

1 Intro

Creating many GPT simulations on a single desktop PC can be tedious and time consuming. The advantage of running GPT on the farm is that you can create many GPT simulations in parallel instead of in series, which can save countless hours (assuming you know what you are doing). The documentation below of basic instructions on how to run GPT simulations on the farm is the culmination of countless hours and countless Advil and countless questions asked to countless people and countless times I've screwed up and had to start all over. I assume that you are either a grad student or an employee looking to use the farm to speed up your GPT simulations. I also assume (and hope) that you have a decent amount of experience with GPT and some experience with Unix shells to understand the example I give below. I hope the below documentation proves useful to you and your research...at least it's much better than the online documentation.

2 Step 1: Are you sure you know what you're doing?

There are countless ways to run on the farm and not get what you want. There are even more ways to have some interesting syntax error that prevents GPT from even running in the first place. Thus, be sure that, at the very least, one of your GPT simulations runs to completion on one of the JLab machines (such as jlab1), not just your desktop PC. Try running a simulation on jlab1. If you get any errors at this point, it's better to deal with them here than on the farm. If it works smoothly, you're most likely good to go.....we'll see.

Note that in order to run GPT on jlab1, you'll need to know how to use GPT on a Unix machine (i.e. with a terminal interface). If you don't have access to a JLab Linux workstation (like an accelerator computer) and mainly use a Windows PC like I do, you can try logging onto jlab1 using PuTTY (see <https://cc.jlab.org/windows/remotedesktop>). You'll only need to do "Stage 1". Once you've logged into login.jlab.org, you can type "ssh jlab1" and enter your CUE password to log onto jlab1. Be sure that *all* of your simulation files (i.e. your input file and fieldmap files) are in an easily accessible place on your J:\ drive on JLab Windows machines (or /u/home/<username>/ directory on Linux machines). You can then try running GPT on jlab1 using your simulation files.

3 Step 2: Are you sure you're allowed?

Once you have a working GPT simulation, the first step to running on the farm is to gain access to the farm. If you don't already have access, you'll need to ask your supervisor about getting access to the farm. In order to use the farm, you need to be a part of one of the groups that is "allowed" to use the farm. I'm in the group called "inj_group", which is an allowed group for using the farm. Note that this has nothing to do with whether or not you are *allowed* to use the farm (Of course you are!), it's to make sure that the farm is only used by those who are authorized to use it (Of course you are, too!). To see what groups you are in, type "groups <your CUE username>" in any JLab terminal.

4 Step 3: Are you even ready?

Once you get access to the farm, you'll need to create an Auger command (batch) file. Auger is the software that manages batch files that are sent to the JLab farm computers to run your jobs. Basically what you'll do is to create a batch file where you'll provide info about your project and what kind of jobs you'd like to run on the farm. Auger will then take that info and send out your jobs to the farm computers (one job per computer). The jobs will be added to the queue with a certain priority (I wouldn't worry about this if you're just starting out. If I were you, I'd be ecstatic if your simulations successfully run on the farm period, regardless of when they actually run.) Once your simulations have finished (successful or not), Auger will then copy the resulting data file(s) back to your directory before deleting them on the farm computer(s).

There are two formats of Auger batch files that you can choose from: one of which you can use and one of which you're gonna use. I'll show how to create the latter below:

4.1 Creating an Auger XML batch file

The xml file consists of two main parts: one for global specifications and one for job specific specifications. To illustrate what each of these sections are and to understand the syntax, I'll show you an example xml batch file below:

```
<Request>
<Email email="yoskowij@jlab.org" request="true" job="true"/>
<Project name="inj_group"/>
<Track name="test"/>
<Name name="GPT_testrun"/>
<TimeLimit time="10"/>

<Command>
use gpt
setenv OMP_WAIT_POLICY PASSIVE
cp /u/group/inj_group/yoskowij/GPT_Farm_Files/* .
gpt -o filename.gdf quads.in eN=${nParticles}
</Command>

<Job>
<Variable name="nParticles" value="100"/>
<Output src="filename.gdf" dest="/u/group/inj_group/yoskowij/GPT_Farm_Files/result_100.gdf"/>
</Job>
<Job>
<Variable name="nParticles" value="200"/>
<Output src="filename.gdf" dest="/u/group/inj_group/yoskowij/GPT_Farm_Files/result_200.gdf"/>
</Job>

</Request>
```

4.1.1 Global Specifications

Everything above `<Command>` are global specifications. They apply to *all* jobs sent to the farm. Some notes about each line:

- The `<Email>` command will tell Auger to email you when the job is completed and tell you whether or not it was successful. It emails the address specified in quotes.
- The `<Project>` command specifies what project your jobs are a part of, assuming it's an allowed project. You can input your project within the quotes. This is a required command.
- The `<Track>` command specifies the type of jobs you are running (see https://scicomp.jlab.org/docs/batch_job_tracks). If you are just starting out, I'd recommend using either the "test" or "debug" tracks. Once you know that your jobs will run smoothly on the farm, you can choose one of the other tracks. This is a required command.
- The `<Name>` command gives your jobs a name so that you can refer to these jobs after they are completed. You can name your jobs whatever you want, so long as you can refer to them easily. This is a required command.
- The `<TimeLimit>` command, as its name suggests, gives a time limit to your jobs. This is useful, as you don't want you jobs to hang on the farm and run forever. You can specify a value and its unit. If you don't specify a unit, the default is minutes. For example, to specify 30 minutes, you can use any of the following: `<TimeLimit time="30"/>`, `<TimeLimit time="30" unit="minutes"/>`, `<TimeLimit time="0.5" unit="hours"/>`.
- I didn't specify it here, but another useful command is `<Memory>` where you can specify the memory usage limit of your jobs. If you're just starting out, this should not be a problem, but it may become a concern for larger simulations. To see if you should be concerned about memory usage, try creating your GPT simulation on a Windows PC and use

Task Manager to see how much memory GPT uses. From there, you can choose a reasonable memory limit for your simulations. You obviously don't want to use too much memory, as it would be unfair to everyone else using the farm.

For more info on the various specifications, refer to https://scicomp.jlab.org/docs/desc_xml_tags.

4.1.2 Global Commands

Between `<Command>` and `</Command>` are the commands you give to each JLab farm computer, just as you would in Terminal. The first line tells the farm computer you'd like to run GPT (specifically it adds the license number to your environment variables so that you have "permission" to run GPT). You are required to have at least one command in your xml file in order to run your job (obviously...).

The second line sets an irrelevant environment variable to passive. If you don't do this, you'll get some warning along the lines of "Warning: Environment variable OMP_WAIT_POLICY=PASSIVE not set," which is strange because the default value *is* passive! This environment variable luckily won't affect your simulations...I just want to reduce the chance that an error, no matter how trivial, screws up my simulations.

The third line copies your simulation files onto each farm computer that you're using (Remember that test run on jlab11 from Step 1 above?) The directory specified in this command is where you have all of your GPT simulation files saved (input file, fieldmap files, etc.). The "*" means copy *all* simulation files in this folder and the "." after it means "copy it here", "here" meaning to the farm computer. If you do not do this, the farm computer will have no files to run your simulations on and you will receive a blank data file (This happened to me many times. Don't be like me!). Note that these files are deleted after your run, so you have to use this command every time you submit your job.

The fourth line is the familiar GPT command line used to run a GPT simulation with a few changes. I'll explain more below:

4.1.3 Job Specifications

The next few lines specify each job that you'd like to run on the farm. In this example, I'm running the Quads simulation from the GPT tutorial for two different numbers of electrons (in this case, 100 electrons and 200 electrons). Each job is specified in the xml file between the two tags `<Job>` and `</Job>`. Any commands that you put in between these two tags gives specifications *for that job only*. Each job has two job-specific commands.

The `<Variable>` command allows you to specify a variable that can be used elsewhere. Remember that GPT allows you to specify a value for a variable used in your input file in the GPT command line. Thus, I can have the first job run the quad input file for 100 electrons and the second one run the input file for 200 electrons. Each job goes to a different farm, which is how you can run many jobs at once. In the quads.in file, I've specified the number of electrons used in the "setparticles" command as "eN". In the command line, I'm setting "eN" to whatever I specify in the variable command within each job. In each variable command, you give your variable a name and a value. It doesn't matter what name you give it. In my case, I'm calling the variable "nParticles" (I'd suggest giving it an easy name to remember what the variable is used for). Once you create the variable, you can reference it elsewhere by putting it between `${<variablenamehere>}`, as I've done in the GPT command line. What this clever use of `<Variable>` does is to have one job run quads.in for 100 electrons and the other for 200 electrons, even though they both use the same input file (which is what I want).

Each job also has an `<Output>` command that allows you to name the resulting GDF data file and give it a destination (outside of the farm computer). Note that in the GPT command line, I've specified the output file as "filename.gdf". This is really a dummy name that will be changed by the `<Output>` commands. Each output command takes "filename.gdf" and puts it in the specified directory with the specified name. What this means is that even though the GPT command line tells each farm computer to call its subsequent GDF file the same name (in this case filename.gdf), these GDF datafiles get returned to me by Auger under the names that I specify in the `<Output>` command, which is how I can distinguish them. Thus, it doesn't really matter what you call the result file in the GPT command line.

Of course, you can run as many jobs as you'd like/need by including more jobs in the xml file. If the number of jobs is on the order of, say, more than 10, I'd recommend using `ForEach`, which will use a `For` loop to create many jobs (see https://scicomp.jlab.org/docs/xml_command_file for more info on this). Obviously I wouldn't start here, but once you get the hang of running simple jobs on the farm, you can try using `ForEach` to simplify your xml file instead of explicitly writing many job commands.

One last note: make sure your job specifications and global specifications are within the `<Request>` and `</Request>` tags. This is important, as your xml file won't get sent to the farm without it.

5 Step 4: Well, we'll see if you're ready

Once you've created your xml file, put it in the same place as your simulation files. Now you are ready to test run your simulations on the farm. First, log onto any JLab computer (either using a terminal on a JLab Linux workstation or using PuTTY on Windows machines). If you use PuTTY, you'll start by logging on to login.jlab.org with your JLab CUE username and password. Once you are logged into a JLab computer, type

```
ssh ifarm.jlab.org
```

and enter your password (again!). This will log you into one of the two interactive nodes: either ifarm1401 or ifarm1402 (it doesn't matter which one you use). You'll have access to all network drives that you would normally have on a Windows machine, except that they'll have different names (see <https://cc.jlab.org/desktopsupport> for more info). If this is your first time running on the farm, you need to get a network certificate by typing in:

```
/site/bin/jcert -create
```

From here, if you haven't already done so, go to your folder on your project's directory here:

```
/u/group/<projectname>/<yourname>/
```

Create a folder for your farm simulation files and copy them there along with the xml file you've created. Once you've double checked your xml file for typos, you can submit your xml file to the farm by typing in

```
jsub -xml <filename>
```

Your xml file will then be parsed. If successful, it will return your job number and your batch will be submitted to the farm. If there are any syntax errors in your xml file, it will let you know then. What it will **NOT** do is tell you any errors or warnings that GPT gives (like it would in the output window of your GPT batch file when run on Windows or within the terminal environment on Linux machines or PuTTY). Thus, it is very important that you are absolutely sure that your GPT program will run correctly and without any non-trivial errors or warnings.

6 Step 5: Thought so...

If you've successfully created an xml file, sent multiple jobs to the farm and got back non-empty datafiles, then congrats! You've done it! You are ready to use the farm. From here, I'd recommend reading https://scicomp.jlab.org/docs/desc_xml_tags to help you customize your xml file. If you can submit a successful job to the farm, you should be able to figure out the rest to meet your needs. Good luck!