### Getting Started with GCHP v11-02c

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### Overview

- 1) What is GCHP and why use it?
- 2) Common Misconceptions
- 3) Useful Tips
- 4) Basic Tutorial
- 5) Introduction to GCHP Source Code
- 6) Resources

### What is GCHP and why use it?

- GCHP features the same science as GEOS-Chem using the standard "classic" capability except:
  - 1. It operates on a cubed-sphere grid
  - 2. It is parallelized using a message-passing interface (MPI)
- GCHP improves upon GCC by:
  - Enabling more accurate transport
  - Providing efficient scaling making finer resolution global simulations possible

### **Common Misconceptions about GCHP**

- I need a high performance compute cluster for GCHP
  - Not true! You can run GCHP on as little as one machine with 6 cores. We are working towards eventually being able to run it on a single core.
- I can only perform high-resolution runs with GCHP
  - Not true! GCHP can run at c24 resolution which is the cubed-sphere equivalent of 4°x5°.
- I need met fields at the same resolution as my run
  - Not true! You may use GCHP with 2°x2.5° meteorology for up to at least c180 (0.5° res), although we recommend keeping the met resolution to no more than twice your run resolution equivalent to ensure quality output. If the met wind fields are too coarse relative to your internal resolution then polar divergence will occur.

### Useful Tip #1: Grid Resolutions

- Cubed-sphere resolution "cN" means each of the six faces are divided into N x N segments.
- An easy rule-of-thumb for resolution mapping is to divide 90 by N to determine the approximate lat-lon degree resolution.

Standard lat-lon resolution	Approximate CS equivalent(s)
4° x 5°	c24
2° x 2.5°	c48
1° x 1.25°	c90
0.5° x 0.625° <sup>1</sup>	c180
0.25° x 0.3125° <sup>2</sup>	c360
0.125° x 0.15625°	c720 <sup>3</sup>

- <sup>1</sup> Native resolution of MERRA-2 product from GMAO
- <sup>2</sup> Native resolution of GEOS-FP product from GMAO
- <sup>3</sup> Native cubed-sphere resolution of GEOS-5

### Useful Tip #2: Resource Allocation

- Number of nodes and number of faces are independent
- Number of cores and number of faces are NOT independent
- Total number of cores must be divisible by 6!
- How does it work?
  - Each NxN face is divided into NX x NY/6 segments, each comprised of approximately N/NX x N\*6/NY cubed-sphere grid cells.
  - Each segment is processed by a single core
  - NX \* NY must therefore equal the total number of cores
  - NX \* NY/6 would ideally be square to minimize required communication between cores
- NX and NY are manually set in config file GCHP.rc but are overwritten by NX and NY of your choosing in utility script runConfig.rc. Setting them will soon be automatic.
- More on this topic later in the presentation.

### **GCHP** Tutorial

- 1. Downloading Source Code
- 2. Create a Run Directory
- 3. Set Link to Source Code
- 4. Load Environment
- 5. Compile GCHP
- 6. Configure Run Settings
- 7. Run a Simulation
- 8. Analyze Output Data
- 9. Reuse a Run Directory

### Step 1: Download Source Code

- You need two repositories for GCHP:
  - 1. GEOS-Chem "classic" (GCC) code as your main directory
  - 2. GCHP code as a subdirectory within it
- Use the GC and GCHP master branches and checkout the version tag for v11-02c

```
git clone -b master https://bitbucket.org/gcst/gc_bleeding_edge Code.v11-02c_gchp
```

```
cd Code.v11-02c_gchp
```

git checkout tags/v11-02c

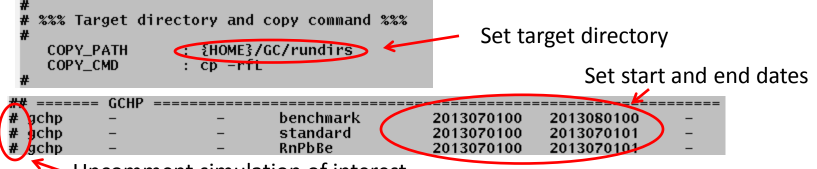
```
git clone -b master https://bitbucket.org/gcst/gchp GCHP
```

```
cd GCHP
```

```
git checkout tags/v11-02c
```

### Step 2: Create a Run Directory

- Download a GCHP run directory from GEOS-Chem Unit Tester
- Use the UT\_Bleeding\_Edge master branch
  - If you have the repository already:
    - git checkout master
    - git pull
    - git checkout tags/v11-02c
  - If you do not have the repository:
    - git clone -b master https:/bitbucket.org/gcst/ut\_bleeding\_edge UT
- To create run directory, modify UT/perl/CopyRunDirs.input:



Uncomment simulation of interest

- Like GCC, each simulation has a different run directory
- Unlike GCC, simulation resolution is configurable at run-time

### Available GCHP Run Directories

- Like GCC, each simulation has a different run directory
- Unlike GCC, simulation resolution is configurable at run-time
- Run directories available in v11-02c:
  - 1. Standard simulation
    - Use for full chemistry science simulations
    - Includes symbolic links to restart files for four resolutions: c24, c48, c90, c180
    - Restart files incorporate UCX initial mixing ratios applied internally in GCC
  - 2. Rn-Pb-Be7 simulation
    - Use for Rn-Pb-Be7 and passive tracer simulations
    - Includes symbolic links to restart files for four resolutions: c24, c48, c90, c180
  - 3. Benchmark simulation
    - Used for benchmarking simulations
    - Like the new GCC benchmark simulation, both simple and complex SOA are on
    - Includes symbolic links to c24 restart file only
    - Restart files does NOT incorporate UCX initial mixing ratios applied internally in GCC

### GCHP Standard Run Directory: Contents

- Configuration files
- Chemistry \*.dat files
- Sample environment files subdirectory
- Sample run scripts subdirectory
- Utility bash scripts

- Output data subdirectory
- README
- Makefile
- Symbolic links to:
  - Restart files
  - Input data directories
  - Cube sphere <-> lat-lon regridding files (tile files)

OD ~/gchp_standard \$ 1s			
bashrcSamples/	h2so4.dat	MetDir@	
brc.dat	HEMCO_Config.rc	org.dat	
build.sh*	HISTORY.rc	OutputDir/	
CAP.rc	<pre>initial_GEOSChem_rst.c180_standard.nc@</pre>	README	
ChemDataDir@	initial_GEOSChem_rst.c24_standard.nc@	runConfig.sh*	
dust.dat	initial_GEOSChem_rst.c48_standard.nc@	runScriptSamples/	
ExtData.rc	initial_GEOSChem_rst.c90_standard.nc@	setCodeDir*	
FJX_j2j.dat	input.geos	so4.dat	
FJX_spec.dat	input.nml	soot.dat	
fvcore_layout.rc	jv_spec_mie.dat	ssa.dat	
GCHP.rc	MainDataDir@	ssc.dat	
gchp.run*	Makefile*	TileFiles@	

# GCHP Run Directory: Environment Setup Files

 Sample environment setup files are stored in the bashrcSamples subdirectory

> OD ~/gchp\_standard \$ cd bashrcSamples/ OD ~/gchp\_standard/bashrcSamples \$ ls gchp.gfortran\_MVAPICH2.bashrc gchp.ifort13\_openmpi\_glooscap.bashrc gchp.ifort15\_mvapich2\_odyssey.bashrc

- The sample files provided are custom for Harvard and Dalhousie Universities
- For your own system you may use these files as examples to build your own
- Please note that the gfortran compiler and the MVAPICH2 and OpenMPI implementations are all open-source
- Contact the GCST if you and your system administrators get stuck

# GCHP Run Directory: Config Files

- GCHP Specific:
  - Cap.rc
    - start/end dates, and more
  - ExtData.rc
    - external data information
  - fvcore\_layout.rc
    - transport-related settings
  - GCHP.rc
    - general settings
  - HISTORY.rc
    - output data settings

- Same as GCC:
  - input.geos
  - HEMCO\_Config.rc

#### • WARNINGS:

- Not all fields in input.geos and HEMCO\_Config.rc are used
- Some settings must be set in multiple files (use runConfig.sh)

	OD ~/gchp_standar	d \$ ls	
	bashrcSamples/	h2so4_dat	MetDir@
	brc.dat	HEMCO_Config.re	org.dat
	build.sh*	HISTORY.rc	OutputDir/
$\boldsymbol{<}$	CAP.rc	<pre>initial_GEOSChem_rst.c180_standard.nc@</pre>	README
	ChemDataDir@	<pre>initial_GEOSChem_rst.c24_standard.nc@</pre>	runConfig.sh*
	duot.dat	initial_GEOSChem_rst.c48_standard.nc@	runScriptSamples/
$\subset$	ExtData.rc	<pre>initial_GEOSChem_rst.c90_standard.nc@</pre>	setCodeDir*
	FJX_j2j.dat	input.geos	so4.dat
	FJX_spec_dat	input.nml	soot.dat
<	fvcore_layout.rc	jv_spec_mie.dat	ssa.dat
	GCHP.rc	MainDataDir@	ssc.dat
	gcnp.run*	Makefile*	TileFiles@

# GCHP Run Directory: Run Scripts and Output Data

#### OutputDir/

- All GCHP output data configured in HISTORY.rc are saved here
- Restart file is not saved here
- Do not remove or rename!
   GCHP will hang without an error message

- runScriptSamples/
  - For Odyssey (Harvard):
    - GCHP\_slurm.run
  - For Glooscap (Dalhousie):
    - GCHP\_gridengine.run
  - For other Systems
    - Use these as examples to build your own

OD ~/gchp_standard \$ ls			
bashrcSamples/	h2so4.dat	MetDir@	
brc.dat	HEMCO_Config.rc	org.dai	
build.sh*	HISTORY.rc	OutputDir/	
CAP.rc	initial_GEOSChem_rst.c180_standard.nc@	README	
ChemDataDir@	initial_GEOSChem_rst.c24_standard.nc@	runConfig.sh*	
dust.dat	initial_GEOSChem_rst.c48_standard.nc@	runScriptSamples/	
ExtData.rc	initial_GEOSChem_rst.c90_standard.nc@	SetCodeDir*	
FJX_j2j.dat	input.geos	so4.dat	
FJX_spec.dat	input.nml	soot.dat	
fvcore_layout.rc	jv_spec_mie.dat	ssa.dat	
GCHP.rc	MainDataDir@	ssc.dat	
gchp.run*	Makefile*	TileFiles@	

# GCHP Run Directory: Utility Scripts

#### • setCodeDir

- creates symbolic link to source code path
- Pass full path (without links) as an argument

### • build.sh

- cleans and compiles code
- executed in Makefile

#### runConfig.sh

- single location to update common run settings
- overwrites config files
- executed in run scripts

> 90% of GCHP errors are due to incorrect or inconsistent config file settings. Use bash script runConfig.sh to avoid common errors.

OD ~/gchp_standar	d \$ ls	
bashrcSamples/	h2so4.dat	MetDir@
brc.dat	HEMCO_Config.rc	org.dat
build.sh*	HISTORY.rc	OutputDir/
CAP.rc	<pre>initial_GEOSChem_rst.c180_standard.nc@</pre>	READILE
ChemDataDir@	initial_GEOSChem_rst.c24_standard.nc@ 🤇	runConfig.sh*
dust.dat	initial_GEOSChem_rst.c48_standard.nc0	runderig Samples/
ExtData.rc	initial_GEOSChem_rst.c90_standard.nc@	setCodeDir*
FJX_j2j.dat	input.geos	304.d3t
FJX_spec.dat	input.nml	soot.dat
fvcore_layout.rc	jv_spec_mie.dat	ssa.dat
GCHP.rc	MainDataDir@	ssc.dat
gchp.run*	Makefile*	TileFiles@

### Step 3: Set Link to Source Code

- GCHP uses a symbolic link to source code called CodeDir
- Run bash shell script **setCodeDir** to set symbolic link:

OD ~/gchp\_standard \$ ./setCodeDir /n/homeO8/elundgren/Code.v11-O2c\_gchp/ CodeDir: symbolic link to `/n/homeO8/elundgren/Code.v11-O2c\_gchp/'

- Things to note:
  - Specify the path to the GEOS-Chem top-level directory, not the GCHP subdirectory
  - Do not include symbolic links in your source code path
  - Unlike GCC, do not edit the Makefile with your source code path

### Step 4: Load GCHP Environment

- Set up your environment prior to compiling and/or running
- On Odyssey:

OD ~ \$ source GCHP.ifort15_mvapich2_odyssey.bashrc Loading modules for GCHP on Odyssey, please wait			
Due to MODULEPATH changes the following have been reloaded: 1) gd/2.0.28-fasrc01			
Currently Loaded Modules: 1) perl/5.10.1-fasrc04	4) intel/15.0.0-fasrc01	7) zlib/1.2.8-fasrc03	
<ol> <li>perl-modules/5.10.1-fasrc12</li> <li>git/2.1.0-fasrc02</li> </ol>	5) gd/2.0.28-fasrc01 6) mvapich2/2.2-fasrc01	<pre>8) hdf5/1.8.12-fasrc12 9) netcdf/4.1.3-fasrc09</pre>	

- Elsewhere:
  - Create a .bashrc file based on sample files in the run directory
  - Using the libraries above is recommended but other combos are possible
    - OpenMPI
    - Intel MPI
    - Gfortran
    - Other NetCDF library versions

### Step 5: Compile GCHP

- Like GCC, compile GCHP from the run directory using the Makefile
- First time compilation (30-60 min): make clean\_compile
  - Warnings, error messages, and pauses are normal
  - Signs of successful compilation:
    - "### GCHP compiled Successfully ###"
    - The following files exist:
      - GCHP/ESMF/esmf.install
      - GCHP/FVdycoreCubed\_GridComp/fvdycore.install
      - GCHP/Shared/mapl.install
- Subsequent compilation: make clean\_standard
  - For updates to GC base code or GCHP top-level directory
  - Not for updates to GCHP subdirectories (e.g. GCHP/Shared)

# Step 6: Configure Run Settings

- Use utility bash script **runConfig.sh** for select config settings:
  - Compute resources (e.g. # nodes and cores)
  - Internal grid resolution
  - Restart file
  - Simulation start/end times
  - Output diagnostic file frequency, duration, and mode
    - e.g. hourly (frequency) that is time-averaged (mode) and contained in daily files (duration)
  - Component on/off switches, including mixing scheme
  - Time-step intervals
  - Debug level for MAPL
- Manually change individual config files for all other settings
- Important things to understand about runConfig.sh
  - Run scripts execute **runConfig.sh** prior to executing **geos**
  - It overwrites input.geos and \*.rc files (BEWARE!!!)
  - Does not updates HEMCO\_Config.rc Or Ext\_Data.rc (yet)

### runConfig.sh: Default Settings Part 1

#### #!/bin/bash runSettings.sh: Update select settings in \*.rc and input.geos config files # Usage: ./runSettings.sh # E. Lundgren, 8/17/2017 Configurables +### #### See Useful Tip #2 at start of slides. Always check that your resources here #### COMPUTE RESOURCE NUM\_NODES=1 match your run script settings! NUM\_CORES\_PER\_NODE=6 NY must be an integer and a multiple of 6 NY=6NX\*NY must equal total number of cores NX=1Choose NX and NY to optimize NX x NY/6 squareness # within contraint of total # of CPUs # e.g., (NX=2,NY=12) if 24 cores, (NX=4,NY=12) if 48 #### INPUT MET RESOLUTION INPUT\_MET\_RES=placeholder # not yet implemented Set simulation resolution **###** INTERNAL CUBED-SPHERE RESOLUTION # Please note that lightning and dust emissions in GCHP are tuned to c24. # Removing the online resolution dependency of lightning and dust by using # offline lightning and dust emissions will be a future update. Until # then, running at resolutions other than c24 will have degraded accuracy # due to the resolution dependency of emissions. Please contact the GCST for more information. CUBE SPHERE RES=24 # 24~4x5, 48~2x2.5, 90~1x1.25, 180~0.5x0.625 ## INITIAL RESTART FILE INITIAL\_RESTART=default )# specify default or your own restart filename

<sup>•</sup>Set restart filename if not using symbolic links included in the run directory

### runConfig.sh: Default Settings Part 2

#### SIMULATION TIMES Start\_Time="20130701 000000" End\_Time="20130701 010000" Puration="00000000 010000"

#### OUTPUT cs\_frequency="010000" cs\_duration="010000" cs\_mode="'time-averaged' T1\_frequency="010000" 11\_duration="010000" 11\_mode="'time averaged'"

#### TURN COMPONENTS N/OFF Turn\_on\_Chemistry=T Turn\_on\_emissions=T Turn\_on\_Dry\_Deposition=T Turn\_on\_Wet\_Deposition=T Turn\_on\_Transport=T Turn\_on\_Cloud\_Conv=T Turn\_on\_PBL\_Mixing=T

#### TIMESTEPS TransConv\_Timestep\_min=10 ChemEmis\_Timestep\_min=20

#### PBL MIXING Use\_nonlocal\_PBL=T

F#### DEBUG OPTIONS

MAPL\_DEBUG\_LEVEL=0

Set simulation start and end times. Set duration to the difference, or multiples of the difference if doing segmented runs.

Set cubed-sphere file duration, data frequency, and mode, either "'timeaveraged'" or "'instantaneous'". These update the "center" collection in HISTORY.rc. Ignore the "II\_" options.

Set high-level input.geos options

For MAPL debugging, set level as high as 20. Be aware this will come at great performance cost! # 0 is none, output increases with higher values (to 20) # requires special handling: omit for now

### Step 7: Run GCHP (single node)

• Two run scripts are provided as examples of how to run GCHP:

OD ~/gchp\_standard \$ cd runScriptSamples/ OD ~/gchp\_standard/runScriptSamples \$ ls gchp\_gridengine.run\* gchp\_slurm.run\* \_

- Both run runConfig.sh prior to executing geos
- runConfig.sh will exit with an error if your settings do not make sense or if your restart file does not exist, thereby preventing GCHP from starting and hanging
- It is up to you, however, to check that your compute resource settings in your run script match those in runConfig.rc
- GCHP standard output is sent to a log called gchp.log
- You may change the log name in your run script.

### Step 8: Analyze Output

- All GCHP output is in netCDF-4 format
- Two outputs:
  - Restart file
    - Stored in top-level of run directory
    - Filename: -gcchem\_internal\_checkpoint\_c24.nc (configured in GCHP.rc)
    - Cubed-sphere grid
  - One or more diagnostic files
    - Output files are stored in OutputDir
    - File format: GCHP. {collection}.YYYYMMDD.nc4
    - Different files for each collection
    - Collections configured in HISTORY.rc
- Regrid from cubed-sphere to lat-lon using tools developed by Seb Eastham:
  - CSRegridTool (FORTRAN): <u>https://bitbucket.org/sdeastham/csregridtool</u>
  - CSRegrid (Matlab): https://bitbucket.org/gcst/csgrid

### Step 9: Reuse a Run Directory

- You can reuse your GCHP run directory but MUST do one of the following prior to rerunning to avoid a seg fault:
  - make cleanup\_output
  - Delete file cap\_restart
- Invoking cleanup\_output will remove all simulation log and output files but will not delete your executable or last build info log
- Sample run scripts include several aliases to reduce typing if running repeatedly
- Experiment with different run settings in runConfig.sh
- If changing # of cores and/or # of nodes:
  - Remember to update **runConfig.sh** as well as your run script
  - Choose NX and NY such that NX by NY/6 is roughly square
  - See next slide for an example

### **GCHP** with Multiple Nodes

- 24 cores across 2 nodes
  - Run script:

#!/bin/bash

#SBATCH -n 24 #SBATCH -N 2

– runConfig.sh:

- 48 cores across 3 nodes
  - Run script:
    - -n 48
    - –N 3
  - runConfig.sh:
    - NUM\_NODES=3
    - NUM\_CORES\_PER\_NODE=16
    - NY=12
    - NX=4

### GCHP Source Code: ESMF, MAPL, FVdycore

ESMF and transport directories: these are compiled once and then you shouldn't need to touch them

	OD ~/Code.v11-02/GCHP \$	ls	
	Chem_OridCompNod.F90	GIGC_Connections.H	HEMCO_Includes_BeforeRun.H
	ESMF/	gigc_diagnostics_mod.F90	Includes_After_Dyn.H
_	FVdycoreCubed_GridComp/	<pre>gigc_finalization_mod.F90</pre>	Includes_After_Run.H
	gchp_utils.F90	GIGC_GridCompMod.F90	Includes_Before_Dyn.H
	gc_land_interface.F90	gigc_initialization_mod.F90	Includes_Before_Run.H
	GCSA-HowTo.docx	GIGC.mk	Makefile
	GEOSChem.F90	gigc_mpi_wrap.F90	Registry/
	GEOS_ctmEnvGridComp.F90	gigc_test_utils.F90	Shared/
	GEOS_HyCoords.H	<pre>gigc_type_mod.F*</pre>	
	gigc_chunk_mod.F90	HEMCO_Includes_AfterRun.H	

This is also compiled once. Most run directory issue errors will point you here.



### GCHP Source Code: MAPL\_Base

OD ~/Code.v11-02/GCHP/Shared/MAPL_Base \$ 1s				
	allgather.H	MAPL_Cap.F90	MAPL_SatVapor.F90	
Error messages	allgatherv.H	MAPL CETTE 90	MAPL_ShmemMod.F90*	
	allreducemax.H*	MAPL_CFIOServer.F90	MAPL_SimpleBundleMod.F90	
may lead you	allreducemin.H*	MAPL_Comms.P90	MAPL_Sort.F90	
here	allreducesup.H*	MAPL_Constants.F90	mapl_stub.pl*	
nere	arraygather.H	MAPL_ErrLog.h	MAPL_stubs.F90	
	arraygatherRcvCnt.H	MAPL_ErrLogMain.h	MAPL_sun_uc.P90	
Posourco sotup	arrayscatter.H	MAPL Exceptions.h	mapl_tree.py	
Resource setup	arrayscatterRcvCnt.H	MAPL_ExtDataGridCompMod.F90	MAPL_VarSpecMod.F90	
or time issues	bcast.H	MAPL_GenericCplComp.F90	mapl_vlist.py*	
	c_mapl_locstream_F.c	MAPL_Generic.F90	memuse.c*	
	CubeIcLatLon.F90	MAPL_Generic.h	overload.macro*	
Input data	evs/	MAPL_Hash.F90	Python/	
	eqsat.H	MAPL_Heapilou.F30	qsatice.H	
issues		MAPL_HistoryGridComp_E90	qsatlqu.H	
	esatice.H	MAPL_HorzTransform.F90	read_parallel.H	
	esatlqu.H	NAPL_InterpMod.F30	recv.H	
Output data 🥢	ESMFL_Mod.P90	MAPL_I0.P90	red_ma.pl*	
	gather.H	MAPL_LoadBalance.F90	Sample_ExtData.rc	
issues	GetPointer.H	MAPL_LocStreamMod.F90	scatter.H	
	getrss.c	MAPL_MaxMinMod.F90	send.H	
	GNUmakefile*	MAPL_MemUtils.F90*	sendrecv.H	
Tile file issues 🧹	hash.c	MAPL_Mod.F90	sort.c	
(lat-lon <-> CS)	hinterp.F	MAPL_NewArthParser.F90*	sun.H	
(10111011 < 200)	HorzBinning.F90	MAPL_NominalOrbitsMod.F90	tests/	
	<pre>mapl_acg.pl*</pre>	MAPL_OrbGridCompMod.F90	TeX/	
	MAPL_Base.F90	MAPL_OrbGridComp.rc	tstqsat.F90	
	MAPL_base.mk*	MAPL_Profiler.F90_	write_parallel.H	

Review your run directory setup before trying to change MAPL code!

### Resources

- GCHP Links:
  - <u>Main Wiki Page</u>
  - Online Tutorial
  - v11-02: new features, benchmarks, open and resolved issues
  - Working Group and Users
  - Timing Tests
- Other Useful Links:
  - Interactive construction of a cubed-sphere grid
  - GMAO MAPL User's Guide (info may be outdated)
  - GEOS-5 wiki page for ExtData (info may be outdated)