

# WorkflowHub: Community Framework for Enabling Scientific Workflow Research and Development

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**Abstract**—Scientific workflows are a cornerstone of modern scientific computing. They are used to describe complex computational applications that require efficient and robust management of large volumes of data, which are typically stored/processed on heterogeneous, distributed resources. The workflow research and development community has employed a number of methods for the quantitative evaluation of existing and novel workflow algorithms and systems. In particular, a common approach is to simulate workflow executions. In previous work, we have presented a collection of tools that have been used for aiding research and development activities in the Pegasus project, and that have been adopted by others for conducting workflow research. Despite their popularity, there are several shortcomings that prevent easy adoption, maintenance, and consistency with the evolving structures and computational requirements of production workflows. In this work, we present *WorkflowHub*, a community framework that provides a collection of tools for analyzing workflow execution traces, producing realistic synthetic workflow traces, and simulating workflow executions. We demonstrate the realism of the generated synthetic traces by comparing simulated executions of these traces with actual workflow executions. We also contrast these results with those obtained when using the previously available collection of tools. We find that our framework not only can be used to generate representative synthetic workflow traces (i.e., with workflow structures and task characteristics distributions that resemble those in traces obtained from real-world workflow executions), but can also generate representative workflow traces at larger scales than that of available workflow traces.

**Index Terms**—Scientific Workflows, Workflow Management Systems, Simulation, Distributed Computing, Workflow Traces

## I. INTRODUCTION

Scientific workflows are relied upon by thousands of researchers [1] for managing data analyses, simulations, and other computations in almost every scientific domain [2]. Scientific workflows have underpinned some of the most significant discoveries of the last decade [3], [4]. These discoveries are in part a result of decades of workflow management system (WMS) research, development, and community engagement to support the sciences [5]. As workflows continue to be adopted by scientific projects and user communities, they are becoming more complex and require more sophisticated workflow management capabilities. Workflows are being designed that can analyze terabyte-scale datasets, be composed of millions of individual tasks that execute for milliseconds up to several hours, process data streams, and process static data in object stores. Catering to these workflow features and demands

requires WMS research and development at several levels, from algorithms and systems all the way to user interfaces.

A traditional approach for testing, evaluating, and evolving WMS is to use full-fledged software stacks to execute applications on distributed platforms and testbeds. Although seemingly natural, this approach has severe shortcomings including lack of reproducible results, limited platform configurations, and time and operational costs. An alternative that reduces these shortcomings is to use simulation, i.e., implement and use a software artifact that models the functional and performance behaviors of software and hardware stacks of interest. Thus, the scientific workflow community has leveraged simulation for the development and evaluation of, for example, novel algorithms for scheduling, resource provisioning, and energy-efficiency, workflow data footprint constraints, exploration of data placement strategies, among others [6], [7].

Studying the execution of workflows in simulation requires sets of workflow applications to be used as benchmarks. This is so that quantitative results are obtained for a range of representative workflows. In [8], we have described a collection of tools and data that together have enabled research and development of the Pegasus [3] WMS. These community resources have enabled over 30 research papers<sup>1</sup> by providing synthetic workflow traces for evaluation via simulation. Despite the extensive usage of this pioneer effort, it lacks (i) a common format for representing workflow execution traces in a way that is agnostic to workflow systems; (ii) structured methods for encoding the workflow design and structure; (iii) robust techniques for generating synthetic workflows in which workflow characteristics conform to the original workflow features; and (iv) a set of tools for analyzing workflow traces, which would support the integration of traces from new application domains.

In this paper, we present the *WorkflowHub project* [9], an open source community framework that provides a collection of structured methods and techniques, implemented as part of usable tools, for analyzing workflow traces and producing synthetic, yet realistic, workflow traces. WorkflowHub mitigates the shortcomings of our previous set of tools by using a common JSON format for representing workflow traces. Any workflow execution log can be captured into this system-agnostic format. In addition, WorkflowHub provides an

<sup>1</sup>Based on records provided by Google Scholar.



statistical distributions (uniform and normal), and as a result workflow performance behavior may not be representative (see results in Section IV).

### III. THE WORKFLOWHUB

The WorkflowHub project (<https://workflowhub.org>) is a community framework for enabling scientific workflow research and development. It provides foundational tools for analyzing workflow execution traces, and generating synthetic, yet realistic, workflow traces. These traces can then be used for experimental evaluation and development of novel algorithms and systems for overcoming the challenge of efficient and robust execution of ever-demanding workflows on increasingly complex distributed infrastructures.

Fig. 1 shows an overview of the workflow research life cycle process that integrates the three axes of the WorkflowHub project: (i) workflow execution traces, (ii) workflow generator, and (iii) workflow simulator.

#### A. Workflow Execution Traces

The first axis of the WorkflowHub project targets the collection and curation of open access production workflow execution traces from various scientific applications, all made available using a common trace format. A workflow execution trace is built based on logs of an actual execution of a scientific workflow on a distributed platform (e.g., clouds, grids, clusters). More specifically, the three main types of information included in the trace are:

- workflow task execution metrics (runtime, input and output data sizes, memory used, energy consumed, CPU utilization, compute resource that was used to execute the task, etc.);
- workflow structure information (inter-task control and data dependencies); and
- compute resource characteristics (CPU speed, available RAM, etc.).

**The WorkflowHub JSON format.** The WorkflowHub project uses a common format for representing collected workflow traces and generated synthetic workflows traces. Workflow simulators and simulation frameworks that support this common format can then use both types of traces interchangeably. This common format uses a JSON specification (publicly available on GitHub [19]), which captures all relevant trace information as listed above. The GitHub repository also provides a Python-based JSON schema validator for verifying the syntax of JSON trace files, as well as their semantics, e.g., whether all files and task dependencies are consistent. Users are encouraged to contribute additional workflow traces for any scientific domain, as long as they conform to the WorkflowHub’s common format.

**Collection of traces.** An integral objective of the WorkflowHub project is to collect and reference open access workflow traces from production workflow systems. Table I summarizes the set of workflow traces currently hosted on WorkflowHub. These traces are from six representative science

domain applications, in which workflows are composed of compute- and/or data-intensive tasks. (Note that although a workflow may be categorized overall as, for example, data-intensive, it may be composed of different kinds of tasks including, e.g., CPU-intensive ones.) We argue that the 101 archived workflow traces form a representative set of small- and large-scale workflow configurations. In addition to consuming/producing large volumes of data processed by thousands of compute tasks, the structures of these workflows are sufficiently complex and heterogeneous to encompass current and emerging large-scale workflow execution models [20].

#### B. Workflow Trace Generator

Workflow execution traces are commonly used to drive experiments for evaluating novel workflow algorithms and systems. It is crucial to run large numbers of such experiments for many different workflow configurations, so as to ensure generality of obtained results. In addition, it is useful to conduct experiments while varying one or more characteristics of the workflow application, so as to study how these characteristics impact workflow execution. For instance, one may wish, for a particular overall workflow structure, to study how the workflow execution scales as the number of tasks increases. And yet, current archives only include traces for limited workflow configurations. And even as efforts are underway, including WorkflowHub, to increase the size of these archives, it is not realistic to expect them to include all relevant workflow configurations for all experimental endeavors. Instead, tools must be provided to generate representative *synthetic* workflow traces. These traces should be generated based on real workflow traces, so as to be representative, while conforming to user-specified characteristics, so as to be useful. The second axis of the WorkflowHub project targets the generation of such realistic synthetic workflow traces with a variety of characteristics.

**The WorkflowHub Python package.** In order to allow users to analyze existing workflow traces and to generate synthetic workflow traces, the WorkflowHub framework provides a collection of tools released as an open source Python package [10], [21]. This package provides several tools for analyzing workflow traces. More specifically, analyses can be performed to produce statistical summaries of workflow performance characteristics, per task type. The package leverages the Python’s SciPy [22] package for performing probability distributions fitting to a series of data to find the best (i.e., minimizes the mean square error) probability distribution that represents the data. In contrast to our previous work [8], which used only two probability distributions for generating workflow performance metrics, the WorkflowHub’s Python package attempts to fit data with 23 probability distributions provided as part of SciPy’s statistics submodule. Fig. 2 shows an example of probability distribution fitting of task runtimes for two task types from different workflow traces, by plotting the cumulative distribution function (CDF) of the data and the best probability distribution found. The outcome of this

TABLE I  
COLLECTION OF WORKFLOW EXECUTION TRACES HOSTED BY WORKFLOWHUB. ALL TRACES WERE OBTAINED USING THE PEGASUS WMS RUNNING ON THE CHAMELEON CLOUD PLATFORM.

Application	Science Domain	Category	# Traces	# Tasks	Runtime and Input/Output Data Sizes Distributions
1000Genome	Bioinformatics	Data-intensive	22	8,844	alpha, chi2, fisk, levy, skewnorm, trapz
Cycles	Agroecosystem	Compute-intensive	24	30,720	alpha, beta, chi, chi2, cosine, fisk, levy, pareto, rdist, skewnorm, triang
Epigenomics	Bioinformatics	Data-intensive	26	15,242	alpha, beta, chi2, fisk, levy, trapz, wald
Montage	Astronomy	Compute-intensive	8	32,606	alpha, beta, chi, chi2, cosine, fisk, levy, pareto, rdist, skewnorm, wald
Seismology	Seismology	Data-intensive	11	6,611	alpha, argus, fisk, levy
SoyKB	Bioinformatics	Data-intensive	10	3,360	argus, dweibull, fisk, gamma, levy, rayleigh, skewnorm, triang, trapz, uniform
6 applications	4 domains	2 categories	101	97,383	18 probability distributions

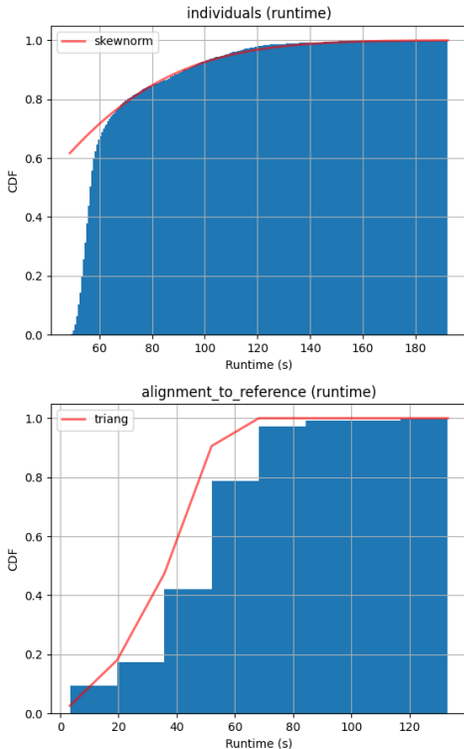


Fig. 2. Example of probability distribution fitting of runtime (in seconds) for *individuals* tasks of the 1000Genome (*top*) and *alignment\_to\_reference* tasks of the SoyKB (*bottom*) workflows.

analysis applied to an entire workflow trace is a summary that includes, for each task type, the best probability distribution fits for runtime, input data size, and output data size. For instance, Table I lists (for each workflow application for which WorkflowHub hosts traces) the probability distributions used for these fits. Listing 1 shows the summary (which is implemented as a Python object) for one particular task type in the 1000Genome workflow application. These summaries can then be used to develop *workflow recipes*, described hereafter.

**Workflow Recipes.** The WorkflowHub Python package also provides a number of workflow “recipes” for generating realistic synthetic workflow traces. Each recipe provides different methods for generating synthetic, yet realistic, workflow traces depending on the properties that define the structure of the

```

1 "individuals": {
2   "runtime": {
3     "min": 48.846,
4     "max": 192.232,
5     "distribution": {
6       "name": "skewnorm",
7       "params": [
8         11115267.652937062,
9         -2.9628504044929433e-05,
10        56.03957070238482
11      ]
12    }
13  },
14  ...
15 }

```

Listing 1. Example of an analysis summary showing the best fit probability distribution for runtime of the *individuals* tasks (1000Genome workflow).

actual workflow. A common method for generating synthetic traces (regardless of the application) is to provide an upper bound for the total number of tasks in the workflow. Although this functionality provides flexibility and control for generating an arbitrary number of synthetic workflows, we have implemented mechanisms that define different lower bound values (for each workflow recipe), so that the workflow structure is guaranteed. The workflow recipe also includes the summaries obtained in the previous steps, so as to generate representative workflow task instances (by sampling task runtimes and input/output data sizes using the probability distribution in the summaries). The current version of the WorkflowHub’s Python package provides recipes for generating synthetic workflows for all 6 applications shown in Table I. Detailed documentation and examples can be found on the project’s website [9] and the online open access package documentation [10].

### C. Workflow Simulator

An alternative to conducting scientific workflow research via real-world experiments is to use simulation. Simulation is used in many computer science domains and can address the limitations of real-world experiments. In particular, real-world experiments are confined to those application and platform configurations that are available to the researcher, and thus typically can only cover a small subset of the relevant scenarios that may be encountered in practice. Furthermore, real-world experiments can be time-, labor-, money-, and energy-intensive, as well as not perfectly reproducible.

The third axis of the WorkflowHub project fosters the use of simulation for scientific workflow research, e.g., the devel-

opment of workflow scheduling and resource provisioning algorithms, the development of workflow management systems, and the evaluation of current and emerging computing platforms for workflow executions. We do not develop simulators as part of the WorkflowHub project. Instead, we catalog open source workflow management systems simulators (such as those developed using the WRENCH framework [23], [24]) that support the WorkflowHub common trace format. In other words, these simulators take as input workflow traces (either from actual workflow executions, or synthetically generated). In the next section, we use one of the simulators cataloged in the WorkflowHub project to quantify the extent to which synthetic traces generated using WorkflowHub tools are representative of real-world traces.

#### IV. CASE STUDY: EVALUATING SYNTHETIC TRACES WITH A SIMULATOR OF A PRODUCTION WMS

In this section, we use a simulator [24] of a state-of-the-art WMS, Pegasus [3], as a case study for evaluation and validation purposes. Pegasus is being used in production to execute workflows for dozens of high-profile applications in a wide range of scientific domains, and is the WMS we used to execute workflows on a cloud environment for the purpose of collecting the traces described in Section III-A.

The simulator is built using WRENCH [23], [24], a framework for implementing simulators of WMSs that are accurate and can run scalably on a single computer, while requiring minimal software development effort. In [24], we have demonstrated that WRENCH achieves these objectives, and provides high simulation accuracy for workflow executions using Pegasus.

##### A. Experimental Scenarios

We consider experimental scenarios defined by particular workflow instances to be executed on particular platforms. To assess the accuracy and scalability of generated synthetic workflows, we have performed real workflow executions with Pegasus and collected raw, time-stamped event traces from these executions. These traces form the ground truth to which we can compare simulated executions.

Actual workflow executions are conducted using the Chameleon Cloud platform, an academic cloud testbed, on which we use homogeneous standard cloud units to run an HTCondor pool with shared file system, a submit node (which runs Pegasus and DAGMan), and a data node placed in the WAN. Each cloud unit consists of a 48-core 2.3GHz processor with 128 GiB of RAM. The bandwidth between the submit node and worker nodes on these instances is about 10Gbps.

Whenever possible, for the experiments conducted in this section, we contrast experimental results obtained with synthetic workflow traces generated with WorkflowHub to results obtained using synthetic workflow traces generated using our previous work [8]. In [24], we have already demonstrated that the simulator framework used in our previous work yields significant discrepancies from actual executions. These

TABLE II  
PARAMETER VALUES USED FOR CALIBRATING THE WORKFLOW GENERATOR FROM OUR PREVIOUS WORK [8].

Application	# Tasks	Runtime Factor	Reference Size
Epigenomics	[125, 263, 405, 559, 713, 803]	0.001	[0.1, 16384]
Montage	[1738, 4846, 7117, 9805]	0.05	[0.1, 113774]

discrepancies mostly stem from the use of a simplistic network simulation model, and from the simulator not capturing relevant details of the system, and thus of the workflow execution. Therefore, to reach fair conclusions regarding the validity of synthetic workflow traces, in this paper we only use the more accurate WRENCH simulator for all experiments. Using this simulator we quantify the extent to which each generated synthetic workflow trace (using our previous work and using WorkflowHub) is representative of the original real-world workflow trace.

The simulator code, details on the calibration procedure, and experimental scenarios used in the rest of this section are all publicly available online [25].

##### B. Evaluating the Accuracy of Synthetic Traces

To evaluate the accuracy of the generated synthetic workflow traces, we consider 2 workflow applications: Montage and Epigenomics. We choose these two applications to allow for comparison with our previous work – the popular generator in [8] can produce synthetic workflow traces for both these applications. For each real-world execution trace of each application, as archived on WorkflowHub, we use WorkflowHub’s Python package for generating a synthetic trace that is similar to the actual execution trace (i.e., we bound the number of tasks to the number of tasks in the actual workflow execution). For comparison purposes, we also generate synthetic traces using the generator from our previous work. To that end, we have calibrated that generator with the parameter values shown in Table II.

**Epigenomics.** Fig. 3 shows simulated empirical cumulative distribution functions (ECDFs) of task submission dates (top) and task completion dates (bottom), for sample runs of real-world and synthetic workflow trace executions of the Epigenomics workflow used on the ILMN dataset. We observe that WorkflowHub’s generated synthetic workflow traces (“synthetic”) yields very similar simulated execution behavior when compared to the real-world execution trace (“real”) – the averaged root mean squared error (RMSE) is 49.02 for task submission and 50.90 for task completion. Recall that small discrepancies in the workflow execution behavior are expected since the workflow task characteristics are sampled from probability distributions. For the synthetic traces generated with the previous generator (“previous”), the averaged RMSEs are 251.87 for task submission and 224.15 for task completion. These substantial discrepancies in execution behavior are mostly due to lack of using accurate probability distributions to model task runtimes, and input and output

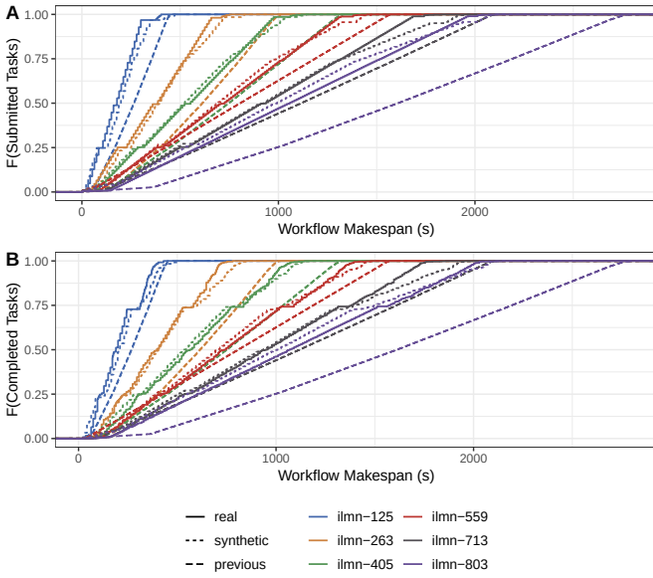


Fig. 3. Empirical cumulative distribution function of task submit times (top) and task completion times (bottom) for sample real-world (“real”) and synthetic (“synthetic” and “previous”) workflow trace executions of Epigenomics using the WRENCH-Pegasus simulator.

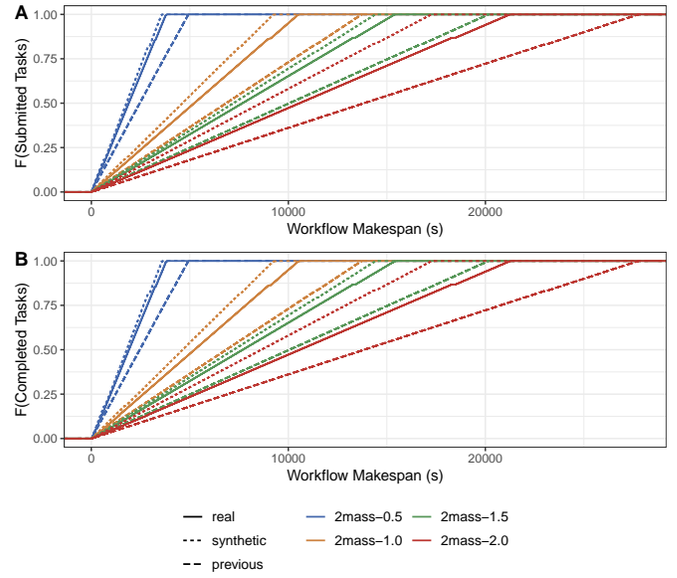


Fig. 4. Empirical cumulative distribution function of task submit times (top) and task completion times (bottom) for sample real-world (“real”) and synthetic (“synthetic” and “previous”) workflow trace executions of Montage using the WRENCH-Pegasus simulator.

file sizes. While WorkflowHub’s generator produces synthetic traces using the best fitted distribution per workflow task type (as described in Section III-A), the previous generator only uses uniform and truncated normal distributions. Moreover, file size generation in that generator uses a random process that leads to inconsistent distribution of file sizes (i.e., the initial reference size seed may shift the density of the distribution toward a lower density area when compared to the empirical distributions obtained from actual workflow executions).

**Montage.** Fig. 4 shows simulated ECDFs for sample runs of real-world and synthetic workflow trace executions of Montage for the 2MASS dataset. Similarly to the Epigenomics results above, WorkflowHub’s generated synthetic workflow traces (“synthetic”) produce workflow execution behaviors close to that of the real-world execution trace (“real”) – average RMSEs are 39.82 for task submission and 46.93 for task completion. When contrasted to synthetic traces generated with the preceding generator (“previous”), RMSEs are 2265.73 and 2253.54 for task submission and completion, respectively. These wide discrepancies are due to the larger number of tasks in the workflow, and Montage’s idiosyncratic workflow structure (extreme fan-in/out pattern). In our previous generator, the generation of file sizes, in particular the larger ones, follows a truncated normal distribution, in which the large variance shortens and broadens the curve, thus generating unbalanced data sizes. As a result, the effect is that tasks in the workflow’s critical path are delayed by data movement operations, causing the workflow makespan to increase. This effect is an artifact of the synthetic trace, and is not seen in real-world executions.

Overall, when using the generated synthetic traces in simulation, we observe large discrepancies when these traces

were generated using our previous work. By contrast, using WorkflowHub’s generation methods produces workflow traces that closely match the real-world execution behaviors (even though some discrepancies necessarily remain due to random sampling effects).

### C. Evaluating the Scaling of Synthetic Traces

In this section, we evaluate the accuracy of the structure of synthetic workflow traces generated based on collected traces at a lower scale, where the scale is the number of workflow tasks. In other words, when generating synthetic workflow traces at various scales, we want to see whether the overall workflow structure is still representative of the workflow application.

We perform experiments using the WRENCH-Pegasus simulator for each workflow application supported by the WorkflowHub project. For each application, we run the simulator for a reference workflow trace (from a real-world execution), and for synthetic traces in which the upper bound limit for the number of tasks are 1K, 5K, 10K, 25K, 50K, and 100K. The goal is to determine whether the simulated execution pattern using these synthetic traces are consistent with that observed when using the reference trace.

Fig. 5 shows ECDFs for task submit times and task completion times for sample runs of these configurations. This figure shows *normalized* makespan values on the horizontal axis. This is to make it possible to use visual inspection for assessing whether the structure of the generated workflows conform with the reference workflow, even though the executions of these workflows have different makespans. In addition, Table III shows the number of tasks that compose the reference workflow trace for each application (“real”) and

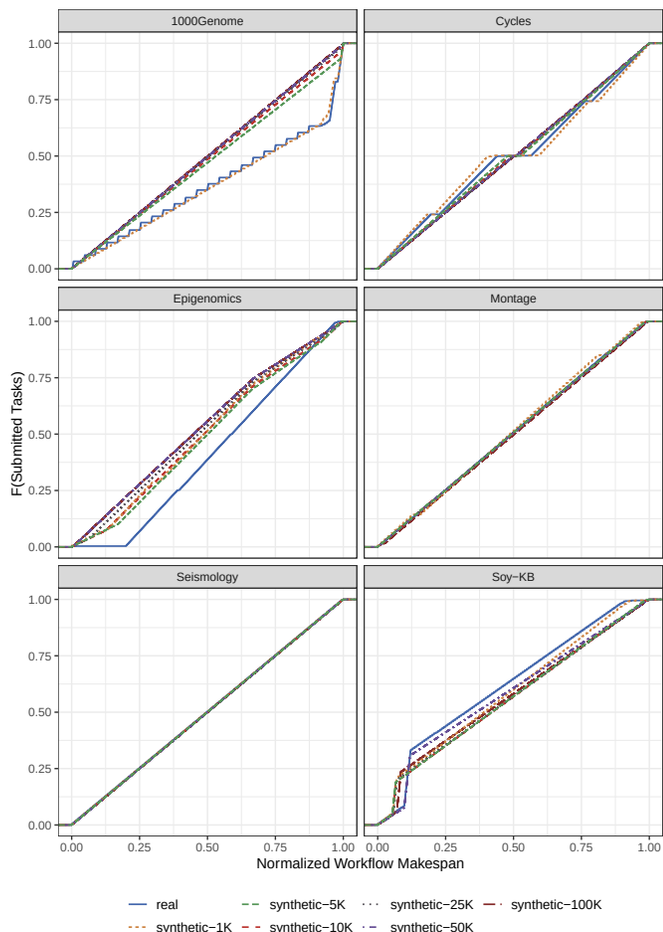


Fig. 5. Empirical cumulative distribution function of task submit times for sample real-world (“real”) and synthetic (“synthetic- $nK$ ”,  $n \in [1, 5, 10, 25, 50, 100]$ ) workflow trace executions using the WRENCH-Pegasus simulator. Workflow makespan has been normalized for comparison purposes.

the RMSE values for the synthetic workflows (“synthetic- $nK$ ”) when contrasted to the reference workflow execution trace.

Overall, generated synthetic traces lead to fairly similar execution patterns when compared to the reference trace. Not surprisingly, the Seismology workflow, due to its simple structure, shows ideal scalability behavior (RMSE values are nearly zero as observed in Table III) – the workflow structure follows a simple *merge* pattern, in which a set of tasks for computing seismogram deconvolutions are followed by a single task that combines all processed fits. Small, yet expected, discrepancies are observed for the Cycles, Montage, and SoyKB workflows. As discussed in Section IV-B, these discrepancies are due to the random generation of workflow task characteristics, which are drawn from various probability distributions.

Visual inspection of Fig. 5 indicates a significant divergence in execution behaviors for the 1000Genome and Epigenomics workflows. For the 1000Genome workflow, the difference between execution behaviors of the reference and *synthetic-1K* workflows is minimum, which is expected as the reference

TABLE III  
ROOT MEAN SQUARE ERRORS (RMSE) FOR LARGE SCALE SYNTHETIC WORKFLOWS. (RMSE VALUES ARE COMPUTED FROM NORMALIZED WORKFLOW MAKESPAN.)

Application	# Tasks (real)	RMSE					
		1K	5K	10K	25K	50K	100K
1000Genome	903	0.03	0.60	0.66	0.69	0.70	0.71
Cycles	331	0.05	0.18	0.26	0.27	0.34	0.36
Epigenomics	125	0.30	0.53	0.49	0.56	0.61	0.61
Montage	1095	0.05	0.06	0.06	0.06	0.07	0.07
Seismology	1738	0.25	0.35	0.16	0.22	0.41	0.48
SoyKB	101	0.05	0.04	0.05	0.05	0.05	0.05
SoyKB	383	0.07	0.22	0.28	0.37	0.37	0.38

workflow is composed of 903 tasks, which is close to the 1,000 tasks in the synthetic workflow (see Table III). However, for synthetic workflows with 5K tasks and higher, the execution pattern does not conform to that with the reference trace (and RMSE values indicate large errors). After carefully inspecting the ECDFs and the workflow structure representation, we noticed that the structure of the 1000Genome workflow, at it scales up, leads to an increase on the number of tasks in the upper levels of the workflow (which are pipelines composed of 6 tasks each), followed by a few tasks that combine the results of these tasks (seen in Fig. 5 as the spike around 90% of the workflow execution time). When increasing the number of tasks for the 1000Genome workflow, this “spike” becomes smoother and fades out in the distribution. A similar structural characteristic is also observed for the Epigenomics workflow. To evaluate this hypothesis, we have also performed runs with an execution trace for the Epigenomics workflow composed of 1095 tasks (also shown in Table III, but not shown in Fig. 5), which produces very similar execution behaviors when compared to the synthetic workflows. We were not able to replicate this experiment for the 1000Genome workflow as we are unable to run the actual application with more than 903 tasks.

Overall, our experiments results show that the tools provided as part of the WorkflowHub framework not only can be used to generate representative synthetic workflow traces (i.e., with workflow structures and task characteristics distributions that resembles those in traces obtained from real-world workflow executions), but can also generate representative workflow traces at larger scales that of available workflow traces. This is crucial for supporting ongoing research that targets large-scale executions of complex scientific applications on emerging platforms.

## V. CONCLUSION

In this paper, we have presented the WorkflowHub project, a community framework for archiving workflow execution traces, analyzing these traces, producing realistic synthetic workflow traces, and simulating workflow executions using all these traces. WorkflowHub provides a collection of resources for developing workflow recipes based on traces collected from the execution of real-world workflow applications. These workflow recipes are then used to produce synthetic, yet

realistic, workflow traces that can enable a variety of novel workflow systems research and development activities. Via a case study using an accurate and scalable simulator of a production WMS, we have demonstrated that WorkflowHub achieves these objectives, and that it favorably compares to a widely used previously developed workflow generator tool. The main finding is that, with WorkflowHub, one can generate representative synthetic workflow traces at various scales in a way it preserves the workflow application’s key features. WorkflowHub is open source and welcomes contributors. It currently provides a collection of 101 traces from actual workflow executions, and can generate synthetic workflows from 6 applications from 4 science domains. Version 0.3 was released in August 2020. We refer the reader to <https://workflowhub.org> for software, documentation, and links to collections of traces and simulators.

A short-term development direction is to use statistical learning methods, such as regression analysis, for automating the process of generating workflow recipes – specifically the description of the workflow structure, i.e., relations between tasks and dependencies. We also intend to provide continuous supported development of novel workflow recipes to broaden the number of science domains in which WorkflowHub can potentially impact research and development efforts. Finally, another future direction is to use synthetic workflows to support the development of simulation-driven pedagogical modules [26], which include targeted activities through which students acquire knowledge by experimenting with various application and platform scenarios in simulation.

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