# Monitor resource usage of an process

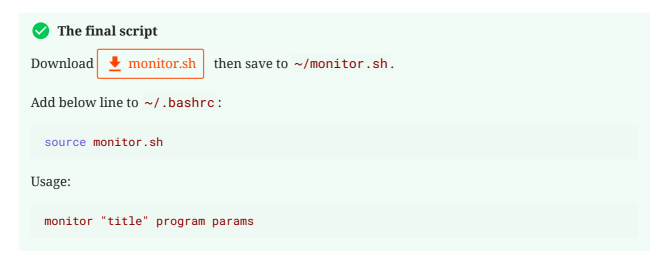
This post presents a custom script to execute a process and monitor the CPU and Memory usage of that process using linux utilities such as top, grep, and awk. The result is exported to a text file, and optionally an image if gnuplot is installed.

#bash #top #grep #awk #performance #gnuplot

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This post is written as a walk through guide, step by step, to help understanding how the script was made.

## 1. Export a function in bash

In a bash file, a function with a name and its body can be defined and then exported with export -f command:

```
myfunc.sh
```

```
#!/bin/bash
myfunc() {
    echo "Params: $@"
}
export -f myfunc
```

To make function available outside of the script, run source command before calling the function:

```
source myfunc.sh
myfunc abc
```

All params are implicit saved into local macros.

Refer to Advanced Bash-Scripting Guide:

Macro	Description
\$BASHPID	Process ID of the current instance of Bash. This is not the same as the \$\$ variable, but it often gives the same result.
\$PPID	Process ID of the parent process
\$\$	Process ID of the script itself

Macro	Description
\$!	Process ID of last job run in background
\$PWD	The current directory that process is in at the time
\$SECONDS	The number of seconds the script has been running
\$1,\$2, \$n	The first, the second and the n-th param
\$#	The number of command-line arguments
\$*	All of the positional parameters, seen as a single word, must be quoted
\$@	Same as \$*, but each parameter is a quoted string, that is, the parameters are passed on intact, without interpretation or expansion. This means, among other things, that each parameter in the argument list is seen as a separate word
\$?	Exit status of a command, function, or the script itself

## 2. Command-Grouping

Refer to Bash Manual - Command Grouping.

Bash provides two ways to group a list of commands to be executed as a unit. When commands are grouped, re-directions may be applied to the entire command list.

### (list)

Placing a list of commands between parentheses causes a sub-shell environment to be created (see Command Execution Environment), and each of the commands in list to be executed in that sub-shell. Since the list is executed in a sub-shell, variable assignments do not remain in effect after the sub-shell completes.

#### { list; }

Placing a list of commands between curly braces causes the list to be executed in the current shell context. No sub-shell is created. The semicolon (or newline) following list is required.

In addition to the creation of a sub-shell, there is a subtle difference between these two constructs due to historical reasons:

- The braces are reserved words, so they must be separated from the list by blanks or other shell meta-characters
- The parentheses are operators, and are recognized as separate tokens by the shell even if they are not separated from the list by whitespace

### 3. Run a process

The basic idea is to run a process in background and while it's running, report its resource usage:

monitor.sh

```
#!/bin/bash
monitor() (
    # run process in background
    echo "Executing $*"
    # get PID of last job in background
    pid=$!
    echo "Executed in PID: $pid"
    ps --no-headers -p $pid
    echo 'CPU MEM'
    # check if a process is running
    while [ -e /proc/$pid ]
        # use ps to get cpu and memory usage
        ps --no-headers -o '%cpu, %mem' -p $pid
        sleep 1
    done
export -f monitor
```

### 4. ps or top

Both ps and top report CPU Usage, but the returned values are different. Let's check the manual of each command:

ps

CPU usage is currently expressed as the percentage of time spent running during the entire lifetime of a process. This is not ideal, and it does not conform to the standards that ps otherwise conforms to. CPU usage is unlikely to add up to exactly 100%. It means ps does not show the instant CPU usage, it shows an average CPU usage over the lifetime of the process.

top

%CPU – CPU Usage, The task's share of the elapsed CPU time since the last screen update, expressed as a percentage of total CPU time. It means if interval is 1 second, top will report CPU usage for the last 1 second. That can be considered as instant report.

Let's check top 's options

### -b: Batch-mode operation

Starts top in Batch mode, which could be useful for sending output from top to other programs or to a file. In this mode, top will not accept input and runs until the iterations limit set with the `-n' command-line option or until killed.

-d : Delay-time interval as: -d ss.t (secs.tenths)

Specifies the delay between screen updates, and overrides the corresponding value in one's personal configuration file or the startup default. Later this can be changed with the d or s interactive commands.

-p: Monitor-PIDs mode as: -pN1 -pN2 ... or -pN1,N2,N3 ...

Monitor only processes with specified process IDs.

Switch to use top get the the process information:

```
#!/bin/bash
monitor() (
    # run process in background
    $* &
    # get PID of last job in background
    pid=$!
    # use top to monitor the process
    top -b -d 1 -p $pid &
    # save top PID to control it
    toppid=$!
    echo 'CPU MEM'
    # check if a process is running
    while [ -e /proc/$pid ]
        sleep 1
    done
    # kill top
    sleep 1
    kill -9 $toppid
export -f monitor
```

#### Run a test:

```
monitor ffmpeg -y -hide_banner -i /dev/video0 -c:v h264_omx -t 10 test.mp4
```

and the result:

```
PID USER PR NI VIRT RES SHR S %CPU %MEM TIME+ COMMAND 2286 pi 20 0 336612 129848 121960 R 34.5 52.5 0:03.36 ffmpeg
```

### 5. grep and awk

Refer to grep manual

Use grep to extract process information lines using pid number as the keyword:

```
top -b -d 1 -p $pid | grep $pid &
```

Refer to awk manual

Use  $\frac{1}{2}$  awk to cut out 2 columns:  $\frac{1}{2}$  and  $\frac{$ 

```
top -b -d 1 -p $pid | grep $pid | awk '{print $9, $10}' &
```

### So, this is a modified version:

```
#!/bin/bash
monitor() (
    # run process in background
    $* &

    # get PID of last job in background
    pid=$!

    # use top to monitor the process
    # use grep to catch useful lines
    # use awk to extract data columns
    top -b -d 1 -p $pid | grep $pid | awk '{print $9, $10}' &

    # save top PID to control it
    toppid=$!

    echo 'CPU MEM'
    # check if a process is running
    while [ -e /proc/$pid ]
```

```
do
    sleep 1
    done

# kill top
    sleep 1
    kill -9 $toppid
)

export -f monitor
```

Run a test command:

```
monitor ffmpeg -y -hide_banner -i /dev/video0 -c:v h264_omx -t 10 test.mp4
```

and the result:

```
CPU MEM
```

### **Bug:** No output value

Surprisingly, there is no output for CPU and MEM usage reported in the output. Search on google, there is a glue of missing data when using grep in a pipeline.

### 6. Line buffered mode

When using pipeline of commands, there is *pipeline buffer* between them.

The output from grep is no longer line buffered but block buffered, usually the block is 4KB,leading to the problem that the next awk command cannot see new data immediately on its input.

Notes from manuals:

```
man grep
--line-buffered
```

Use line buffering on output. This can cause a performance penalty.

### man awk

```
-W interactive
```

sets un-buffered writes to stdout and line buffered reads from stdin. Records from stdin are lines regardless of the value of RS.

Combining them together and testing again:

#### monitor.sh

```
#!/bin/bash
monitor() (
    # run process in background
    # get PID of last job in background
    pid=$!
    # use top to monitor the process
    # use grep to catch useful lines, use line buffered mode
    # use awk to extract data column, read input in line buffered mode
    top -b -d 1 -p $pid | grep --line-buffered $pid | awk -W interactive '{print
$9, $10}' &
    # save top PID to control it
    toppid=$!
    echo 'CPU MEM'
    # check if a process is running
    while [ -e /proc/$pid ]
    do
        sleep 1
    done
    # kill top
    sleep 1
    kill -9 $toppid
export -f monitor
```

#### Run a test:

```
monitor ffmpeg -y -hide_banner -i /dev/video0 -c:v h264_omx -t 10 test.mp4
```

### and the result comes with expected values:

```
CPU MEM
20.0 0.8
21.0 3.5
67.3 5.1
89.1 6.0
77.2 9.4
```

### 7. Save log with tee

Use **tee** to read from the standard input and write to the standard output and a file. That is simple enough to clone the output to a log file:

```
$* | tee log.txt &
top -b -d 1 -p $pid | grep --line-buffered $pid | awk -W interactive '{print $9,
$10}' | tee usage.txt &
```

### Then modify the script:

monitor.sh

```
#!/bin/bash
monitor() (
    # run process in background
    $* | tee log.txt &
    # get PID of last job in background
    pid=$!
    # use top to monitor the process
    # use grep to catch useful lines, use line buffered mode
    # use awk to extract data columns, read input in line buffered mode
    top -b -d 1 -p $pid | grep --line-buffered $pid | awk -W interactive '{print
$9, $10}' | tee usage.txt &
    # save top PID to control it
    toppid=$!
    echo 'CPU MEM'
    # check if a process is running
    while [ -e /proc/$pid ]
        sleep 1
    done
    # kill top
    sleep 1
    kill -9 $toppid
export -f monitor
```

### Run a test:

```
monitor ffmpeg -y -hide_banner -i /dev/video0 -c:v h264_omx -t 10 test.mp4
```

### 🏮 Bug: Empty log and invalid output

After using tee, there are twp issues happened:

- an empty *log.txt* file
- the usage.txt content is invalid

### 7.1. Fix empty log

When making pipeline to tee, only the STDOUT (1) is forwarded, while ffmpeg prints output on the STDERR (2) not on the STDOUT (1).

Fix it by redirect ffmpeg STDERR to STDOUT:

```
$* 2>&1 | tee log.txt &
```

### 7.2. Fix wrong data

Add some debug lines ps -p \$pid to check the process ID after creating the processes:

```
#!/bin/bash
monitor() (
    # run process in background
    $* | tee log.txt &
    # get PID of last job in background
    pid=$!
    ps -p $pid
    # use top to monitor the process
    # use grep to catch useful lines, use line buffered mode
    # use awk to extract data columns, read input in line buffered mode
    top -b -d 1 -p $pid | grep --line-buffered $pid | awk -W interactive '{print
$9, $10}' | tee > usage.txt &
    # save top PID to control it
    toppid=$!
    ps -p $toppid
    echo 'CPU MEM'
    # check if a process is running
    while [ -e /proc/$pid ]
        sleep 1
    done
    # kill top
    sleep 1
    kill -9 $toppid
```

```
export -f monitor
```

Then, it prints out the PID of tee, not the PID of ffmpeg or top.

```
PID TTY TIME CMD
647 pts/0 00:00:00 tee
PID TTY TIME CMD
652 pts/0 00:00:00 tee
```

### Get PID of a process in pipeline

In bash, pipeline cause commands to run in a sub-shell!

For example, \$\* | tee > log.txt & will run \$\* in a sub-shell, and tee > log.txt will run in current shell, therefore tee's PID will be saved in the macro \$!.

The solution is to save the PID in the newly created sub-shell in which \$! returns correct PID, then load that PID later:

```
# save to pid.txt
($* 2>&1 & echo $! > pid.txt) | tee > log.txt &
# load from pid.txt
pid=$(<pid.txt)</pre>
```

### Then modify the script:

```
#!/bin/bash
monitor() (
    # run process in background
    ($* 2>&1 & echo $! > pid.txt) | tee log.txt &
    # get PID of last job in background
    pid=$(<pid.txt)</pre>
    ps -p $pid
    # use top to monitor the process
    # use grep to catch useful lines, use line buffered mode
    # use awk to extract data columns, read input in line buffered mode
    (top -b -d 1 -p $pid & echo $! > pid.txt) | grep --line-buffered $pid | awk -
W interactive '{print $9, $10}' | tee usage.txt &
    # save top PID to control it
    toppid=$(<pid.txt)</pre>
    ps -p $toppid
    echo 'CPU MEM'
```

Run a test:

```
monitor ffmpeg -y -hide_banner -i /dev/video0 -c:v h264_omx -t 10 test.mp4
```

and got correct PIDs for ffmpeg and top:

```
PID TTY TIME CMD

2352 pts/0 00:00:00 ffmpeg

PID TTY TIME CMD

2360 pts/0 00:00:00 top
```

## 8. Graph with gnuplot

Gnuplot is a portable command-line graph utility for Linux, OS/2, MS Windows, OSX, VMS, and many other platforms. It can produce many different types of output, including terminal and file.

#### Terminal output

```
gnuplot -e " \
    set term dumb; \
    plot \
        'usage.txt' using 1 title '%CPU' with lines, \
        '' using 2 title 'MEM' with lines \
"
```

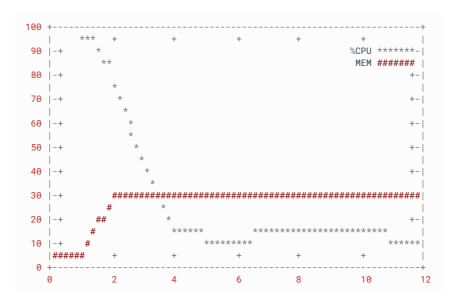
### **PNG Image output**

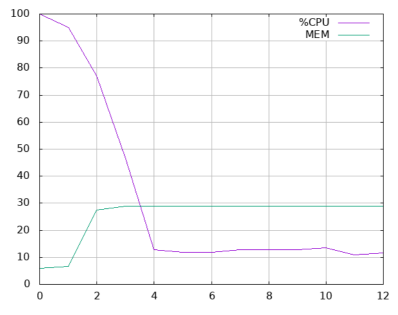
```
gnuplot -e " \
  set term png size 640, 480; \
```

Run a test:

```
monitor "test" "ffmpeg -y -hide_banner -i /dev/video0 -c:v h264_omx -t 10
test.mp4"
```

It prints out a good graph in the terminal as well as in a PNG image:





Resource Usage

### 9. Some enhancements

Some small modifications to make script run in multiple sessions:

#### 1. Set title for a new session

Let's modify the script to accept params in this format:

```
monitor "title" command
```

by extracting those params at the beginning of the script:

```
monitor() (
    # extract params
    title=$1
    command=${@:2} # get params from the 2nd one
    ...
)
```

The title will be used to name the session, to create a folder to save log files.

#### 2. Save output to a separated folder

```
# create result folder if not existed
[ ! -d $title ] && mkdir $title
```

### 3. Change output format to CPU= X MEM= Y

It needs to change the data column index in gnuplot:

```
# *-usage.txt content:
# CPU= X MEM= Y
# X is at 2nd column,
# Y is at 4th column
gnuplot -e " \
    set term dumb; \
    plot \
        '${title}/${title}-usage.txt' using 2 title '%CPU' with lines, \
        '' using 4 title 'MEM' with lines \
"
```

## 10. The final script

Download the final script in the top of this page.

Monitor resource usage of an process - Code Inside Out