Cloud Computing Architecture

Semester project

May 13, 2023

Overview

The semester project consists of four parts, two of which are described in detail in this handout. In this project, you will explore how to schedule latency-sensitive and batch applications in a cloud cluster. You will deploy applications inside containers and gain experience using a popular container orchestration platform, Kubernetes. Containers are a convenient and lightweight mechanism for packaging code and all its dependencies so that applications can run quickly and reliably from one computing environment to another.

You will work in groups of three students and submit a single report per group. Please submit your report in the format of the project report template. We will be assigning groups for the project, however you will have a chance to optionally let us know your preferences for teammates. If you know one or two other students in the class that you would like to work with on the project, please submit your group preference by March 9th, 2023. To do so, each student in your preferred group should sign up for the same group number in the Project Group Selection page on Moodle. We will notify you about final group assignments on March 13th and then you may redeem your cloud credits and begin working on the project.

Important Dates

March 9th, 2023: Deadline to submit group preferences. Remember that you must either subscribe to a group or join the general group (Group 1) to be assigned randomly by us.

March 13th, 2023: Groups are assigned and announced. Start working on project.

April 6th, 2023 at 13:00: Deadline to submit Part 1 and 2 of the project. May 25th, 2023 at 13:00: Deadline to submit Part 3 and 4 of the project.

We will release Part 3 and 4 of the project mid-April. Parts 3 and 4 are more open-ended and will require more time to complete than Part 1 and 2. Please plan your time accordingly.

Cloud Environment and Credits

To run experiments for the project, you will use Google Cloud. We will provide you with Google Cloud credits for your project. To redeem your cloud credits, please follow the steps in Part 1 (Section 1.1), when your project group assignment is confirmed. Each group member should create

a Google Cloud account at ${\tt https://accounts.google.com}$. Please use your ETH email address to create the account.

1 Part 1

In Part 1 of this project, you will run a latency-critical application, memcached, inside a container. Memcached is a distributed memory caching system that serves requests from clients over the network. A common performance metric for memcached is the tail latency (e.g., 95th percentile latency) under a desired query rate. You will measure tail latency as a function of queries per second and explore the impact of hardware resource interference. To add different types of hardware resource contention, you will use the iBench microbenchmark suite to apply different sources of interference (e.g., CPU, caches, memory bandwidth).

Follow the setup instructions below to deploy a Google Cloud cluster using the **kops** tool. Your cluster will consist of four virtual machines (VMs). One VM will serve as the Kubernetes cluster master, one VM will be used to run the memcached server application and iBench workloads, and two VMs will be used to run a client program that generates load for the memcached server.

This document contains setup instructions. Answer and submit the questions for Part 1 of the project in the report **template**.

1.1 Setup Instructions

Installing necessary tools

For the setup of the project, you will need to install **kubernetes**, **google-tools** and **kops**. Instructions based on the operating system on your local machine are provided in the links above. Having installed all the tools successfully, the following three commands should return output in your terminal (for the rest of the document the \$ symbol is there to declare a bash command and you shouldn't type it explicitly):

- 1. \$ kubectl --help
- 2. \$ kops --help
- 3. \$./google-cloud-sdk/bin/gcloud --help

Note that the final command is relative to where you have downloaded the google cloud tools. If you have installed via a package manager or have added the gcloud tools to your \$PATH you don't need the prefix and you can just type gcloud. Note that you have to open a new terminal or refresh your shell using source for your \$PATH to be updated.

All the scripts that you will need for both parts of the project are available here:

git clone https://github.com/eth-easl/cloud-comp-arch-project.git

Redeeming cloud credits and creating Google Cloud project

Each group member should create a Google Cloud account at https://accounts.google.com. Use your ETH email address to create the account. Each group will receive a \$50 Google Cloud coupon code. Select **one** group member to enter their name and ETH email address at the link you will receive when the groups have been assigned. Only redeem one coupon per group. If you need more credits you can get in touch with the TA team.

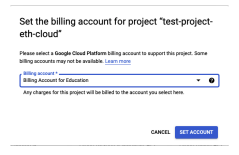
After installing kubernetes tools, connect your local client to your google cloud account using:

gcloud init

A browser window will open and you will have to login in with your ETH address. Afterwards, you will give <code>google-cloud-sdk</code> permissions to your account and then in the command line you will pick a name for the project. When creating the project name use <code>cca-eth-2023-group-XXX</code> (where XXX is your group number). Only one group member (who also redeemed the cloud credit coupon) should create the Google Cloud project. This person will add other group members as Project Owners (see instructions below). After the other group members are added as Project Owners, they will simply select the existing project name when they run the <code>gcloud init</code> command. All group members will have access to the project and share the cloud credits.

Do not configure any default computer region and zone. For deploying a cluster on Google Cloud we will modify some of the instructions listed here, which will be given below.

After creating the project you can log into the google cloud console and will be prompted to select a billing account for the project. In the pop up choose Billing account for education as below and click Set account:



Afterwards, you can try the command **gcloud compute zones list**. The first time you should get a prompt to enable the compute engine API that looks like this:

API [compute.googleapis.com] not enabled on project [project number]. Would you like to enable and retry (this will take a few minutes)? (y/N)?

After the API is enabled you can repeat the command which should now yield the following output:

\$ gcloud compute zones list

| + gerous compare remes rrs | - | |
|----------------------------|-------------|--------|
| NAME | REGION | STATUS |
| us-east1-b | us-east1 | UP |
| us-east1-c | us-east1 | UP |
| us-east1-d | us-east1 | UP |
| us-east4-c | us-east4 | UP |
| us-east4-b | us-east4 | UP |
| us-east4-a | us-east4 | UP |
| us-central1-c | us-central1 | UP |
| us-central1-a | us-central1 | UP |

| us-central1-f | us-central1 | UP |
|----------------|--------------|----|
| us-central1-b | us-central1 | UP |
| us-west1-b | us-west1 | UP |
| us-west1-c | us-west1 | UP |
| us-west1-a | us-west1 | UP |
| europe-west4-a | europe-west4 | UP |
| europe-west4-b | europe-west4 | UP |
| europe-west4-c | europe-west4 | UP |
| europe-west1-b | europe-west1 | UP |
| europe-west1-d | europe-west1 | UP |
| europe-west1-c | europe-west1 | UP |
| europe-west3-c | europe-west3 | UP |
| europe-west3-a | europe-west3 | UP |
| europe-west3-b | europe-west3 | UP |
| europe-west2-c | europe-west2 | UP |
| europe-west2-b | europe-west2 | UP |
| europe-west2-a | europe-west2 | UP |
| | | |

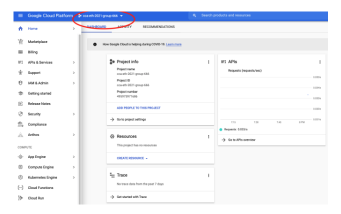
Then you will need to configure your default credentials using:

\$ gcloud auth application-default login

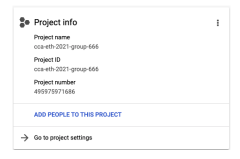
This will redirect you to a browser window where you will login with the same account you used when you setup the <code>gloud init</code> command.

Giving your teammates owner permission to the project

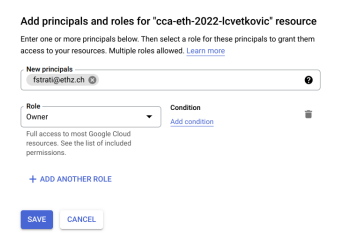
After creating the cca-eth-2023-group-XXX project on Google Cloud, give your group members access to the project and cloud credits by navigating to the Google Cloud console menu. Make sure your project is properly displayed on the top left as below:



In the project info click Add people to this project.



Type the email addresses of your teammates, select **Owner** as a role and click **Save**. **Note that** your teammates should have created a google cloud account with their ETH address in advance to put them as project owners.



Deploying a cluster using kops

At this point you will deploy a cluster using **kops**. First of all you will need to create an empty bucket to store the configuration for your clusters. Do this by running:

- \$ gsutil mb gs://cca-eth-2023-group-XXX-ethzid/
- ... where XXX is your group number and ethzid is your ETH username. Then run the following command to have the KOPS_STATE_STORE command to your environment for the subsequent steps:
- \$ export KOPS_STATE_STORE=gs://cca-eth-2023-group-XXX-ethzid/

If you open another terminal this and other environmental variables will not be preserved. You can preserve it by adding it with an **export** command to your .bashrc. You should substitute the number of your group and your ETH username as before.

Small Hint: Since Windows users cannot use the "export" command, you can manually add this environment variable. This tip also applies to the following "export" commands and "PROJECT='gcloud config get-value project'" commands.

For the first part of the exercise you will need a 3 node cluster. Two VMs will have 2 cores. One of these VMs will be the node where memcached and iBench will be deployed and another will be used for for the mcperf memcached client which will measure the round-trip latency of memcached requests. The third VM will have 8 cores and hosts the mcperf client which generates the request load for the experiments.

Before you deploy the cluster with **kops** you will need an ssh key to login to your nodes once they are created. Execute the following commands to go to your **.ssh** folder and create a key:

```
$ cd ~/.ssh
$ ssh-keygen -t rsa -b 4096 -f cloud-computing
```

Once you have created the key, go to lines 16 and 43 of the part1.yaml file (provided in the github link above) and substitute the placeholder values with your group number and ethzid. Then run the following commands to create a kubernetes cluster with 1 master and 2 nodes.

- \$ PROJECT=`gcloud config get-value project`
- \$ kops create -f part1.yaml

We will now add the key as a login key for our nodes. Type the following command:

\$ kops create secret --name part1.k8s.local sshpublickey admin -i ~/.ssh/cloud-computing.pub

We are ready now to deploy the cluster by typing:

\$ kops update cluster --name part1.k8s.local --yes --admin

Your cluster should need around 5-10 minutes to be deployed. You can validate this by typing:

\$ kops validate cluster --wait 10m

The command will terminate when your cluster is ready to use. If you get a **connection refused** or **cluster not yet healthy** messages, wait while the previous command automatically retries. When the command completes, you can type:

\$ kubectl get nodes -o wide

... to get the status and details of your nodes as follows:

| NAME | STATUS | ROLES | AGE | VERSION | INTERNAL-IP | EXTERNAL-IP |
|----------------------------|--------|--------|------|---------|-------------|----------------|
| master-europe-west3-a-2s21 | Ready | master | 3m2s | v1.19.7 | 10.156.0.63 | 34.107.107.152 |
| memcache-server-jrk4 | Ready | node | 102s | v1.19.7 | 10.156.0.61 | 34.107.94.26 |
| client-agent-vg5v | Ready | node | 98s | v1.19.7 | 10.156.0.62 | 34.89.236.52 |
| client-measure-ngwk | Ready | node | 102s | v1.19.7 | 10.156.0.60 | 35.246.185.27 |

You can connect to any of the nodes by using your generated ssh key and the node name. For example to connect to the client-agent node, you can type:

Running memcached and the mcperf load generator

To launch memcached using Kubernetes, run the following:

Then run the following:

\$ kubectl get pods -o wide

The output should look like:

```
NAME READY STATUS RESTARTS AGE IP NODE some-memcached 1/1 Running 0 42m 100.96.3.3 memcache-server-zns8
```

Use the IP address above (100.96.3.3 in this example) as the MEMCACHED_IP in the remaining instructions. Now ssh into both the client-agent and client-measure VMs and run the following commands to compile the mcperf memcached load generator:

```
$ sudo apt-get update
$ sudo apt-get install libevent-dev libzmq3-dev git make g++ --yes
$ sudo cp /etc/apt/sources.list /etc/apt/sources.list~
$ sudo sed -Ei 's/^# deb-src /deb-src /' /etc/apt/sources.list
$ sudo apt-get update
$ sudo apt-get build-dep memcached --yes
$ cd && git clone https://github.com/shaygalon/memcache-perf.git
$ cd memcache-perf
$ git checkout Oafbe9b
$ make
```

On the client-agent VM, you should now run the following command to launch the mcperf memcached client load agent with 16 threads:

```
$ ./mcperf -T 16 -A
```

On the client-measure VM, run the following command to first load the memcached database with key-value pairs and then query memcached with throughput increasing from 30000 queries per second (QPS) to 110000 QPS in increments of 5000:

... where MEMCACHED_IP is from the output of kubectl get pods -o wide above and INTERNAL_AGENT_IP is from the Internal IP of the client-agent node from the output of kubectl get nodes -o wide. You should look at the output of ./mcperf -h to understand the different flags in the above commands.

Introducing Resource Interference

Now we are going to introduce different types of resource interference with iBench microbenchmarks. Run the following commands:

\$ kubectl create -f interference/ibench-cpu.yaml

This will launch a CPU interference microbenchmark. You can check it is running correctly with:

\$ kubectl get pods -o wide

(wait until READY 1/1 and STATUS Running shows before starting a run).

When you have finished collecting memcached performance measurements with CPU interference, you should kill the job by running:

\$ kubectl delete pods ibench-cpu

You can apply the above three steps for any of the six ibench-cpu, ibench-lld, ibench-lli, ibench-lli, ibench-lli, and ibench-membw interference microbenchmarks. For Part 1 you will perform experiments to investigate the effect of the different types of interference. After now having followed this tutorial, you are able to run those experiments. First, start with reading the information of what to run for Part 1 in the project report template.

Deleting your cluster

<u>IMPORTANT</u>: you must delete your cluster when you are not using it! Otherwise, you will easily use up all of your cloud credits! When you are ready to work on the project, you can easily re-launch the cluster with the instructions above.

To delete your cluster, run on your local machine the command:

\$ kops delete cluster part1.k8s.local --yes

1.2 Notes

- Writing a script to automatize the data collection for Parts 1 and 2 is not mandatory and will not affect the grading. However, using automation scripts will be required for Parts 3 and 4, thus we encourage you to practice this approach in order to save time in the future.
- Parts 3 and 4 of the project are more resource-demanding and more costly in comparison to Parts 1 and 2 so make sure to plan your budget (usage of redeemed cloud credits) accordingly.

2 Part 2

In Part 2 of this project, you will run eight different throughput-oriented ("batch") workloads from the PARSEC (and SPLASH2x) benchmark suite: blackscholes, canneal, dedup, ferret, freqmine, radix and vips. You will first explore each workload's sensitivity to resource interference using iBench on a small 2 core VM (e2-standard-2). This is somewhat similar to what you did in Part 1 for memcache. Next, you will investigate how each workload benefits from parallelism by measuring the performance of each job with 1, 2, 4, 8 threads on a large 8 core VM (e2-standard-8). In the latter scenario, no interference is used.

Follow the setup instructions below to deploy a Google Cloud cluster and run the batch applications. Answer and submit the questions for Part 2 of the project in the report **template**.

2.1 Setup

In order to complete this Part of the project, we will have to study the behavior of PARSEC in two different contexts. For both, we will require that kubectl, kops and gcloud sdk are set up. This should already be the case if you have completed Part 1.

We have provided you with a set of yaml files which are useful towards spawning kubectl jobs for workloads and interference. The interference files are the same as in Part 1, but you must change the nodetype from memcached to parsec. The workloads are in the parsec-benchmarks folder in the github repo. All these files cover the workloads in the PARSEC suite, as well as the iBench interference sources relevant for this part: cpu, l1d, l1i, l2, l1c, memBW.

2.1.1 PARSEC Behavior with Interference

For the first half of Part 2, you will have to set up a single node cluster consisting of a VM with 2 CPUs. For this, we will employ **kops** and make use of the **part2a.yaml** file (make sure to update the file with values for your GCP project and configBase):

```
$ export KOPS_STATE_STORE=<your-gcp-state-store>
$ PROJECT=`gcloud config get-value project`
$ kops create -f part2a.yaml
$ kops update cluster part2a.k8s.local --yes --admin
$ kops validate cluster --wait 10m
$ kubectl get nodes -o wide
```

If successful, you should see something like this:

```
NAME
                               STATUS
                                        ROLES
                                                  AGE
                                                         VERSION
                                                                    INTERNAL-IP
                                                                                   EXTERNAL-IP
master-europe-west3-a-9nx1
                                                  3m2s
                                                         v1.19.7
                                                                    10.156.0.46
                                                                                   34.107.0.118
                              Ready
                                        master
                                                                                   35.234.110.58
parsec-server-s28x
                              Ready
                                        node
                                                  104s
                                                         v1.19.7
                                                                    10.156.0.47
```

Now you should be able to connect to the parsec-server VM using either ssh:

```
$ ssh -i ~/.ssh/cloud-computing ubuntu@35.234.110.58
```

Or by using gcloud:

To make sure that the jobs can be scheduled successfully, run the following command in order to assign the appropriate label to the parsec node (replace the cparsec-server-name> with the name of the parsec server observed in the output of the kubectl get nodes command):

\$ kubectl label nodes <parsec-server-name> cca-project-nodetype=parsec

For this part of the study we will sometimes require to set up some form of interference, and also deploy a job. For this example, we will use the PARSEC dedup job together with iBench CPU interference. Here is where we will use kubectl together with some of the yaml files we provide. The following code snippet spins up the interference, and runs the PARSEC dedup job:

```
$ kubectl create -f interference/ibench-cpu.yaml # Wait for interference to start
$ kubectl create -f parsec-benchmarks/part2a/parsec-dedup.yaml
```

Please note that, for Part 2a, you should use the job templates contained in the parsec-benchmarks/part2a folder. blackscholes, canneal, ferret and freqmine use the *simlarge* dataset, while dedup, radix, and vips use the *native* dataset. This is specified in the startup command for the container in the template file.

Make sure that the interference has properly started **before** running the **PARSEC** job. One way to see if the interference and the **PARSEC** job has started refers to **ssh**-ing into the VM and using the **htop** command to inspect running processes. You should see an image like below:

You can get information on submitted jobs using:

\$ kubectl get jobs

In order to get the output of the PARSEC job, you will have to collect the logs of its pods. To do so, you will have to run the following commands.

Note that the job name needs to match the one you get from kubectl get jobs.

Run experiments sequentially and wait for one benchmark to finish before you spin up the next one. Once you are done with running one experiment, make sure to terminate the started jobs. You can terminate them all together using:

```
$ kubectl delete jobs --all
$ kubectl delete pods --all
```

Alternatively, you can do so one-by-one using the following command:

```
$ kubectl delete job <job_name>
```

<u>IMPORTANT</u>: you must delete your cluster when you are not using it! Otherwise, you will easily use up all of your cloud credits! When you are ready to work on the project, you can easily re-launch the cluster with the instructions above. To delete your cluster, use the command:

\$ kops delete cluster part2a.k8s.local --yes

2.1.2 PARSEC Parallel Behavior

For the second half of Part 2, you will have to look into the parallel behavior of PARSEC, more specifically, how does the performance of various jobs in PARSEC change as more threads are added (more specifically 1, 2, 4 and 8 threads). For this part of the study, no interference is used.

You will first have to spawn a cluster as in section 2.1.1, however, this time use the part2b.yaml file we provided (make sure to update the file with values for your GCP project and configBase). Once more, this will be a single node cluster with an 8 CPU VM. You will have to vary the number of threads for each PARSEC job. To do so, change the value of the -n parameter in the relevant yaml files. The corresponding .yaml files are in parsec-benchmarks/part2b folder of the GitHub repo. Note that, for Part 2b, all of the jobs use the native dataset.

Other relevant instructions for this task can be found in section 2.1.1.

<u>IMPORTANT</u>: you must delete your cluster when you are not using it! Otherwise, you will easily use up all of your cloud credits! When you are ready to work on the project, you can easily re-launch the cluster with the instructions above. To delete your cluster, use the command:

\$ kops delete cluster part2b.k8s.local --yes

2.2 Notes

- Writing a script to automatize the data collection for Parts 1 and 2 is not mandatory and will not affect the grading. However, using automation scripts will be required for Parts 3 and 4, thus we encourage you to practice this approach in order to save time in the future.
- Parts 3 and 4 of the project are more resource-demanding and more costly in comparison to Parts 1 and 2 so make sure to plan your budget (usage of redeemed cloud credits) accordingly.

3 Part 3

In Part 3, you will put the previous two parts of the project together. You will now co-schedule the latency-critical memcached application from Part 1 and all seven batch applications from Part 2 in a heterogeneous cluster consisting of VMs with different numbers of cores. Your cluster will consist of a VM for the Kubernetes master (same as in Part 1), 3 VMs for the memcperf clients (2 agents and 1 measure machine), and 3 heterogeneous VMs (each with 2, 4, and 8 cores) which will be labeled node-a-2core, node-b-4core, node-c-8core, respectively, and used to run memcached and the PARSEC batch applications. Note that these VMs also have different configurations (as you can see in the part3.yaml file): node-a-2core is of type t2d-standard-2, node-b-4core is of type n2d-highcpu-4, and node-c-8core is of type e2-standard-8. The number of CPUs, the CPU platform, and the amount of memory differ in these VMs, which is something that you should take into account in your scheduling policy.

Your goal is to design a scheduling policy to minimize the time it takes for all seven batch workloads to complete while guaranteeing a tail latency service level objective (SLO) for the long-running memcached service. It might be helpful to take into account the characteristics of the PARSEC jobs that you noted in Part 2 (e.g. speedup across cores, total running time). For this part of the project, the memcached service will receive requests from the client at a steady rate and you will measure the request tail latency. Your scheduling policy should minimize the total time to complete all the PARSEC apps, without violating a strict service level objective for memcached of 1 ms 95th percentile latency at 30K QPS. You must ensure that all seven PARSEC apps succeed because a job may stop due to errors (e.g., out of memory). Use the native dataset size for all PARSEC jobs. At every point, you must use as many resources of your cluster as possible.

To design and implement your scheduling policy, you will experiment with different job collocations and resource management strategies using mechanisms in Kubernetes. Use what you learned about the performance characteristics of each application in Parts 1 and 2 to decide with which degree of parallelism to run each workload and which applications to collocate on shared resources.

The template with questions and space to record your results can be found here: template.

Hint: You may modify the YAML files provided, write a script for launching the batch jobs, or apply any other techniques you choose, as long as you describe them clearly in your report. You can choose which jobs to collocate, which degree of parallelism to use for the batch and memcached workloads, and when to launch particular batch jobs. Use any Kubernetes mechanism you wish to implement a scheduling policy. You may find node/pod affinity and/or resource requests/limits particularly useful. You may also want to use taskset in the container command arguments to pin containers to certain CPU cores on a node. Keep in mind that a job may fail due to a lack of resources. Use kubectl describe jobs to monitor jobs.

3.1 Setup

Run the following command to create a Kubernetes cluster with 1 master and 6 nodes. Make sure to update the part3.yaml file with the name of your project and your ConfigBase.

```
$ export KOPS_STATE_STORE=<your-gcp-state-store>
```

^{\$} PROJECT='gcloud config get-value project'

^{\$} kops create -f part3.yaml

We are ready now to deploy the cluster by typing:

\$ kops update cluster --name part3.k8s.local --yes --admin

Your cluster should need around 5-10 minutes to be deployed. You can validate the cluster with the command:

\$ kops validate cluster --wait 10m

The command will terminate when your cluster is ready to use. Afterward you can type

kubectl get nodes -o wide

to get the status and details of your nodes as follows:

| NAME | STATUS | ROLES | AGE | VERSION | INTERNAL-IP | EXTERNAL-IP |
|----------------------------|--------|--------|-----|---------|---------------|----------------|
| client-agent-a-d81z | Ready | node | 23m | v1.19.7 | 10.156.15.222 | 35.234.120.124 |
| client-agent-b-xpt7 | Ready | node | 23m | v1.19.7 | 10.156.15.224 | 34.107.4.82 |
| client-measure-x1xw | Ready | node | 23m | v1.19.7 | 10.156.15.223 | 35.242.212.158 |
| master-europe-west3-a-cdp2 | Ready | master | 24m | v1.19.7 | 10.156.15.225 | 34.89.196.131 |
| node-a-2core-qtrb | Ready | node | 23m | v1.19.7 | 10.156.15.221 | 34.89.217.203 |
| node-b-4core-gq6s | Ready | node | 23m | v1.19.7 | 10.156.15.220 | 34.107.20.21 |
| node-c-8core-3kz9 | Ready | node | 23m | v1.19.7 | 10.156.15.226 | 34.107.23.202 |

To connect to any of the machines you can run:

Modify the memcached and PARSEC batch job YAML files from Parts 1 and 2 of the project and use the **kubectl create** commands to launch the workloads in the cluster. You may want (and we encourage you, since it will be compulsory in the last part) to write scripts to launch the jobs. The memcached job should start first and continue running throughout the whole experiment while receiving a constant load of 30K QPS from the **mcperf** client. After you have started memcached and the client load, you can start the PARSEC batch jobs in whichever order you choose. Your goal is to minimize the time from the start of the first PARSEC job to the time the last PARSEC job completes while ensuring that the 95th percentile latency for memcached remains below 1ms.

For part 3 and 4, you must use a custom version of mcperf. It provides two features: it adds two columns that contain start and end time for each measurement, and it will allow variable traces (for the last part). To install the augmented version of mcperf on the client-agent-* and client-measure, follow the instructions below:

```
$ sudo sh -c "echo deb-src http://europe-west3.gce.archive.ubuntu.com/ubuntu/ bionic main \
    restricted >> /etc/apt/sources.list"
$ sudo apt-get update
$ sudo apt-get install libevent-dev libzmq3-dev git make g++ --yes
$ sudo apt-get build-dep memcached --yes
```

- \$ git clone https://github.com/eth-easl/memcache-perf-dynamic.git
- \$ cd memcache-perf-dynamic
- \$ make

Instead of sweeping request throughput as you did in Part 1, you now want to generate load at a constant rate of approximately 30K QPS while reporting latency periodically (e.g., every 10 seconds). To do this, run the following command on the client-agent-a machine:

\$./mcperf -T 2 -A

and the following command on the client-agent-b machine:

 $\ \ ./mcperf -T 4 -A$

and the following command on the client-measure VM:

You can get the execution time of the batch jobs by parsing the JSON output of the **kubectl** command that returns information about each job, including its start and completion time. To do that, after all jobs have been completed run:

```
$ kubectl get pods -o json > results.json
$ python3 get_time.py results.json
```

where **get_time.py** is a python script that you can find here.

<u>IMPORTANT</u>: you must delete your cluster when you are not using it! Otherwise, you will easily use up all of your cloud credits! When you are ready to work on the project, you can easily re-launch the cluster with the instructions above.

To delete your cluster, use the command:

```
$ kops delete cluster --name part3.k8s.local --yes
```

3.2 Questions

Answer and submit the questions for Part 3 in the report template.

3.3 Submission

For part 3, is expected that you submit:

- The answers to the questions
- Any modified or added yaml file for the jobs
- Any script, if you automated that part
- Any further script or file that you consider useful for the understanding of your scheduling policy
- Your measurement output files, as explained below

Your submission **must** contain the measurement results that you are describing in the report. Place in the root of the archive a directory called $part_3_results_group_XXX$, with 6 files inside, 3.json and 3.txt. XXX is the group number as 3 digits (e.g. group 1 should put 001). For i = 1..3, they should be named $pods_i.json$ and $mcperf_i.txt$. The JSON file should be the entire output of the get pods command as described above, while the txt is the output of mcperf execution. You can find an example of the expected mcperf output format here. Usually, copying from the

console is sufficient. Trailing new lines and white spaces are ignored, you can use either Unix-like line endings (\n) or Windows-like line endings (\n) .

Please follow that request, your group may lose points if the files are not present or with invalid contents.

There are no requirements regarding the structure of the other requested files.

4 Part 4

In Part 4, you will co-schedule PARSEC batch jobs on a single 4-core server running memcached. In contrast to Part 3, we will now dynamically vary the load on the long-running memcached service, such that the number of cores needed by the memcached server to meet the tail latency service level objective (SLO) ranges from 1 to 2 cores. You will design a scheduling policy that grows and shrinks the resource allocation of memcached and opportunistically uses temporarily available cores to complete the PARSEC jobs as quickly as possible. Your scheduling policy must guarantee a memcached tail latency SLO of 1ms 95th percentile latency. For this part you will be using a cluster consisting of 4 nodes: a 2 core VM cluster master, a 4 core high memory VM for the memcached server and PARSEC jobs, a 16 core VM for the mcperf agent, and a 2 core VM for the mcperf measurement machine.

You will need to implement your own controller to launch jobs and dynamically adjust their available resources, based on your scheduling policy. In this part of the project, we will not use Kubernetes because Kubernetes does not provide an API to change a container's resource allocation during run time. Instead, you will use Docker to launch containers to run the PARSEC benchmarks and dynamically adjust their resources. For memcached, we provide instructions for installing and running memcached directly on the VM (rather than in a Docker container) and using the taskset command to dynamically adjust resources. The reason we do not use Docker to run memcached in this part of the assignment is because we have observed that memcached's resources are not effectively constrained with docker --cpuset-cpus, since most of the processing in memcached application is network packet processing, which executes in kernel threads. Your controller should monitor CPU utilization and/or other types of resources and metrics to decide if resources need to be adjusted to meet the SLO. Your controller should make dynamic resource allocation decisions, such that the PARSEC batch jobs are completed as quickly as possible while still enforcing memcached's SLO.

For this part we also provide an augmented version of mcperf which is capable of generating random loads on the memcached server, as well as specific load traces. Refer to the instructions provided in part 3 to install this version.

Implementing the controller and scheduling policy

We recommend implementing your controller in python and using the Docker Python SDK to manage containers. Alternatively, you may implement the controller in Go using the Docker Go SDK. You can find examples of managing containers using the Docker SDK, for both Python and Go. If you plan on using such an SDK, you might find it useful to use the shell command sudo usermod -a -G docker <your-username>. This will allow you to use the SDK programmatically without encountering permission errors. You will also be able to run docker commands without using sudo.

In addition to running containers, you will need to update containers while they are running. Updating a container refers to dynamically adjusting properties of the container, such as the CPU allocation. You can read more about updating containers in the Docker update command documentation. You can update docker containers using Docker SDK commands. In case you find it helpful for your scheduling policy (though it is not required), you can also pause and unpause containers. Pausing a container has the effect of temporarily stopping the execution of the processes in the container (i.e., releasing CPU resources) while retaining the container's state (i.e., keeping the container's memory resources). Unpausing containers resumes the execution of processes in the

container.

Your controller should run on the 4-highmem core memcached server and monitor CPU utilization. The controller should use CPU utilization statistics to make dynamic scheduling decisions. You can monitor CPU utilization on the server by reading and post-processing data from /proc/stat files on the VM. There are also language specific options for monitoring metrics, such as psutil for Python.

In addition to CPU utilization, you may use other inputs for your scheduling policy, if you wish. This is not required, but may let you implement an even better policy. Please explain in your report any additional inputs you choose to consider in your scheduling policy at the controller.

Evaluating the scheduling policy

You will evaluate your scheduling policy with a dynamic mcperf load trace that we provide (see instructions below). You should use mcperf to investigate the performance of your scheduling policy with various load traces (e.g., try different random seeds and time intervals). Experimenting with various load traces will allow you to analyze when and why your policy performs well and understand in which scenarios the policy does not adapt appropriately.

Generating the plots

In this part of the project, you will be asked to generate some plots which often require that you aggregate data gathered from different VMs. This can be challenging since you'll need to temporally correlate this data across the VMs. A straight forward way to do this is to save the Unix time whenever you log an event, as this time is roughly synchronized across VMs. You can further use other information such as dynamic mcperf's --qps_interval or -t parameter (see documentation here). Our dynamic mcperf version should also by default print the simulation's start and end Unix times in the output logs. Another alternative is to use the shell command date +%s. These times can then be used when generating the plots to synchronize events that take place on different VMs.

4.1 Setup

4.1.1 Installation

Run the following command to create a kubernetes cluster with 1 master and 3 nodes.

```
$ export KOPS_STATE_STORE=<your-gcp-state-store>
$ PROJECT='gcloud config get-value project'
$ kops create -f part4.yaml
```

We are ready now to deploy the cluster by typing:

```
$ kops update cluster --name part4.k8s.local --yes --admin
```

Your cluster should need around 5-10 minutes to be deployed. You can validate the cluster with the command:

```
$ kops validate cluster --wait 10m
```

The command will terminate when your cluster is ready to use. Afterwards you can type:

kubectl get nodes -o wide

To get the status and details of your nodes as follows:

| NAME | STATUS | ROLES | AGE | VERSION | INTERNAL-IP | EXTERNAL-IP |
|------------------------|--------|--------|-------|---------|-------------|----------------|
| client-agent-bf7q | Ready | node | 111s | v1.19.7 | 10.138.0.33 | 35.230.78.193 |
| client-measure-5v6m | Ready | node | 116s | v1.19.7 | 10.138.0.32 | 35.227.161.236 |
| master-us-west1-a-kh69 | Ready | master | 3m23s | v1.19.7 | 10.138.0.34 | 35.247.63.197 |
| memcache-server-qmql | Ready | node | 111s | v1.19.7 | 10.138.0.31 | 34.83.56.78 |

You will first need to manually install memcached on the memcache-sever VM. To do so, you must first use the following commands:

```
$ sudo apt update
$ sudo apt install -y memcached libmemcached-tools
```

To make sure the installation succeeded, run the following command:

\$ sudo systemctl status memcached

you should see and ouput similar to the following being produced:

You will need to expose the service to the outside world, and increase its default starting memory. To do so, open memcached's configuration file using the command:

\$ sudo vim /etc/memcached.conf

To update memcached's memory limit, look for the line starting with -m and update the value to 1024. Similarly, to expose the memcached server to external requests, locate the line starting with -1 and replace the localhost address with the internal IP of the memcache-server VM. You can also specify the number of memcached threads here by introducing a line starting with -t followed by the number of threads. Save the file, then execute the next command to restart memcached with the new configuration:

\$ sudo systemctl restart memcached

Running sudo systemctl status memcached again should yield an output similar as before, however, you should see the updated parameters in the command line. If you completed these steps successfully, memcached should be running and listening for requests on the VMs internal IP on port 11211.

Install on client-agent and client-measure the augmented version of mcperf following the Part 3 instructions.

On the client-agent VM, you should now run the following command to launch the mcperf memcached client load agent with 16 threads:

\$./mcperf -T 16 -A

On the client-measure VM, run the following command to first load the memcached database with key-value pairs and then query memcached with a dynamic load generator which will produce a random throughput between 5000 and 100000 queries per second during each interval. The throughput target will change and will be assigned to another QPS for the next interval. Note that, differently from before, the output appears only at the end of the measurement. In the example the interval duration is set to 2 seconds, whilst the overall execution time is 10 seconds, which will result in five different QPS intervals:

The INTERNAL_MEMCACHED_IP and INTERNAL_AGENT_IP are the internal IPs of the memcache-sever and client-agent retrieved from the output of kubectl get nodes -o wide.

For more information on the dynamic load generator, and the available options it provides, check the guide in the README.md of the public repository.

PARSEC jobs can be started using Docker. For instance, one can start the **blackscholes** job on core 0 (--cpuset-cpus="0" parameter) and with 2 threads (-n 2 parameter) using the following command:

```
docker run --cpuset-cpus="0" -d --rm --name parsec \
     anakli/cca:parsec_blackscholes \
     ./run -a run -S parsec -p blackscholes -i native -n 2
```

You can also inspect the yaml files for the PARSEC jobs from the previous parts to further understand their command lines. You can find the rest of the docker images here. Make sure to use the native datasets for the jobs and the following image versions:

- blackscholes: anakli/cca:parsec_blackscholes
- canneal: anakli/cca:parsec_canneal
- dedup: anakli/cca:parsec_dedup
- ferret: anakli/cca:parsec_ferret
- freqmine: anakli/cca:parsec_freqmine
- radix: anakli/cca:splash2x_radix
- vips: anakli/cca:parsec_vips

<u>IMPORTANT</u>: you must delete your cluster when you are not using it! Otherwise, you will easily use up all of your cloud credits! When you are ready to work on the project, you can easily re-launch the cluster with the instructions above.

To delete your cluster, use the command:

```
$ kops delete cluster --name part4.k8s.local --yes
```

4.1.2 Setting resource limits

taskset is an essential command for setting the CPU affinity of processes. For instance, running taskset -a -cp 0-2 <pid> will bind all threads (-a switch) of the running process indicated by <pid> (-p parameter) to the CPUs 0, 1 and 2 (-c parameter). One can also use this command when starting up processes. More information on taskset can be obtained here.

For Docker, the --cpuset-cpus parameter is used to set the cores a container is able to use. This parameter can be set both when spinning up a container (e.g. sudo docker run --cpuset-cpus="0-2" ...) or updated when a container is already running (e.g. docker container update --cpuset-cpus="0-2" CONTAINER).

You are also free to use other means to dynamically adjust resource allocation for your running jobs. This can refer to resources other than CPU cores.

4.2 Questions

Answer and submit the questions for Part 4 in the report template.

4.3 Submission

For part 4, is expected that you submit:

- The answers to the questions
- The script you used to automate the scheduler
- Any other script or file that you consider needed/useful for the script above
- Your measurement output files, as explained below

You should have at least 6 scheduler runs in your report (3 for question 4.3, 3 for question 4.4). For each of them, you have to produce a couple of files, namely the mcperf output (similar to what explained in part 3, and with an example here) and the container execution log.

Since you are not expected to use k8s for this part, you have to produce a text-based log. We provide you with a utility class in Python that does exactly that. Feel free to re-implement it in any language that you decide to use, but the output must adhere to its format. Each line represents an event, and starts with a date in ISO format (e.g. 2023-04-12T09:52:37.019688), followed by the event name (start, end, update_cores, pause, unpause, and custom) and the job name (memcached, blackscholes, canneal, dedup, ferret, freqmine, radix, vips). start event must be followed by two further elements that represent the list of CPU cores (0..3) the process has assigned at the beginning and the number of (software) threads that is started with. update_cores event has an additional argument that represents the new list of assigned cores. custom event has an arbitrary string (that is URL-encoded) as the last parameter; use this event if you are applying different techniques that are not supported by the logger, or if you want to add comments to the trace. Trailing whitespaces and newlines are ignored, you can use either Unix-like line endings (\n) or Windows-like line endings (\r\n).

The file must start and end with start and end events for scheduler, with no core assignment specified. Remember that each PARSEC job that you start must eventually end. Remember that also memcached needs a start event, but you don't need to end it; if memcached is already running, log start memcached just after the start scheduler event. Refer to this file for an example.

Then, create two directories in the root of the archive, $part_4_3_results_group_XXX$ and $part_4_4_results_group_XXX$, each with 6 files inside. For measurement i=1..3, they should be named $jobs_i.txt$ and $mcperf_i.txt$. The jobs file should contain the log as explained above, while the mcperf one represents the mcperf output.

If the file or directory format does not adhere to this description, or if the files are not complete or do not match the charts, you may have points subtracted from the related questions.

There are no requirements regarding the structure of the other requested files.

5 FAQ

- When running kops create:
 - if you get the following error: failed to create file as already exists: gs://cca-eth-2023-group-XXX-ethzid/part1.k8s.local/config. error: error creating cluster: file already exists, you need to delete the contents of your Google Cloud storage bucket, the recreate it with the following commands:

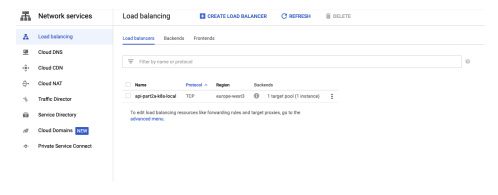
```
$ gsutil rm -r gs://cca-eth-2023-group-XXX-ethzid/
$ gsutil mb gs://cca-eth-2023-group-XXX-ethzid/
```

- if you get the following error: Error: error creating cluster: error writing Cluster "part1.k8s.local": error from acl provider "k8s.io/kops/acl/gce": error querying bucket "...": googleapi: Error 404: The requested project was not found., notFound, make sure you have set the credentials correctly:
 - \$ gcloud auth application-default login
- When ssh-ing into a cluster node, if you get an error like WARNING: REMOTE HOST IDENTIFICATION HAS CHANGED!
 Offending ED25519 key in /Users/username/.ssh/known_hosts:9

Host key verification failed

then you need to run ssh-keygen -R <host> where <host> is the IP address of the server you want to access.

- If kubectl commands prompt you for a username and password, or if kops validate says Unauthorized, first try to re-export the k8s credentials configuration using kops export kubecfg --admin. If it still does not work, delete the cluster and recreate it from scratch.
- If for any reason you cannot delete the cluster with the kops command do the following:
 - Go to console.cloud.google.com
 - Type in the search bar the term "Load balancers". You should be redirected to a page similar to the one below:



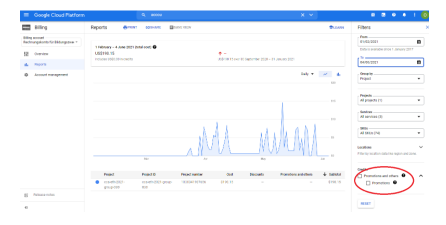
- Select and delete the load balancer.
- Then type in the search bar the term "Instance groups". You should be redirected to a page similar to the one below:



- Select and delete all the instance groups.
- Delete your Google Cloud storage bucket by typing:
 - \$ gsutil rm -r gs://cca-eth-2023-group-XXX-ethzid/
- Also under "External IP addresses" check there are no charges for left over static IPs.
- If your Google Cloud Credits are disappearing even though no charges appear on your Billing Overview, make sure you have unselected "Promotions"
 - Go to console.cloud.google.com
 - Type in the search bar the term "Account Overview". Select "Go to linked billing account" if prompted. You should be redirected to a page similar to the one below:



- Click on "View report".
- Make sure you unclick "Promotions and Other" as shown below and select a reasonable To/From time range:



| • If you run out of credits for your project, please email cloud-arch-ta@lists.inf.ethz.ch to request additional cloud credits. |
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