Reproducible Workflow on a Public Cloud for Computational Fluid Dynamics

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Abstract—In a new effort to make our research transparent and reproducible by others, we developed a workflow to run and share computational studies on the public cloud Microsoft Azure. It uses Docker containers to create an image of the application software stack. We also adopt several tools that facilitate creating and managing virtual machines on compute nodes and submitting jobs to these nodes. The configuration files for these tools are part of an expanded “reproducibility package” that includes workflow definitions for cloud computing, in addition to input files and instructions. This facilitates recreating the cloud environment to rerun the computations under the same conditions. Although cloud providers have improved their offerings, many researchers using high-performance computing (HPC) are still skeptical about cloud computing. Thus, we ran benchmarks for tightly coupled applications to confirm that the latest HPC nodes of Microsoft Azure are indeed a viable alternative to traditional on-site HPC clusters. We also show that cloud offerings are now adequate to complete computational fluid dynamics studies with in-house research software that uses parallel computing with GPUs. Finally, we share with the community what we have learned from nearly two years of using Azure cloud to enhance the transparency and reproducibility in our computational simulations.

Reproducible research and replication studies are essential components for the progress of evidence-based science, even more now when nearly all fields advance via computation. We use computer simulations and data models to
create new knowledge, but how do we provide evidence that this new knowledge is justified? Traditional journal publications exclude software and data products from the peer-review process, yet reliance on ever more complex computational artifacts and methods is the norm. Lacking standards for documenting, reporting and reviewing the computational facets of research, it becomes difficult to verify and corroborate the findings presented in journals and conferences.

The literature is cluttered with confused and sometimes contradictory definitions for reproducible research, reproducibility, replicability, repetition, etc. It is thus worth clarifying how we use these terms. “Reproducible research” was used by geophysics professor Jon Claerbout in the 1990s to mean computational studies that are published with sufficient transparency so other scientists can recreate the results. His research group at Stanford created a reproducible-research environment whose goal was complete documentation of scientific computations, in such a way that a reader could reproduce all the results and figures in a paper using the author-provided computer programs and raw data. This requires open data and open source software and, for this reason, the reproducibility movement is closely linked with the open science movement. The term “replication” has been adopted to refer to an independent study generating new data which, when analyzed, lead to the same findings. We follow this convention, adopted also recently in reports from the National Academies of Sciences, Engineering, and Medicine.

Many efforts to develop cyberinfrastructure that supports reproducible research have been launched in past years. They address concerns like automatic capture of changes to software (version control systems), persistent data archival, global registration of data identifiers, workflow management, etc. But capturing the whole computational environment used in a research project remains one of the most difficult problems. Computational researchers often use a multilayer stack of software applications that can be laborious to build from scratch. Container technology like Docker is a recent addition to the reproducibility toolbox. In this article, we develop and assess a workflow for reproducible research on the public cloud provider Microsoft Azure, adopting Docker containers and several other tools to automate and fully document scientific computations. The workflow follows the idea of using container technology to enhance transparency and reproducibility, and to reduce the burden of installing our application software stack. We go one step further by moving the computational work to Azure cloud resources using tools that programatically capture the virtual environment configuration. The configuration files become part of expanded “reproducibility packages” for the study. This paper does not target portability of our software and workflow to diverse high-performance computing (HPC) environments. Our goal is, rather, to publish our computational research with deep transparency. The workflow we developed and document here is expressly designed for Azure cloud offerings. An interested reader could thus rerun our simulations under exactly the same conditions as we used, provided they are willing and able to create an Azure account.

Universities and national laboratories spend millions of dollars to deploy and maintain on-site HPC clusters. At the George Washington University, we have access to a cluster called Colonial One. The cluster is now 6 years old and approaching its end-of-life. On average, its computational resources are idle 9% of the time and unavailable to users for roughly five days a year (due to maintenance). Administrators of the cluster have been recently considering integrating cloud-computing platforms in the research-computing portfolio. Over the past decade, cloud-computing platforms have rapidly evolved, now offering solutions for scientific applications. From a user’s point of view, a cloud platform offers great flexibility (hardware and virtual machines) with instantaneous availability of infinite (in appearance) computational resources. No more waiting time in job-submission queues! This promises to greatly facilitate code development, debugging, and testing. Resources
allocated on the cloud are released as soon as the job is done, avoiding paying for idle time. On a public-cloud platform—such as Microsoft Azure, Google Cloud, Amazon AWS—with a “pay-as-you-go” type of subscription, a user directly sees how much it costs to run a scientific application, while such information usually remains obscure to the end-user on university-managed clusters. Cost models that make sense for researchers, labs, and universities are still unclear. Yet, this information is key when making a decision to adopt a cloud workflow for research.

Until recently, public cloud offerings were inadequate to the needs of computational scientists using HPC, mainly due to performance overhead of virtualization or lack of support for fast networking and hardware accelerators. Freniere and co-authors (2016) used the OSU microbenchmark suite to report performance degradation in networking with Amazon AWS, compared to their on-site local HPC cluster. We take the same approach with Microsoft Azure to verify that their improved services are now suitable for HPC applications. We also share what we have learned from nearly two years of using cloud computing for our computational fluid dynamics (CFD) simulations, with the hope that it will shed some light on the costs and benefits, and help others considering using cloud computing for their research.

REPRODUCIBLE CLOUD-BASED WORKFLOW

Scientific publications reporting computational results often lack sufficient details to reproduce the researcher’s computational environment; e.g., they may miss to mention external libraries used along with the main computational code. We have learned the hard way how different versions of the same external library can alter the numerical results and even the scientific findings of a computational study. This section presents an overview and mini-tutorial of the workflow we developed to aim for the highest level of reproducibility of our computational research, with the best available technology solutions. In the process of creating this reproducible workflow, we also evaluated the suitability of public cloud offerings by Microsoft Azure for our research computing needs. The tools we adopted for computing on cloud resources are specific for this provider.

Use of Container Technology

To overcome the so-called “dependency hell” and facilitate reproducibility and portability, we use the container technology provided by the open-source project Docker. A container represents an isolated user space where application programs run directly on the operating system’s kernel of the host (with limited access to its resources). In contrast with virtual machines, containers do not include a full operating system, making them lighter and faster. Containers allow recreating the same runtime environment of an application (including all its dependencies) from one machine to another. They empower researchers to share pre-built images of their software stack, ameliorating one of the biggest pain points for reproducible research: building the software on a different machine. The majority of enterprise developers today are familiar with Docker container technology (first released six years ago). But in academic settings, many are still unaware of its value. We present this section as an overview for research software developers unfamiliar with Docker, but comfortable with scripting, version control, and distributed collaboration.

A container is an instance of an image. The developer builds this image on a local machine and pushes it to a public registry to share it with other users. Users pull the image from the public registry—in our case, DockerHub—and create containers out of it. To create a Docker image, the developer writes a Dockerfile: an ASCII file containing instructions that tell Docker what to do. For example, we start building a new image from a base image using the keyword FROM. We then write the different shell instructions (prefixed with the keyword RUN) to build a multilayered runtime environment of the application that includes all its dependencies. Once we have built and tested the image on the local machine, we push it to a repository on the DockerHub registry: a place to store and retrieve Docker images. Listing 1 provides the command lines to build (docker build) and push (docker push) an image of our CFD software (barbagroup/petibm:0.4-GPU-IntelMPI-ubuntu) that we used to obtain some of the results presented in the following section. Here, CLOUDREPRO is an environment variable set to the local path of the GitHub repository for this paper, cloud-repro,
Listing 1. Build and push a Docker image.

```bash
$ cd $CLOUDREPRO/docker/petibm
$ docker build
    -- tag=barbagroup/petibm:0.4-GPU-IntelMPI-ubuntu
    -- file=Dockerfile.
$ docker push
    barbagroup/petibm:0.4-GPU-IntelMPI-ubuntu
```

A reader interested in reproducing the computational results can now pull the application image from DockerHub, and create a Docker container to run the CFD application software in a faithfully reproduced local environment. Our objective is to create and run containers on the public cloud provider Microsoft Azure, and make it possible for the readers to also run these containers to reproduce our results. Figure 1 shows a graphical representation of the workflow we developed using Docker and various tools for running CFD simulations on Microsoft Azure. The following section explains these tools.

Use of Public Cloud Resources

To run computational jobs on Microsoft Azure, we use several tools that facilitate creating and managing an HPC-capable cluster of cloud nodes, including virtual machines, virtual networks, job and task scheduling infrastructure. Azure Batch works with both embarrassingly parallel workloads and tightly coupled MPI jobs (the latter being the case of our CFD software). To use Azure Batch, we first need to configure a workspace on Microsoft Azure. This can be done either via the Azure Portal in a web browser or from a local terminal using the open-source tool Azure CLI (version 2.0.57, https://github.com/Azure/azure-cli). We prefer to use the command-line solution (program `az`), as it allows us to keep track of the steps taken to configure the cloud workspace (see Listing 2). First, we set the Azure subscription we want to use (let’s call it `reprosubscription`). Next, we create a resource group (`reprorg`) located in this case in the East US region, which will contain all the Azure resources. We create an

```
Listing 2. Configure the workspace on Microsoft Azure.

```bash
$ az account set -- subscription reprosubscription
$ az group create -- name reprogr -- location eastus
$ az storage account create -- name reprostorage
    -- resource- group reprogr -- sku Standard_LRS
    -- location eastus
$ az batch account create -- name reprobatch
    -- resource- group reprogr -- location eastus
    -- storage- account reprostorage
$ az storage share create -- name fileshare
    -- account -- name reprostorage -- account- key
    storagekey -- quota 100
```
Azure Storage account \((\text{reprostorage})\) in the resource group, as well as an Azure Batch account \((\text{reprobatch})\) associated with the storage account. Finally, we create a fileshare (in this case of size 100 GB) in the storage.

To create computational nodes and submit container-based jobs to Azure Batch, we use the open-source command-line utility Batch Shipyard (version 3.6.1: https://github.com/Azure/batch-shipyard). Batch Shipyard is entirely driven by configuration files: the utility parses user-written YAML files to automatically create pools of compute nodes on Azure Batch and to submit jobs to those pools. Typically, we need to provide four configuration files:

- `config.yaml` contains information about the Azure Storage account and Docker images to use.
- `credentials.yaml` stores the necessary credentials to use the different Microsoft Azure service platforms (e.g., Azure Batch and Azure Storage).
- `pool.yaml` is where the user configures the pool of virtual machines to create.
- `jobs.yaml` details the configuration of the jobs to submit to the pool.

Once the configuration files are written, we invoke Batch Shipyard (program `shipyard`) on our local machine. The folder `examples/snake2d2k35/config_shipyard` in the repository accompanying this paper contains an example YAML file to create a pool of two NC24r compute nodes (featuring K80 GPUs and using InfiniBand network). Listing 3 shows the commands to run in your local terminal to create a pool of compute nodes on Azure Batch and submit jobs to it. The Docker image of our CFD application is pulled from the registry to the virtual machines during the pool creation (`shipyard pool add`). We then upload the input files to the compute nodes (`shipyard data ingress`) and submit jobs to the pool (`shipyard jobs add`). The tasks for a job will start automatically upon submission.

**Listing 3. Create a pool and submit jobs to it.**

```bash
$ cd $CLOUDREPRO/examples/snake2d2k35
$ az storage directory create --name snake2d2k35
    --share-name fileshare --account-name reprostorage
$ export SHIPYARD_CONFIGDIR=config_shipyard
$ shipyard pool add
$ shipyard data ingress
$ shipyard jobs add
```
Once the simulations are done (i.e., the job tasks are complete), we delete the jobs and the pool (Listing 4). The output of the computation is now stored in the fileshare in our Azure Storage account. We can download the data to our local machine to perform additional postprocessing steps (such as flow visualizations).

Reproducible research requires authors to make their code and data available. Thus, the Dockerfile and YAML configuration files should be made part of an extended reproducibility package that includes workflow instructions for cloud computing, in addition to other input files. Such a reproducibility package facilitates recreating the cloud environment to run the simulations under the same conditions. The reproducibility packages of the examples showcased in the following section are available in the GitHub repository cloud-repro, which includes instructions on how to reproduce the results.

Listing 4. Delete the pool and jobs, and download to output to a local machine.

$ shipyard pool del
$ shipyard jobs del
$ mkdir output
$ az storage file download-batch --source fileshare/snake2d2k25 --destination output --account-name reprostorage

RESULTS

The top concerns of researchers considering cloud computing are performance and cost. Until just a few years ago, the products offered by cloud providers were unsuitable to the needs of computational scientists using HPC, due to performance overhead of virtualization or lack of support for fast networking. Azure only introduced nodes with GPU devices during late 2016 and Infiniband support for Linux virtual machines on the NC-series in 2017. Our first objective was to assess the performance on cloud nodes for the type of computations in our research workflows with tightly coupled parallel applications. We present results from benchmarks and test-cases showing that we are able to obtain similar performance in terms of latency and bandwidth using the Azure virtual network, comparing to a traditional university-managed HPC cluster (Colonial One). Table 1 lists the hardware specifications of the nodes used on Microsoft Azure and Colonial One. Our target research application relies on three-dimensional CFD simulations with our in-house research software. We include here a sample of the types of results needed to answer our research question, obtained by running on the public cloud using the reproducible workflow described in the previous section. The goal is to showcase the potential of cloud computing for CFD, share the lessons we learned in the process, as well as analyze the cost scenarios for full applications.

### MPI Communication Benchmarks

We ran point-to-point MPI benchmarks from the Ohio State University Micro-Benchmarks suite (version 5.6, http://mvapich.cse.ohio-state.edu/benchmarks/) on Microsoft Azure and Colonial One, to investigate the performance in terms of latency and bandwidth. The latency test is carried out in a ping-pong fashion and measures the time elapsed to get a response; the sender sends a message with a certain data size and waits for the receiver to send back the message with the same data size. The bandwidth test measures the maximum sustained rate that can be achieved on the network; the sender sends a fixed number of messages to a receiver that replies only after receiving all of them. The tests ran on NC24r nodes on Azure and Ivygpu nodes on Colonial One, all of them featuring a network

| Platform | Node | Intel Xeon CPU | # threads | NVIDIA GPU | RAM (GiB) | SSD Storage (GiB) |
|----------|------|----------------|-----------|------------|----------|------------------|
| Azure    | NC24r| Dual 12-Core E5-2690v3 (2.60 GHz) | 24 | 2 × K80 | 224 | 1440 |
|          | H16r | Dual 8-Core E5-2667v3 (3.20 GHz) | 16 | – | 112 | 2000 |
| Colonial One | Ivygpu | Dual 6-Core E5-2620v2 (2.10 GHz) | 12 | 2 × K20 | 120 | 93 |
|          | Short | Dual 8-Core E5-2650v2 (2.60 GHz) | 16 | – | 120 | 93 |

On both platforms, MPI applications take advantage of RDMA (remote direct memory access) network with FDR InfiniBand and ECC is enabled for GPU computing.
interface for RDMA (Remote Direct Memory Access) connectivity to communicate over InfiniBand. (RDMA allows direct access to a remote’s memory without involving the operating system of the host and remote.) Figure 2 reports the mean latencies and bandwidths obtained over five repetitions on both platforms. For small message sizes, the average latencies on Colonial One and Azure are 1.25 and 1.95 μs, respectively.

For all message sizes, the latency reported on Azure is slightly higher than the value obtained on Colonial One. The maximum sustained bandwidth rates for Colonial One and Azure are on average 6.2 and 5.2 GB/s, respectively. For all message sizes, a similar bandwidth rate was achieved on Azure and Colonial One. Over the last few years, Microsoft Azure has indeed improved its HPC solutions to provide networking capabilities that are comparable or even better than our 6-year-old university-managed cluster.

Poisson Benchmarks

CFD algorithms often require solving linear systems with iterative methods. For example, the Navier–Stokes solver implemented in our software requires the solution of a Poisson system at every time step to project the velocity field onto the divergence-free space (to satisfy the incompressibility constraint). We investigated the time-to-solution for a 3-D Poisson system (obtained with a 7-point stencil central-difference scheme) on different nodes of Microsoft Azure and Colonial One. The solution method was a conjugate-gradient (CG) method with a classical algebraic multigrid (AMG) preconditioner using an exit criterion set to an absolute tolerance of $10^{-12}$. The iterative solver ran on CPU nodes (H16r instances on Azure and Short nodes on Colonial One) using the CG algorithm from the PETSc library and an AMG preconditioner from Hypre BoomerAMG. Figure 3 (top) reports the mean runtimes (averaged over five repetitions) to iteratively solve the system, on a uniform grid of 50 million cells ($1000 \times 1000 \times 50$), as we increase the number of nodes in the pool (strong scaling). Runtimes obtained on Colonial One and Azure are similar. We also solved the Poisson system with the NVIDIA AmgX library on multiple GPU devices using NC24r instances on Azure and Ivygpu nodes on Colonial One. The Poisson system for a base mesh of 6.25 million cells ($500 \times 500 \times 25$) was solved on a single compute node using 12 MPI processes and 2 GPU devices; we then doubled the mesh size as we doubled the number of nodes, keeping the same number of MPI processes and GPUs per node (weak scaling). The number of iterations to reach convergence increases with the size of the system, so we normalize the runtimes by the number of iterations. Figure 3 (bottom left) shows the normalized mean runtimes (five repetitions) obtained on Azure and Colonial One. The smaller runtimes on Azure are explained by the fact that the NC-series of Microsoft Azure features NVIDIA K80 GPU devices with a higher compute capability than the K20 GPUs on Colonial One. The bottom-right panel of the figure reports the normalized mean runtimes obtained on Azure when we load the bandwidth with a larger problem; the base mesh now contains 25 millions cells ($1000 \times 500 \times 50$). We
observe larger variations in the time-to-solution for the Poisson system on Microsoft Azure with GPU computing; runtimes are more uniform on Colonial One.

Flow Around a Flying Snake Cross Section

Our research lab is interested in understanding the aerodynamics of flying animals via CFD simulations. One of our applications deals with the aerodynamics of a snake species, *Chrysopelea paradisi*, that lives in Southeast Asia. This arboreal reptile has the remarkable capacity to turn its entire body into a wing and glide over several meters. The so-called “flying snake” jumps from tree branches, undulates in the air, and is able to produce lift by expanding its ribcage to flatten its ventral surface (morphing its normally circular cross section into a triangular shape).

To study the flow around the flying snake, we developed a CFD software called PetIBM, an open-source toolbox that solves the 2-D and 3-D incompressible Navier–Stokes equations using a projection method (seen as an approximate block-LU decomposition of the fully discretized equations) and an immersed-boundary method (IBM). Within this framework, the fluid equations are solved over an extended grid that does not conform to the surface of the body immersed in the computational domain. To model the presence of the body, the momentum equation is augmented with a forcing term that is activated in the vicinity of the immersed boundary. This technique allows the use of simple fixed structured Cartesian grids to solve the equations. PetIBM implements immersed boundary methods (IBMs) that fit into the projection method of Perot. In the present study, we use the IBM scheme proposed by Li et al. PetIBM runs on distributed-memory architectures using the efficient data structures and parallel routines from the PETSc library. The software also implements the possibility to solve linear systems on multiple GPU devices distributed across the nodes with the NVIDIA AmgX library and our AmgXWrapper. One of the requirements for reproducible computational results is to make the code available under a public license (ideally allowing reuse and modification by others). In that regard, PetIBM is open source, version-controlled on GitHub (version 0.4, https://github.com/barbagroup/PetIBM), and shared under the permissive (non copy-left) BSD-3 clause license. We also provide a Docker image of PetIBM on DockerHub and its Dockerfile is available in the GitHub repository of the software.

2-D Flow Around a Snake Cross Section

We submitted a job on Azure Batch (with Batch Shipyard) to compute the 2-D flow around an anatomically accurate sectional shape of the gliding snake. The cross section has a chord-length $c = 1$ and forms a 35° angle of attack with the incoming freestream flow. The Reynolds number, based on the freestream speed, the body
chord-length, the kinematic viscosity, is set to $Re = 2000$. The immersed boundary is centered in a $30c \times 30c$ computational domain that contains just over 2.9 million cells. The grid is uniform in the vicinity of the body and stretched to the external boundaries with a constant ratio (see Table 2 for details about the grid). The discretization of the immersed boundary has the same resolution as the background fluid grid. A convective condition was used at the outlet boundary while freestream conditions were enforced on the three other boundaries. The job was submitted to a pool of two NC24r nodes, using 12 MPI processes and 2 GPU devices per node. It completed 200 000 time steps (i.e., 80 time units of flow simulation with time-step size $\Delta t = 0.0004c/U_\infty$) in just above 7 h (wall-clock time).

Figure 5 shows the history of the force coefficients on the 2-D cross section. The lift coefficient only maintains its maximum mean value during the early stage of the simulation (up to 40 time units). Between 40 and 50 time units, the mean value starts to drop. The time-averaged force coefficients (between 40 and 80 time units) are reported in Table 3. Figure 4 shows snapshots of the vorticity field after 20, 44, 45, and 80 time units of flow simulation. After 20 time units, the vortices shed from the snake section are almost aligned in the near wake (with a slight deflection toward the lower part of the domain). Snapshots of the vorticity at time units 44 and 45 show that the initial alignment is perturbed by vortex-merging events (same-sign vortices merging together to form a stronger one). Following that, the wake signature is altered for the rest of the simulation: vortices are not aligned anymore, the wake becomes wider (leading to lower aerodynamic forces) with a 1S+1P vortex signature (a single clockwise-rotating vortex on the upper part and a vortex pair on the lower part).

### Table 2. Details about the computational grids used for the snake simulations with PetIBM.

| Case | Domain | Uniform region | Smallest cell-width | Stretching ratio | Size           |
|------|--------|----------------|---------------------|------------------|----------------|
| 2-D  | $30 \times 30$ | $([-0.52, 3.48] \times [-2, 2]$ | $0.004 \times 0.004$ | 1.01 | $1704 \times 1706$ |
| 3-D  | $30 \times 30 \times 3.2$ | $([-0.52, 3.48] \times [-2, 2] \times [0, 3.2]$ | $0.008 \times 0.008 \times 0.08$ | 1.01 | $1071 \times 1072 \times 40$ |

Distances are expressed in terms of chord-length units.

**Figure 4.** Filled contour of the vorticity field ($-5 \leq \omega_zc/U_\infty \leq 5$) after 20, 44, 45, and 80 time units of flow simulation with PetIBM for the snake cross section at a $35^\circ$ angle of attack and Reynolds number 2000. Vortex merging events trigger a change in the wake signature causing the drop in the mean value of the lift coefficient.

**3-D Flow Around a Snake Cylinder**

Although the 2-D simulations can give us some insights into flow dynamics happening in the wake behind the snake, we know that at this Reynolds number ($Re = 2000$), 3-D structures will develop in the wake. We submitted jobs...
on Azure Batch to perform direct numerical simulation of the 3-D flow around a snake model: a cylinder with the same anatomically accurate cross section. The computational domain extends in the $z$-direction over a length of $3.2c$ and the grid now contains about 46 million cells (with a uniform discretization in the $z$-direction and periodic boundary conditions; see Table 2). The job was submitted to a pool of two NC24r nodes using 24 MPI processes and 4 GPU devices per node. The task completed 100 000 time steps (100 time units with a time-step size $\Delta t = 0.01c/U_\infty$) in about five days and 16 h.

Figure 5 compares the history of the force coefficients between 2-D and 3-D configurations. The force coefficients resulting from the 2-D simulation of the snake are higher than those obtained for the snake cylinder; we computed a relative difference of $+37.9\%$ and $+14.4\%$ for the time-averaged drag and lift coefficient, respectively (see Table 3). Two-dimensional computational simulations of fundamentally 3-D flows lead to incorrect estimation of the force coefficients, as is well known. The grid used for the 3-D simulation contains about 46 million cells and is somewhat coarse for direct numerical simulation at this Reynolds number, but acceptable for our exploratory analysis. A finer grid would be preferable to capture the flow dynamics in detail, but the purpose of this article is not targeting the physics. We did however run a simulation on a finer grid (about 233 million cells) resulting in a relative difference (with respect to the “coarse”-grid simulation) of $+6.5\%$ and $+3\%$ for the time-averaged drag and lift coefficients, respectively. We decided to continue this paper with the coarser grid due to the large cost difference (see the next section, Cost Analysis and User Experience). Details about this fine-grid simulation are available as supplementary material on the GitHub repository for this study (https://nbviewer.jupyter.org/github/barbagroup/cloud-repro/blob/master/misc/independence/grid.ipynb).

Table 3. Time-averaged force coefficients on the snake model at Reynolds number 2000 and angle of attack $35^\circ$ for the 2-D and 3-D configurations.

| Case | $C_D$     | $C_L$     |
|------|-----------|-----------|
| 3D   | 0.8390    | 1.5972    |
| 2D   | 1.1567 (+37.9%) | 1.8279 (+14.4%) |

(We average the force coefficients between 40 and 80 time units of flow simulation and report the relative difference of the 2-D values with respect to the 3-D ones.)

Figure 6 shows the instantaneous spanwise vorticity averaged along the $z$-direction after 80 and 100 time units. Compared to the snapshots from the 2-D simulation, we observe that free-shear layers roll up into vortices further away from the snake body and that the von Kármán street exhibits a narrower wake than in the 2-D simulations. We also note the presence of an unsteady recirculation region just behind the

\[ \text{Table 3. Time-averaged force coefficients on the snake model at Reynolds number 2000 and angle of attack $35^\circ$ for the 2-D and 3-D configurations.} \]

\begin{tabular}{|c|c|c|}
\hline
Case & $(C_D)$ & $(C_L)$ \\
\hline
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\hline
\end{tabular}

(We average the force coefficients between 40 and 80 time units of flow simulation and report the relative difference of the 2-D values with respect to the 3-D ones.)
snake cylinder, and alternating regions of positive and negative cross-flow velocity showing the presence of von Karman vortices (see Figure 7). Figure 8 shows a side-view of the isosurfaces of the Q-criterion after 100 time units and highlights the complexity of the 3-D turbulent wake generated by the snake model.

The selection of results presented here corresponds to a typical set of experiments included in a CFD study. A comprehensive study might include several similar sets, leading to additional insights about the flow dynamics, and clues to future avenues of research. We include this selection here to represent a typical workflow, and combine our discussion with a meaningful analysis of the costs associated with running CFD studies in a public cloud.

Table 4. Charges incurred for the usage of different services on Microsoft Azure.

| Service name          | Cost (USD) | % of total cost |
|-----------------------|------------|-----------------|
| Bandwidth             | 46.85      | 0.23            |
| Data management       | 0.56       | 0.003           |
| Networking            | 1.38       | 0.007           |
| Storage               | 25.93      | 0.16            |
| Virtual machines      | 20,582.64  | 99.6            |
| **Total**             | **20,614** |                 |

COST ANALYSIS AND USER EXPERIENCE

For the year 2018, Microsoft Azure granted our research lab a sponsorship (in terms of cloud credits) to run CFD simulations with our in-house software. With a “pay-as-you-go” subscription, users can immediately see how much it costs to run a scientific application on a public cloud; such information is often hidden to end-users on university-managed clusters. From May to December, we spent a total of 20,614 USD to run CFD simulations, including a couple dozen snake simulations. (The output of those simulations is now being processed to further analyze the complex flow dynamics generated behind the snake model.) During that period, we have been charged for data management, networking, storage, bandwidth, and virtual machines (see Table 4). More than 99% of the charges incurred were for the usage of virtual machines, mainly instances from the NC-series (see Table 5). To run PetIBM simulations of the snake model, we used the NC24r virtual machines to get access to NVIDIA K80 GPUs and InfiniBand networking. For example, the 2-D snake run reported in this paper cost 55.4 USD (2 NC24r

Figure 7. Filled contour of the streamwise velocity (top) and cross-flow velocity (bottom) behind the snake cylinder in the x/z plane at y/c = −0.2 in the wake of the snake cylinder with a 35° angle of attack at Reynolds number 2000 after 100 time-units of flow simulations. There are 52 contours from −1.0 to 1.0. The gray area shows a projection of the snake cylinder in the x/z plane. The solid black line defines the contour with u_x = 0 (top) and u_y = 0 (bottom).

Figure 8. Lateral view of the isosurfaces of the Q-criterion (Q = 1) in the wake of the snake cylinder (with a 35° angle of attack) at Reynolds number 2000. The isosurfaces are colored with the streamwise vorticity (−5 ≤ w_z/c/U_∞ ≤ 5). The figure was generated using the visualization software VisIt.¹⁷
instances with a hourly price of 3.96 USD for about 7 h). The three-dimensional computation cost 1077.1 USD (2 NC24r instances for about 136 h). (Note that with 3-year-reserved instances, the 2-D and 3-D snake simulations would have only cost 24.6 and 478.1 USD, respectively.) The 3-D simulation on the finer grid (233 million cells) ran on 6 NC24r instances and cost about 7965 USD to compute 100,000 time steps (cost not included in Table 4). This is more than seven times the cost we paid for the “coarse”-grid run! It would have been too expensive to run a batch of simulations on the finer grid for our exploratory analysis of the flow around the snake cylinder.

Running CFD simulations on Microsoft Azure was a first for our research lab. As novices in cloud computing, it took us several months to become familiar with the technical vocabulary and the infrastructure of Microsoft Azure before we could submit our first simulation of the flow around a snake profile.

Command-line utilities such as Azure CLI and Batch Shipyard were of great help to create and manage resources on Microsoft Azure for executing our reproducible research workflow. (In our lab, we tend to avoid interacting directly with graphical user interfaces to keep a trace of the commands run and make the workflow more reproducible.) Azure CLI helped us to setup the Azure Batch and Storage accounts as well as moving data between the cloud platform and our local machines. Thanks to Batch Shipyard, we did not have to dig into the software development kit of Azure Batch to use the service. Writing YAML configuration files was all we needed to do to create a pool of virtual machines with Batch Shipyard and to submit jobs to it. With Batch Shipyard, we have painlessly submitted jobs on Azure Batch to run multi-instance tasks in Docker containers, everything from the command-line terminal on a local machine. Note that Batch Shipyard also supports Singularity containers. Singularity is an open source container platform designed for HPC workloads. Singularity containers can be used on traditional HPC clusters such as Colonial One; Docker containers usually cannot. Indeed, running a Docker container involves running a Docker daemon (a background process) which requires root privileges (that users do not and should not have on production clusters). The container technology from Singularity was designed from the ground up to prevent escalation in user privileges.

Our in-house CFD software, PetIBM, relies on MPI to run applications on distributed-memory architectures. The Poisson system is solved on distributed GPU devices using the NVIDIA AmgX library. Thus, we used Azure instances of the NC-series, which feature a network interface for remote direct memory access (RDMA) connectivity, allowing nodes in the same pool to communicate over InfiniBand network. The Docker images were specifically built to be able to run MPI applications on RDMA-capable virtual machines (such as the NC24r instance with a CentOS-based 7.3 HPC image) with multiple-instance tasks on Azure Batch service. As of this writing, only Intel MPI 5.x versions are supported with the Azure Linux RDMA drivers. Both the Docker image and the configurations files for deploying the cloud resources give an interested reader the opportunity to reproduce our computations in an identical environment. Even if this opportunity is not accompanied by portability—e.g., one may depend on a different MPI library to use a different system—the human-readable Docker files and configuration files do offer transparency.

Microsoft Azure offers the possibility of taking advantage of surplus capacity with “low-priority”

| Instance | cores | RAM (GiB) | disk sizes (GiB) | GPU | pay-as-you-go (dedicated) ($/hr) | pay-as-you-go (low-priority) ($/hr) | 1-year reserved ($/hr) | 3-year reserved ($/hr) |
|----------|-------|----------|------------------|-----|-------------------------------|---------------------------------|-------------------------|------------------------|
| NC6      | 6     | 56       | 340              | 1 × K80 | 0.90                           | 0.18                            | 0.5733                  | 0.3996                 |
| NC12     | 12    | 112      | 680              | 2 × K80 | 1.80                           | 0.36                            | 1.1466                  | 0.7991                 |
| NC24     | 24    | 224      | 1,440            | 4 × K80 | 3.60                           | 0.72                            | 2.2932                  | 1.5981                 |
| NC24r    | 24    | 224      | 1,440            | 4 × K80 | 3.96                           | 0.792                           | 2.5224                  | 1.7578                 |

(Prices as of March 24, 2019, for CentOS or Ubuntu Linux Virtual Machines in the East US region.)
* The NC24r configuration provides a low latency, high throughput network interface optimized for tightly coupled parallel applications.
virtual machines that are substantially cheaper than “dedicated” ones (see Table 5 for pricing options on the NC series with Linux-based virtual machines). For example, the low-priority NC24r instance costs 0.792 USD/h, five times cheaper than its dedicated counterpart. The reader should keep in mind that low-priority virtual machines may not be available for allocation or may be preempted at any time. Thus, low-priority instances should be avoided for long-running MPI jobs in pools, where internode communication is enabled. We only used dedicated virtual machines to run our CFD simulations. Moreover, our job tasks used a shared filesystem (GlusterFS on compute nodes) for I/O operations and Batch Shipyard would fail to create the pool if low-priority nodes were requested.

Microsoft Azure implements quotas and limits on resources with Azure Batch service, such as the maximum number of dedicated cores that can be used in a certain region. To be able to run our simulations, we had to contact Azure Support through Azure Portal to request a quota increase for a given type of instance in a specific region and for a given subscription. We had to go through this process at least five times during our sponsorship period. Readers should be aware of these quotas and limits before scaling up workloads on Azure Batch.

CONCLUSION

Exploring public cloud services as an alternative (or complementary) solution to university-managed HPC clusters to run CFD simulations with our in-house software, we used Microsoft Azure to generate reproducible computational results. Our reproducible workflow makes use of Docker containers to faithfully port the runtime environment of our CFD application to Azure, and the command-line utilities Azure CLI and Batch Shipyard to run multi-instance tasks within Docker containers on the Azure Batch platform. We ran CFD simulations on instances of the NC-series that feature NVIDIA K80 GPU devices and have access to InfiniBand network. Latency and bandwidth microbenchmarks show that our university-managed HPC system and Microsoft Azure can deliver similar performances. Thanks to a Microsoft Azure sponsorship, we were able to run dozens of CFD simulations of a gliding snake’s model. Our study of the 3-D flow around a cylinder with an anatomically accurate cross section of the flying snake is currently ongoing. We plan to analyze the output data from these runs, and publish them in a later article.

In this article, we show that public cloud resources are today able to deliver similar performances to a university-managed cluster, and thus can be regarded as a suitable solution for research computing. But in addition to performance, researchers also worry about cost. We share the actual costs incurred to run 2-D and 3-D CFD simulations using cloud services with a pay-as-you-go model, and report on reduced costs possible with reserved instances. Universities (and possibly funding agencies) may obtain even more favorable pricing through bids or medium-term contracts. Researchers thinking of adding cloud computing to proposal budgets might encounter other barriers, however. For example, some universities exempt large equipment purchases from indirect (facilities and administrative, F&A) costs, while they may add these overhead rates to cloud purchases. Until these internal policies are adjusted, cloud computing may not be adopted widely.

In a heightened effort to make our research transparent and reproducible by others, we maintain in a public version-controlled repository all the files necessary to re-run the examples highlighted in this paper. The repository is found at https://github.com/barbagroup/cloud-repro and is also permanently archived in Zenodo.\(^{18}\) Our expanded reproducibility packages contain input files, Batch Shipyard configuration files, command-line instructions to set up and submit jobs to Microsoft Azure, as well as the postprocessing scripts to reproduce the figures of this paper. We included in the Zenodo archive all secondary data required to reproduce the figures without running the simulations again. Although they are not required to reproduce the computations, the Dockerfiles to build Docker images are also made available. We even include a Dockerfile to reproduce the local environment with the visualization tools needed for postprocessing. These files contain details about all the libraries needed to create the computational runtime environment. Both the Dockerfiles and the...
workflow files may not execute as they were designed to do in a period of a few years, due to technology changes. Obsolescence of digital research objects, including software libraries and computational environments, is an acknowledged source of non-reproducibility, in the long run. The recent report of the National Academies highlights this [6, p. 57]. Indeed, tools in our workflow that we have no control over (Azure CLI, Batch Shipyard) will likely change, perhaps breaking backward compatibility. That is inevitable. But when they do, our fully documented workflow (via human-readable configuration files) will continue to offer transparency of what we did and how we did it.

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