# **GenChem Documentation**

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### Introduction

GenChem is a system to generate and test chemical mechanisms for the EMEP MSC-W model [Simpson2012] and 1-D canopy model, ESX [SimpsonTuovinen2014]. GenChem consists of two main directories, **chem** and **box**.

The **chem** directory contains several chemical mechanisms written in chemist-friendly format (e.g. k1 NO2 + OH = HNO3). A python script *GenChem.py* can be used to convert these files to fortran friendly input files for the EMEP model, usually with the help of some wrapper script, either *do.GenChem*, *do.testChems*, or *emep\_setup.py*. The fortran files produced by these scripts have the prefix "CM\_" or "CMX\_", where CM denotes Chemical Mechanism.

Although GenChem can be run directly from within the **chem** directory, the strongly recommended approach is to use the scripts available in the **box** directory. In this approach GenChem is first applied, and then the resulting CM files are compiled and run as box-model simulations. Once all looks okay, a final script can be run to add additional code, and provide an EMEP-ready set of fortran files. This approach ensures that the CM files compile as they should, and allows rapid testing of several chemical mechanisms alongside each other.

#### 1.1 Code structure

The directory structure for GenChem can be summarised as:

```
# GenChem's mechanism tree
XXX/chem/scripts
                        # scripts, including do.GenChem and GenChem.py
XXX/chem/base_mechanisms # collection of main chemical schemes
XXX/chem/extra_mechanisms # collection of extra reactions for chemical schemes
                         # emissplit files, see ...
XXX/chem/inputs
XXX/box
                       # Top of ESX directory tree
XXX/box/src
                       # source files
                       # scripts
XXX/box/scripts
                    # documentation, as .rst files plus sphinx conf system
XXX/doc
XXX/doc/_build
                    # processed documentation, as .pdf and html
XXX/doc/_build/html # .. as .html (aim your browser at index.html here)
XXX/doc/_build/latex # .. as .pdf (aim your pdfreader at GenChemDoc.pdf here)
```

(where XXX could any suitable user-directory into which GenChem was unpacked, e.g. /home/fred/chemwork/GenChem.)

# 1.2 Requirements

GenChem has been developed on Ubuntu linux systems, and should work on any modern linux/unix computer. The code has also been run on Windows via a virtual ubuntu environment. The minumum requirements are a modern fortran compiler and python3 (probably 3.5 or higher).

We have used for example

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- gfortran (gcc 4.6.1) on Linux Xubuntu PC system
- gfortran (gcc 4.4.7) on HP supercomputer
- ifort 13.0.1

**Warning:** \*\* NOTE !! This user-guide is a work-in-progress manual on the GenChem system, with this interim version produced for interested users, Jan. 2020. \*\*

**TWO** 

#### **Quick start**

We will proceed directly to a run of the box-model system, to show how chemical schemes are normally compiled into CM\_ and CMX\_ files, and used in box-model simulations. This is actually the normal and recommended way to prepare files for the EMEP model, but also provides a good environment for comparing chemical mechanisms.

### 2.1 Step 1: initial setup

If not run previously, some preliminary steps are needed to set up a working directory. From the **GenChem/box** directory (cd GenChem/box), do:

```
cd somepath/GenChem-xxx/box
mkdir tmp_work

cd tmp_work/

cp ../src/*.f90 .
    cp ../src/Makefile .
    cp ../src/config_box.nml .
    cp ../scripts/do.testChems .
    cp ../scripts/emep_setup.py .
```

If you want, the script box\_setup.sh from the box/scripts directory can also be used to perform the above steps:

```
cd somepath/GenChem-xxx/box
scripts/box_setup.sh tmp_work
```

# 2.2 Step 2: do.testChems

At this stage, one can try compiling a chemical scheme. With the example of EmChem19, and now from our tmp\_work directory, try:

```
./do.testChems EmChem19a
```

This script will run GenChem.py on the EmChem19a scheme (also adding a few extra reactions from a helper BoxAero mechanism), run "make", and then run the resulting box-model code. Results will appear in one log-file (RES.EmChem19a, way too wordy!), and as comma-separated results in the Output directory: file boxEmChem19a.csv. This file is readable with e.g. libreoffice. Plot scripts are also available (see BELOW), for easy visualisation and comparison of these csv results.

The CM\_ and CMX\_ fortran files produced by this process are saved in directories, e.g. here in ZCMBOX\_EmChem19a. These files could be used in the EMEP model if wanted, but usually the more complex script emep\_setup.py (described below) is used for that. (Hence we reserve the prefix ZCMBOX for files created by do.testChems and ZCM for those created with emep\_setup.py, see below.)

Now, if one wants to compare several schemes, one can do e.g.:

```
./do.testChems EmChem19a CRI-R5-emep MCM_v3.3
```

This would produce 3 output .csv files, which again are easily plotted against each other.

Technical comments:

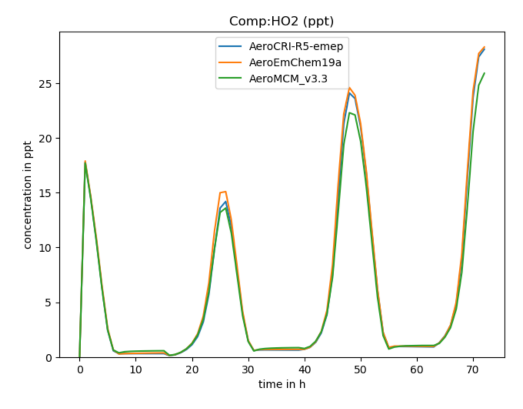
- do.testChems is just a simple wrapper, which cleans up files, runs another script (do.GenChem), and runs the box model.
- MCM is a very large scheme and this can take a while, or stress your PC's memory! Try with the smaller schemes first.

# 2.3 2a. Plotting?

If one has run say 3 chemical schemes using Step 2 above, the results are easily plotted from the box/tmp\_work/Output directory:

```
../../scripts/boxplots.py -h for help!
../../scripts/boxplots.py -v 03 -i boxEmChem19a.csv boxChem1.csv boxChem2.csv -p
```

Using 'ALL' or 'DEF' with -v results in all or many common species being plotted at once (-p is assumed in this case. For example, here we can see a comparison of three schemes produced with this script:



Another crude+helpful script just grabs the concentrations:

```
../../scripts/getboxconcs.py O3 boxEmChem19a.csv
```

which results in ResConcs\_boxEmChem19a\_O3\_ppb.txt

### 2.4 2b. Box-config

The script do.testChems above compiles the executable boxChem for each mechanism in turn, and by default runs this using some settings from the default config\_box.nml file. This file contains a number of important settings which by deault run a 24-hour simulation (starting at 12:00 GMT), with set emissions, temperature of 298.15 K, mixing height of 1000 m, and some boundary conditions. Default outputs are also given.

The user can of course change these settings (do this in your working directory, not in *src*). We explain the variables and choices here.

Note these config files follow fortran namelist conventions. Text following an exclamation mark (!) is ignored.

#### 2.4.1 Time-related variables

#### 2.4.2 Geographical location

```
lat = 45.05, ! degrees N
lon = 15.06, ! degrees E
```

#### 2.4.3 Emissions

# 2.5 Step 3: emep\_setup.py

The do.testChems script described above is best for quickly testing and comparing different mechanisms. Usually these comparisons only involve gas-phase mechanisms such as EmChem19a or MCM\_v3.3. However, the EMEP model usually requires a host of extra species and reactions to accommodate sea-salt, dust, organic aerosols, and pollen. It also requires files to specify how emissions and boundary conditions should be distributed among specific species, e.g. how a VOC emission should be split into C2H6, C2H4, nC4H10 etc.

In fact, for the EMEP model, GenChem produces many files which are copied into ZCM\_XXX directories for the scheme XXX you wish to use:

```
$ls -x ZCM_EmChem19a/
```

```
CM_ChemDims_mod.f90
                         CM_ChemGroups_mod.f90
                                                    CM_ChemRates_mod.f90
                         CM DryDep.inc
                                          CM EmisFile.inc
                                                           CM emislist.csv
CM_ChemSpecs_mod.f90
CM EmisSpecs.inc
                   CM Reactions1.inc
                                      CM Reactions2.inc
                                                          CM Reactions.log
CM WetDep.inc
                                  CMX_BiomassBurningMapping_FINNv1.5.txt
CMX_BiomassBurningMapping_GFASv1.txt
                                         CMX_BoundaryConditions.txt
fig_box.nml run_emislist/ (with emislist.defaults.sox etc..)
```

The recommended way to get this directory is to use the script *emep\_setup.py* from your temporary work directory within the **box** system. So, from e.g. box/tmp\_work, do:

./emep\_setup.py EmChem19a

or just:

```
./emep_setup.py
```

and this will provide a list of options.

You can edit the *emep\_setup.py* scripts, maybe renaming it as *my\_setup.py* directory. If selecting from the provided base\_mechanisms and extra\_mechanisms you only need to extend the possible command lines as provided by the *cmdx* dictionary:

```
cmdx['EmChem19a'] = '-b EmChem19a -e PM_VBS_EmChem19 BVOC_MTERP1_EmChem19'+common
cmdx['CRI-R5-emep'] = '-b CRI-R5-emep -e common'
```

The '-b' argument gives the base mechanism, and then you can have any number of compatable extra mechanisms (-e argument).

Any keys from *cmdx* can be used by *emep\_setup.py*. For example, if the user builds a new base scheme *usersChem* and some OA scheme, *usersSOA*, then *emep\_setup.py* can be edited to add these as a new option:

```
cmdx['usersChem'] = '-b usersChem -e usersSOA'+common
```

you could do:

```
do.testChems usersChem  # GOOD TO CHECK FIRST

emep_setup.py usersChem  # Creates ZCM_usersChem
```

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### Formatting of GenChem files

#### 3.1 Reactions files

#### Example:

```
* Some simple lines

1.4e-12*EXP(-1310.*TINV) : 03 + N0 = N02 + <02> ; acp2004

5.681e-34*EXP(-2.6*LogTdiv300) : OP + <02> + <M> = 03 ; acp2004

2.15e-11*EXP(110.*TINV) : OD + <N2> = OP ; Updated (IUPAC 2009)

emisfiles:sox,nox,co,voc,nh3
rcemis(NO,KDIM) : = NO ;
```

- END-OF-LINE is ";". Text after this (e.g. references, or unused "products") will be ignored.
- Separator between rate coefficient and reaction is ":".
- lines beginning with "\*" are comments (no ";" needed here)
- lines beginning with "rcemis" are emission terms
- lines beginning with "emisfiles" give name of emission files, e.g. nox
- Some coefficients are defined in GenIn.shorthand, e.g. TINV, LogTdiv300
- Anything else is simply used as the rate coefficient. (Do not add spaces!)

Four types of tracers/catalysts/yields are allowed, denoted by different types of parentheses:

- e.g. [OH] + VOC -> SOA will put xnew(OH) into the loss rate of VOC, but will not change the loss rate of OH.
- 2) e.g. {O2} + OD -> OP will ignore the O2 term. Make sure it is in the reaction rate though if needed!
- 3) e.g. OP + <O2> + <M> -> O3 will ignore the O2 and M term AND add their concentrations to the reaction rate (multiply it). This system is only used for these "special" species (O2, N2, M) as they must be predefined, e.g. O2(k), in boxChem and/or EMEP codes.
- 4) e.g. 1.36e-11: [OXYL] + [OH] = |YCOXY(0)| ASOC\_ug1 + ... will replace the contents of the || term with yield coefficients which will be updated each time-step in the EMEP model. These variables (here YCOXY(0)) must be predefined in order for emep\_setup.py and the emep model to compile.

# 3.2 Species files

The input to the GenChem.py script is GenIn\_Species.csv, but this is assembled by do.GenChem from all needed \_Species.csv files from the base\_mechanisms and extra\_mechanisms sub-directories. For For example, for a typical emep run with base EmChem19a, do.GenChem appends EmChem19a\_Species.csv, SeaSalt\_Species.csv, and many more into one GenIn\_Species.csv. The file contains columns with species name, type, formula, and various settings related to dry and wet deposition

#### Example lines:

Further details can be found in the documentation article ...

#### 3.3 Shorthands file

Shorthands are text-strings used in the Reactions.txt file, usually to represent commonly used rate-coefficients. The meaning of the text-string is given in \_Shorthand.txt file, e.g.

In these examples, XT is just a character-saving replacement for temp, FH2O gives a more complex expression, which also uses the pre-defined variable TINV = 1/temp. KHO2RO2 is a common rate-coefficient, but here we see that comments are allowed - anything afer the 2nd term. FInally, the KMT12 term shows that complex fuction calls are also allowed. IMPORTANT - avoid white space in any terms!

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### **FOUR**

#### **Contributors**

The GenChem system was created over many years:

David Simpson, Norwegian Meteorological Institute & Chalmers, 1998-2019: wrote original GenChem.pl scripts, the boxChem system, assorted helper scripts (do.GenChem, boxplots.py, etc.), and python3 conversion.

Alan Briolat, Stockholm Environment Institute at York, 2013: wrote the first python version: GenChem.py.

Hannah Imhof, Chalmers, 2016: added extra flexibility and types of arrays (e.g. factor groups), plus further scripts. Added CRI and MCM chemical mechanisms.

John Johansson, Chalmers, 2017-2019: improved organisation and flexibility of GenChem system.

Robert Bergström, Chalmers and SMHI, 2017-2019: development of chemical mechanisms (e.g. EmChem19 family, VBS schemes) for gas and aerosols.

Alvaro Valdebenito, Norwegian Meteorological Institute, 2018-2019, various bug-fixes and updates. Added pollen.

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# References

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