces are ignored
- Mathematical operations have precedence over string concatenation

```
$ echo "1 5" | awk '{print $1 "+" $2 "=" $1 + $2}
1+5=6
```

String concatenation

- Adjacent strings are concatenated
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```
$ echo "1 5" | awk '{print $1 "+" $2 "=" $1 + $2}
1+5=6
```

Exercise

Follow the instructions in script.sh

Workshop Outline

- 1. Creating shell script
- 2. Exploring data with grep, sort, uniq and wc
- 3. sed search/replace and line deletion
- 4. awk advanced data processing language

grep, sed, and awk

```
# Use grep to get lines that match a pattern
grep 'Fred' m.tab
# Use sed to search and replace
sed 's/Fred/George/g' m.tab
# Use awk with columnar data
awk '{print $2,$1,$4}' m.tab
```

Pipelines

```
# All of these commands can be piped together.
grep 'Fred' m.tab | sed 's/Fred/George/g' | awk '{print $2,$4}'
```

Advanced Unix Workshop

BCBGSO 2016 Workshop Series

Python Workshop --- 2-5pm @MBB