



# EXECUTING WORKFLOWS

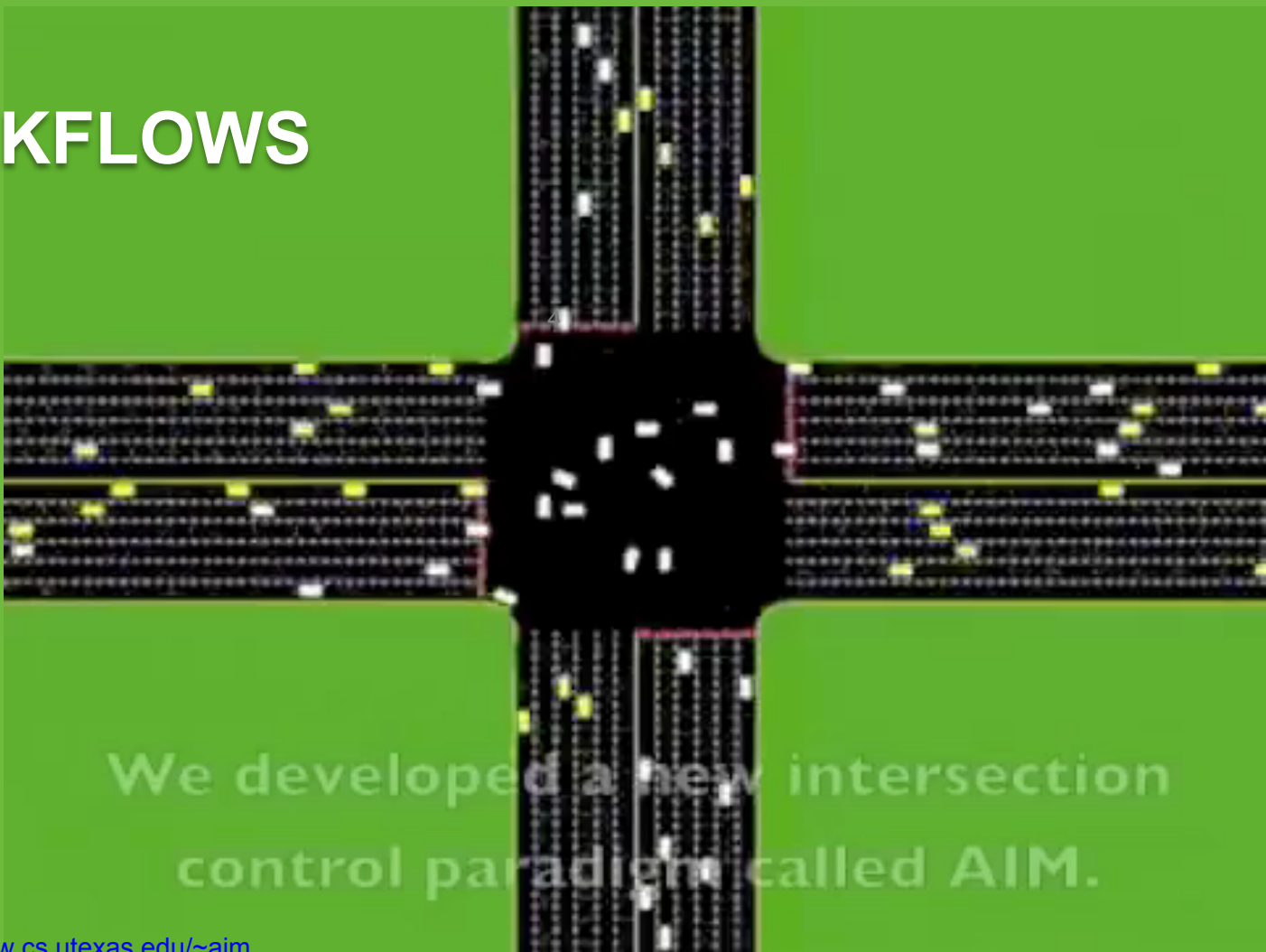
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# WHY SHOULD YOU CARE ABOUT WORKFLOWS?

- You came to the SDL workshop to learn to use large-scale parallel systems for simulation, data, and learning:
  - architectures
  - programming models
  - communication
  - solvers
  - visualization
  - profiling
  - optimizing for node-level performance
  - scaling parallel applications to tens of thousands of nodes
- Once you've written your application and achieved near-peak node-level performance and scalability, what comes next?



# WORKFLOWS



We developed a new intersection control paradigm called AIM.

# ALLOCATION EXAMPLES

- Small number of large jobs running on Mira over course of the year followed by some analysis
  - 40 jobs x 16K nodes \* 16 cores/node \* 12 hours  $\approx$  100M core-hours
  - ~Reasonable to manage manually
- Large number of simulation+analysis jobs running across multiple facilities, requires coordination of jobs submitted to multiple schedulers, large/long data transfers, and interaction with project storage and archival storage
  - Programmatic management would be a clear benefit

Workflows can save you!

...but you might have to code them yourself!

# WHY USE WORKFLOWS?

- Automate job submission
- Simplify computational campaigns
- Increase concurrency by disentangling data dependencies
- Robustness: Improve error handling and recovery (retries)
- Coscheduling of multiple resources
- Systematize data management
- Provenance/Metadata tracking
  - Validation
  - Reuse

# WHAT IS A WORKFLOW?

- It depends who you ask
- Basically a collection of jobs to be run
  - Could be a sequence of individual jobs
  - Could be a sequence of varying numbers of jobs
- Available means of describing and running workflows
  - Script jobs
  - Job dependencies
  - Ensemble jobs
  - In situ
  - Custom workflows
  - Workflow software

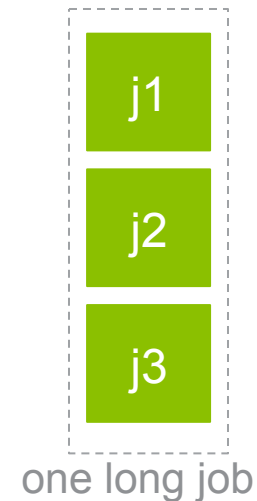
# WORKFLOW VIA SCRIPT JOB

- You can submit a job that executes your application
  - `qsub -q cache-quad -n 512 -t 60 -A yourproject application.exe`
- Alternatively, you can submit a script job that executes multiple applications sequentially
  - `qsub -q cache-quad -n 512 -t 60 -A yourproject script.sh`

## *script.sh*

```
aprun -n 512 -N 1 application.exe  
aprun -n 512 -N 1 application.exe
```

- With this approach, you wait in the queue one time to run your application multiple times
- However, small long jobs tend to stay in the queue longer than small short jobs
  - It may be better to submit individual jobs with dependencies





# WORKFLOW VIA COBALT JOB DEPENDENCIES

- A simple way to achieve a linear workflow is simply to set dependencies between your jobs
  - `qsub -q cache-quad -n 512 -t 60 -A yourproject a.out`  
Job 12345 submitted
  - `qsub -q cache-quad -n 512 -t 60 -A yourproject --dependencies 12345 a.out`
  - `qsub -q cache-quad -n 512 -t 60 -A yourproject --dependencies 12345:12346 a.out`
- Is there an advantage to setting job dependencies?
  - Dependent jobs accumulate score more quickly
- How many jobs can be submitted at once?

j1

j2

j3

multiple  
independent  
short  
jobs

```
qstat -Q
```

Name	Users	Groups	MinTime	MaxTime	MaxRunning	MaxQueued	MaxUserNodes	MaxNodeHours
cache-quad	None	None	None	06:00:00	10	20	None	None
flat-quad	None	None	None	06:00:00	10	20	None	None

- On Theta, max queued is set to 20

# ENSEMBLE JOBS ON THETA

```
for x in {0..100}
do
  # background one aprun call per job
  aprun -n 1 -N 1 myapp.py --args $x &
  sleep 1
done
wait
```

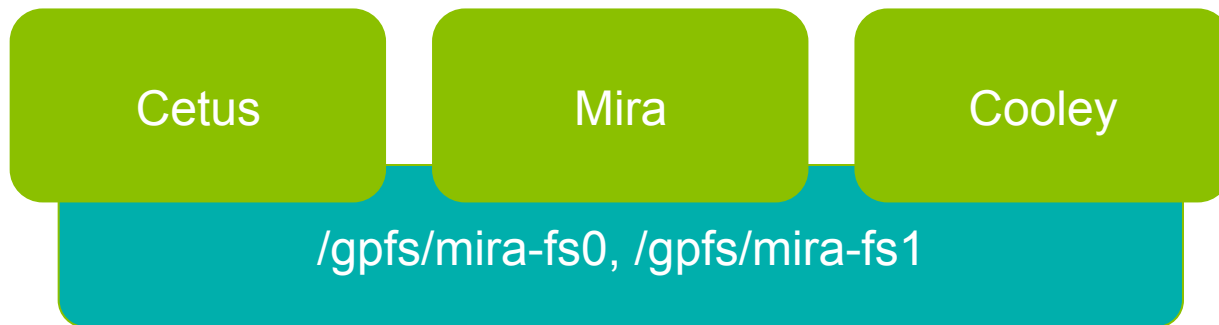
## Note:

- There is a system limitation of 1000 simultaneous apruns per Cobalt script job
- You should include a sleep call between aprun invocations

For more details of running ensemble jobs on Theta, see Paul Rich's slides  
[https://www.alcf.anl.gov/files/rich-ensemble-jobs-2017\\_1.pdf](https://www.alcf.anl.gov/files/rich-ensemble-jobs-2017_1.pdf)

# WORKFLOW ACROSS ALCF SYSTEMS

- How can I set job dependencies across multiple resources (e.g. Mira and Cooley)?
  - ALCF does not provide a means of doing this currently
  - However, Cetus and Cooley mount the Mira filesystems. Analysis jobs run on Cooley without needing to transfer the data.



# SOFTWARE FOR MANAGING WORKFLOWS

This manual effort can be ameliorated by scripting your job submission so that new jobs are automatically submitted as you drop below the `max_queued` limit. Multiple toolkits are available for this purpose.

## **Balsam** (<https://www.alcf.anl.gov/balsam>)

Balsam is an ALCF project for managing jobs and workflows on our systems. Balsam allows users to define campaigns of many jobs with interdependencies and control how they execute on the systems, either by submitting them to the job scheduler, or running them within a single job for higher throughput.

## **Swift** (<http://swift-lang.org>)

Swift provides a C-like language for expressing workflows that consist of command-line invocations, and an engine for managing their execution. For a recent Python implementation of Swift concepts, also see Parsl (<http://parsl-project.org>).

# SOFTWARE FOR MANAGING WORKFLOWS

## **Fireworks** (<https://pythonhosted.org/FireWorks>)

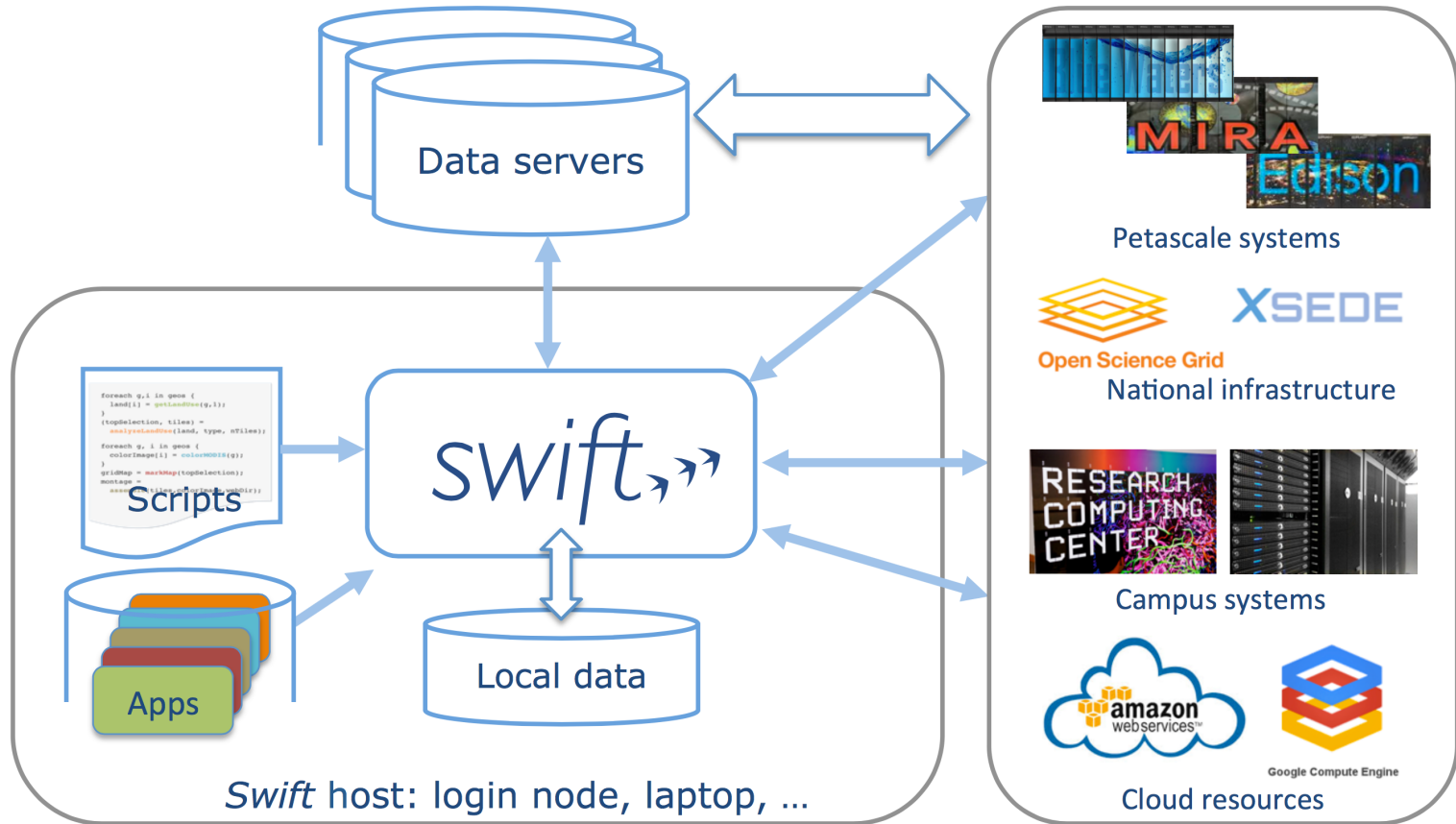
FireWorks is a free, open-source code for defining, managing, and executing workflows. Complex workflows can be defined using Python, JSON, or YAML, are stored using MongoDB, and can be monitored through a built-in web interface. Workflow execution can be automated over arbitrary computing resources, including those that have a queueing system. FireWorks has been used to run millions of workflows encompassing tens of millions of CPU-hours across diverse application areas and in long-term production projects over the span of multiple years.

## **Pegasus/Condor** (<https://pegasus.isi.edu>)

Pegasus bridges the scientific domain and the execution environment by automatically mapping high-level workflow descriptions onto distributed resources. It automatically locates the necessary input data and computational resources necessary for workflow execution. Pegasus enables scientists to construct workflows in abstract terms without worrying about the details of the underlying execution environment or the particulars of the low-level specifications required by the middleware

**Many others...**

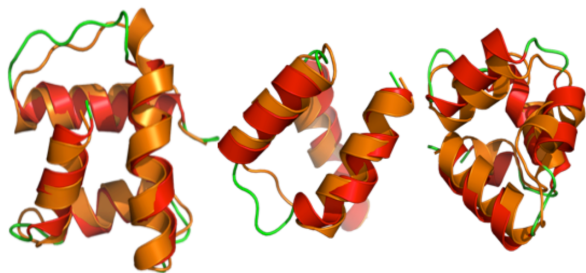
# Swift enables execution of simulation campaigns across multiple HPC and cloud resources



# When do you need HPC workflow?

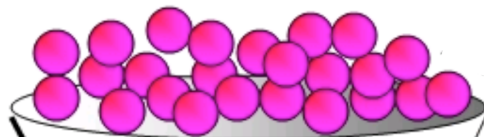
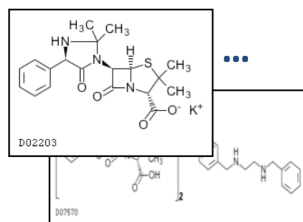
Example application: protein-ligand docking for drug screening

$O(10)$  proteins  
implicated in a disease



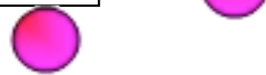
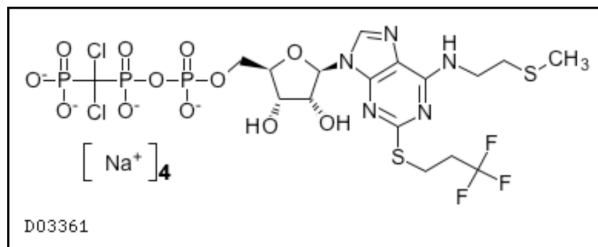
X

$O(100K)$   
drug  
candidates



= 1M  
docking  
tasks...

...then hundreds of  
detailed MD  
models to find  
10-20 fruitful  
candidates for  
wetlab & APS  
crystallography



# Expressing this many task workflow in Swift

*For protein docking workflow:*

```
foreach p, i in proteins {  
    foreach c, j in ligands {  
        (structure[i,j], log[i,j]) =  
            dock(p, c, minRad, maxRad);  
    }  
scatter_plot = analyze(structure)
```

*To run:*

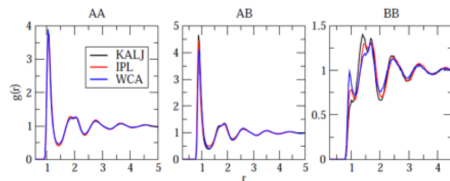
```
swift -site cooley,blues dock.swift
```



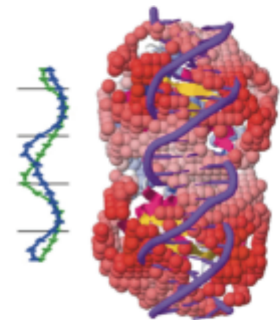
# Large-scale applications using Swift

- A Simulation of super-cooled glass materials
- B Protein and biomolecule structure and interaction
- C Climate model analysis and decision making for global food production & supply
- D Materials science at the Advanced Photon Source
- E Multiscale subsurface flow modeling
- F Modeling of power grid for OE applications

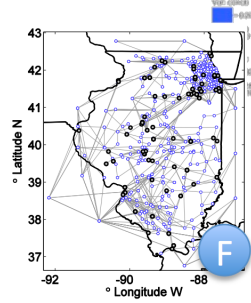
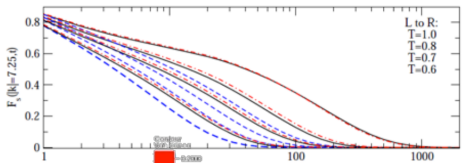
All have published science results obtained using Swift



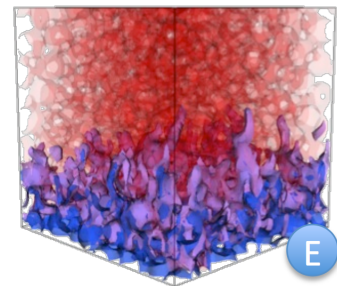
A



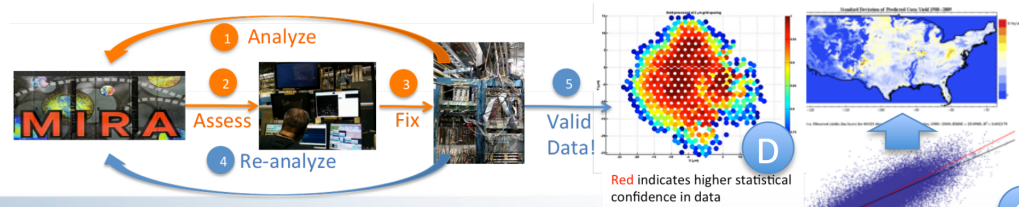
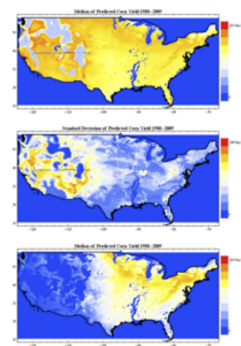
B



F



E



D

C

# BALSAM

We are developing Balsam to manage workflows on systems at ALCF (and elsewhere). What is Balsam?

- Workflow system for managing large campaigns of interdependent jobs (unlimited queue depth)
- Service for managing flow of jobs into scheduler (automated job submission)
- Flexible launcher for high throughput execution by side-stepping the queue (user control)
- Built on Python and Django
  - Powerful Django object-relational mapper (ORM)
  - Support for many databases (sqlite, MySQL, Postgres)
- Handles data transfers for stage-in and stage-out
- Modular interface to job sources, schedulers, and transfer utilities

# BALSAM INSTALLATION ON THETA

```
export PATH=$PATH:$HOME/bin:/opt/intel/python/2017.0.035/intelpython35/bin
conda config --add channels intel
conda create --name balsam_env intelpython3_full python=3
source activate balsam_env
cp /opt/cray/pe/mpt/7.6.0/gni/mpich-intel-abi/16.0/lib/libmpi* ~/.conda/envs/
balsam_env/lib/
export LD_LIBRARY_PATH=~/.conda/envs/balsam_env/lib:$LD_LIBRARY_PATH

git clone git@xgitlab.cels.anl.gov:datascience/balsam.git
cd balsam
pip install -e .
```

# BALSAM USE CASES

Add an application to the database

```
balsam app --name myapp --desc "my application" --exec myapp.py
```

Add a job to the database (options mirror qsub/aprun options)

```
balsam job --name myjob --workflow balsam_tutorial --application myapp \  
--wall-min 60 --num-nodes 1 --ranks-per-node 1
```

Launch the Balsam service to automatically flow jobs from the database to the scheduler (subject to user-defined limits)

```
balsam service
```

Submit Balsam launcher jobs manually to run jobs from the database continuously in the context of a single Cobalt job

```
qsub balsam launcher --consume-all --max-ranks 4
```

# BALSAM COMMAND LINE INTERFACE (CLI)

```
$ balsam ls
```

job_id	name	workflow	application	state
43e4d1be-fc30-4618-91e0-65b100bc18a8	myjob	balsam_tutorial	myapp	CREATED
77d5e32e-8595-4162-ac21-1b86835a6cd0	myjob	balsam_tutorial	myapp	CREATED
ebcdc382-d548-4d1d-8c6a-53f3ac59d64e	myjob	balsam_tutorial	myapp	CREATED
e705d917-b196-4cf4-9b6f-c2ea453e14ae	myjob	balsam_tutorial	myapp	CREATED
35a570d9-2120-42a8-9984-387285827606	myjob	balsam_tutorial	myapp	CREATED
50d8bea1-977a-4028-84ac-022686c7949b	myjob	balsam_tutorial	myapp	CREATED
eb613637-efc2-49c0-9105-bac8cd48fdd8	myjob	balsam_tutorial	myapp	CREATED

```
$ balsam rm jobs --id 43e4d1be-fc30-4618-91e0-65b100bc18a8
```

```
$ balsam --help
```

app	add a new application definition
job	add a new Balsam job
dep	add a dependency between two existing jobs
ls	list jobs, applications, or jobs-by-workflow
modify	alter job or application
rm	remove jobs or applications from the database
qsub	add a one-line bash command or script job
killjob	Kill a job without removing it from the DB
mkchild	Create a child job of a specified job
launcher	Start an instance of the balsam launcher
dbserver	Start/stop database server process

# MORE INFO

- Code available at [xgitlab.cels.anl.gov/datascience/balsam](https://xgitlab.cels.anl.gov/datascience/balsam)
- Documentation
  - Generated docs exist in repo under balsam/docs
  - Online documentation at [balsam.alcf.anl.gov](https://balsam.alcf.anl.gov) (soon)
- Misha Salim will present use of Balsam for hyperparameter optimization on Thursday

## Balsam - HPC Workflow and Edge Service

Balsam is a Python-based service that handles the cumbersome process of running many jobs across one or more HPC resources. It runs on the login nodes, keeping track of all your jobs and submitting them to the local scheduler on your behalf.

### Why do I want this?

Whereas a local batch scheduler like Cobalt runs on behalf of **all users**, with the goals of fair resource sharing and maximizing overall utilization, Balsam runs on **your** behalf, interacting with the scheduler to check for idle resources and sizing jobs to minimize time-to-solution.

You could use Balsam as a drop-in replacement for `qsub`, simply using `balsam qsub` to submit your jobs with absolutely no restrictions. Let Balsam throttle submission to the local queues, package jobs into ensembles for you, and dynamically size these packages to exploit local scheduling policies.

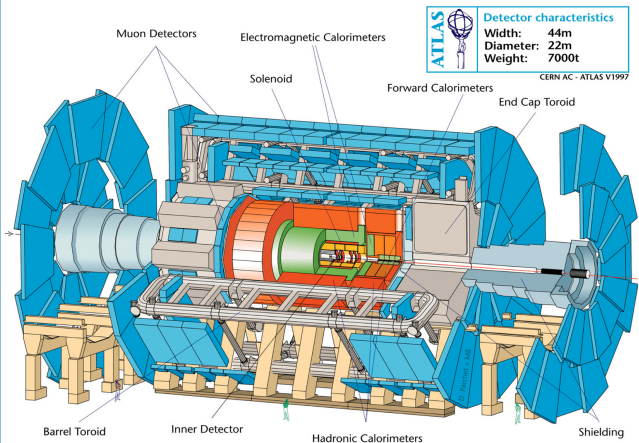
There is **much more** to Balsam, which is a complete service for managing complex workflows and optimized scheduling across multiple HPC resources.

### Quickstart

- [Install Balsam](#)
  - [Prerequisites](#)
  - [Environment](#)
  - [Get Balsam](#)
  - [Quick Tests](#)
- [The Balsam Database](#)
  - [Creating a new Balsam DB](#)
  - [Starting up the Balsam DB Server](#)
  - [Specifying which DB to use](#)
- [Hello World and Testing](#)

# HIGH ENERGY PHYSICS EXAMPLE

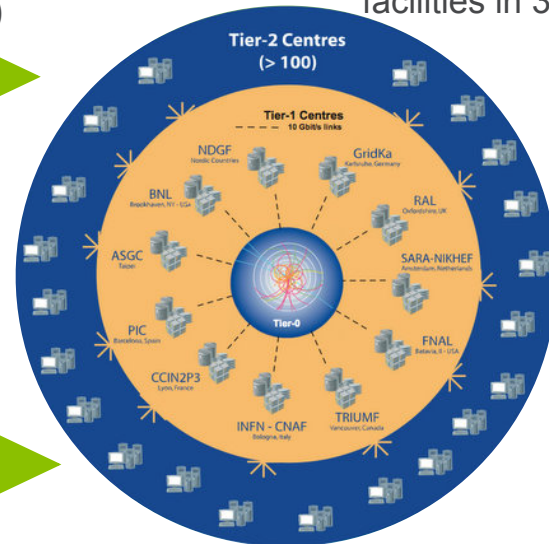
ATLAS detector at LHC



Experimental data  
(>25PB/year)



LHC Computing Grid  
(analysis)  
>170 computing facilities in 36 countries



Simulation data



ALCF



# HIGH ENERGY PHYSICS EXAMPLE

- The ATLAS experiment uses more than 1 billion compute hours per year
- This consists of integration, event generation, showering, simulation, reconstruction, and analysis
- This (very loosely coupled) workflow propagates step-by-step through the ATLAS production system based on requests from users.
- Simulation with Geant4 accounts for 60% of ATLAS's computing. We wanted to leverage Mira to offload some of this computing, and chose to target event generation (not simulation) as a first step. This task required coordination of multiple workflow-like steps:
  - fetching job descriptions from ATLAS (manual process)
  - serial phase-space grid integration on local cluster
  - transfer of data from local cluster to ALCF
  - large-scale event generation on Mira, based on incoming integration grids
  - transfer of data from ALCF to local cluster
  - post-processing of output data to exportable format
  - transfer of data from local cluster to WLCG

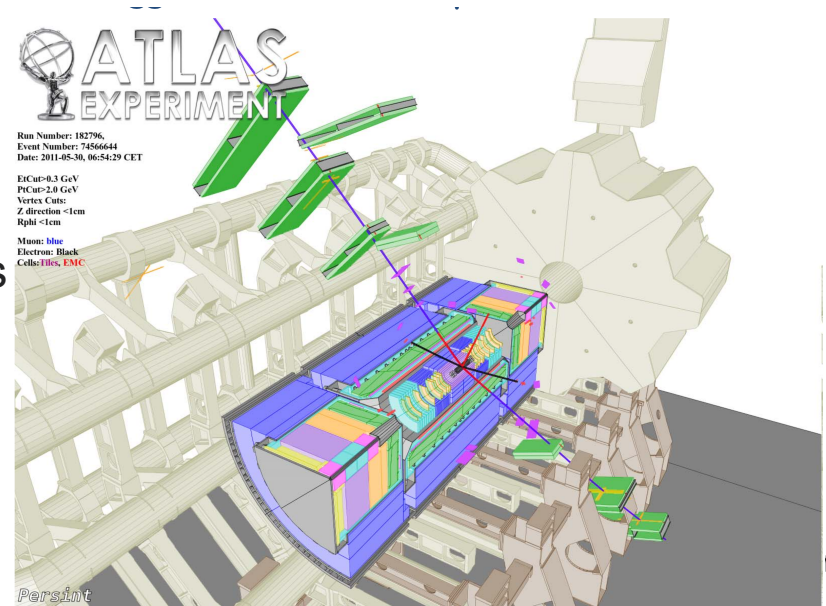


# EXAMPLE: HEP EVENT GENERATION

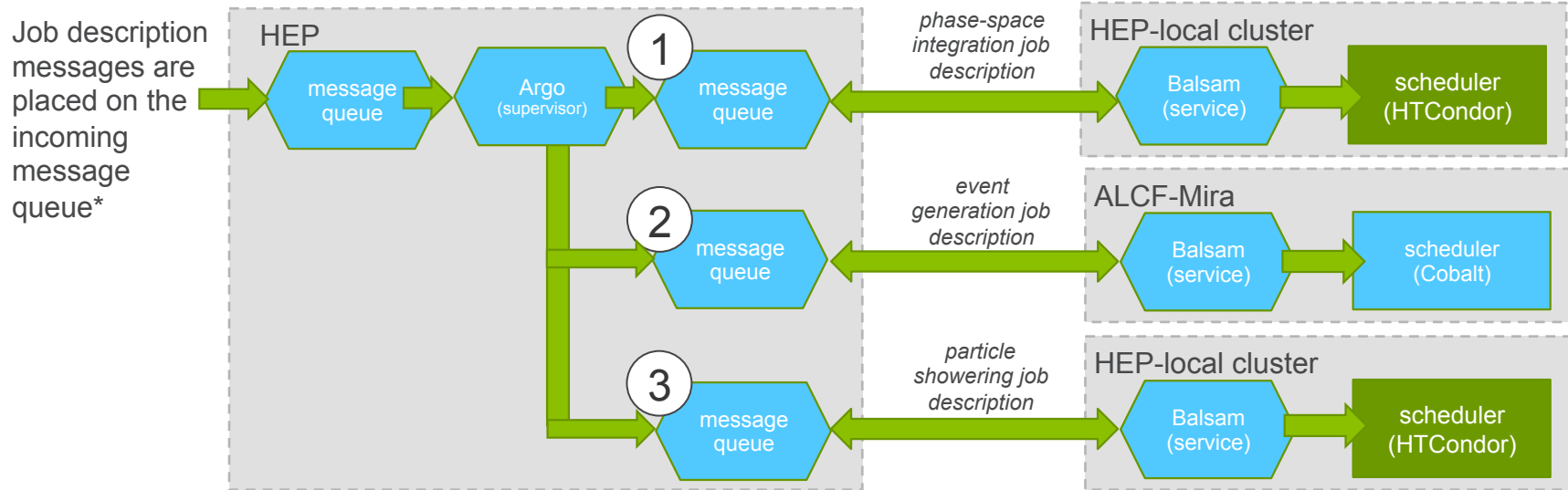
Monte Carlo-based generation of particle collision events such as occur in the ATLAS detector at the Large Hadron Collider, using Alpgen.

Consists of three stages:

1. Generation of phase-space integration grid
2. Generation of weighted events
3. Unweighting of events



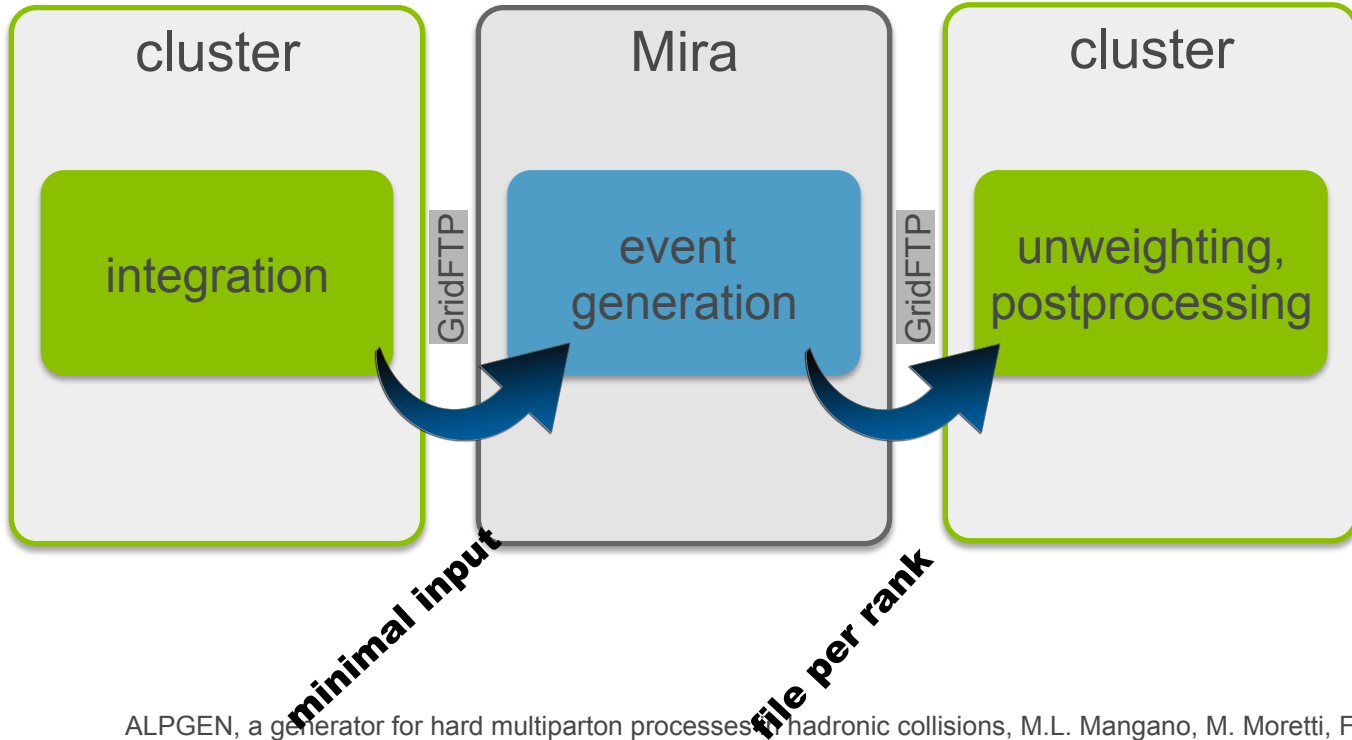
# MULTI-RESOURCE EVENT GENERATION WORKFLOW



Final events are transferred manually to ATLAS; this could be automated in future

\* Note: To date, all jobs have been injected manually; integration with external services (e.g. PanDA) is incomplete

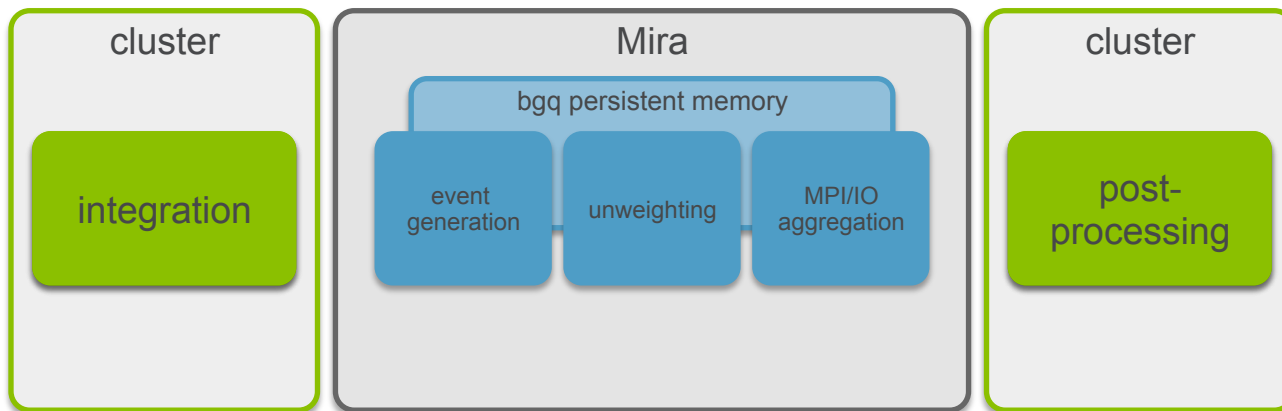
# EVENT GENERATION ON MIRA



ALPGEN, a generator for hard multiparton processes in hadronic collisions, M.L. Mangano, M. Moretti, F. Piccinini, R. Pittau, A. Polosa, JHEP 0307:001,2003

# EVENT GENERATION ON MIRA

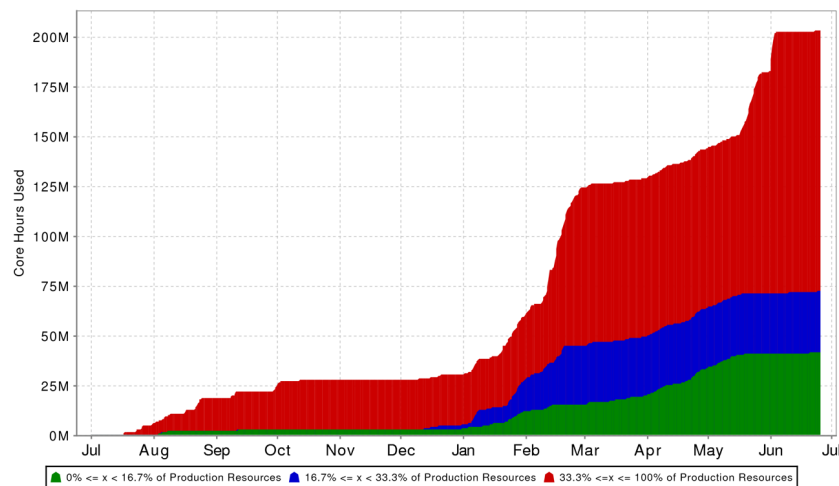
- Combine multiple application invocations in single script job
- Use persistent memory for exchanging data between invocations
- Aggregate data from persistent memory to filesystem



# RESULTS

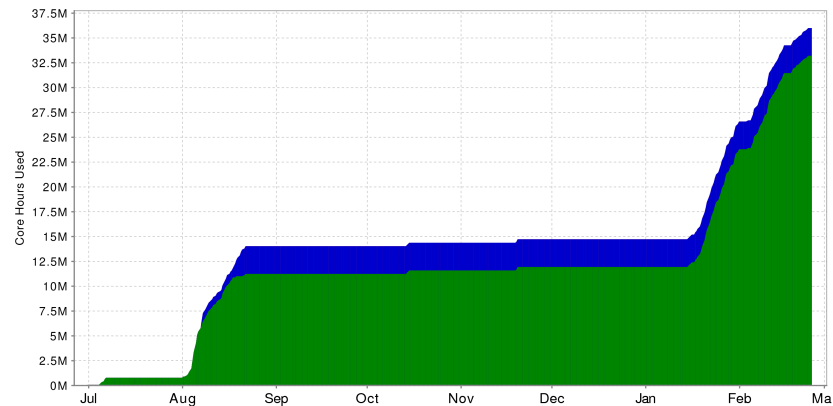
- **100M+ hours used for ATLAS event generation through Balsam**
- Unprecedented rates of event generation (largest event generation jobs on Mira requested >1T events)
- More complex/rare events than could be produced on the Grid
- Mira has become the primary site for ATLAS event generation
- 100 million compute hours have been offloaded from the WLCG to Mira, freeing this time to be used for other purposes
- Events that would have taken years to produce on the Grid were generated within a couple months, accelerating the simulation and analysis pipeline and therefore publications

**EnergyFEC**  
Machine: MIRA  
Total core hours used per category  
2015-07-01 to 2016-06-26

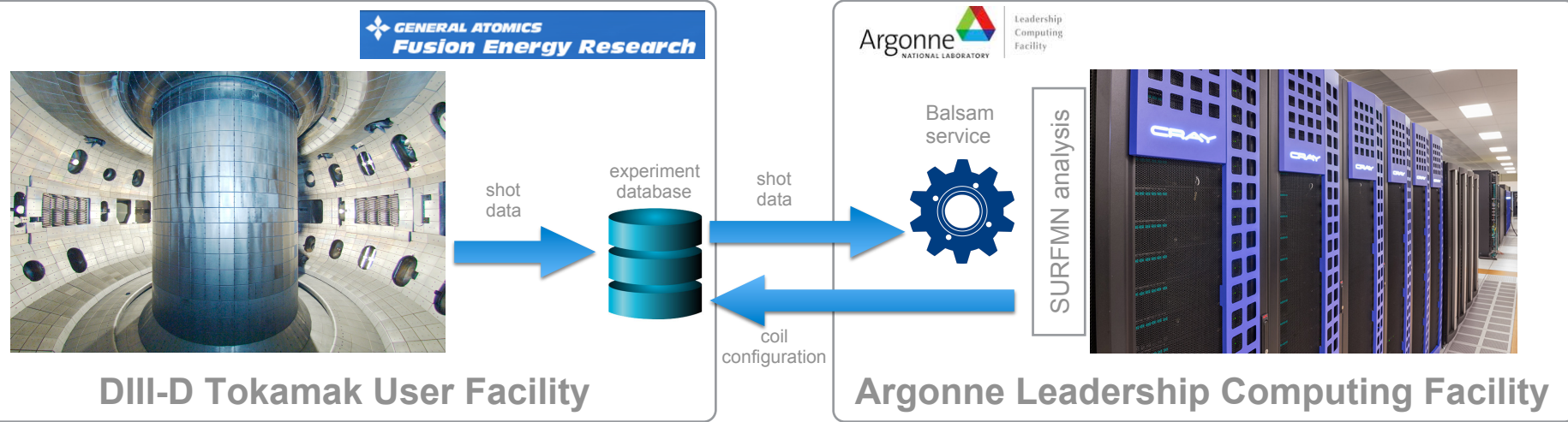


**EnergyFEC\_3 Machine: THETA**

Allocation Core Hours: 40,000,000 Data Dates: 2017-07-01 through 2018-02-25  
Usage Core Hours: 36,051,682 (90.1%) Allocation Dates: 2017-07-01 through 2018-06-30



# AUTOMATIC BETWEEN-SHOT ANALYSIS OF FUSION EXPERIMENT DATA



- ▶ Scientists configure experimental “shots” every 15 minutes
  - A shot is an attempt to magnetically confine high temperature plasma
  - The timing/current of magnetic coils are configured to control the plasma during a disruption to avoid damage to the containing vessel (applicable to DIII-D and future reactors)
  - Analyses indicate how to optimize coil configuration for confinement
- ▶ Each shot triggers an automatic, real-time analysis job at ALCF
- ▶ GA scientists integrate analysis results into configuration for next shot
- ▶ Analysis at ALCF enables more complex analyses ( $128^2$  FFT vs  $32^2$ ) to be completed faster, improving the accuracy of results and allowing analyses to inform every shot instead of every other

## original timeline



## new timeline



*Faster analysis time allows analysis results to be integrated into magnet configuration for subsequent shots. Higher resolution analyses improve configuration accuracy.*

# SUMMARY

- Workflows are inevitable in computational science
- Small-scale workflow management should be easy and is normally managed using scripting
- Large-scale workflow management can benefit from a workflow system
  - In the transition from small to large workflows, you might build your own workflow system
- The Data Science group at ALCF can help scale your workflows on our systems

