GEOS-Chem Reference, Vol. 1: Makefiles

GEOS-CHEM SUPPORT TEAM

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1 Routine/Function Prologues

1.1 Module Interface Makefile (Main-level)

This is a "router" makefile. It calls the main GEOS-Chem Makefile (in the GeosCore subdirectory) to direct the Unix "make" utility how to build the GEOS-Chem source code.

REMARKS:

```
To build the programs, call "make" with the following syntax:
    make TARGET [ OPTIONAL-FLAGS ]
  To display a complete list of options, type "make help".
  Makefile uses the following variables:
  Variable
           Description
  GEOSDIR
            Specifies the directory where GEOS-Chem "core" routines are found
  GEOSTOM
            Specifies the directory where GEOS-Chem + TOMAS routines are found
REVISION HISTORY:
   16 Sep 2009 - R. Yantosca - Initial version
   24 Nov 2009 - R. Yantosca - Now call libbpch and libcore targets in
                            the Makefile in the GeosCore sub-directory
   11 Dec 2009 - R. Yantosca - Now get SHELL from Makefile_header.mk
   25 Jan 2010 - R. Yantosca - Added Makefile targets for TOMAS microphysics
   16 Feb 2011 - R. Yantosca - Added Makefile targets for APM microphysics
   04 Nov 2011 - R. Yantosca - Remove ESMF targets, those are not needed
   24 Jan 2012 - R. Yantosca - Also add libnc target to build netCDF utils
   11 May 2012 - R. Yantosca - Now make sure that all targets of the
                            GeosCore/Makefile are pointed to properly
# Get the Unix shell definition
include ./Makefile_header.mk
# Define variables
GEOSAPM = GeosApm
GEOSDIR = GeosCore
GEOSTOM = GeosTomas
GTMM = GTMM
# Makefile targets: type "make help" for a complete list!
```

.PHONY: all lib libkpp libnc libutil exe clean realclean doc docclean help

```
all:
@$(MAKE) -C $(GEOSDIR) all
lib:
@$(MAKE) -C $(GEOSDIR) lib
libcore:
@$(MAKE) -C $(GEOSDIR) libcore
libiso:
@$(MAKE) -C $(GEOSDIR) libiso
libkpp:
@$(MAKE) -C $(GEOSDIR) libkpp
libnc:
@$(MAKE) -C $(GEOSDIR) libnc
ncdfcheck:
@$(MAKE) -C $(GEOSDIR) ncdfcheck
libutil:
@$(MAKE) -C $(GEOSDIR) libutil
libheaders:
@$(MAKE) -C $(GEOSDIR) libheaders
exe:
@$(MAKE) -C $(GEOSDIR) exe
clean:
@$(MAKE) -C $(GEOSDIR) clean
distclean:
@$(MAKE) -C $(GEOSDIR) distclean
realclean:
@$(MAKE) -C $(GEOSDIR) realclean
doc:
@$(MAKE) -C $(GEOSDIR) doc
docclean:
@$(MAKE) -C $(GEOSDIR) docclean
help:
@$(MAKE) -C $(GEOSDIR) help
```

```
#-----
# Targets for mercury simulation (ccc, 6/7/10)
.PHONY: allhg libhg libgtmm exehg
allhg:
@$(MAKE) -C $(GEOSDIR) allhg
libhg:
@$(MAKE) -C $(GEOSDIR) libhg
ligbtmm:
@$(MAKE) -C $(GEOSDIR) libgtmm
exehg:
@$(MAKE) -C $(GEOSDIR) exehg
# Targets for TOMAS aerosol microphysics code (win, bmy, 1/25/10)
.PHONY: tomas libtomas exetomas cleantomas
tomas:
@$(MAKE) -C $(GEOSTOM) TOMAS=yes all
libtomas:
@$(MAKE) -C $(GEOSTOM) TOMAS=yes lib
exetomas:
@$(MAKE) -C $(GEOSTOM) TOMAS=yes exe
cleantomas:
@$(MAKE) -C $(GEOSTOM) TOMAS=yes clean
# Targets for APM aerosol microphysics code (bmy, 2/16/11)
.PHONY: apm libapm exeapm cleanapm
apm:
@$(MAKE) -C $(GEOSAPM) APM=yes all
libapm:
@$(MAKE) -C $(GEOSAPM) APM=yes lib
```

```
exeapm:
```

@\$(MAKE) -C \$(GEOSAPM) APM=yes exe

cleanapm:

@\$(MAKE) -C \$(GEOSAPM) APM=yes clean

1.1.1 Makefile_header.mk

This sub-makefile defines the variables which specify compilation options for the different supported compiler/platform combinations. Also, the default makefile compilation rules are specified here.

REMARKS:

To build the programs, call "make" with the following syntax:

```
make TARGET [ OPTIONAL-FLAGS ]
To display a complete list of options, type "make help".
```

The following variables are exported to the main-level Makefile:

Variable Description CCContains the default C compilation commands (for PGI only) Contains the Fortran compilation commands F90 Contains the command to force F90 "free format" compilation FREEFORM LD Contains the command to link to libraries & make executable LINK Contains the commands to link to GEOS-Chem built libraries R8 Contains the command to force REAL -> REAL*8 SHELL Contains the default Unix shell to use when building code

FFLAGS is a local variable that is not returned to the "outside world", but is only used locally. COMPILER, HDF5, and OMP are all input via the command line or via environment variables.

REVISION HISTORY:

- 16 Sep 2009 R. Yantosca Initial version
- 22 Sep 2009 R. Yantosca Bug fix, added -I\$(HDR) to F90 compilation lines
- 24 Sep 2009 R. Yantosca added NONUMA option for PGI compiler
- 07 Oct 2009 R. Yantosca Replaced .SUFFIXES section w/ pattern rules
- 19 Nov 2009 R. Yantosca Now use OMP variable to determine whether to turn on OpenMP parallelization options
- 23 Nov 2009 R. Yantosca Now use -module \$(MOD) instead of -I\$(MOD) to specify the directory for *.mod files on both IFORT and PGI compilers.
- 23 Nov 2009 R. Yantosca Now use -moddir=\$(MOD) and -M\$(MOD) instead of -I\$(MOD) to specify the directory for *.mod files on the SunStudio compiler.
- 23 Nov 2009 R. Yantosca Change DEBUG to allow for new version of
 Totalview which doesn't choke when debugging
 parallel code (Totalview 8.6.1-1)
- 02 Dec 2009 R. Yantosca Added SUN32 switch for building 32-bit executbable on the SunStudio compiler
- 11 Dec 2009 R. Yantosca Now define SHELL here and export to other

 Makefiles, so as to have a single place where
 the Unix shell name is defined.
- 21 Dec 2009 R. Yantosca Add H5I and H5L variables to specify the HDF5 library and include paths. Also set the default to not link to the HDF5 libraries.
- 21 Dec 2009 R. Yantosca Now pass LINK back to the outside world, so that the Makefile that builds the executable can reference it.
- 19 Jan 2010 R. Yantosca Minor fix, add -m64 if SUN32 is not defined.
- 25 Jan 2010 R. Yantosca Now add -DTOMAS to FFLAGS if necessary
- 28 Jan 2010 C. Carouge Add -lIsoropia to LINK, for ISORROPIA II
- 16 Feb 2011 R. Yantosca Now add -DAPM to FFLAGS if necessary
- 25 Aug 2011 R. Yantosca Add "-fp-model source" to FFLAGS for IFORT compiler. This will prevent aggressive optimizations from changing numerical results.
- 25 Aug 2011 R. Yantosca Add -CU (check for uninit'd variables) to FFLAGS when using IFORT w/ the DEBUG option.
- 26 Aug 2011 R. Yantosca Allow for deactivation of the "-fp-model source" option by using the PRECISE=no env variable
- 24 Jan 2012 R. Yantosca If NETCDF=yes, GEOS-Chem will link and include to the netCDF dir paths that are specified
- 24 Jan 2012 R. Yantosca Now use := for makefile assignment statements
- 10 Feb 2012 R. Yantosca When compiling with NETCDF=yes or HDF5=yes, we must also add the flags -mcmodel=medium -i-dynamic to FFLAGS in order to avoid memory errors (for IFORT only)
- 10 Feb 2012 R. Yantosca Remove -CU from the DEBUG option (IFORT only)
- 19 Mar 2012 R. Yantosca Add optional NO_ISO switch, which will turn off the ISORROPIA ATE package for testing
- 05 Apr 2012 R. Yantosca Now assume netCDF is always used

05 Apr 2012 - R. Yantosca - Change BL_INC_NETCDF to INC_NETCDF

```
05 Apr 2012 - R. Yantosca - Change BL_INC_HDF5 to INC_HDF5
   05 Apr 2012 - R. Yantosca - Change BL_LIB_NETCDF to LIB_NETCDF
   05 Apr 2012 - R. Yantosca - Change BL_LIB_HDF5
                                           to LIB_HDF5
   30 Apr 2012 - R. Yantosca - Add NETCDF3=[yes|no] makefile option
   30 Apr 2012 - R. Yantosca - Use separate netCDF link and include paths
                          for netCDF3 and for netCDF4
   30 Apr 2012 - R. Yantosca - Also add -mcmodel=medium flag for PGI compiler
   09 May 2012 - R. Yantosca - Now try to get the proper linking sequence
                          for netCDF etc w/ nf-config and nc-config.
   11 May 2012 - R. Yantosca - Now export NCL (netCDF linking sequence)
# Default settings for Makefile options
# IFORT is default compiler
ifndef COMPILER
COMPILER := ifort
endif
# Get Operating System (Linux = Linux; Darwin = MacOSX)
ifndef UNAME
UNAME
          := $(shell uname)
endif
# OpenMP is turned on by default
ifndef OMP
OMP
        := yes
endif
# HDF5 I/O is turned off by default
ifndef HDF5
HDF5
       := no
endif
# Use precise FP math optimization (i.e. to avoid numerical noise)
ifndef PRECISE
PRECISE
       := yes
endif
# TOMAS runs on single processor (at least for now!)
ifeq ($(TOMAS), yes)
OMP
        := no
endif
```

```
# Default values for variables
# If your system uses "/bin/sh", then uncomment this line!
SHELL
         := /bin/sh
# If your system uses "/bin/bash", then uncomment this line!
#SHELL
         := /bin/bash
# Library include path
         := -I$(GC_INCLUDE)
NCI
# Library link path: first try to get the list of proper linking flags
# for this build of netCDF with nf-config and nc-config.
         := $(shell $(GC_BIN)/nf-config --flibs)
NCL
        += $(shell $(GC_BIN)/nc-config --libs)
NCL
NCL
        := $(filter -1%,$(NCL))
#%%%% NOTE TO GEOS-CHEM USERS: If you do not have netCDF-4.2 installed
#%%%% Then you can add/modify the linking sequence here. (This sequence
#%%%% is a guess, but is probably good enough for other netCDF builds.)
ifeq ($(NCL),)
NCL
         := -lnetcdf -lhdf5_hl -lhdf5 -lz
endif
# Prepend the library directory path to the linking sequence
         := -L\$(GC\_LIB) \$(NCL)
NCL
# Command to link to the various library files (-lHeaders should be last!)
        := -L$(LIB) -lKpp -lIsoropia -lGeosUtil -lHeaders
LINK
         := $(LINK) -1NcUtils $(NCL)
# Commands to link to libraries, for GTMM code (-lHeaders should be last!)
LHG
         := -L$(LIB) -lKpp -lIsoropia -lHg -lGeosUtil -lHeaders
LHG
        := $(LINK) -1NcUtils $(NCL)
# Add the HDF5 library link commands (optional)
ifeq ($(HDF5),yes)
LINK
        := \$(LINK) - L\$(H5L)
LHG
         := \$(LINK) - L\$(H5L)
endif
# For ESMF development
ifeq ($(ESMF),yes)
LINK
        += -lesmf $(Lib_chem_base) $(Lib_chem_shared) $(Lib_pilgrim)
                 $(LIB_MAPL_BASE) $(LIB_CFIO) $(LIB_GFIO) $(LIB_MPEU) \
```

```
$(LIB_ESMF) $(LIB_SDF) $(LIB_SYS) $(LIB_MPI)
                  $(ESMF_LDFLAGS) -lmpi_cxx -lstdc++ -limf -lrt -ldl
         += -1ESMF
T.HG
endif
#-----
# IFORT compilation options (default)
#-----
ifeq ($(COMPILER),ifort)
# Turn on -traceback option by default for debugging runs
ifdef DEBUG
TRACEBACK := yes
endif
# Pick compiler options for debug run or regular run
ifdef DEBUG
FFI.AGS
        := -cpp -w -00 -auto -noalign -convert big_endian -g
else
        := -cpp -w -02 -auto -noalign -convert big_endian -vec-report0
FFLAGS
endif
# OSX compilation options
ifeq ($(UNAME), Darwin)
FFLAGS += -Wl,-stack_size,0x2cb410000 # Allow 12GB of stack space
ifdef DEBUG
FFLAGS += -g0 -debug -save-temps -fpic -W1,-no_pie
endif
endif
# Add options for medium memory model. This is to prevent G-C from
# running out of memory at hi-res, especially when using netCDF I/O.
ifneq ($(UNAME),Darwin)
FFLAGS
        += -mcmodel=medium -i-dynamic
endif
# Prevent any optimizations that would change numerical results
# This is needed to prevent numerical noise from ISORROPIA (bmy, 8/25/11)
ifeq ($(PRECISE),yes)
FFLAGS
         += -fp-model source
endif
# Turn on OpenMP parallelization
ifeq ($(OMP),yes)
FFLAGS
         += -openmp -Dmultitask
endif
# Also add TOMAS aerosol microphysics option
```

```
ifeq ($(TOMAS),yes)
FFLAGS
         += -DTOMAS
endif
# Also add APM aerosol microphysics option
ifeq ($(APM),yes)
FFLAGS
         += -DAPM
endif
# Add special IFORT optimization commands
ifdef IPO
FFLAGS
          += -ipo -static
endif
# Add option for "array out of bounds" checking
ifdef BOUNDS
FFLAGS
          += -CB
endif
# Also add traceback option
ifdef TRACEBACK
FFLAGS
       += -traceback
endif
# Option to turn off ISORROPIA for testing
ifdef NO_ISO
FFLAGS
         += -DNO_ISORROPIA
endif
# Include options (i.e. for finding *.h, *.mod files)
INCLUDE := -I$(HDR) -module $(MOD) $(NCI)
# Also append HDF5 include commands (optional)
ifeq ($(HDF5),yes)
INCLUDE += -DUSE_HDF5 -I$(H5I)
endif
# Also add ESMF linking option
ifeq ($(ESMF),yes)
FFLAGS
         += -DESMF_
endif
ifeq ($(ESMF_TESTBED),yes)
       += -DESMF_TESTBED_
FFLAGS
INCLUDE += -I\$(HDR)
endif
# DEVELOPMENT FLAG - MSL
```

```
ifeq ($(DEVEL),yes)
FFLAGS += -DDEVEL
endif
CC
F90
       = ifort $(FFLAGS) $(INCLUDE)
       = ifort $(FFLAGS)
LD
FREEFORM = -free
R8
      = -r8
endif
# Portland Group (PGI) compilation options
#-----
ifeq ($(COMPILER),pgi)
# Pick compiler options for debug run or regular run
ifdef DEBUG
FFLAGS
        := -byteswapio -Mpreprocess -Bstatic -g -00
else
        := -byteswapio -Mpreprocess -Bstatic -fast
FFLAGS
endif
# Add options for medium memory model. This is to prevent G-C from
# running out of memory at hi-res, especially when using netCDF I/O.
        += -mcmodel=medium
FFLAGS
# Turn on OpenMP parallelization
ifeq ($(OMP),yes)
FFLAGS
        += -mp -Mnosgimp -Dmultitask
endif
# Add option for suppressing PGI non-uniform memory access (numa) library
ifeq ($(NONUMA),yes)
FFLAGS
        += -mp=nonuma
endif
# Also add TOMAS aerosol microphysics option
ifeq ($(TOMAS),yes)
FFLAGS
        += -DTOMAS
endif
# Also add APM aerosol microphysics option
ifeq ($(APM),yes)
FFLAGS
        += -DAPM
endif
```

```
# Add option for "array out of bounds" checking
ifdef BOUNDS
FFI.AGS
        +=-C
endif
# Option to turn off ISORROPIA for testing
ifdef NO_ISO
FFLAGS
        += -DNO_ISORROPIA
endif
# Include options (i.e. for finding *.h, *.mod files)
       := -I$(HDR) -module $(MOD) $(NCI)
INCLUDE
# Also append HDF5 include commands (optional)
ifeq ($(HDF5),yes)
       += -DUSE_HDF5 -I$(H5I)
INCLUDE
endif
CC
       := gcc
F90
        := pgf90 $(FFLAGS) $(INCLUDE)
        := pgf90 $(FFLAGS)
LD
FREEFORM := -Mfree
R8
       := -Mextend -r8
endif
# Prior to 4/30/12:
# For now, remove SunStudio compiler option. Pretty much everyone now is using
# a Linux-like O/S and can use either IFORT or PGI compilers. (bmy, 4/30/12)
## SunStudio compilation options
-----
#ifeq ($(COMPILER),sun)
## Pick compiler options for debug run or regular run
## NOTE: -native builds in proper options for whichever chipset you have!
#ifdef DEBUG
        := -fpp -g -00 -stackvar -xfilebyteorder=big16:%all -native
#FFLAGS
#else
#FFLAGS
         := -fpp -fast -stackvar -xfilebyteorder=big16:%all -native
#endif
## Build Sun for 32-bit platform
#ifdef SUN32
#FFLAGS
        += -m32
#else
\#FFLAGS += -m64
```

```
#endif
## Turn on OpenMP parallelization
#ifeq ($(OMP),yes)
#FFLAGS
       += -openmp=parallel -Dmultitask
#endif
## Also add TOMAS aerosol microphysics option
#ifeq ($(TOMAS),yes)
#FFLAGS
        += -DTOMAS
#endif
## Also add APM aerosol microphysics option
#ifeq ($(APM),yes)
#FFLAGS
        += -DAPM
#endif
## Add option for "array out of bounds" checking
#ifdef BOUNDS
#FFLAGS
       += -C
#endif
## Option to turn off ISORROPIA for testing
#ifdef NO_ISO
#FFLAGS += -DNO_ISORROPIA
#endif
## Include options (i.e. for finding *.h, *.mod files)
#INCLUDE := -I$(HDR) -moddir=$(MOD) -M$(MOD) $(NCI)
## Also append HDF5 include commands (optional)
#ifeq ($(HDF5),yes)
#INCLUDE += -DUSE_HDF5 -I$(H5I)
#endif
#CC
      :=
## If your compiler is under the name "f90", use these lines!
#F90
       := f90 $(FFLAGS) $(INCLUDE)
         := f90 $(FFLAGS)
##-----
## If your compiler is under the name "sunf90", use these lines!
##F90
         := sunf90 $(FFLAGS) $(INCLUDE)
         := sunf90 $(FFLAGS)
##LD
##-----
#FREEFORM := -free
#R8
        := -xtypemap=real:64
#
```

```
#endif
# Prior to 4/30/12:
# For now, remove IBM/XLF compiler option. Pretty much everyone now is using
# a Linux-like O/S and can use either IFORT or PGI compilers. (bmy, 4/30/12)
## IBM/XLF compilation options
## NOTE: someone who runs on IBM compiler should check this !!!
##-----
#ifeq ($(COMPILER),xlf)
## Default compilation options
#FFLAGS = -bmaxdata:0x80000000 -bmaxstack:0x80000000 -qfixed -qsuffix=cpp=f -q64
## Add optimization options
#FFLAGS += -03 -qarch=auto -qtune=auto -qcache=auto -qmaxmem=-1 -qstrict
## Turn on OpenMP parallelization
#ifeq ($(OMP),yes)
#FFLAGS += -qsmp=omp:opt -WF,-Dmultitask -qthreaded
#endif
## Prior to 11/19/09:
### Add more options for parallel run
##ifndef DEBUG
##FFLAGS += -qsmp=omp:opt -WF,-Dmultitask -qthreaded
##endif
## Also add TOMAS aerosol microphysics option
#ifeq ($(TOMAS),yes)
#FFLAGS += -DTOMAS
#endif
## Also add APM aerosol microphysics option
#ifeq ($(APM),yes)
#FFLAGS += -DAPM
#endif
## Add option for "array out of bounds" checking
#ifdef BOUNDS
#FFLAGS += -C
#endif
## Option to turn off ISORROPIA for testing
#ifdef NO_ISO
#FFLAGS
        += -DNO_ISORROPIA
#endif
```

```
#
## Include options (i.e. for finding *.h, *.mod files)
\#INCLUDE = -I\$(HDR) -I \$(MOD) \$(NCI)
## Also append HDF5 include commands if necessary
#ifeq ($(HDF5),yes)
#INCLUDE += -DUSE_HDF5 -I$(H5I)
#endif
#
#CC
#F90
     = xlf90_r $(FFLAGS) $(INCLUDE)
    = xlf90_r $(FFLAGS)
#FREEFORM = -qrealsize=8
\#R8 = -r8
#endif
# Specify pattern rules for compiliation
# (i.e. tell "make" how to compile different types of source code files)
%.o: %.f
$(F90) -c $<
%.o: %.F
$(F90) -c $<
%.o: %.f90
$(F90) -c $(FREEFORM) $<
%.o: %.F90
$(F90) -c $(FREEFORM) $<
%.o : %.c
$(CC) -c $*.c
# Export global variables so that the main Makefile will see these
export CC
export F90
export FREEFORM
export LD
export LINK
export R8
export SHELL
export NCL
```

1.2 Module Interface Makefile (in the GeosUtil subdirectory)

This makefile compiles the various GEOS-Chem utility modules, which provide basic functionality for:

- Collapsing vertical levels in the stratosphere
- Date and time computations
- Defining data directories
- Defining the GEOS-Chem horizontal grid
- Defining the GEOS-Chem pressure coordinate grid
- Defining the logical units for GEOS-Chem file I/O
- Defining various Unix commands
- Platform-specific error handling
- Manipulating string variables
- Regridding data (horizontally) from fine to coarse resolution

REMARKS:

To build the programs, call "make" with the following syntax:

```
make TARGET [ OPTIONAL-FLAGS ]
```

To display a complete list of options, type "make help".

Makefile uses the following variables:

Variable	Description
SHELL	Specifies the shell for "make" to use (usually SHELL=/bin/sh)
ROOTDIR	Specifies the root-level directory of the GEOS-Chem code
HDR	Specifies the directory where GEOS-Chem include files are found
LIB	Specifies the directory where library files (*.a) are stored
MOD	Specifies the directory where module files (*.mod) are stored
AR	Sys var w/ name of library creator program (i.e., "ar", "ranlib")
MAKE	Sys var w/ name of Make command (i.e, "make" or "gmake")

REVISION HISTORY:

```
19 Nov 2009 - R. Yantosca - Initial version
   23 Nov 2009 - R. Yantosca - Now don't copy module files; they will be
                            automatically written to the mod directory
   11 Dec 2009 - R. Yantosca - Now get SHELL from Makefile_header.mk
   21 Dec 2009 - R. Yantosca - If HDF5=yes, then look for hdf5.mod in the
                            HDF5 include path $(HDF5_INC).
   01 Mar 2012 - R. Yantosca - Replace grid_mod.F with grid_mod.F90, to
                            facilitate work on the GI model
   03 Apr 2012 - M. Payer - Add new module regrid_a2a_mod.F90 (M. Cooper)
# Define variables
ROOTDIR := ...
HDR
      := $(ROOTDIR)/Headers
HELP
      := $(ROOTDIR)/help
      := $(ROOTDIR)/lib
LIB
MOD
      := \$(ROOTDIR)/mod
# Include header file. This returns CC, F90, FREEFORM, LD, R8, SHELL,
# as well as the default Makefile compilation rules for source code files.
include $(ROOTDIR)/Makefile_header.mk
# List of files to compile. Here the order is not important,
# as we will explicity define the dependencies listing below.
# List of source files
SRC := $(wildcard *.F) $(wildcard *.F90)
# Skip these files
SRC := $(filter-out grid_mod.F,$(SRC))
# Replace .f and .f90 extensions with *.o
TMP := \$(SRC:.F=.o)
OBJ := (TMP:.F90=.0)
# Special files just for IFORT
ifeq ($(COMPILER),ifort)
OBJ += ifort_errmsg.o
endif
# Special files just for PGI
ifeq ($(COMPILER),pgi)
OBJ += linux_err.o
endif
```

```
# Makefile targets: type "make help" for a complete listing!
.PHONY: clean help
lib: $(OBJ)
$(AR) crs libGeosUtil.a $(OBJ)
mv libGeosUtil.a $(LIB)
clean:
rm -f *.o *.mod
help:
@$(MAKE) -C $(HELP)
#-----
# Dependencies listing (grep "USE " to get the list of module references!)
# From this list of dependencies, the "make" utility will figure out the
# correct order of compilation (so we don't have to do that ourselves).
# This also allows us to compile on multiple processors with "make -j".
#
# NOTES:
# (1) Only specify object-file dependencies that are within this directory.
      Object files in other directories will be referenced at link-time.
# (2) For "make -j" to work, all files in this directory must have a
      listed dependency.
: bpch2_mod.F
bpch2_mod.o
                                    error_mod.o
                                                  file_mod.o
                                    julday_mod.o
                : charpak_mod.F
charpak_mod.o
directory_mod.o : directory_mod.F
error_mod.o
                : error_mod.F
file_mod.o
                : file_mod.F
                                    error_mod.o
global_grid_mod.o : global_grid_mod.F90 error_mod.o
                                                  grid_mod.o
grid_mod.o
               : grid_mod.F90
                                    error_mod.o
ifort_errmsg.o
                : ifort_errmsg.F
julday_mod.o
               : julday_mod.F
linux_err.o
                : linux_err.c
                : pressure_mod.F
                                    error_mod.o
pressure_mod.o
regrid_1x1_mod.o : regrid_1x1_mod.F
                                    charpak_mod.o
                                                  error_mod.o \
                                    grid_mod.o
regrid_a2a_mod.o : regrid_a2a_mod.F90
                                    file_mod.o
                                                  grid_mod.o
                                    charpak_mod.o error_mod.o \
time_mod.o
                : time_mod.F
                                    grid_mod.o
                                                  julday_mod.o
transfer_mod.o
                : transfer_mod.F
                                    error_mod.o
unix_cmds_mod.o : unix_cmds_mod.F
```

```
# NOTE: if HDF5=yes then we need to also look for the hdf5.mod
# in the HDF5 include path (bmy, 12/21/09)
hdf_mod.o : hdf_mod.F error_mod.o grid_mod.o
ifeq ($(HDF5),yes)
$(F90) -DUSE_HDF5 -I$(HDF_INC) -c $<
endif</pre>
```

1.3 Module Interface Makefile (in the ISOROPIA/ subdirectory)

This makefile compiles the ISOROPIA code. Object files (*.o) are bundled into the libIsoropia.a library (located in the LIB directory). Module files (*.mod) are copied to the MOD directory.

REMARKS:

To build the programs, call "make" with the following syntax:

```
make TARGET [ OPTIONAL-FLAGS ]
```

To display a complete list of options, type "make help".

Makefile uses the following variables:

Variable Description SHELL Specifies the shell for "make" to use (usually SHELL=/bin/sh) Specifies the root-level directory of the GEOS-Chem code ROOTDIR HDR Specifies the directory where GEOS-Chem include files are found Specifies the directory where library files (*.a) are stored LIB Specifies the directory where module files (*.mod) are stored MOD Sys var w/ name of library creator program (i.e., "ar", "ranlib") AR Sys var w/ name of Make command (i.e, "make" or "gmake") MAKE R8 Specifies the c

REVISION HISTORY:

```
21 Dec 2009 - C. Carouge - Initial version
22 Aug 2011 - R. Yantosca - Add "-fp-model source" flag for IFORT compiler,
which prevents random numerical noise
```

25 Aug 2011 - R. Yantosca - Remove -fp-model source flag here, as this is now added to FFLAGS in Makefile_header.mk

```
# Define variables
SHELL = /bin/sh
ROOTDIR = ...
    = $(ROOTDIR)/Headers
HDR.
HELP
     = $(ROOTDIR)/help
     = $(ROOTDIR)/lib
LIB
     = $(ROOTDIR)/mod
MOD
# Include header file. This returns variables CC, F90, FREEFORM, LD, R8,
# as well as the default Makefile compilation rules for source code files.
include $(ROOTDIR)/Makefile_header.mk
# List of files to compile (the order is important!). We specify these as
# a list of object files (*.o). For each object file, the "make" utility
# will find the corresponding source code file (*.F) and compile it.
# List of source files
SRC = $(wildcard *.F) $(wildcard *.F90)
# Replace .F and .F90 extensions with *.o
TMP = \$(SRC:.F=.o)
OBJ = (TMP:.F90=.0)
#-----
# Makefile targets: type "make help" for a complete listing!
.PHONY: clean help
lib: $(OBJ)
$(AR) crs libIsoropia.a $(OBJ)
mv libIsoropia.a $(LIB)
clean:
rm -f *.o *.mod
help:
@$(MAKE) -C $(HELP)
# Dependencies listing (grep "USE " to get the list of module references!)
```

1.4 Module Interface Makefile (in the GeosCore subdirectory)

This is the main GEOS-Chem makefile. It compiles the GEOS-Chem core source code files and bundles all of the object files (*.o) into the libGeosCore.a library (located in the LIB directory). Module files (*.mod) are copied to the MOD directory.

REMARKS:

To build the programs, call "make" with the following syntax:

```
make TARGET [ OPTIONAL-FLAGS ]
```

To display a complete list of options, type "make help".

Makefile uses the following variables:

Variable	Description
SHELL ROOTDIR BIN BPCH DOC	Specifies the shell for "make" to use (usually SHELL=/bin/sh) Specifies the root directory for the GEOS-Chem code Specifies the directory where executable files are stored Specifies the directory where the G-C bpch routines are stored Specifies the directory for generating desumentation w/ ProTeV
EXE	Specifies the directory for generating documentation w/ ProTeX Specifies the name of the executable file
HDR	Specifies the directory where include files are found
LIB	Specifies the directory where library files (*.a) are stored
LINK	Specifies the link commands to the GEOS-Chem library files
KPP	Specifies the directory where th KPP solver files reside
MOD	Specifies the directory where module files (*.mod) are stored
NCDF	Specifies the directory where netCDF utilities are stored
OBJ	Specifies the list of object files (*.o) to be created.
UTIL	Specifies the directory where the G-C utility modules are found

```
AR
             Sys var w/ name of library creator program (i.e., "ar", "ranlib")
  MAKE
             Sys var w/ name of Make command (i.e, "make" or "gmake")
             Cmd line argument; specifies either 43 or 54 tracer simulation
  NTRAC
  KPPSOLVER Cmd line argument; specifies the type of integrator to use
  NOTE: CC, F90, FREEFORM, LD, R8 are included from "Makefile_header.mk".
  %%% You can compile GEOS-Chem in parallel using the "make -jN" option!
                                                                       %%%
  %%%
                                                                       %%%
                                                                       %%%
  %%% N = number of proceses that you want to run simultaneously (i.e.
  %%% (when one file is finished compiling, "make" will immediately start
                                                                       %%%
                                                                       %%%
  %%% on the next one). Usually N is the # of processors on your system.
  REVISION HISTORY:
   16 Sep 2009 - R. Yantosca - Initial version
   18 Sep 2009 - P. Le Sager - Removed - LKppInt
   21 Sep 2009 - R. Yantosca - Now call Makefile in help directory to
                              display the help screen options
   19 Nov 2009 - R. Yantosca - Now compile the various GEOS-Chem utility
                              modules in the GeosUtil subdirectory
   19 Nov 2009 - R. Yantosca - Now compile the GEOS-Chem bpch module
                              separately in the GeosBpch subdirectory
   19 Nov 2009 - R. Yantosca - Now list all object dependencies explicitly,
                              to be able to use "make -j" (parallel make)
   23 Nov 2009 - R. Yantosca - Remove "main.o" explicitly form the "exe"
                              makefile target. This will be now compiled in
                              the proper sequence given the dependency
                              ordering. This allows "make -j" to work.
   23 Nov 2009 - R. Yantosca - Now don't copy module files; they will be
                              automatically written to the mod directory
   23 Nov 2009 - R. Yantosca - Removed libbpch; bundled that into libutil
   23 Nov 2009 - R. Yantosca - Added separate target libcore.
                                                           lib is now
                              a synonym for "libkpp libutil libcore"
   01 Dec 2009 - R. Yantosca - Modified the "exe" target for SunStudio compiler
                              which chokes at link time if the list of object
                              files is not explicitly passed
   02 Dec 2009 - R. Yantosca - Add conditional statements in dependencies list
                              for the SunStudio compiler
   11 Dec 2009 - R. Yantosca - Now get SHELL from Makefile_header.mk
   21 Dec 2009 - R. Yantosca - Now get LINK from Makefile_header.mk
   25 Jan 2010 - R. Yantosca - Whem making "realclean", also call "clean" in
                              the $(GEOSTOM)/Makefile. Also make sure to
                              remove executables in the $(BIN) directory.
   28 Jan 2010 - C. Carouge - Modifications for ISORROPIA II
   08 Feb 2010 - C. Carouge - Modifications for F. Paulot's isoprene scheme
```

10 May 2010 - R. Yantosca - Add dependency for RD_AOD.f

SRC := \$(wildcard *.F) \$(wildcard *.F90)

```
14 May 2010 - C. Carouge - Updates for mercury simulation
   20 Aug 2010 - R. Yantosca - Modifications for MERRA met fields
   16 Feb 2011 - R. Yantosca - Add modifications for APM (G. Luo)
   05 Aug 2011 - M. Long
                         - Now compile module files in Headers/ directory
   04 Nov 2011 - R. Yantosca - Remove references to ESMF subdirectory
   08 Dec 2011 - M. Payer - Remove obsolete GEIA biogenic emissions routines
   24 Jan 2012 - R. Yantosca - Also add libnc target to build netCDF utils
   25 Jan 2012 - R. Yantosca - Add ncdfcheck target to check netCDF install
   05 Apr 2012 - R. Yantosca - Now assume netCDF will always be used
   05 Apr 2012 - R. Yantosca - Now retire rdlai.F, readlai.F
   11 Apr 2012 - R. Yantosca - Now retire obsolete lai_mod.F
   11 Apr 2012 - R. Yantosca - Reference modis_lai_mod.F90 in timeseries diag
   11 Apr 2012 - R. Yantosca - Remove all references to obsolete lai_mod.F
   12 Apr 2012 - R. Yantosca - Remove reference to findmon.F
   19 Apr 2012 - R. Yantosca - Remove reference to rd_prof.F
# Define variables
ROOTDIR := ..
APM
       := $(ROOTDIR)/GeosApm
BIN
       := \$(ROOTDIR)/bin
BPCH
      := $(ROOTDIR)/GeosBpch
DOC
      := \$(ROOTDIR)/doc
EXE
      := geos
HDR
      := $(ROOTDIR)/Headers
HELP
      := $(ROOTDIR)/help
ISO
      := $(ROOTDIR)/ISOROPIA
LIB
      := $(ROOTDIR)/lib
KPP
      := $(ROOTDIR)/KPP
MOD
      := \$(ROOTDIR)/mod
NCDF
      := $(ROOTDIR)/NcdfUtil
TOM
      := $(ROOTDIR)/GeosTomas
GTMM
      := $(ROOTDIR)/GTMM
UTIL
       := $(ROOTDIR)/GeosUtil
# Include header file. This returns CC, F90, FREEFORM, LD, R8, SHELL,
# as well as the default Makefile compilation rules for source code files.
include $(ROOTDIR)/Makefile_header.mk
# List of files to compile (the order is important!). We specify these as
# a list of object files (*.o). For each object file, the "make" utility
# will find the corresponding source code file (*.f, *.f90) and compile it.
# List of source files
```

```
# Replace .f and .f90 extensions with *.o
TMP := \$(SRC:.F=.o)
OBJ := \$(TMP:.F90=.o)
# Makefile targets: type "make help" for a complete listing!
.PHONY: clean realclean doc docclean help distclean
                                            # Build libraries
all:
@$(MAKE) lib
                                     # and the executable
ifeq ($(UNAME),Darwin)
ranlib -c ../lib/*
endif
@$(MAKE) exe
lib:
                                            # Build all G-C libraries
@$(MAKE) libnc
@$(MAKE) libheaders
@$(MAKE) libkpp
@$(MAKE) libutil
@$(MAKE) libiso
@$(MAKE) libcore
libcore: $(OBJ)
                                            # Build code in GeosCore/
libiso:
                                            # Build code in ISOROPIA/
@$(MAKE) -C $(ISO)
libkpp:
                                            # Build code in KPP/
@$(MAKE) -C $(KPP)
libnc:
                                            # Build code in NcdfUtil/
@$(MAKE) -C $(NCDF) lib
ncdfcheck:
                                            # Check netCDF library
@$(MAKE) libnc
@$(MAKE) -C $(NCDF) check
libutil:
                                            # Build code in GeosUtil/
@$(MAKE) -C $(UTIL)
                                            # Build code in Headers/
libheaders:
@$(MAKE) -C $(HDR)
                                            # Build executable
exe:
$(LD) $(OBJ) $(LINK) -o $(EXE)
```

```
cp -f $(EXE) $(BIN)
                                            # Remove files here
clean:
rm -f *.o *.mod geos geosapm geostomas
distclean:
                                            # Synonym for "realclean"
@$(MAKE) realclean
realclean:
                                            # Remove files everywhere
@$(MAKE) clean
@$(MAKE) -C $(NCDF) clean
@$(MAKE) -C $(APM) clean
@$(MAKE) -C $(ISO) clean
@$(MAKE) -C $(KPP) realclean
@$(MAKE) -C $(TOM) clean
@$(MAKE) -C $(GTMM) clean
@$(MAKE) -C $(UTIL) clean
@$(MAKE) -C $(HDR) clean
@$(MAKE) docclean
rm -f $(LIB)/*.a
rm -f $(MOD)/*.mod
rm -f $(BIN)/geos*
                                            # Build documentation
doc:
@$(MAKE) -C $(DOC) all
docclean:
                                            # Remove documentation
@$(MAKE) -C $(DOC) clean
help:
                                            # Show help screen
@$(MAKE) -C $(HELP)
# Targets for mercury simulation (ccc, 6/7/10)
# Build Hg/GTMM executable
allhg:
@$(MAKE) libhg
@$(MAKE) exehg
libhg:
                                            # Compile HG/GTMM code
@$(MAKE) libgtmm
@$(MAKE) lib
libgtmm:
                                            # Compile GTMM model code
@$(MAKE) -C $(GTMM)
                                            # Create Hg/GTMM exec file
exehg:
```

\$(LD) \$(OBJ) \$(LHG) -o \$(EXE)

arsl1k.o

```
cp -f $(EXE) $(BIN)
# Dependencies listing (grep "USE " to get the list of module references!)
# From this list of dependencies, the "make" utility will figure out the
# correct order of compilation (so we don't have to do that ourselves).
# This also allows us to compile on multiple processors with "make -j".
# NOTES:
# (1) Only specify object-file dependencies that are within this directory.
      Object files in other directories will be referenced at link-time.
# (2) For "make -j" to work, all files in this directory must have a
      listed dependency.
#-----
                          : a3_read_mod.F
a3_read_mod.o
                                                   dao_mod.o
                            diag_mod.o
                                                   logical_mod.o
a6_read_mod.o
                          : a6_read_mod.F
                                                   dao_mod.o
                                                   logical_mod.o
                            diag_mod.o
acetone_mod.o
                          : acetone_mod.F
                                                   dao_mod.o
                            diag_mod.o
                                                   megan_mod.o
aerosol_mod.o
                          : aerosol_mod.F
                                                   comode_mod.o
                            dao_mod.o
                                                   diag_mod.o
                            logical_mod.o
                                                   tracerid_mod.o
                            tracer_mod.o
                                                   tropopause_mod.o
aircraft_nox_mod.o
                          : aircraft_nox_mod.F
                            dao_mod.o
                                                   diag_mod.o
                          : airmas.F
airmas.o
                          : anthroems.F
anthroems.o
                            future_emissions_mod.o edgar_mod.o
                            geia_mod.o
                                                   logical_mod.o
                            scale_anthro_mod.o
                                                   tracer_mod.o
                            tracerid_mod.o
arctas_ship_emiss_mod.o
                          : arctas_ship_emiss_mod.F
                            logical_mod.o
                                                   scale_anthro_mod.o
                            tracerid_mod.o
                                                   tracer_mod.o
```

: arsl1k.F

backsub.o	: backsub.F		
benchmark_mod.o	<pre>: benchmark_mod.F tracerid_mod.o</pre>	tracer_mod.o	\
biofuel_mod.o	<pre>: biofuel_mod.F dao_mod.o epa_nei_mod.o logical_mod.o tracerid_mod.o</pre>	<pre>diag_mod.o future_emissions_mod.o streets_anthro_mod.o tracer_mod.o</pre>	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
biofit.o	: biofit.F		
biomass_mod.o	<pre>: biomass_mod.F diag_mod.o gfed2_biomass_mod.o logical_mod.o tracerid_mod.o</pre>	<pre>gc_biomass_mod.o gfed3_biomass_mod.o tracer_mod.o</pre>	\\\\
BLKSLV.o	: BLKSLV.F		
boxvl.o	: boxvl.F	dao_mod.o	
bravo_mod.o	<pre>: bravo_mod.F future_emissions_mod.o scale_anthro_mod.o</pre>	<pre>logical_mod.o tracerid_mod.o</pre>	\
bromocarb_mod.o	<pre>: bromocarb_mod.F dao_mod.o logical_mod.o tracer_mod.o</pre>	<pre>diag_mod.o pbl_mix_mod.o</pre>	\ \
c2h6_mod.o	<pre>: c2h6_mod.F biofuel_mod.o dao_mod.o geia_mod.o logical_mod.o tracer_mod.o</pre>	biomass_mod.o diag_mod.o global_oh_mod.o tracerid_mod.o	////
cac_anthro_mod.o	<pre>: cac_anthro_mod.F future_emissions_mod.o scale_anthro_mod.o tracer_mod.o</pre>	<pre>logical_mod.o tracerid_mod.o</pre>	\ \
calcrate.o	<pre>: calcrate.F comode_mod.o diag_mod.o drydep_mod.o</pre>	dao_mod.o diag63_mod.o emissions_mod.o	\ \ \ \

	<pre>logical_mod.o planeflight_mod.o cldice_HBrHOBr_rxn.o</pre>	<pre>pbl_mix_mod.o tracerid_mod.o</pre>	\ \
carbon_mod.o	<pre>: carbon_mod.F biomass_mod.o dao_mod.o drydep_mod.o gfed2_biomass_mod.o global_no3_mod.o global_o3_mod.o logical_mod.o pbl_mix_mod.o vdiff_pre_mod.o</pre>	comode_mod.o diag_mod.o future_emissions_mod.o gfed3_biomass_mod.o global_oh_mod.o megan_mod.o tracerid_mod.o tropopause_mod.o meganut_mod.o	///////////////////////////////////////
ch3i_mod.o	<pre>: ch3i_mod.F biofuel_mod.o dao_mod.o diag_pl_mod.o tracerid_mod.o uvalbedo_mod.o</pre>	<pre>biomass_mod.o diag_mod.o logical_mod.o tracer_mod.o</pre>	\ \ \ \ \ \ \
chemdr.o	<pre>: chemdr.F aerosol_mod.o dao_mod.o diag_pl_mod.o dust_mod.o planeflight_mod.o tracer_mod.o tropopause_mod.o</pre>	comode_mod.o diag_oh_mod.o future_emissions_mod.o logical_mod.o restart_mod.o tracerid_mod.o uvalbedo_mod.o	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
<pre>chemistry_mod.o</pre>	: chemistry_mod.F acetone_mod.o c2h6_mod.o ch3i_mod.o dao_mod.o dust_mod.o h2_hd_mod.o isoropiaII_mod.o mercury_mod.o rpmares_mod.o seasalt_mod.o sulfate_mod.o tagged_ox_mod.o tracer_mod.o	aerosol_mod.o carbon_mod.o comode_mod.o drydep_mod.o global_ch4_mod.o hcn_ch3cn_mod.o logical_mod.o optdepth_mod.o RnPbBe_mod.o strat_chem_mod.o tagged_co_mod.o tracerid_mod.o	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
cleanup.o	: cleanup.F		\

	acetone_mod.o	aerosol_mod.o	\
	arctas_ship_emiss_mod.o		\
	biomass_mod.o	biofuel_mod.o	\
	bravo_mod.o	c2h6_mod.o	\
	cac_anthro_mod.o	carbon_mod.o	\
	co2_mod.o	comode_mod.o	\
	dao_mod.o	depo_mercury_mod.o	\
	diag_mod.o	diag03_mod.o	\
	diag04_mod.o	diag41_mod.o	\
	diag50_mod.o	diag51_mod.o	\
	diag51b_mod.o	diag_oh_mod.o	\
	diag_pl_mod.o	drydep_mod.o	\
	dust_mod.o	dust_dead_mod.o	\
	edgar_mod.o	emep_mod.o	\
	epa_nei_mod.o	gc_biomass_mod.o	\
	isoropiaII_mod.o	<pre>gfed2_biomass_mod.o</pre>	\
	gfed3_biomass_mod.o		\
	global_ch4_mod.o	global_hno3_mod.o	\
	global_no3_mod.o	global_nox_mod.o	\
	global_o1d_mod.o	global_oh_mod.o	\
	h2_hd_mod.o	hcn_ch3cn_mod.o	\
	modis_lai_mod.o	land_mercury_mod.o	\
	lightning_nox_mod.o	linoz_mod.o	\
	megan_mod.o	mercury_mod.o	\
	ocean_mercury_mod.o	pbl_mix_mod.o	\
	pjc_pfix_mod.o	planeflight_mod.o	\
	seasalt_mod.o	sulfate_mod.o	\
	tagged_co_mod.o	toms_mod.o	\
	tracer_mod.o	transport_mod.o	\
	tropopause_mod.o	uvalbedo_mod.o	\
	vistas_anthro_mod.o	wetscav_mod.o	\
	icoads_ship_mod.o	tpcore_fvdas_mod.o	\
	tpcore_geos5_window_mod.	_	\
	tpcore_geos57_window_mod		\
	retro_mod.o	strat_chem_mod.o	
	_		
	cldice_HBrHOBr_rxn.F		
	CLDSRF.F		
	co2_mod.F	biomass_mod.o	\
	diag04_mod.o	tracer_mod.o	\
Ċ	l_mod.o		

cldice_HBrHOBr_rxn.o

CLDSRF.o

co2_mod.o

tracerio

CO_strat_pl.o : CO_strat_pl.F

> dao_mod.o tracer_mod.o tracerid_mod.o tropopause_mod.o

comode_mod.o	: comode_mod.F	tracer_mod.o
convection_mod.o	<pre>: convection_mod.F dao_mod.o depo_mercury_mod.o gc_type_mod.o logical_mod.o tracer_mod.o wetscav_mod.o</pre>	<pre>diag_mod.o fvdas_convect_mod.o gcap_convect_mod.o mercury_mod.o tracerid_mod.o \</pre>
<pre>dao_mod.o gc_type_mod.o</pre>	: dao_mod.F logical_mod.o	tracer_mod.o \
decomp.o	: decomp.F	
depo_mercury_mod.o	<pre>: depo_mercury_mod.F dao_mod.o logical_mod.o</pre>	<pre>diag_mod.o \ tracerid_mod.o</pre>
diag03_mod.o	: diag03_mod.F	tracerid_mod.o
diag04_mod.o	: diag04_mod.F	
diag1.o	<pre>: diag1.F dao_mod.o tracer_mod.o tropopause_mod.o</pre>	<pre>diag_mod.o tracerid_mod.o ocean_mercury_mod.o</pre>
diag3.o	: diag3.F biofuel_mod.o diag_mod.o diag04_mod.o diag42_mod.o diag_pl_mod.o drydep_mod.o tracer_mod.o wetscav_mod.o	biomass_mod.o diag03_mod.o diag41_mod.o diag56_mod.o depo_mercury_mod.o logical_mod.o tracerid_mod.o
diag41_mod.o	: diag41_mod.F	pbl_mix_mod.o
diag42_mod.o	<pre>: diag42_mod.F dao_mod.o tracerid_mod.o</pre>	logical_mod.o \ tracer_mod.o
diag48_mod.o	<pre>: diag48_mod.F dao_mod.o tracerid_mod.o</pre>	<pre>pbl_mix_mod.o \ tracer_mod.o</pre>

diag49_mod.o	<pre>: diag49_mod.F dao_mod.o pbl_mix_mod.o tracer_mod.o</pre>	<pre>modis_lai_mod.o tracerid_mod.o</pre>	\ \
diag50_mod.o	<pre>: diag50_mod.F comode_mod.o logical_mod.o tracerid_mod.o tropopause_mod.o</pre>	<pre>dao_mod.o pbl_mix_mod.o tracer_mod.o</pre>	\ \ \
diag51_mod.o	<pre>: diag51_mod.F dao_mod.o logical_mod.o tracerid_mod.o tropopause_mod.o</pre>	<pre>modis_lai_mod.o pbl_mix_mod.o tracer_mod.o</pre>	\ \ \
diag51b_mod.o	<pre>: diag51b_mod.F dao_mod.o logical_mod.o tracerid_mod.o tropopause_mod.o</pre>	<pre>modis_lai_mod.o pbl_mix_mod.o tracer_mod.o</pre>	\ \ \
diag56_mod.o	: diag56_mod.F		
diag63_mod.o	<pre>: diag63_mod.F dao_mod.o tracerid_mod.o</pre>	<pre>pbl_mix_mod.o tracer_mod.o</pre>	\
diag_2pm.o	: diag_2pm.F diag_mod.o	tropopause_mod.o	\
diag_mod.o	: diag_mod.F		
diag_oh_mod.o	<pre>: diag_oh_mod.F comode_mod.o tracerid_mod.o</pre>	<pre>logical_mod.o tracer_mod.o</pre>	\
diag_pl_mod.o	<pre>: diag_pl_mod.F comode_mod.o tracerid_mod.o</pre>	<pre>logical_mod.o tracer_mod.o</pre>	\
drydep_mod.o	<pre>: drydep_mod.F dao_mod.o logical_mod.o tracerid_mod.o</pre>	<pre>comode_mod.o diag_mod.o pbl_mix_mod.o tracer_mod.o</pre>	\ \ \

meganut_mod.o

diagoh.o	: diagoh.F	diag_mod.o
dust_dead_mod.o	: dust_dead_mod.F	dao_mod.o
dust_mod.o	<pre>: dust_mod.F comode_mod.o diag_mod.o dust_dead_mod.o tracerid_mod.o</pre>	dao_mod.o \ drydep_mod.o \ logical_mod.o \ tracer_mod.o
edgar_mod.o	<pre>: edgar_mod.F future_emissions_mod.o scale_anthro_mod.o</pre>	<pre>logical_mod.o tracerid_mod.o</pre>
EFOLD.0	: EFOLD.F	
emep_mod.o	<pre>: emep_mod.F future_emissions_mod.o scale_anthro_mod.o tracer_mod.o</pre>	<pre>logical_mod.o \ tracerid_mod.o \</pre>
emf_scale.o	: emf_scale.F	tracerid_mod.o
emfossil.o retro_mod.o	<pre>: emfossil.F bravo_mod.o dao_mod.o edgar_mod.o epa_nei_mod.o logical_mod.o streets_anthro_mod.o tracerid_mod.o c2h6_mod.o</pre>	<pre>cac_anthro_mod.o diag_mod.o diag_mod.o emep_mod.o icoads_ship_mod.o nei2005_anthro_mod.o tracer_mod.o vistas_anthro_mod.o \</pre>
<pre>emissdr.o lightning_nox_mod.o bromocarb_mod.o</pre>	<pre>: emissdr.F acetone_mod.o biofuel_mod.o diag_mod.o logical_mod.o megan_mod.o tracerid_mod.o</pre>	aircraft_nox_mod.o \ dao_mod.o \ emissions_mod.o \ tracer_mod.o \ meganut_mod.o \
emissions_mod.o	<pre>: emissions_mod.F arctas_ship_emiss_mod.o bravo_mod.o cac_anthro_mod.o</pre>	biomass_mod.o \ c2h6_mod.o \ carbon_mod.o \

ch3i_mod.o

co2_mod.o

	dust_mod.o emep_mod.o global_ch4_mod.o hcn_ch3cn_mod.o logical_mod.o nei2005_anthro_mod.o retro_mod.o seasalt_mod.o sulfate_mod.o tracer_mod.o	edgar_mod.o epa_nei_mod.o h2_hd_mod.o icoads_ship_mod.o mercury_mod.o paranox_mod.o RnPbBe_mod.o streets_anthro_mod.o vistas_anthro_mod.o	`\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
epa_nei_mod.o	<pre>: epa_nei_mod.F future_emissions_mod.c scale_anthro_mod.o tracer_mod.o</pre>	logical_mod.o tracerid_mod.o	\ \
fast_j.o	<pre>: fast_j.F dao_mod.o</pre>	toms_mod.o	\
fertadd.o	: fertadd.F	logical_mod.o	
fjfunc.o	: fjfunc.F		
fjx_acet_mod.o	: fjx_acet_mod.F		
future_emissions_mod.o	: future_emissions_mod.F	,	
fvdas_convect_mod.o	<pre>: fvdas_convect_mod.F dao_mod.o depo_mercury_mod.o tracerid_mod.o</pre>	<pre>diag_mod.o logical_mod.o tracer_mod.o</pre>	\ \
fyhoro.o	: fyhoro.F		
fyrno3.o	: fyrno3.F		
<pre>gamap_mod.o ifeq (\$(COMPILER),sun)</pre>	: gamap_mod.F diag03_mod.o diag41_mod.o diag48_mod.o diag50_mod.o diag51b_mod.o diag_pl_mod.o logical_mod.o tracer_mod.o diag63_mod.o	diag04_mod.o diag42_mod.o diag49_mod.o diag51_mod.o diag56_mod.o drydep_mod.o tracerid_mod.o wetscav_mod.o	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
• • • •			

\$(F90) -00 -c \$<

endif			
gasconc.o	<pre>: gasconc.F comode_mod.o drydep_mod.o logical_mod.o</pre>	dao_mod.o tropopause_mod.o	\ \
GAUSSP.o	: GAUSSP.F		
gc_biomass_mod.o	<pre>: gc_biomass_mod.F future_emissions_mod.o tracerid_mod.o</pre>	<pre>logical_mod.o tracer_mod.o</pre>	\
gc_type_mod.o	: gc_type_mod.F		
<pre>gcap_convect_mod.o</pre>	<pre>: gcap_convect_mod.F dao_mod.o</pre>	diag_mod.o	\
gcap_read_mod.o	<pre>: gcap_read_mod.F diag_mod.o</pre>	dao_mod.o logical_mod.o	\
geia_mod.o	: geia_mod.F		
GEN. o	: GEN.F		
geos57_read_mod.o	: geos57_read_mod.F90 dao_mod.o		\
get_global_ch4.o	<pre>: get_global_ch4.F future_emissions_mod.o</pre>	logical_mod.o	\
getifsun.o	: getifsun.F	comode_mod.o	
gfed2_biomass_mod.o	<pre>: gfed2_biomass_mod.F future_emissions_mod.o tracer_mod.o</pre>	logical_mod.o tracerid_mod.o	\
gfed3_biomass_mod.o	<pre>: gfed3_biomass_mod.F future_emissions_mod.o tracer_mod.o</pre>	<pre>logical_mod.o tracerid_mod.o</pre>	\
global_br_mod.o	: global_br_mod.F	tropopause_mod.o	
global_ch4_mod.o	<pre>: global_ch4_mod.F dao_mod.o diag_oh_mod.o global_oh_mod.o</pre>	<pre>diag_mod.o diag_pl_mod.o logical_mod.o</pre>	\ \ \

	tracer_mod.o	vdiff_pre_mod.o	
global_hno3_mod.o	: global_hno3_mod.F dao_mod.o	tracer_mod.o	\
global_no3_mod.o	: global_no3_mod.F		
global_nox_mod.o	: global_nox_mod.F		
global_o1d_mod.o	: global_o1d_mod.F		
global_o3_mod.o	: global_o3_mod.F		
global_oh_mod.o	: global_oh_mod.F dao_mod.o		\
h2_hd_mod.o	: h2_hd_mod.F biofuel_mod.o dao_mod.o drydep_mod.o global_nox_mod.o global_oh_mod.o scale_anthro_mod.o tracerid_mod.o tropopause_mod.o	biomass_mod.o diag_mod.o geia_mod.o global_o1d_mod.o logical_mod.o tagged_co_mod.o tracer_mod.o meganut_mod.o	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
hcn_ch3cn_mod.o	<pre>: hcn_ch3cn_mod.F biomass_mod.o diag_mod.o global_oh_mod.o pbl_mix_mod.o</pre>	<pre>dao_mod.o geia_mod.o logical_mod.o tracerid_mod.o</pre>	\ \ \ \
i6_read_mod.o	: i6_read_mod.F dao_mod.o logical_mod.o	diag_mod.o	\
icoads_ship_mod.o	<pre>: icoads_ship_mod.F future_emissions_mod.o scale_anthro_mod.o tracer_mod.o</pre>	<pre>logical_mod.o tracerid_mod.o</pre>	\ \
initialize.o	: initialize.F diag_mod.o diag04_mod.o diag42_mod.o diag_pl_mod.o	diag03_mod.o diag41_mod.o diag56_mod.o logical_mod.o	\\\\

inphot.o : inphot.F

<pre>input_mod.o diag51_mod.o diag56_mod.o diag_pl_mod.o</pre>	: input_mod.F benchmark_mod.o depo_mercury_mod.o diag04_mod.o diag42_mod.o diag49_mod.o diag51b_mod.o diag_oh_mod.o drydep_mod.o emissions_mod.o gamap_mod.o logical_mod.o ocean_mercury_mod.o restart_mod.o tracerid_mod.o diag63_mod.o	biofuel_mod.o
isoropiaII_mod.o	<pre>: isoropiaII_mod.F dao_mod.o logical_mod.o tracer_mod.o</pre>	global_hno3_mod.o \ tracerid_mod.o \ tropopause_mod.o
jsparse.o	: jsparse.F	comode_mod.o
JRATET.o	: JRATET.F	fjx_acet_mod.o
JVALUE. o	: JVALUE.F	
jv_index.o	: jv_index.F	
ksparse.o	: ksparse.F	
land_mercury_mod.o	<pre>: land_mercury_mod.F biomass_mod.o depo_mercury_mod.o logical_mod.o</pre>	dao_mod.o \ modis_lai_mod.o \ tracerid_mod.o \
LEGNDO.o	: LEGNDO.F	
lightning_nox_mod.o	<pre>: lightning_nox_mod.F dao_mod.o diag_mod.o</pre>	<pre>diag56_mod.o logical_mod.o</pre>
linoz_mod.o	<pre>: linoz_mod.F dao_mod.o tracer_mod.o</pre>	tracerid_mod.o \ tropopause_mod.o

logical_mod.o	: logical_mod.F		
lump.o	: lump.F comode_mod.o	tracerid_mod.o	\
main.o	: main.F a3_read_mod.o benchmark_mod.o convection_mod.o diag_mod.o diag42_mod.o diag49_mod.o diag51_mod.o diag_oh_mod.o depo_mercury_mod.o emissions_mod.o gcap_read_mod.o lightning_nox_mod.o logical_mod.o pbl_mix_mod.o planeflight_mod.o transport_mod.o transport_mod.o vdiff_mod.o merra_cn_mod.o gc_environment_mod.o diag63_mod.o mapping_mod.o	a6_read_mod.o chemistry_mod.o comode_mod.o diag41_mod.o diag48_mod.o diag50_mod.o diag51b_mod.o dao_mod.o drydep_mod.o global_ch4_mod.o i6_read_mod.o modis_lai_mod.o linoz_mod.o megan_mod.o ocean_mercury_mod.o soaprod_mod.o tracer_mod.o tropopause_mod.o uvalbedo_mod.o merra_a1_mod.o merra_i6_mod.o olson_landmap_mod.o	///////////////////////////////////////
mapping_mod.o	: mapping_mod.F90 logical_mod.o		\
MATIN4.o	: MATIN4.F		
megan_mod.o	<pre>: megan_mod.F modis_lai_mod.o meganut_mod.o geos57_read_mod.o</pre>	a3_read_mod.o logical_mod.o merra_a1_mod.o	\ \
meganut_mod.o	: meganut_mod.F	dao_mod.o	
mercury_mod.o	<pre>: mercury_mod.F dao_mod.o diag03_mod.o</pre>	<pre>depo_mercury_mod.o diag_mod.o</pre>	\ \

	<pre>drydep_mod.o global_o3_mod.o land_mercury_mod.o logical_mod.o pbl_mix_mod.o tracerid_mod.o tropopause_mod.o</pre>	<pre>global_br_mod.o global_oh_mod.o ocean_mercury_mod.o RnPbBe_mod.o tracer_mod.o vdiff_pre_mod.o</pre>	\ \ \ \
merra_a1_mod.o	<pre>: merra_a1_mod.F logical_mod.o</pre>	dao_mod.o	\
merra_a3_mod.o	<pre>: merra_a3_mod.F logical_mod.o</pre>	dao_mod.o	\
merra_cn_mod.o	<pre>: merra_cn_mod.F logical_mod.o</pre>	dao_mod.o	\
merra_i6_mod.o	<pre>: merra_i6_mod.F logical_mod.o</pre>	dao_mod.o	\
MIESCT.o	: MIESCT.F		
modis_lai_mod.o	<pre>: modis_lai_mod.F90 mapping_mod.o</pre>		\
ndxx_setup.o	<pre>: ndxx_setup.F biofuel_mod.o diag_oh_mod.o logical_mod.o tracer_mod.o wetscav_mod.o</pre>	<pre>diag_mod.o drydep_mod.o planeflight_mod.o tracerid_mod.o diag63_mod.o</pre>	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
nei2005_anthro_mod.o	<pre>: nei2005_anthro_mod.F future_emissions_mod.o scale_anthro_mod.o tracer_mod.o</pre>	<pre>logical_mod.o tracerid_mod.o</pre>	\ \
NOABS.o	: NOABS.F		
ocean_mercury_mod.o	<pre>: ocean_mercury_mod.F dao_mod.o diag03_mod.o tracerid_mod.o</pre>	depo_mercury_mod.o logical_mod.o tracer_mod.o	\ \
ohsave.o	: ohsave.F diag_mod.o	<pre>comode_mod.o tracerid_mod.o</pre>	\
olson_landmap_mod.o	: olson_landmap_mod.F90		\

mapping_mod.o

optdepth_mod.o : optdepth_mod.F diag_mod.o

OPMIE.o : OPMIE.F

paranox_mod.o : paranox_mod.F

dao_mod.o tracer_mod.o \

tracerid_mod.o

partition.o : partition.F

comode_mod.o tracerid_mod.o

pbl_mix_mod.o : pbl_mix_mod.F \

dao_mod.o diag_mod.o

logical_mod.o tracer_mod.o

pderiv.o : pderiv.F

photoj.o : photoj.F

physproc.o : physproc.F \

comode_mod.o logical_mod.o

chemistry_mod.o

 $\verb|pjc_pfix_geos5_window_mod.o|: \verb|pjc_pfix_geos5_window_mod.F||$

pjc_pfix_geos57_window_mod.o: pjc_pfix_geos57_window_mod.F

pjc_pfix_mod.o : pjc_pfix_mod.F

planeflight_mod.o : planeflight_mod.F

comode_mod.o dao_mod.o

precipfrac.o : precipfrac.F dao_mod.o

pulsing.o : pulsing.F

RD_AOD.o : RD_AOD.F

rd_js.o : rd_js.F

rdsoil.o : rdsoil.F

RD_TJPL.o : RD_TJPL.F

readchem.o : readchem.F

	<pre>diag_pl_mod.o logical_mod.o</pre>	drydep_mod.o	\
reader.o	: reader.F		
read_jv_atms_dat.o	: read_jv_atms_dat.F90		
restart_mod.o	: restart_mod.F		\
	comode_mod.o	dao_mod.o	\
	logical_mod.o	tracer_mod.o	
retro_mod.o	: retro_mod.F		\
	future_emissions_mod.o	· ·	\
	scale_anthro_mod.o	tracerid_mod.o	\
	tracer_mod.o		
RnPbBe_mod.o	: RnPbBe_mod.F		\
	dao_mod.o	diag_mod.o	\
	logical_mod.o	tracer_mod.o	\
	tropopause_mod.o		
rpmares_mod.o	: rpmares_mod.F		\
	dao_mod.o	<pre>global_hno3_mod.o</pre>	\
	tracerid_mod.o	tracer_mod.o	\
	tropopause_mod.o		
ruralbox.o	: ruralbox.F		\
	comode_mod.o	tropopause_mod.o	
scale_anthro_mod.o	: scale_anthro_mod.F		
seasalt_mod.o	: seasalt_mod.F		\
	dao_mod.o	diag_mod.o	\
	drydep_mod.o	logical_mod.o	\
	pbl_mix_mod.o	tracerid_mod.o	\
	<pre>tracer_mod.o ssa_bromine_mod.o</pre>	vdiff_pre_mod.o	\
	ssa_bromine_mod.o		
set_aer.o	: set_aer.F		
set_prof.o	: set_prof.F		\
	dao_mod.o	toms_mod.o	
setemdep.o	: setemdep.F	drydep_mod.o	\
	tracer_mod.o	tracerid_mod.o	
actomia o	. gotomia E		`
setemis.o	<pre>: setemis.F aircraft_nox_mod.o</pre>	biofuel_mod.o	\
	arrerare_moa.u	DIOI del_mod.O	`

		<pre>biomass_mod.o diag_mod.o lightning_nox_mod.o pbl_mix_mod.o tropopause_mod.o</pre>	<pre>comode_mod.o emissions_mod.o logical_mod.o tracerid_mod.o</pre>	\ \ \
setmodel.o	:	setmodel.F		
sfcwindsqr.o	:	sfcwindsqr.F	dao_mod.o	
smvgear.o	:	smvgear.F	comode_mod.o	
soaprod_mod.o	:	<pre>soaprod_mod.F carbon_mod.o logical_mod.o</pre>	dao_mod.o tracer_mod.o	\
soilbase.o	:	soilbase.F		
soilcrf.o	:	soilcrf.F		
soilnoxems.o	:	<pre>soilnoxems.F dao_mod.o future_emissions_mod.o meganut_mod.o</pre>	<pre>diag_mod.o logical_mod.o</pre>	\ \
soiltemp.o	:	soiltemp.F		
soiltype.o	:	soiltype.F		
SPHERE.o	:	SPHERE.F		
ssa_bromine_mod.o	:	<pre>ssa_bromine_mod.F bromocarb_mod.o logical_mod.o tracerid_mod.o</pre>	<pre>comode_mod.o pbl_mix_mod.o tropopause_mod.o</pre>	\ \
streets_anthro_mod.o	:	streets_anthro_mod.F future_emissions_mod.o scale_anthro_mod.o tracer_mod.o	<pre>logical_mod.o tracerid_mod.o</pre>	\ \
strat_chem_mod.o	:	strat_chem_mod.F90 dao_mod.o linoz_mod.o tracer_mod.o tropopause_mod.o	<pre>logical_mod.o tagged_ox_mod.o tracerid_mod.o</pre>	\ \ \ \

subfun.o : subfun.F

<pre>sulfate_mod.o</pre>	: sulfate_mod.F arctas_ship_emiss_mod.o bravo_mod.o comode_mod.o diag_mod.o edgar_mod.o epa_nei_mod.o gfed2_biomass_mod.o global_hno3_mod.o icoads_ship_mod.o nei2005_anthro_mod.o scale_anthro_mod.o streets_anthro_mod.o tracer_mod.o uvalbedo_mod.o wetscav_mod.o	biomass_mod.o cac_anthro_mod.o dao_mod.o drydep_mod.o emep_mod.o future_emissions_mod.o gfed3_biomass_mod.o global_oh_mod.o logical_mod.o pbl_mix_mod.o seasalt_mod.o tracerid_mod.o tropopause_mod.o vdiff_pre_mod.o	///////////////////////////////////////
sunparam.o	: sunparam.F		
<pre>gc_type2_mod.o gc_environment_mod.o gc_type_mod.o</pre>	<pre>: gc_type2_mod.F90 : gc_environment_mod.F90</pre>	<pre>tracer_mod.o gc_type2_mod.o</pre>	\
tagged_co_mod.o	<pre>: tagged_co_mod.F biofuel_mod.o dao_mod.o diag_pl_mod.o global_oh_mod.o megan_mod.o pbl_mix_mod.o tracer_mod.o</pre>	biomass_mod.o diag_mod.o global_nox_mod.o logical_mod.o meganut_mod.o tracerid_mod.o tropopause_mod.o	\\\\\\
tagged_ox_mod.o	<pre>: tagged_ox_mod.F dao_mod.o diag_pl_mod.o logical_mod.o pbl_mix_mod.o tracer_mod.o</pre>	<pre>diag_mod.o drydep_mod.o meganut_mod.o tracerid_mod.o tropopause_mod.o</pre>	\ \ \ \ \ \
tcorr.o	: tcorr.F		
toms_mod.o	: toms_mod.F		
tpcore_bc_mod.o	<pre>: tpcore_bc_mod.F logical_mod.o</pre>	tracer_mod.o	\

tpcore_geos5_window_mod.o : tpcore_geos5_window_mod.F90 \$(F90) -c \$(FREEFORM) \$(R8) \$< tpcore_geos57_window_mod.o : tpcore_geos57_window_mod.F90 \$(F90) -c \$(FREEFORM) \$(R8) \$< : tpcore_mod.F tpcore_mod.o dao_mod.o diag_mod.o global_ch4_mod.o tracer_mod.o \$(F90) -c \$(R8) \$< tpcore_window_mod.o : tpcore_window_mod.F dao_mod.o diag_mod.o global_ch4_mod.o tracer_mod.o \$(F90) -c \$(R8) \$< : tracer_mod.F tracer_mod.o tracerid_mod.o : tracerid_mod.F gc_type2_mod.o logical_mod.o tracer_mod.o gc_environment_mod.o transport_mod.o : transport_mod.F dao_mod.o diag_mod.o pjc_pfix_mod.o logical_mod.o tpcore_mod.o tpcore_bc_mod.o tpcore_fvdas_mod.o tpcore_window_mod.o tracer_mod.o tpcore_geos5_window_mod.o tpcore_geos57_window_mod.o pjc_pfix_geos5_window_mod.o pjc_pfix_geos57_window_mod.o : tropopause.F tropopause.o dao_mod.o diag_mod.o logical_mod.o tropopause_mod.o tropopause_mod.o : tropopause_mod.F comode_mod.o dao_mod.o diag_mod.o logical_mod.o : update.F update.o : vdiff_pre_mod.F vdiff_pre_mod.o tracer_mod.o ifeq (\$(COMPILER),sun) \$(F90) -03 -c \$< endif

vdiff_mod.o : vdiff_mod.F90 comode_mod.o dao_mod.o diag_mod.o depo_mercury_mod.o drydep_mod.o logical_mod.o ocean_mercury_mod.o pbl_mix_mod.o tracer_mod.o tracerid_mod.o vdiff_pre_mod.o vistas_anthro_mod.o : vistas_anthro_mod.F future_emissions_mod.o epa_nei_mod.o logical_mod.o scale_anthro_mod.o tracerid_mod.o tracer_mod.o : wetscav_mod.F wetscav_mod.o dao_mod.o diag_mod.o depo_mercury_mod.o logical_mod.o mercury_mod.o ocean_mercury_mod.o tracerid_mod.o tracer_mod.o : XSEC1D.F XSEC1D.o XSECO2.o : XSECO2.F XSECO3.o : XSECO3.F

1.5 Module Interface Makefile (in the KPP subdirectory)

This is main "router" makefile for the KPP solver. It compiles the KPP code for one of the following types of GEOS-Chem simulations:

- 1. GEOS-Chem "standard" simulation (43 tracers)
- 2. GEOS-Chem "secondary organic aerosol" simulation (54 tracers)

The KPP code will be compiled using one of the following numerical solvers:

- 1. rosenbrock (This is the default option.)
- 2. lsodes
- 3. radau5
- 4. runge_kutta

REMARKS:

To build the programs, call "make" with the following syntax:

```
make TARGET [ OPTIONAL-FLAGS ]
```

To display a complete list of options, type "make help".

Makefile uses the following variables:

```
Variable
          Description
_____
          -----
SHELL
          Specifies the shell for "make" to use (usually SHELL=/bin/sh)
ROOTDIR
          Specifies the root-level directory of the GEOS-Chem code
DOC
          Specifies the directory where GEOS-Chem documentation is found
HDR
          Specifies the directory where GEOS-Chem include files are found
LIB
          Specifies the directory where library files (*.a) are stored
MOD
          Specifies the directory where module files (*.mod) are stored
AR
          Sys var w/ name of library creator program (i.e., "ar", "ranlib")
          Sys var w/ name of Make command (i.e, "make" or "gmake")
MAKE
          Cmd line argument; specifies either 43 or 54 tracer simulation
NTRAC
KPPSOLVER Cmd line argument; specifies the type of integrator to use
```

REVISION HISTORY:

```
16 Sep 2009 - R. Yantosca - Initial version
18 Sep 2009 - P. Le Sager - Added kppintegrator target & commented
"make -C int" calls
20 Nov 2009 - P. Le Sager - Added CHEM option
23 Nov 2009 - R. Yantosca - Added realclean target
11 Dec 2009 - R. Yantosca - Now get SHELL from Makefile_header.mk
```

Define variables

ROOTDIR = ...

DOC = \$(ROOTDIR)/doc HDR = \$(ROOTDIR)/Headers HELP = \$(ROOTDIR)/help LIB = \$(ROOTDIR)/lib MOD = \$(ROOTDIR)/mod

Include header file. This returns CC, F90, FREEFORM, LD, R8, SHELL,
as well as the default Makefile compilation rules for source code files.
include \$(ROOTDIR)/Makefile_header.mk

```
# Make the standard 43-tracer simulation the default
ifndef CHEM
CHEM = standard
endif
# Check if NTRAC option is used
ifdef NTRAC
ifeq ($(NTRAC),43)
CHEM = standard
endif
ifeq ($(NTRAC),54)
CHEM = SOA
endif
endif
# Make rosenbrock the default solver
ifndef KPPSOLVER
KPPSOLVER = rosenbrock
endif
# solver (S=Source, T=Target)
SOLVER_SFILE=./int/gckpp_Integrator_$(KPPSOLVER).F90
SOLVER_TFILE=./$(CHEM)/gckpp_Integrator.F90
# Makefile targets: type "make help" for a complete listing!
.PHONY: all lib kppintegrator clean realclean doc help
all: lib
lib: kppintegrator
@$(MAKE) -C $(CHEM)
kppintegrator:
@diff $(SOLVER_SFILE) $(SOLVER_TFILE) ;\
if [ $$? == 1 ] ; then
echo " copy $(SOLVER_SFILE) --> $(SOLVER_TFILE)";\
cp $(SOLVER_SFILE) $(SOLVER_TFILE) ;
fi
clean:
```

```
@$(MAKE) -C $(CHEM) clean
realclean:
@$(MAKE) -C standard clean
@$(MAKE) -C SOA clean
@$(MAKE) -C isoprene clean
help:
@$(MAKE) -C $(HELP)
```

1.6 Module Interface Makefile (in the KPP/43t subdirectory)

This makefile compiles the KPP solver code for the GEOS-Chem 43 tracer simulation (i.e. without secondary organic aerosol tracers). Object files (*.o) are bundled into the libKpp.a library (located in the LIB directory). Module files (*.mod) are copied to the MOD directory.

REMARKS:

```
To build the programs, call "make" with the following syntax:
```

```
make TARGET [ OPTIONAL-FLAGS ]
```

To display a complete list of options, type "make help".

Makefile uses the following variables:

```
Variable
           Description
           Specifies the shell for "make" to use (usually SHELL=/bin/sh)
SHELL
           Specifies the root-level directory of the GEOS-Chem code
ROOTDIR
           Specifies the directory where GEOS-Chem include files are found
HDR
           Specifies the directory where library files (*.a) are stored
LIB
MOD
           Specifies the directory where module files (*.mod) are stored
           Sys var w/ name of library creator program (i.e., "ar", "ranlib")
AR
MAKE
           Sys var w/ name of Make command (i.e, "make" or "gmake")
```

REVISION HISTORY:

```
16 Sep 2009 - R. Yantosca - Initial version
```

```
21 Sep 2009 - R. Yantosca - Now call Makefile in help directory to
                       display the help screen options
  23 Nov 2009 - R. Yantosca - Now don't copy module files; they will be
                       automatically written to the mod directory
   11 Dec 2009 - R. Yantosca - Now get SHELL from Makefile_header.mk
# Define variables
ROOTDIR = ../..
HDR
     = $(ROOTDIR)/Headers
     = $(ROOTDIR)/help
HELP
LIB
     = (ROOTDIR)/lib
MOD
     = \$(ROOTDIR)/mod
# Include header file. This returns CC, F90, FREEFORM, LD, R8, SHELL,
# as well as the default Makefile compilation rules for source code files.
include $(ROOTDIR)/Makefile_header.mk
# List of files to compile. Here the order is not important,
# as we will explicity define the dependencies listing below.
# Source code files
SRC = $(wildcard gckpp*.F90)
# Object files
OBJ = \$(SRC:.F90=.o)
# Makefile targets: type "make help" for a complete listing!
.PHONY: clean help
lib: $(OBJ)
$(AR) crs libKpp.a $(OBJ)
mv libKpp.a $(LIB)
clean:
rm -f *.o *.mod geos
help:
@$(MAKE) -C $(HELP)
# Dependencies listing (grep "USE " to get the list of module references!)
```

From this list of dependencies, the "make" utility will figure out the

```
# correct order of compilation (so we don't have to do that ourselves!)
gckpp_Function.o
                      : gckpp_Parameters.o
gckpp_Global.o
                      : gckpp_Parameters.o
gckpp_Hessian.o
                      : gckpp_Parameters.o
                        gckpp_HessianSP.o
gckpp_Initialize.o
                      : gckpp_Parameters.o
                        gckpp_Global.o
                        gckpp_Util.o
                        gckpp_Monitor.o
gckpp_Integrator.o
                      : gckpp_Parameters.o
                        gckpp_Global.o
                        gckpp_Function.o
                        gckpp_Rates.o
                        gckpp_Jacobian.o
                        gckpp_LinearAlgebra.o
gckpp_Jacobian.o
                      : gckpp_Parameters.o
                        gckpp_JacobianSP.o
gckpp_LinearAlgebra.o : gckpp_Parameters.o
                        gckpp_JacobianSP.o
gckpp_Model.o
                      : gckpp_Precision.o
                        gckpp_Parameters.o
                        gckpp_Global.o
                        gckpp_Function.o
                        gckpp_Integrator.o
                        gckpp_Rates.o
                        gckpp_Jacobian.o
                        gckpp_Hessian.o
                        gckpp_Stoichiom.o
                        gckpp_Monitor.o
                        gckpp_Util.o
                        gckpp_LinearAlgebra.o
                      : gckpp_Precision.o
gckpp_Parameters.o
#gckpp_Rates.o
                       : gckpp_Parameters.o
                         gckpp_Global.o
#
                         gckpp_Monitor.o
#
                         gckpp_comode_mod.o
```

1.7 Module Interface Makefile (in the KPP/SOA subdirectory)

This makefile compiles the KPP solver code for the GEOS-Chem SOA simulation (with aromatic formation of secondary organic aerosol tracers). Object files (*.o) are bundled into the libKpp.a library (located in the LIB directory). Module files (*.mod) are copied to the MOD directory.

REMARKS:

To build the programs, call "make" with the following syntax:

Makefile uses the following variables:

Variable	Description
SHELL	Specifies the shell for "make" to use (usually SHELL=/bin/sh)
ROOTDIR	Specifies the root-level directory of the GEOS-Chem code
HDR	Specifies the directory where GEOS-Chem include files are found
LIB	Specifies the directory where library files (*.a) are stored
MOD	Specifies the directory where module files (*.mod) are stored
AR	Sys var w/ name of library creator program (i.e., "ar", "ranlib")
MAKE	Sys var w/ name of Make command (i.e, "make" or "gmake")

REVISION HISTORY:

```
16 Sep 2009 - R. Yantosca - Initial version
   21 Sep 2009 - R. Yantosca - Now call Makefile in help directory to
                         display the help screen options
   23 Nov 2009 - R. Yantosca - Now don't copy module files; they will be
                         automatically written to the mod directory
   11 Dec 2009 - R. Yantosca - Now get SHELL from Makefile_header.mk
# Define variables
ROOTDIR = ../..
     = $(ROOTDIR)/Headers
HDR
HELP
      = $(ROOTDIR)/help
      = (ROOTDIR)/lib
MOD
      = \$(ROOTDIR)/mod
# Include header file. This returns CC, F90, FREEFORM, LD, R8, SHELL,
# as well as the default Makefile compilation rules for source code files.
include $(ROOTDIR)/Makefile_header.mk
# List of files to compile. Here the order is not important,
# as we will explicity define the dependencies listing below.
# Source code files
SRC = $(wildcard gckpp*.F90)
# Object files
OBJ = \$(SRC:.F90=.0)
# Makefile targets: type "make help" for a complete listing!
.PHONY: clean help
lib: $(OBJ)
$(AR) crs libKpp.a $(OBJ)
mv libKpp.a $(LIB)
clean:
rm -f *.o *.mod geos
help:
@$(MAKE) -C $(HELP)
# Dependencies listing (grep "USE " to get the list of module references!)
```

```
#
# From this list of dependencies, the "make" utility will figure out the
# correct order of compilation (so we don't have to do that ourselves!)
gckpp_Function.o
                      : gckpp_Parameters.o
gckpp_Global.o
                      : gckpp_Parameters.o
gckpp_Hessian.o
                      : gckpp_Parameters.o
                        gckpp_HessianSP.o
gckpp_Initialize.o
                      : gckpp_Parameters.o
                        gckpp_Global.o
                        gckpp_Util.o
                        gckpp_Monitor.o
gckpp_Integrator.o
                      : gckpp_Parameters.o
                        gckpp_Global.o
                        gckpp_Function.o
                        gckpp_Rates.o
                        gckpp_Jacobian.o
                        gckpp_LinearAlgebra.o
                      : gckpp_Parameters.o
gckpp_Jacobian.o
                        gckpp_JacobianSP.o
gckpp_LinearAlgebra.o : gckpp_Parameters.o
                        gckpp_JacobianSP.o
gckpp_Model.o
                      : gckpp_Precision.o
                        gckpp_Parameters.o
                        gckpp_Global.o
                        gckpp_Function.o
                        gckpp_Integrator.o
                        gckpp_Rates.o
                        gckpp_Jacobian.o
                        gckpp_Hessian.o
                        gckpp_Stoichiom.o
                        gckpp_Monitor.o
                        gckpp_Util.o
                        gckpp_LinearAlgebra.o
gckpp_Parameters.o
                      : gckpp_Precision.o
gckpp_Rates.o
                      : gckpp_Parameters.o
                        gckpp_Global.o
                        gckpp_Monitor.o
```

```
gckpp_Stoichiom.o
                      : gckpp_Parameters.o
                        gckpp_StoichiomSP.o
gckpp_Util.o
                      : gckpp_Parameters.o
                        gckpp_Global.o
                        gckpp_Monitor.o
```

Module Interface Makefile (in the GeosTomas subdirectory) 1.8

This is the main makefile for GEOS-Chem + TOMAS aerosol microphysics. It compiles the GEOS-Chem core source code files and into object files (*.o). Module files (*.mod) are copied to the MOD directory.

REMARKS:

To build the programs, call "make" with the following syntax:

```
make TARGET [ OPTIONAL-FLAGS ]
To display a complete list of options, type "make help".
```

%%% NOTE: Most of the time this Makefile will be called automatically %%% ""," from the router Makefile in the top-level directory. However, if %%% %% you are in the ./GeosCore directory, then you can call this Makefile %%%%%% to build the GEOS-Chem source code, libraries, and executables. %%%

Makefile uses the following variables:

Variable	Description
SHELL	Specifies the shell for "make" to use (usually SHELL=/bin/sh)
ROOTDIR	Specifies the root directory for the GEOS-Chem code
BIN	Specifies the directory where executable files are stored
BPCH	Specifies the directory where the G-C bpch routines are stored
DOC	Specifies the directory for generating documentation w/ ProTeX
EXE	Specifies the name of the executable file
HDR	Specifies the directory where include files are found
LIB	Specifies the directory where library files (*.a) are stored
LINK	Specifies the link commands to the GEOS-Chem library files
KPP	Specifies the directory where th KPP solver files reside
MOD	Specifies the directory where module files (*.mod) are stored
OBJ	Specifies the list of object files (*.o) to be created.

```
UTIL
            Specifies the directory where the G-C utility modules are found
  AR
            Sys var w/ name of library creator program (i.e., "ar", "ranlib")
            Sys var w/ name of Make command (i.e, "make" or "gmake")
  MAKE.
  NTRAC
            Cmd line argument; specifies either 43 or 54 tracer simulation
  KPPSOLVER Cmd line argument; specifies the type of integrator to use
  NOTE: CC, F90, FREEFORM, LD, R8 are included from "Makefile_header.mk".
  %%% You can compile GEOS-Chem in parallel using the "make -jN" option!
                                                                     %%%
                                                                     %%%
  %%%
  %%% N = number of proceses that you want to run simultaneously (i.e.
                                                                     %%%
  %%% (when one file is finished compiling, "make" will immediately start
                                                                    %%%
  %%% on the next one). Usually N is the # of processors on your system.
                                                                     %%%
  REVISION HISTORY:
   25 Jan 2010 - R. Yantosca - Initial version for TOMAS microphysics
   28 Jan 2010 - C. Carouge - Modifications for ISORROPIA II
   10 May 2010 - R. Yantosca - Add dependency for RD_AOD.f
   26 Aug 2010 - R. Yantosca - Modifications for MERRA met fields
   23 Sep 2010 - R. Yantosca - Removed "tropopause.f" from compile list
   23 Sep 2010 - R. Yantosca - Updated dependencies for v9-01-01
   06 Dec 2011 - R. Yantosca - Remove #ifdef blocks for GI model code
                           - Remove references to ESMF directory
   16 Feb 2012 - M. Payer
   05 Apr 2012 - R. Yantosca - Always assume netCDF will be used
   05 Apr 2012 - R. Yantosca - Added c2h6_mod.o to emfossil.F dependency
   05 Apr 2012 - R. Yantosca - Now compile the NcdfUtil routines
```

05 Apr 2012 - R. Yantosca - Remove reference to rdland.F, rdlai.F, readlai.F

12 Apr 2012 - R. Yantosca - Remove reference to findmon.F 19 Apr 2012 - R. Yantosca - Remove reference to rd_prof.F

Define variables

ROOTDIR := ...

APM:= \$(ROOTDIR)/APM := \$(ROOTDIR)/binBIN **BPCH** := \$(ROOTDIR)/GeosBpch CORE := \$(ROOTDIR)/GeosCore DOC := \$(ROOTDIR)/docEXE := geostomas HDR := \$(ROOTDIR)/Headers

HELP := \$(ROOTDIR)/help ISO := \$(ROOTDIR)/ISOROPIA := \$(ROOTDIR)/lib LIB KPP := \$(ROOTDIR)/KPPMOD := \$(ROOTDIR)/mod:= \$(ROOTDIR)/NcdfUtil NCDF

UTIL := \$(ROOTDIR)/GeosUtil GTMM := \$(ROOTDIR)/GTMM

This directory only contains files that are different for the TOMAS # aerosol microphysics. For files that are the same as for the regular # GEOS-Chem code, look in the GeosCore directory. := ../GeosCore VPATH

Include header file. This returns CC, F90, FREEFORM, LD, R8, SHELL, # as well as the default Makefile compilation rules for source code files. include \$(ROOTDIR)/Makefile_header.mk

List of files to compile. We need to manually list all object files, since # some of these will be referenced from the ../GeosCore directory. OB.J =BLKSLV.o drydep_mod.o partition.o CLDSRF.o dust_dead_mod.o pbl_mix_mod.o CO_strat_pl.o dust_mod.o pderiv.o EFOLD.o edgar_mod.o photoj.o FLINT.o emep_mod.o physproc.o GAUSSP.o emf_scale.o pjc_pfix_geos5_window_mod.o \ GEN.o emfossil.o pjc_pfix_mod.o JRATET.o planeflight_mod.o geos57_read_mod.o JVALUE.o precipfrac.o olson_landmap_mod.o LEGNDO.o pulsing.o mapping_mod.o MATIN4.o emissdr.o rd_js.o MIESCT.o emissions_mod.o NOABS.o modis_lai_mod.o OPMIE.o epa_nei_mod.o RD TJPL.o fast_j.o RnPbBe_mod.o fertadd.o SPHERE. o XSEC1D.o fjfunc.o rdsoil.o XSECO2.o fjx_acet_mod.o readchem.o XSECO3.o future_emissions_mod.o reader.o a3_read_mod.o fvdas_convect_mod.o a6_read_mod.o fyhoro.o restart_mod.o acetone_mod.o fyrno3.o rpmares_mod.o aerosol_mod.o gamap_mod.o ruralbox.o aircraft_nox_mod.o gasconc.o scale_anthro_mod.o airmas.o gc_biomass_mod.o anthroems.o gcap_convect_mod.o seasalt_mod.o arctas_ship_emiss_mod.o gcap_read_mod.o set_aer.o arsl1k.o geia_mod.o set_prof.o backsub.o gfed3_biomass_mod.o

get_global_ch4.o

setemdep.o

getifsun.o

benchmark_mod.o

biofit.o	gfed2_biomass_mod.o	setemis.o	\		
biofuel_mod.o	global_ch4_mod.o	setmodel.o	ί.		
biomass_mod.o	global_hno3_mod.o	sfcwindsqr.o	ĺ		
boxv1.o	global_no3_mod.o	smvgear.o	Ì		
bravo_mod.o	global_nox_mod.o	soaprod_mod.o	Ì		
c2h6_mod.o	global_o1d_mod.o	soilbase.o	Ì		
cac_anthro_mod.o	global_o3_mod.o	soilcrf.o	Ì		
calcrate.o	global_oh_mod.o	soilnoxems.o	ĺ		
carbon_mod.o	soiltemp.o		ĺ		
ch3i_mod.o	h2_hd_mod.o	soiltype.o	ĺ		
chemdr.o	hcn_ch3cn_mod.o	streets_anthro_mod.o	ί.		
chemistry_mod.o	i6_read_mod.o	subfun.o	ĺ		
cleanup.o	icoads_ship_mod.o	sulfate_mod.o	ĺ		
co2_mod.o	initialize.o	sunparam.o	ĺ		
comode_mod.o	inphot.o	tagged_co_mod.o	΄,		
convection_mod.o	input_mod.o	tagged_ox_mod.o	΄,		
dao_mod.o	isoropiaII_mod.o	odggod_on_mou.o	`		
decomp.o	jsparse.o	toms_mod.o	`		
diag03_mod.o	jv_index.o	tpcore_bc_mod.o	′		
diag04_mod.o	ksparse.o	tpcore_fvdas_mod.o	′		
diag1.o	tpcore_geos5_window_mod	-	`		
diag3.o	lightning_nox_mod.o	tpcore_mod.o	`		
diag41_mod.o	linoz_mod.o	tpcore_window_mod.o	`		
diag42_mod.o	logical_mod.o	tracer_mod.o	ĺ		
diag48_mod.o	lump.o	tracerid_mod.o	`		
diag49_mod.o	main.o	transport_mod.o	′		
diag50_mod.o	megan_mod.o	retro_mod.o	`		
diag51_mod.o	meganut_mod.o	tropopause_mod.o	`		
diag51b_mod.o	mercury_mod.o	oropopudas_mod.c	`		
diag56_mod.o	mmran_16.o	update.o	`		
diag_2pm.o	ndxx_setup.o	uvalbedo_mod.o	ĺ		
diag_mod.o	nei2005_anthro_mod.o	vdiff_mod.o	`		
diag_oh_mod.o	ocean_mercury_mod.o	vdiff_pre_mod.o	′		
diag_pl_mod.o	ohsave.o	vistas_anthro_mod.o	′		
diagoh.o	optdepth_mod.o	wetscav_mod.o	′		
aero_drydep.o	tomas_mod.o	webbeav_mod.o	′		
tomas_tpcore_mod.o	RD_AOD.o	depo_mercury_mod.o	′		
land_mercury_mod.o	global_br_mod.o	merra_a1_mod.o	′		
merra_a3_mod.o	merra_cn_mod.o	merra_i6_mod.o	′		
diag63_mod.o	paranox_mod.o		'		
bromocarb_mod.o	cldice_HBrHOBr_rxn.o	<pre>read_jv_atms_dat.o ssa_bromine_mod.o</pre>	'		
strat_chem_mod.o	gc_type2_mod.o		'		
		gc_environment_mod.o	\		
tpcore_geos57_window_mod					
pjc_pfix_geos57_window_m	Ju. 0				
#===========	=======================================				

#-----

[#] Makefile targets: type "make help" for a complete listing!

^{#-----}

.PHONY: clean realclean doc docclean help all: @\$(MAKE) TOMAS=yes lib @\$(MAKE) TOMAS=yes exe # Build normal GEOS-Chem lib: @\$(MAKE) libnc **Q\$(MAKE)** libheaders @\$(MAKE) libkpp @\$(MAKE) libutil @\$(MAKE) libiso @\$(MAKE) libtomas # Build code in GeosTomas/ libtomas: \$(OBJ) libiso: # Build code in ISOROPIA/ @\$(MAKE) -C \$(ISO) libkpp: # Build code in KPP/ @\$(MAKE) -C \$(KPP) # Build code in NcdfUtil/ libnc: @\$(MAKE) -C \$(NCDF) lib ncdfcheck: # Check netCDF library @\$(MAKE) libnc @\$(MAKE) -C \$(NCDF) check # Build code in GeosUtil/ libutil: @\$(MAKE) -C \$(UTIL) libheaders: # Build code in Headers/ @\$(MAKE) -C \$(HDR) # Build executable exe: \$(LD) \$(OBJ) \$(LINK) -o \$(EXE) cp -f \$(EXE) \$(BIN) clean: # Remove files here rm -f *.o *.mod geos geosapm geostomas realclean: # Remove files everywhere @\$(MAKE) clean @\$(MAKE) -C \$(ISO) clean @\$(MAKE) -C \$(KPP) realclean @\$(MAKE) -C \$(UTIL) clean

```
@$(MAKE) -C $(CORE) clean
@$(MAKE) -C $(GTMM) clean
@$(MAKE) -C $(HDR) clean
@$(MAKE) docclean
rm -f $(LIB)/*.a
rm -f $(MOD)/*.mod
rm -f $(BIN)/geos*
                                                # Build documentation
doc:
@$(MAKE) -C $(DOC) all
docclean:
                                                # Remove documentation
@$(MAKE) -C $(DOC) clean
                                                # Show help screen
help:
@$(MAKE) -C $(HELP)
# Dependencies listing (grep "USE " to get the list of module references!)
# From this list of dependencies, the "make" utility will figure out the
# correct order of compilation (so we don't have to do that ourselves).
# This also allows us to compile on multiple processors with "make -j".
# NOTES:
# (1) Only specify object-file dependencies that are within this directory.
      Object files in other directories will be referenced at link-time.
# (2) For "make -j" to work, all files in this directory must have a
      listed dependency.
: a3_read_mod.F
a3_read_mod.o
                                                   dao_mod.o
                            diag_mod.o
                                                  logical_mod.o
                         : a6_read_mod.F
a6_read_mod.o
                                                  dao_mod.o
                            diag_mod.o
                                                  logical_mod.o
acetone_mod.o
                          : acetone_mod.F
                                                   dao_mod.o
                            diag_mod.o
                                                  megan_mod.o
aerosol_mod.o
                          : aerosol_mod.F
                                                  comode_mod.o
                            dao_mod.o
                                                  diag_mod.o
                            logical_mod.o
                                                  tracerid_mod.o
                            tracer_mod.o
                                                   tropopause_mod.o
aircraft_nox_mod.o
                         : aircraft_nox_mod.F
                            dao_mod.o
                                                   diag_mod.o
```

airmas.o	: airmas.F	
anthroems.o	geia_mod.o	edgar_mod.o \ logical_mod.o \ tracer_mod.o \
arctas_ship_emiss_mod.o	· ·	scale_anthro_mod.o \ tracer_mod.o
arsl1k.o	: arsl1k.F	
backsub.o	: backsub.F	
benchmark_mod.o	<pre>: benchmark_mod.F tracerid_mod.o</pre>	tracer_mod.o
biofuel_mod.o	<pre>epa_nei_mod.o logical_mod.o</pre>	<pre>diag_mod.o</pre>
biofit.o	: biofit.F	
biomass_mod.o	<pre>gfed2_biomass_mod.o logical_mod.o</pre>	<pre>gc_biomass_mod.o \ gfed3_biomass_mod.o \ tracer_mod.o</pre>
BLKSLV.o	: BLKSLV.F	
boxvl.o	: boxvl.F	dao_mod.o
bravo_mod.o		<pre>logical_mod.o tracerid_mod.o</pre>
bromocarb_mod.o		<pre>diag_mod.o \ pbl_mix_mod.o \</pre>
c2h6_mod.o	: c2h6_mod.F biofuel_mod.o	biomass_mod.o \

	<pre>dao_mod.o geia_mod.o logical_mod.o tracer_mod.o</pre>	<pre>diag_mod.o global_oh_mod.o tracerid_mod.o</pre>	\ \
cac_anthro_mod.o	<pre>: cac_anthro_mod.F future_emissions_mod.o scale_anthro_mod.o tracer_mod.o</pre>	<pre>logical_mod.o tracerid_mod.o</pre>	\ \
calcrate.o	<pre>: calcrate.F comode_mod.o diag_mod.o drydep_mod.o logical_mod.o planeflight_mod.o cldice_HBrHOBr_rxn.o</pre>	dao_mod.o diag63_mod.o emissions_mod.o pbl_mix_mod.o tracerid_mod.o	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
carbon_mod.o	<pre>: carbon_mod.F biomass_mod.o dao_mod.o drydep_mod.o gfed2_biomass_mod.o global_no3_mod.o global_o3_mod.o logical_mod.o pbl_mix_mod.o tracer_mod.o vdiff_pre_mod.o tomas_mod.o</pre>	comode_mod.o diag_mod.o future_emissions_mod.o gfed3_biomass_mod.o global_oh_mod.o megan_mod.o tracerid_mod.o tropopause_mod.o meganut_mod.o	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
ch3i_mod.o	<pre>: ch3i_mod.F biofuel_mod.o dao_mod.o diag_pl_mod.o tracerid_mod.o uvalbedo_mod.o</pre>	<pre>biomass_mod.o diag_mod.o logical_mod.o tracer_mod.o</pre>	\ \ \ \
chemdr.o	<pre>: chemdr.F aerosol_mod.o dao_mod.o diag_pl_mod.o dust_mod.o planeflight_mod.o tracer_mod.o tropopause_mod.o</pre>	comode_mod.o diag_oh_mod.o future_emissions_mod.o logical_mod.o restart_mod.o tracerid_mod.o uvalbedo_mod.o	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
chemistry_mod.o	: chemistry_mod.F		\

cleanup.o

acetone_mod.o	aerosol_mod.o	\
c2h6_mod.o	carbon_mod.o	\
ch3i_mod.o	comode_mod.o	\
dao_mod.o	drydep_mod.o	\
dust_mod.o	global_ch4_mod.o	\
h2_hd_mod.o	hcn_ch3cn_mod.o	\
isoropiaII_mod.o	logical_mod.o	\
mercury_mod.o	optdepth_mod.o	\
rpmares_mod.o	RnPbBe_mod.o	\
seasalt_mod.o	strat_chem_mod.o	\
sulfate_mod.o	tagged_co_mod.o	\
tagged_ox_mod.o	tracerid_mod.o	\
tracer_mod.o	tomas_mod.o	
51 G 5 5 2 2 2 3 G 7 5		
cleanup.F		\
acetone_mod.o	aerosol_mod.o	\
_	aircraft_nox_mod.o	\
biomass_mod.o	biofuel_mod.o	\
bravo_mod.o	c2h6_mod.o	\
cac_anthro_mod.o	carbon_mod.o	\
co2_mod.o	comode_mod.o	\
dao_mod.o	depo_mercury_mod.o	\
diag_mod.o	diag03_mod.o	\
diag04_mod.o	diag41_mod.o	\
diag50_mod.o	diag51_mod.o	'
diag51b_mod.o	diag_oh_mod.o	'
_	•	'
<pre>diag_pl_mod.o dust_mod.o</pre>	<pre>drydep_mod.o dust_dead_mod.o</pre>	'
		'
edgar_mod.o	emep_mod.o	'
epa_nei_mod.o	gc_biomass_mod.o	'
isoropiaII_mod.o	gfed2_biomass_mod.o	,
gfed3_biomass_mod.o	111101	,
global_ch4_mod.o	global_hno3_mod.o	,
global_no3_mod.o	global_nox_mod.o	'
global_o1d_mod.o	global_oh_mod.o	,
h2_hd_mod.o	hcn_ch3cn_mod.o	١.
modis_lai_mod.o	<pre>land_mercury_mod.o</pre>	
lightning_nox_mod.o	linoz_mod.o	\
megan_mod.o	mercury_mod.o	\
ocean_mercury_mod.o	pbl_mix_mod.o	\
pjc_pfix_mod.o	planeflight_mod.o	\
seasalt_mod.o	sulfate_mod.o	\
tagged_co_mod.o	toms_mod.o	\
tracer_mod.o	transport_mod.o	\
tropopause_mod.o	uvalbedo_mod.o	\
vistas_anthro_mod.o	wetscav_mod.o	\
icoads_ship_mod.o	tpcore_fvdas_mod.o	\
tpcore_geos5_window_mod	. 0	\

		<pre>tpcore_geos57_window_mo retro_mod.o</pre>	d.o tomas_mod.o	\
cldice_HBrHOBr_rxn.o	:	cldice_HBrHOBr_rxn.F		
CLDSRF.o	:	CLDSRF.F		
co2_mod.o	:	co2_mod.F	biomass_mod.o	\
tra	ceri	diag04_mod.o d_mod.o	tracer_mod.o	\
CO_strat_pl.o	:	CO_strat_pl.F		\
		<pre>dao_mod.o tracerid_mod.o</pre>	<pre>tracer_mod.o tropopause_mod.o</pre>	\
comode_mod.o	:	comode_mod.F	tracer_mod.o	
convection_mod.o	:	convection_mod.F		\
		dao_mod.o	diag_mod.o	\
		depo_mercury_mod.o	fvdas_convect_mod.o	\
		gc_type_mod.o	<pre>gcap_convect_mod.o</pre>	\
		logical_mod.o	mercury_mod.o	\
		tracer_mod.o	tracerid_mod.o	\
		wetscav_mod.o		
dao_mod.o	:	dao_mod.F		\
		logical_mod.o	tracer_mod.o	\
gc_type_mod.o				
decomp.o	:	decomp.F		
depo_mercury_mod.o	:	depo_mercury_mod.F		\
		dao_mod.o	diag_mod.o	\
		logical_mod.o	tracerid_mod.o	
diag03_mod.o	:	diag03_mod.F	tracerid_mod.o	
diag04_mod.o	:	diag04_mod.F		
diag1.o	:	diag1.F		\
		dao_mod.o	diag_mod.o	\
		tracer_mod.o	tracerid_mod.o	\
		tropopause_mod.o	ocean_mercury_mod.o	
diag3.o	:	diag3.F		\
_		biofuel_mod.o	biomass_mod.o	\
		diag_mod.o	diag03_mod.o	\
		diag04_mod.o	diag41_mod.o	\
		5 –	3 -	•

	<pre>diag42_mod.o diag_pl_mod.o drydep_mod.o tracer_mod.o wetscav_mod.o</pre>	<pre>diag56_mod.o depo_mercury_mod.o logical_mod.o tracerid_mod.o tomas_mod.o</pre>	\ \ \
diag41_mod.o	: diag41_mod.F	pbl_mix_mod.o	
diag42_mod.o	<pre>: diag42_mod.F dao_mod.o tracerid_mod.o</pre>	<pre>logical_mod.o tracer_mod.o</pre>	\
diag48_mod.o	<pre>: diag48_mod.F dao_mod.o tracerid_mod.o</pre>	<pre>pbl_mix_mod.o tracer_mod.o</pre>	\
diag49_mod.o	<pre>: diag49_mod.F dao_mod.o pbl_mix_mod.o tracer_mod.o</pre>	<pre>modis_lai_mod.o tracerid_mod.o</pre>	\ \
diag50_mod.o	<pre>: diag50_mod.F comode_mod.o logical_mod.o tracerid_mod.o tropopause_mod.o</pre>	<pre>dao_mod.o pbl_mix_mod.o tracer_mod.o</pre>	\ \ \
diag51_mod.o	<pre>: diag51_mod.F dao_mod.o logical_mod.o tracerid_mod.o tropopause_mod.o</pre>	<pre>modis_lai_mod.o pbl_mix_mod.o tracer_mod.o</pre>	\ \ \
diag51b_mod.o	<pre>: diag51b_mod.F dao_mod.o logical_mod.o tracerid_mod.o tropopause_mod.o</pre>	<pre>modis_lai_mod.o pbl_mix_mod.o tracer_mod.o</pre>	\ \ \
diag56_mod.o	: diag56_mod.F		
diag63_mod.o	<pre>: diag63_mod.F dao_mod.o tracerid_mod.o</pre>	<pre>pbl_mix_mod.o tracer_mod.o</pre>	\
diag_2pm.o	: diag_2pm.F diag_mod.o	tropopause_mod.o	\

diag_mod.o	: diag_mod.F		
diag_oh_mod.o	<pre>: diag_oh_mod.F comode_mod.o tracerid_mod.o</pre>	<pre>logical_mod.o tracer_mod.o</pre>	\
diag_pl_mod.o	<pre>: diag_pl_mod.F comode_mod.o tracerid_mod.o</pre>	<pre>logical_mod.o tracer_mod.o</pre>	\
drydep_mod.o	<pre>: drydep_mod.F dao_mod.o logical_mod.o tracerid_mod.o meganut_mod.o</pre>	<pre>comode_mod.o diag_mod.o pbl_mix_mod.o tracer_mod.o tomas_mod.o</pre>	\ \ \ \
diagoh.o	: diagoh.F	diag_mod.o	
dust_dead_mod.o	: dust_dead_mod.F	dao_mod.o	
dust_mod.o	<pre>: dust_mod.F comode_mod.o diag_mod.o dust_dead_mod.o tracerid_mod.o</pre>	<pre>dao_mod.o drydep_mod.o logical_mod.o tracer_mod.o</pre>	\ \ \
edgar_mod.o	<pre>: edgar_mod.F future_emissions_mod.o scale_anthro_mod.o</pre>	<pre>logical_mod.o tracerid_mod.o</pre>	\
EFOLD.o	: EFOLD.F		
emep_mod.o	<pre>: emep_mod.F future_emissions_mod.o scale_anthro_mod.o tracer_mod.o</pre>	<pre>logical_mod.o tracerid_mod.o</pre>	\ \
emf_scale.o	: emf_scale.F	tracerid_mod.o	
emfossil.o	<pre>: emfossil.F bravo_mod.o dao_mod.o edgar_mod.o epa_nei_mod.o logical_mod.o streets_anthro_mod.o tracerid_mod.o</pre>	cac_anthro_mod.o diag_mod.o emep_mod.o icoads_ship_mod.o nei2005_anthro_mod.o tracer_mod.o vistas_anthro_mod.o	\ \ \ \ \

retro_mod.o		c2h6_mod.o		
emissdr.o		emissdr.F acetone_mod.o biofuel_mod.o diag_mod.o logical_mod.o megan_mod.o	aircraft_nox_mod.o dao_mod.o emissions_mod.o tracer_mod.o	\ \ \ \ \ \
bromocarb_mc	od.o	tracerid_mod.o	meganut_mod.o	\
emissions_mod.o	:	emissions_mod.F arctas_ship_emiss_mod.o bravo_mod.o cac_anthro_mod.o ch3i_mod.o dust_mod.o emep_mod.o global_ch4_mod.o hcn_ch3cn_mod.o logical_mod.o nei2005_anthro_mod.o retro_mod.o seasalt_mod.o sulfate_mod.o tracer_mod.o	biomass_mod.o c2h6_mod.o carbon_mod.o co2_mod.o edgar_mod.o epa_nei_mod.o h2_hd_mod.o icoads_ship_mod.o mercury_mod.o paranox_mod.o RnPbBe_mod.o streets_anthro_mod.o vistas_anthro_mod.o	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
epa_nei_mod.o	:	<pre>epa_nei_mod.F future_emissions_mod.o scale_anthro_mod.o tracer_mod.o</pre>	<pre>logical_mod.o tracerid_mod.o</pre>	\ \
fast_j.o	:	<pre>fast_j.F dao_mod.o</pre>	toms_mod.o	\
fertadd.o	:	fertadd.F	logical_mod.o	
fjfunc.o	:	fjfunc.F		
fjx_acet_mod.o	:	fjx_acet_mod.F		
future_emissions_m	nod.o :	future_emissions_mod.F		
fvdas_convect_mod.	o :	<pre>fvdas_convect_mod.F dao_mod.o depo_mercury_mod.o tracerid_mod.o</pre>	<pre>diag_mod.o logical_mod.o tracer_mod.o</pre>	\ \

fyhoro.o	: fyhoro.F	
fyrno3.o	: fyrno3.F	
gamap_mod.o	: gamap_mod.F diag03_mod.o diag41_mod.o diag48_mod.o diag50_mod.o diag51b_mod.o diag_pl_mod.o togical_mod.o tomas_mod.o	diag04_mod.o \ diag42_mod.o \ diag49_mod.o \ diag51_mod.o \ diag56_mod.o \ drydep_mod.o \ tracerid_mod.o \ wetscav_mod.o \ diag63_mod.o
<pre>ifeq (\$(COMPILER),sun) \$(F90) -00 -c \$< endif</pre>		
gasconc.o	<pre>: gasconc.F comode_mod.o drydep_mod.o logical_mod.o</pre>	dao_mod.o \ tropopause_mod.o \
GAUSSP.o	: GAUSSP.F	
gc_biomass_mod.o	<pre>: gc_biomass_mod.F future_emissions_mod.o tracerid_mod.o</pre>	logical_mod.o \ tracer_mod.o
gc_type_mod.o	: gc_type_mod.F	
<pre>gcap_convect_mod.o</pre>	<pre>: gcap_convect_mod.F dao_mod.o</pre>	diag_mod.o
gcap_read_mod.o	<pre>: gcap_read_mod.F diag_mod.o</pre>	dao_mod.o \ logical_mod.o
geia_mod.o	: geia_mod.F	
GEN. o	: GEN.F	
geos57_read_mod.o	: geos57_read_mod.F90 dao_mod.o	\
get_global_ch4.o	<pre>: get_global_ch4.F future_emissions_mod.o</pre>	\ logical_mod.o
getifsun.o	: getifsun.F	comode_mod.o

<pre>gfed2_biomass_mod.o</pre>	gfed2_biomass_mod.F future_emissions_mod.o tracer_mod.o	<pre>logical_mod.o tracerid_mod.o</pre>	\
<pre>gfed3_biomass_mod.o</pre> :	gfed3_biomass_mod.F future_emissions_mod.o tracer_mod.o	<pre>logical_mod.o tracerid_mod.o</pre>	\
global_br_mod.o :	global_br_mod.F	tropopause_mod.o	
global_ch4_mod.o :	global_ch4_mod.F dao_mod.o diag_oh_mod.o global_oh_mod.o tracer_mod.o	<pre>diag_mod.o diag_pl_mod.o logical_mod.o vdiff_pre_mod.o</pre>	\ \ \
global_hno3_mod.o :	global_hno3_mod.F dao_mod.o	tracer_mod.o	\
global_no3_mod.o :	global_no3_mod.F		
global_nox_mod.o :	global_nox_mod.F		
global_o1d_mod.o :	global_o1d_mod.F		
global_o3_mod.o :	global_o3_mod.F		
global_oh_mod.o :	global_oh_mod.F dao_mod.o		\
h2_hd_mod.o :	h2_hd_mod.F biofuel_mod.o dao_mod.o drydep_mod.o global_nox_mod.o global_oh_mod.o scale_anthro_mod.o tracerid_mod.o tropopause_mod.o	biomass_mod.o diag_mod.o geia_mod.o global_o1d_mod.o logical_mod.o tagged_co_mod.o tracer_mod.o meganut_mod.o	\ \\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
hcn_ch3cn_mod.o :	hcn_ch3cn_mod.F biomass_mod.o diag_mod.o global_oh_mod.o pbl_mix_mod.o	<pre>dao_mod.o geia_mod.o logical_mod.o tracerid_mod.o</pre>	\ \ \
i6_read_mod.o :	i6_read_mod.F		\

	<pre>dao_mod.o logical_mod.o</pre>	diag_mod.o	\
icoads_ship_mod.o	<pre>: icoads_ship_mod.F future_emissions_mod.o scale_anthro_mod.o tracer_mod.o</pre>	<pre>logical_mod.o tracerid_mod.o</pre>	\ \
<pre>initialize.o inphot.o</pre>	<pre>: initialize.F diag_mod.o diag04_mod.o diag42_mod.o diag_pl_mod.o</pre> <pre>: inphot.F</pre>	diag03_mod.o diag41_mod.o diag56_mod.o logical_mod.o	\\\\
input_mod.o	_	biofuel_mod.o diag03_mod.o diag41_mod.o ag48_mod.o \ ag50_mod.o	\ \ \ \ \
diag51_mod.o diag56_mod.o diag_pl_mod.o	<pre>diag51b_mod.o diag_oh_mod.o drydep_mod.o emissions_mod.o gamap_mod.o</pre>	future_emissions_mod.o land_mercury_mod.o ercury_mod.o planeflight_mod.o tpcore_bc_mod.o tracer_mod.o wetscav_mod.o diag63_mod.o	// ////
isoropiaII_mod.o	<pre>: isoropiaII_mod.F dao_mod.o logical_mod.o tracer_mod.o</pre>	<pre>global_hno3_mod.o tracerid_mod.o tropopause_mod.o</pre>	\ \
jsparse.o	: jsparse.F	comode_mod.o	
JRATET.0	: JRATET.F	fjx_acet_mod.o	
JVALUE.0	: JVALUE.F		
jv_index.o	: jv_index.F		

ksparse.o	: ksparse.F		
land_mercury_mod.o	<pre>: land_mercury_mod.F biomass_mod.o depo_mercury_mod.o logical_mod.o</pre>	<pre>dao_mod.o modis_lai_mod.o tracerid_mod.o</pre>	\ \ \
LEGNDO.o	: LEGNDO.F		
lightning_nox_mod.o	<pre>: lightning_nox_mod.F dao_mod.o diag_mod.o</pre>	diag56_mod.o logical_mod.o	\
linoz_mod.o	: linoz_mod.F dao_mod.o tracer_mod.o	<pre>tracerid_mod.o tropopause_mod.o</pre>	\
logical_mod.o	: logical_mod.F		
lump.o	: lump.F comode_mod.o	tracerid_mod.o	\
main.o	: main.F a3_read_mod.o benchmark_mod.o convection_mod.o diag_mod.o diag42_mod.o diag49_mod.o diag51_mod.o diag51_mod.o depo_mercury_mod.o emissions_mod.o gcap_read_mod.o input_mod.o lightning_nox_mod.o pbl_mix_mod.o pbl_mix_mod.o ptpcore_bc_mod.o transport_mod.o vetscav_mod.o merra_a1_mod.o merra_i6_mod.o o olson_landmap_mod.o	a6_read_mod.o chemistry_mod.o comode_mod.o diag41_mod.o diag48_mod.o diag50_mod.o diag51b_mod.o dao_mod.o drydep_mod.o global_ch4_mod.o i6_read_mod.o modis_lai_mod.o linoz_mod.o megan_mod.o ocean_mercury_mod.o soaprod_mod.o tracer_mod.o tracer_mod.o vdiff_mod.o merra_cn_mod.o gc_environment_mod.o diag63_mod.o mapping_mod.o	///////////////////////////////////////

mapping_mod.o	: mapping_mod.F90 logical_mod.o		\
MATIN4.o	: MATIN4.F		
megan_mod.o	<pre>: megan_mod.F modis_lai_mod.o meganut_mod.o geos57_read_mod.o</pre>	a3_read_mod.o logical_mod.o merra_a1_mod.o	\ \
meganut_mod.o	: meganut_mod.F	dao_mod.o	
mercury_mod.o	<pre>: mercury_mod.F dao_mod.o diag03_mod.o drydep_mod.o global_o3_mod.o land_mercury_mod.o logical_mod.o pbl_mix_mod.o tracerid_mod.o tropopause_mod.o</pre>	depo_mercury_mod.o diag_mod.o global_br_mod.o global_oh_mod.o ocean_mercury_mod.o RnPbBe_mod.o tracer_mod.o vdiff_pre_mod.o	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
merra_a1_mod.o	<pre>: merra_a1_mod.F logical_mod.o</pre>	dao_mod.o	\
merra_a3_mod.o	<pre>: merra_a3_mod.F logical_mod.o</pre>	dao_mod.o	\
merra_cn_mod.o	<pre>: merra_cn_mod.F logical_mod.o</pre>	dao_mod.o	\
merra_i6_mod.o	: merra_i6_mod.F logical_mod.o	dao_mod.o	\
MIESCT.o	: MIESCT.F		
modis_lai_mod.o	<pre>: modis_lai_mod.F90 mapping_mod.o</pre>		\
ndxx_setup.o	<pre>: ndxx_setup.F biofuel_mod.o diag_oh_mod.o logical_mod.o tracer_mod.o wetscav_mod.o diag63_mod.o</pre>	<pre>diag_mod.o drydep_mod.o planeflight_mod.o tracerid_mod.o tomas_mod.o</pre>	\ \ \ \

nei2005_anthro_mod.o	:	<pre>nei2005_anthro_mod.F future_emissions_mod.o scale_anthro_mod.o tracer_mod.o</pre>	<pre>logical_mod.o tracerid_mod.o</pre>	\ \
NOABS.o	:	NOABS.F		
ocean_mercury_mod.o	:	ocean_mercury_mod.F dao_mod.o diag03_mod.o tracerid_mod.o	<pre>depo_mercury_mod.o logical_mod.o tracer_mod.o</pre>	\ \
ohsave.o	:	ohsave.F diag_mod.o	<pre>comode_mod.o tracerid_mod.o</pre>	\
olson_landmap_mod.o	:	olson_landmap_mod.F90 mapping_mod.o		\
optdepth_mod.o	:	optdepth_mod.F	diag_mod.o	
OPMIE.o	:	OPMIE.F		
paranox_mod.o	:	<pre>paranox_mod.F dao_mod.o tracerid_mod.o</pre>	tracer_mod.o	\
partition.o	:	partition.F comode_mod.o	tracerid_mod.o	\
pbl_mix_mod.o	:	<pre>pbl_mix_mod.F dao_mod.o logical_mod.o</pre>	diag_mod.o tracer_mod.o	\
pderiv.o	:	pderiv.F		
photoj.o	:	photoj.F		
physproc.o	:	<pre>physproc.F comode_mod.o chemistry_mod.o</pre>	logical_mod.o	\
pjc_pfix_geos5_window_mod.o	:	pjc_pfix_geos5_window_m	od.F	
pjc_pfix_geos57_window_mod.	o:	pjc_pfix_geos57_window_	mod.F	
pjc_pfix_mod.o	:	pjc_pfix_mod.F		

planeflight_mod.o	<pre>: planeflight_mod.F comode_mod.o tracer_mod.o</pre>	dao_mod.o tropopause_mod.o	\
precipfrac.o	: precipfrac.F	dao_mod.o	
pulsing.o	: pulsing.F		
RD_AOD.o	: RD_AOD.F		
rd_js.o	: rd_js.F		
rdsoil.o	: rdsoil.F		
RD_TJPL.o	: RD_TJPL.F		
readchem.o	<pre>: readchem.F diag_pl_mod.o logical_mod.o</pre>	drydep_mod.o	\
reader.o	: reader.F		
read_jv_atms_dat.o	: read_jv_atms_dat.F90		
restart_mod.o	<pre>: restart_mod.F comode_mod.o logical_mod.o</pre>	dao_mod.o tracer_mod.o	\
retro_mod.o	<pre>: retro_mod.F future_emissions_mod.o scale_anthro_mod.o tracer_mod.o</pre>	<pre>logical_mod.o tracerid_mod.o</pre>	\ \
RnPbBe_mod.o	: RnPbBe_mod.F dao_mod.o logical_mod.o tropopause_mod.o	diag_mod.o tracer_mod.o	\ \
rpmares_mod.o	<pre>: rpmares_mod.F dao_mod.o tracerid_mod.o tropopause_mod.o</pre>	<pre>global_hno3_mod.o tracer_mod.o</pre>	\ \
ruralbox.o	<pre>: ruralbox.F comode_mod.o</pre>	tropopause_mod.o	\
scale_anthro_mod.o	: scale_anthro_mod.F		

seasalt_mod.o	: seasalt_mod.F		\
	dao_mod.o	diag_mod.o	\
	drydep_mod.o	logical_mod.o	\
	pbl_mix_mod.o	tracerid_mod.o	\
	tracer_mod.o	vdiff_pre_mod.o	\
	tomas_mod.o	ssa_bromine_mod.o	
		224_220	
set_aer.o	: set_aer.F		
set_prof.o	: set_prof.F		\
_	dao_mod.o	toms_mod.o	
setemdep.o	: setemdep.F	drydep_mod.o	\
	tracer_mod.o	tracerid_mod.o	
setemis.o	: setemis.F		\
Setemis.0		1. d - f 1 1	`
	aircraft_nox_mod.o	biofuel_mod.o	,
	biomass_mod.o	comode_mod.o	\
	diag_mod.o	emissions_mod.o	\
	lightning_nox_mod.o	logical_mod.o	\
	pbl_mix_mod.o	tracerid_mod.o	\
	tropopause_mod.o		
setmodel.o	: setmodel.F		
sfcwindsqr.o	: sfcwindsqr.F	dao_mod.o	
	B		
smvgear.o	: smvgear.F	comode_mod.o	
soaprod_mod.o	: soaprod_mod.F		\
<u> </u>	carbon_mod.o	dao_mod.o	ĺ
	logical_mod.o	tracer_mod.o	`
	logical_mod.o	tracer_mod.o	
soilbase.o	: soilbase.F		
soilcrf.o	: soilcrf.F		
soilnoxems.o	: soilnoxems.F		\
2022220	dao_mod.o	diag_mod.o	``
	future_emissions_mod.o	•	`
	meganut_mod.o	logical_mou.o	`
soiltemp.o	: soiltemp.F		
soiltype.o	: soiltype.F		
SPHERE.o	: SPHERE.F		

<pre>ssa_bromine_mod.o streets_anthro_mod.o</pre>	<pre>: ssa_bromine_mod.F bromocarb_mod.o logical_mod.o tracerid_mod.o : streets_anthro_mod.F</pre>	<pre>comode_mod.o pbl_mix_mod.o tropopause_mod.o</pre>	\ \ \
	<pre>future_emissions_mod.o scale_anthro_mod.o tracer_mod.o</pre>	<pre>logical_mod.o tracerid_mod.o</pre>	\
strat_chem_mod.o	: strat_chem_mod.F90 dao_mod.o linoz_mod.o tracer_mod.o tropopause_mod.o	<pre>logical_mod.o tagged_ox_mod.o tracerid_mod.o</pre>	\ \ \ \
subfun.o	: subfun.F		
<pre>sulfate_mod.o</pre>	: sulfate_mod.F arctas_ship_emiss_mod.o bravo_mod.o comode_mod.o diag_mod.o edgar_mod.o epa_nei_mod.o gfed2_biomass_mod.o global_hno3_mod.o icoads_ship_mod.o logical_mod.o pbl_mix_mod.o seasalt_mod.o tracerid_mod.o vdiff_pre_mod.o	biomass_mod.o cac_anthro_mod.o dao_mod.o drydep_mod.o emep_mod.o future_emissions_mod.o gfed3_biomass_mod.o global_oh_mod.o tomas_mod.o nei2005_anthro_mod.o scale_anthro_mod.o streets_anthro_mod.o tracer_mod.o uvalbedo_mod.o wetscav_mod.o	///////////////////////////////////////
sunparam.o	: sunparam.F		
<pre>gc_type2_mod.o gc_environment_mod.o gc_type_mod.o</pre>	<pre>: gc_type2_mod.F90 : gc_environment_mod.F90</pre>	<pre>tracer_mod.o gc_type2_mod.o</pre>	\
tagged_co_mod.o	<pre>: tagged_co_mod.F biofuel_mod.o dao_mod.o diag_pl_mod.o global_oh_mod.o megan_mod.o</pre>	biomass_mod.o diag_mod.o global_nox_mod.o logical_mod.o meganut_mod.o	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\

	<pre>pbl_mix_mod.o tracer_mod.o</pre>	<pre>tracerid_mod.o tropopause_mod.o</pre>	\
tagged_ox_mod.o	<pre>: tagged_ox_mod.F dao_mod.o diag_pl_mod.o logical_mod.o pbl_mix_mod.o tracer_mod.o</pre>	<pre>diag_mod.o drydep_mod.o meganut_mod.o tracerid_mod.o tropopause_mod.o</pre>	\ \ \ \ \
tcorr.o	: tcorr.F		
toms_mod.o	: toms_mod.F		
tpcore_bc_mod.o	<pre>: tpcore_bc_mod.F logical_mod.o</pre>	tracer_mod.o	\
<pre>tpcore_geos5_window_mod.o \$(F90) -c \$(FREEFORM) \$(R8)</pre>	<pre>: tpcore_geos5_window_mod \$<</pre>	.F90	
<pre>tpcore_geos57_window_mod.o \$(F90) -c \$(FREEFORM) \$(R8)</pre>		d.F90	
<pre>tpcore_mod.o \$(F90) -c \$(R8) \$<</pre>	<pre>: tpcore_mod.F diag_mod.o tracer_mod.o</pre>	<pre>dao_mod.o global_ch4_mod.o tomas_tpcore_mod.o</pre>	\
<pre>tpcore_window_mod.o \$(F90) -c \$(R8) \$<</pre>	<pre>: tpcore_window_mod.F diag_mod.o tracer_mod.o</pre>	<pre>dao_mod.o global_ch4_mod.o tomas_mod.o</pre>	\
tracer_mod.o	: tracer_mod.F		
<pre>tracerid_mod.o gc_environment_mod.o</pre>	<pre>: tracerid_mod.F logical_mod.o</pre>	<pre>gc_type2_mod.o tracer_mod.o</pre>	\
transport_mod.o	: transport_mod.F dao_mod.o logical_mod.o tpcore_mod.o tpcore_fvdas_mod.o tracer_mod.o tpcore_geos5_window_mod tpcore_geos57_window_mod pjc_pfix_geos5_window_mod	d.o	\\\\\\\\\\

pjc_pfix_geos57_window_mod.o

tropopause.o	<pre>: tropopause.F dao_mod.o logical_mod.o</pre>	diag_mod.o tropopause_mod.o	\
tropopause_mod.o	<pre>: tropopause_mod.F comode_mod.o diag_mod.o</pre>	dao_mod.o logical_mod.o	\
update.o	: update.F		
<pre>vdiff_pre_mod.o ifeq (\$(COMPILER),sun) \$(F90) -03 -c \$< endif</pre>	: vdiff_pre_mod.F	tracer_mod.o	
vdiff_mod.o	<pre>: vdiff_mod.F90 comode_mod.o depo_mercury_mod.o drydep_mod.o ocean_mercury_mod.o tracer_mod.o vdiff_pre_mod.o</pre>	<pre>dao_mod.o diag_mod.o logical_mod.o pbl_mix_mod.o tracerid_mod.o</pre>	\ \ \ \
vistas_anthro_mod.o	<pre>: vistas_anthro_mod.F future_emissions_mod.o logical_mod.o tracerid_mod.o</pre>	<pre>epa_nei_mod.o scale_anthro_mod.o tracer_mod.o</pre>	\ \
wetscav_mod.o	<pre>: wetscav_mod.F dao_mod.o depo_mercury_mod.o mercury_mod.o tracerid_mod.o tomas_mod.o</pre>	<pre>diag_mod.o logical_mod.o ocean_mercury_mod.o tracer_mod.o</pre>	\ \ \ \
XSEC1D.o	: XSEC1D.F		
XSECO2.o	: XSECO2.F		
XSEC03.o	: XSECO3.F		
# # Dependencies of files specific to TOMAS microphysics (bmy, 1/25/10) #			

```
tomas_mod.o
                             : tomas_mod.F
                      dao_mod.o
                                               diag_mod.o
                              diag_pl_mod.o
                                                       logical_mod.o
                                                       tracer_mod.o
                               tropopause_mod.o
                               tracerid_mod.o
                             : tomas_tpcore_mod.F90
tomas_tpcore_mod.o
                               tomas_mod.o
                                                        tracerid_mod.o
aero_drydep.o
                             : aero_drydep.F
                               dao_mod.o
                                                        diag_mod.o
                               drydep_mod.o
                                                        dust_mod.o
                               logical_mod.o
                                                       pbl_mix_mod.o
                               tracer_mod.o
                                                       tracerid_mod.o
                               tropopause_mod.o
                                                        tomas_mod.o
```

1.9 Module Interface Makefile (in the GTMM subdirectory)

This is main "router" makefile for the GTMM model. It compiles the GTMM code for GEOS-Chem mercury simulations. **REMARKS**:

To build the programs, call "make" with the following syntax:

```
make TARGET [ OPTIONAL-FLAGS ]
```

To display a complete list of options, type "make help".

Makefile uses the following variables:

Variable	Description
SHELL	Specifies the shell for "make" to use (usually SHELL=/bin/sh)
ROOTDIR	Specifies the root-level directory of the GEOS-Chem code
DOC	Specifies the directory where GEOS-Chem documentation is found
HDR	Specifies the directory where GEOS-Chem include files are found
LIB	Specifies the directory where library files (*.a) are stored
MOD	Specifies the directory where module files (*.mod) are stored
AR	Sys var w/ name of library creator program (i.e., "ar", "ranlib")

MAKE

```
REVISION HISTORY:
   16 Sep 2009 - R. Yantosca - Initial version
   18 Sep 2009 - P. Le Sager - Added kppintegrator target & commented
                        "make -C int" calls
  21 Sep 2009 - C. Carouge - Adapted to use with GTMM model.
# Define variables
SHELL
     = /bin/sh
ROOTDIR = ...
HDR
     = $(ROOTDIR)/Headers
HELP
     = $(ROOTDIR)/help
     = (ROOTDIR)/lib
LIB
MOD
     = \$(ROOTDIR)/mod
LGTMM = -L\$(LIB) -1Hg
# Include header file. This returns variables CC, F90, FREEFORM, LD, R8,
# as well as the default Makefile compilation rules for source code files.
include $(ROOTDIR)/Makefile_header.mk
# List of files to compile. Here the order is not important,
# as we will explicity define the dependencies listing below.
# Source code files
SRC = \$(wildcard *.F90)
# Object files
OBJ = \$(SRC:.F90=.0)
# Makefile targets: type "make help" for a complete listing!
#-----
.PHONY: clean help gtmm
lib: $(OBJ)
$(AR) crs libHg.a $(OBJ)
mv libHg.a $(LIB)
gtmm:
@$(MAKE) lib
@$(MAKE) exe
```

Sys var w/ name of Make command (i.e, "make" or "gmake")

exe:

```
$(LD) $(OBJ) $(LGTMM) -o gtmm
clean:
rm -f *.o *.mod gtmm
help:
@$(MAKE) -C $(HELP)
# Dependencies listing (grep "USE " to get the list of module references!)
# From this list of dependencies, the "make" utility will figure out the
# correct order of compilation (so we don't have to do that ourselves!)
: CleanupCASAarrays.F90 defineConstants.o
CleanupCASAarrays.o
                             loadCASAinput.o
                                                 defineArrays.o
GTMM.o
                            : GTMM.F90
                                                 defineConstants.o
                                                 defineArrays.o
                             loadCASAinput.o
                             dorestart_mod.o
                                                 input_gtmm_mod.o
GTMM_coupled.o
                            : GTMM_coupled.F90
                                                 defineConstants.o
                             defineArrays.o
                                                 dorestart_mod.o
                      loadCASAinput.o
                                          input_gtmm_mod.o
HgOutForGEOS.o
                            : HgOutForGEOS.F90
                                                 defineConstants.o
                             loadCASAinput.o
                                                 defineArrays.o
                             CasaRegridModule.o
assignAgeClassToRunningPool.o : assignAgeClassToRunningPool.F90
                             defineConstants.o
                                                 loadCASAinput.o
                             defineArrays.o
assignRanPoolToAgeClass.o
                            : assignRanPoolToAgeClass.F90
                             defineConstants.o
                                                 loadCASAinput.o
                             defineArrays.o
                            : defineArrays.F90
                                                 defineConstants.o
defineArrays.o
doFPARandLAI.o
                            : doFPARandLAI.F90
                                                 defineConstants.o
                             loadCASAinput.o
                                                 defineArrays.o
doHerbCarbon.o
                            : doHerbCarbon.F90
                                                 defineConstants.o
                                                 defineArrays.o
                             loadCASAinput.o
# $(F90) -00 -c $(FREEFORM) doHerbCarbon.F90
```

<pre>doHerbCarbonHg.o # \$(F90) -00 -c \$(FREEFORM)</pre>	loadCASAinput.o	defineConstants.o defineArrays.o	\
doHerbivory.o	: doHerbivory.F90	defineConstants.o defineArrays.o	\
doHgDeposition.o	0 1	defineConstants.o defineArrays.o	\
doLatitude.o		defineConstants.o defineArrays.o	\
doLeafRootShedding.o	: doLeafRootShedding.F90 loadCASAinput.o	defineConstants.o defineArrays.o	\
doMaxHg.o	0	defineConstants.o defineArrays.o	\
doNPP.o		defineConstants.o defineArrays.o	\
doOptimumTemperature.o	: doOptimumTemperature.FS loadCASAinput.o	90 defineConstants.o defineArrays.o	\
doPET.o		defineConstants.o defineArrays.o	\
doSoilMoisture.o		defineConstants.o defineArrays.o	\
<pre>doTreeCarbon.o # \$(F90) -00 -c \$(FREEFORM)</pre>	loadCASAinput.o	defineConstants.o defineArrays.o	\
doTreeCarbonHg.o	loadCASAinput.o	defineConstants.o defineArrays.o	\
# \$(F90) -00 -c \$(FREEFORM)	-		
getAgeClassBF.o	0 0	defineConstants.o defineArrays.o	\
getFireParams.o	0	defineConstants.o defineArrays.o	\
getFuelWood.o	0	defineConstants.o defineArrays.o	\

getSoilMoistParams.o	<pre>: getSoilMoistParams.F90 defineConstants.o loadCASAinput.o defineArrays.o</pre>	\
getSoilParams.o	: getSoilParams.F90 defineConstants.o loadCASAinput.o defineArrays.o	\
input_gtmm_mod.o	<pre>: input_gtmm_mod.F90 defineConstants.o defineArrays.o</pre>	\
load_GC_data.o	: load_GC_data.F90 defineConstants.o loadCASAinput.o CasaRegridModule.o	\
loadCASAinput.o	: loadCASAinput.F90 defineConstants.o defineArrays.o CasaRegridModule.o	\
loadHgDeposition.o	: loadHgDeposition.F90 defineConstants.o loadCASAinput.o defineArrays.o CasaRegridModule.o	\
organizeAgeClasses.o	: organizeAgeClasses.F90 defineConstants.o loadCASAinput.o defineArrays.o	\
processData.o	: processData.F90 defineConstants.o loadCASAinput.o defineArrays.o	\
# \$(F90) -00 -c \$(FREEFORM)		
sort_pick_veg.o	: sort_pick_veg.F90 defineConstants.o	
dorestart_mod.o	<pre>: dorestart_mod.F90 defineConstants.o defineArrays.o</pre>	\

1.10 Module Interface Makefile (in doc subdirectory)

Makefile for building the documentation (in PDF and PostScript formats) for the GEOS-Chem model source code, makefiles, and related scripts.

REMARKS:

To build the documentation, call "make" with the following syntax:
 make TARGET [OPTIONAL-FLAGS]

To display a complete list of options, type "make help".

You must have the LaTeX utilities (latex, dvips, dvipdf) installed on your system in order to build the documentation.

16 Sep 2009 - R. Yantosca - Initial version

```
21 Sep 2009 - R. Yantosca - Now call Makefile in help directory to
                          display the help screen options
   19 Nov 2009 - R. Yantosca - Now build documentation for Makefiles in the
                          GeosUtil and GeosCore subdirectories
   11 Dec 2009 - R. Yantosca - Now get SHELL from Makefile_header.mk
   28 Jan 2010 - R. Yantosca - Now add the Makefiles in the GeosTomas and
                          ISOROPIA subdirs to the documentation
   27 Aug 2010 - R. Yantosca - Brought up to date for newly documented code
   31 Aug 2010 - R. Yantosca - Add documentation for GTMM routines
   14 Sep 2010 - R. Yantosca - Now split make commands into include files
# Initialization
# Define variables
ROOTDIR := ..
BIN
     := \$(ROOTDIR)/bin
BPCH
     := $(ROOTDIR)/GeosBpch
CORE := $(ROOTDIR)/GeosCore
DOC
     := \$(ROOTDIR)/doc
GTMM
     := $(ROOTDIR)/GTMM
     := $(ROOTDIR)/Headers
HDR
HELP := $(ROOTDIR)/help
ISO
     := $(ROOTDIR)/ISOROPIA
KPP
      := $(ROOTDIR)/KPP
MOT
      := $(ROOTDIR)/GeosTomas
UTIL
      := $(ROOTDIR)/GeosUtil
# Get the Unix shell in SHELL from the Makefile_header.mk
include $(ROOTDIR)/Makefile_header.mk
# Makefile targets
.PHONY: all docclean help
all: srcdoc utildoc gtmmdoc makedoc
clean:
rm -f *.tex *.ps *.pdf
help:
@$(MAKE) -C $(HELPDIR)
```

1.10.1 Makefile_SrcDoc.mk (in doc subdirectory)

This Makefile fragment contains commands to build the documentation for the GEOS-Chem Source Code. It is inlined into the Makefile (in the doc subdirectory) by an "include" command.

REMARKS:

```
To build the documentation, call "make" with the following syntax: make TARGET [ OPTIONAL-FLAGS ]
To display a complete list of options, type "make help".
You must have the LaTeX utilities (latex, dvips, dvipdf) installed on your system in order to build the documentation.
```

REVISION HISTORY:

include ./Makefile_GtmmDoc.mk

```
14 Sep 2010 - R. Yantosca - Initial version, split off from Makefile
14 Sep 2010 - R. Yantosca - Added optdepth_mod.f to list
15 Sep 2010 - R. Yantosca - Added diag_2pm, diag_56, diagoh, ohsave
16 Sep 2010 - R. Yantosca - Added diag_pl_mod
04 Nov 2010 - R. Yantosca - Added acetone_mod
10 Nov 2010 - R. Yantosca - Added lightning_nox_mod
19 Nov 2010 - R. Yantosca - Added anthroems, RnPbBe_mod, tagged_ox_mod
19 Nov 2010 - R. Yantosca - Added tcorr, emfossil, emf_scale
01 Dec 2010 - R. Yantosca - Added global_br_mod, global_no3_mod
01 Dec 2010 - R. Yantosca - Added global_nox_mod, global_old_mod
01 Dec 2010 - R. Yantosca - Added global_oh_mod, toms_mod
02 Dec 2010 - R. Yantosca - Added upbdflx_mod, diag41_mod, diag42_mod
02 Dec 2010 - R. Yantosca - Added diag03_mod, diag49_mod, diag50_mod
```

```
02 Dec 2010 - R. Yantosca - Added diag51_mod, diag51b_mod, boxvl, rdmonot
   02 Dec 2010 - R. Yantosca - Added rdlight, rdland, rdsoil, emmonot
    16 Dec 2010 - R. Yantosca - Renamed output files to "GC_Ref_Vol_3.*"\
   21 Dec 2010 - R. Yantosca - Added comode_mod
    11 Jul 2011 - R. Yantosca - Added restart_mod
    19 Jul 2011 - R. Yantosca - Changed *.f* to *.F* for ESMF compatibility
    29 Jul 2011 - R. Yantosca - Added planeflight_mod
    22 Aug 2011 - R. Yantosca - Added retro_mod
   07 Sep 2011 - R. Yantosca - Added gfed3_biomass_mod, *jv*_mod files
   22 Dec 2011 - M. Payer
                             - Added aerosol_mod, drydep_mod, seasalt_mod,
                               and sulfate_mod
   07 Feb 2012 - M. Payer - Added paranox_mod, diag63_mod
   08 Feb 2012 - R. Yantosca - Added geos57_read_mod.F90
   28 Feb 2012 - R. Yantosca - Added pbl_mix_mod
   05 Mar 2012 - M. Payer
                           - Added tracer_mod
   06 Mar 2012 - R. Yantosca - Added photoj.F and set_prof.F
   07 Mar 2012 - M. Payer - Added global_ch4_mod
   22 Mar 2012 - M. Payer - Added c2h6_mod, olson_landmap_mod
   29 Mar 2012 - R. Yantosca - Added lai_mod
   29 Mar 2012 - R. Yantosca - Added modis_lai_mod and mapping_mod
   09 Apr 2012 - R. Yantosca - Added modules from Headers/ directory
    13 Apr 2012 - R. Yantosca - Removed findmon.F, rdlai.F, lai_mod.F
    19 Apr 2012 - R. Yantosca - Added read_jv_atms_dat.F90
    15 May 2012 - R. Yantosca - Added tpcore_bc_mod.F
    22 May 2012 - M. Payer
                           - Add bromocarb_mod.F, cldice_HBrHOBr_rxn.F,
                               and ssa_bromine_mod.F
# List of source code files (order is important)
```

```
SRC1 :=
./intro.geos-chem
./headers.geos-chem
$(HDR)/define.h
$(HDR)/CMN_SIZE_mod.F
$(HDR)/CMN_DEP_mod.F
$(HDR)/CMN_DIAG_mod.F
$(HDR)/CMN_GCTM_mod.F
$(HDR)/CMN_NOX_mod.F
$(HDR)/CMN_O3_mod.F
$(HDR)/CMN_mod.F
$(HDR)/cmn_fj_mod.F
$(HDR)/commsoil_mod.F
$(HDR)/comode_loop_mod.F
$(HDR)/jv_cmn_mod.F
$(HDR)/jv_mie_mod.F
$(HDR)/smv_dimension_mod.F
$(HDR)/smv_errcode_mod.F
$(HDR)/smv_physconst_mod.F
```

\$(CORE)/main.F	\
\$(CORE)/acetone_mod.F	\
\$(CORE)/aerosol_mod.F	\
\$(CORE)/arctas_ship_emiss_mod.F	\
\$(CORE)/bravo_mod.F	\
\$(CORE)/bromocarb_mod.F	\
\$(CORE)/c2h6_mod.F	\
\$(CORE)/cac_anthro_mod.F	`
\$(CORE)/chemistry_mod.F	,
\$(CORE)/co2_mod.F	``
\$(CORE)/comode_mod.F	`
\$(CORE)/convection_mod.F	`
\$(CORE)/dao_mod.F	`
\$(CORE)/depo_mercury_mod.F	`
\$(CORE)/diag03_mod.F	\
\$(CORE)/diag04_mod.F	\
_	\
\$(CORE)/diag41_mod.F	\
\$(CORE)/diag42_mod.F	\
\$(CORE)/diag49_mod.F	\
\$(CORE)/diag50_mod.F	\
\$(CORE)/diag51b_mod.F	,
\$(CORE)/diag56_mod.F	,
\$(CORE)/diag63_mod.F	,
\$(CORE)/diag_pl_mod.F	\
\$(CORE)/diag_oh_mod.F	///////////////////////////////////////
\$(CORE)/diag_mod.F	\
\$(CORE)/drydep_mod.F	\
\$(CORE)/dust_mod.F	\
\$(CORE)/emep_mod.F	\
\$(CORE)/emissions_mod.F	\
\$(CORE)/fjx_acet_mod.F	\
\$(CORE)/gamap_mod.F	\
\$(CORE)/geos57_read_mod.F90	\
\$(CORE)/gfed3_biomass_mod.F	\
\$(CORE)/global_br_mod.F	\
\$(CORE)/global_ch4_mod.F	\
\$(CORE)/global_no3_mod.F	\
\$(CORE)/global_nox_mod.F	\
\$(CORE)/global_o1d_mod.F	\
\$(CORE)/global_o3_mod.F	,
\$(CORE)/global_oh_mod.F	Ì
\$(CORE)/h2_hd_mod.F	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
\$(CORE)/icoads_ship_mod.F	Ì
\$(CORE)/input_mod.F	`\
\$(CORE)/isoropiaII_mod.F	\ \
\$(CORE)/land_mercury_mod.F	\
\$(CORE)/lightning_nox_mod.F	\
	\
\$(CORE)/linoz_mod.F	\

\$(CORE)/logical_mod.F	\
\$(CORE)/mapping_mod.F90	\
\$(CORE)/megan_mod.F	Ì
\$(CORE)/meganut_mod.F	Ì
\$(CORE)/merra_a1_mod.F	ì
\$(CORE)/merra_a3_mod.F	`
\$(CORE)/merra_cn_mod.F	`
\$(CORE)/merra_i6_mod.F	\ \ \ \
\$(CORE)/meira_io_mod.r \$(CORE)/modis_lai_mod.F90	'
	'
\$(CORE)/nei2005_anthro_mod.F	'
\$(CORE)/olson_landmap_mod.F90	,
\$(CORE)/optdepth_mod.F	\ \ \
\$(CORE)/paranox_mod.F	\
\$(CORE)/pbl_mix_mod.F	\
<pre>\$(CORE)/pjc_pfix_mod.F</pre>	\
<pre>\$(CORE)/planeflight_mod.F</pre>	\
\$(CORE)/retro_mod.F	\
\$(CORE)/RnPbBe_mod.F	\
\$(CORE)/scale_anthro_mod.F	\
<pre>\$(CORE)/seasalt_mod.F</pre>	\
\$(CORE)/ssa_bromine_mod.F	\ \ \ \
\$(CORE)/strat_chem_mod.F90	\
\$(CORE)/sulfate_mod.F	\
\$(CORE)/tagged_ox_mod.F	\
\$(CORE)/toms_mod.F	
\$(CORE)/tpcore_bc_mod.F	\
\$(CORE)/tracer_mod.F	`
\$(CORE)/tropopause_mod.F	
\$(CORE)/tpcore_fvdas_mod.F90	/
\$(CORE)/tpcore_geos5_window_mod.F90	\
	\
\$(CORE)/transport_mod.F	'
\$(CORE)/upbdflx_mod.F	,
\$(CORE)/vdiff_mod.F90	,
\$(CORE)/vdiff_pre_mod.F	\
\$(CORE)/vistas_anthro_mod.F	\
./subs.geos-chem	\
\$(CORE)/anthroems.F	/
\$(CORE)/boxvl.F	\
\$(CORE)/cldice_HBrHOBr_rxn.F	\
\$(CORE)/diag1.F	\
\$(CORE)/diag3.F	\
\$(CORE)/diag_2pm.F	\
\$(CORE)/diagoh.F	\
\$(CORE)/emfossil.F	///////////////////////////////////////
\$(CORE)/emf_scale.F	\
\$(CORE)/fast_j.F	\
\$(CORE)/photoj.F	`
\$(CORE)/set_prof.F	\
# (0010T) \ D00 - b1 01 .1	`

```
$(CORE)/initialize.F
$(CORE)/ndxx_setup.F
$(CORE)/ohsave.F
$(CORE)/rdsoil.F
$(CORE)/read_jv_atms_dat.F90
$(CORE)/ruralbox.F
$(CORE)/setemis.F
$(CORE)/sfcwindsqr.F
# Output file names
TEX1 := GC_Ref_Vol_3.tex
DVI1 := GC_Ref_Vol_3.dvi
PDF1 := GC_Ref_Vol_3.pdf
PS1 := GC_Ref_Vol_3.ps
# Make commands
srcdoc:
rm -f $(TEX1)
protex -sf $(SRC1) > $(TEX1)
latex $(TEX1)
latex $(TEX1)
latex $(TEX1)
dvipdf $(DVI1) $(PDF1)
dvips $(DVI1) -o $(PS1)
rm -f *.aux *.dvi *.log *.toc
```

1.10.2 Makefile_UtilDoc.mk (in doc subdirectory)

This Makefile fragment contains commands to build the documentation for the GEOS-Chem utility modules. It is inlined into the Makefile (in the doc subdirectory) by an "include" command.

REMARKS:

```
To build the documentation, call "make" with the following syntax:
   make TARGET [ OPTIONAL-FLAGS ]

To display a complete list of options, type "make help".

You must have the LaTeX utilities (latex, dvips, dvipdf) installed on your system in order to build the documentation.
```

```
14 Sep 2010 - R. Yantosca - Initial version, split off from Makefile 16 Dec 2010 - R. Yantosca - Renamed output files to "GC_Ref_Vol_2.*"
```

```
19 Jul 2011 - R. Yantosca - Changed *.f* to *.F* for ESMF compatibility
     3 Apr 2012 - M. Payer
                              - Added *.F90 so that grid_mod.F90 and
                                regrid_a2a_mod.F90 are included
# List of source code files
SRC3 :=
./intro.util
                            \
$(wildcard $(UTIL)/*.F)
$(wildcard $(UTIL)/*.F90)
# Output file names
TEX3 := GC_Ref_Vol_2.tex
DVI3 := GC_Ref_Vol_2.dvi
PDF3 := GC_Ref_Vol_2.pdf
PS3 := GC_Ref_Vol_2.ps
# Make commands
utildoc:
rm -f $(TEX3)
protex -sf $(SRC3) > $(TEX3)
latex $(TEX3)
latex $(TEX3)
latex $(TEX3)
dvipdf $(DVI3) $(PDF3)
dvips $(DVI3) -o $(PS3)
rm -f *.aux *.dvi *.log *.toc
```

1.10.3 Makefile_GtmmDoc.mk (in doc subdirectory)

This Makefile fragment contains commands to build the documentation for the Global Terrestrial Mercury Model (GTMM). It is inlined into the Makefile (in the doc subdirectory) by an "include" command.

REMARKS:

```
To build the documentation, call "make" with the following syntax:
   make TARGET [ OPTIONAL-FLAGS ]

To display a complete list of options, type "make help".

You must have the LaTeX utilities (latex, dvips, dvipdf) installed on your system in order to build the documentation.
```

14 Sep 2010 - R. Yantosca - Initial version, split off from Makefile 16 Dec 2010 - R. Yantosca - Renamed output files to "GC_Ref_Vol_4.*"

```
19 Jul 2011 - R. Yantosca - Changed *.f* to *.F* for ESMF compatibility
# List of source code files (order is important)
SRC4 :=
./intro.gtmm
$(GTMM)/GTMM.F90
$(GTMM)/CasaRegridModule.F90
$(GTMM)/defineArrays.F90
$(GTMM)/defineConstants.F90
$(GTMM)/dorestart_mod.F90
$(GTMM)/input_gtmm_mod.F90
$(GTMM)/loadCASAinput.F90
./subs.gtmm
$(GTMM)/CleanupCASAarrays.F90
$(GTMM)/GTMM_coupled.F90
$(GTMM)/HgOutForGEOS.F90
$(GTMM)/assignAgeClassToRunningPool.F90
$(GTMM)/assignRanPoolToAgeClass.F90
$(GTMM)/doFPARandLAI.F90
$(GTMM)/doHerbCarbon.F90
$(GTMM)/doHerbCarbonHg.F90
$(GTMM)/doHerbivory.F90
$(GTMM)/doHgDeposition.F90
$(GTMM)/doLatitude.F90
$(GTMM)/doLeafRootShedding.F90
$(GTMM)/doMaxHg.F90
$(GTMM)/doNPP.F90
$(GTMM)/doOptimumTemperature.F90
$(GTMM)/doPET.F90
$(GTMM)/doSoilMoisture.F90
$(GTMM)/doTreeCarbon.F90
$(GTMM)/doTreeCarbonHg.F90
$(GTMM)/getAgeClassBF.F90
$(GTMM)/getFireParams.F90
$(GTMM)/getFuelWood.F90
$(GTMM)/getSoilMoistParams.F90
$(GTMM)/getSoilParams.F90
$(GTMM)/loadHgDeposition.F90
$(GTMM)/load_GC_data.F90
$(GTMM)/organizeAgeClasses.F90
$(GTMM)/processData.F90
$(GTMM)/sort_pick_veg.F90
```

[#] Output file names

```
TEX4 := GC_Ref_Vol_4.tex
DVI4 := GC_Ref_Vol_4.dvi
PDF4 := GC_Ref_Vol_4.pdf
PS4 := GC_Ref_Vol_4.ps

# Make commands
gtmmdoc:
rm -f $(TEX4)
protex -sf $(SRC4) > $(TEX4)
latex $(TEX4)
latex $(TEX4)
latex $(TEX4)
dvipdf $(DVI4) $(PDF4)
dvips $(DVI4) -o $(PS4)
rm -f *.aux *.dvi *.log *.toc
```

1.10.4 Makefile_MakeDoc.mk (in doc subdirectory)

This Makefile fragment contains commands to build the documentation for the GEOS-Chem Makefiles It is inlined into the Makefile (in the doc subdirectory) by an "include" command.

REMARKS:

```
To build the documentation, call "make" with the following syntax:
   make TARGET [ OPTIONAL-FLAGS ]

To display a complete list of options, type "make help".

You must have the LaTeX utilities (latex, dvips, dvipdf) installed on your system in order to build the documentation.
```

```
14 Sep 2010 - R. Yantosca - Initial version, split off from Makefile 16 Dec 2010 - R. Yantosca - Renamed output files to "GC_Ref_Vol_1.*"
```

```
$(KPP)/standard/Makefile
$(KPP)/SOA/Makefile
$(TOM)/Makefile
$(GTMM)/Makefile
$(DOC)/Makefile
$(DOC)/Makefile_SrcDoc.mk
$(DOC)/Makefile_UtilDoc.mk
$(DOC)/Makefile_GtmmDoc.mk
$(DOC)/Makefile_MakeDoc.mk
$(HELP)/Makefile
# Output file names
TEX2 := GC_Ref_Vol_1.tex
DVI2 := GC_Ref_Vol_1.dvi
PDF2 := GC_Ref_Vol_1.pdf
PS2 := GC_Ref_Vol_1.ps
# Make command
makedoc:
rm -f $(TEX2)
protex -fS $(SRC2) > $(TEX2)
latex $(TEX2)
latex $(TEX2)
latex $(TEX2)
dvipdf $(DVI2) $(PDF2)
dvips $(DVI2) -o $(PS2)
rm -f *.aux *.dvi *.log *.toc
```

1.11 Module Interface Makefile (in the help subdirectory)

Displays the makefile help screen for GEOS-Chem.

REMARKS:

```
To build the programs, call "make" with the following syntax:

make TARGET [ OPTIONAL-FLAGS ]

To display a complete list of options, type "make help".

Makefile uses the following variables:

Variable Description
```

SHELL Specifies the shell for "make" to use (usually SHELL=/bin/sh)

```
21 Sep 2009 - R. Yantosca - Initial version
   24 Sep 2009 - R. Yantosca - Added info about NONUMA option for PGI
   24 Sep 2009 - R. Yantosca - Now list rosenbrock as default solver
   19 Nov 2009 - R. Yantosca - Updated comments
   23 Nov 2009 - R. Yantosca - Updated comments
   11 Dec 2009 - R. Yantosca - Now get SHELL from Makefile_header.mk
   21 Dec 2009 - R. Yantosca - Added info about HDF5 option
   25 Jan 2010 - R. Yantosca - Added info about TOMAS option
   10 Mar 2010 - C. Carouge - Remove info about TOMAS option. Keep info
                               about tomas target.
   26 Aug 2011 - R. Yantosca - Added info about APM targets
   26 Aug 2011 - R. Yantosca - Add info about the PRECISE=no option
   11 May 2012 - R. Yantosca - Updated to include info about new make options
# Get the Unix shell (in SHELL variable) from Makefile_header.mk
ROOTDIR = ...
include $(ROOTDIR)/Makefile_header.mk
help:
@echo '%%%
               GEOS-Chem Help Screen
@echo
@echo 'Usage: make TARGET [ OPTIONAL-FLAGS ]'
@echo ''
@echo 'TARGET may be one of the following:'
                    Default target (synonym for "lib exe")'
@echo 'all
@echo 'lib
                     Builds GEOS-Chem source code'
@echo 'libcore
                    Builds GEOS-Chem objs & libs only in GeosCore/'
                     Builds GEOS-Chem objs & libs only in GeosHeaders/'
@echo 'libheaders
@echo 'libiso
                     Builds GEOS-Chem objs & libs only in ISOROPIA/'
                     Builds GEOS-Chem objs & libs only in KPP/'
@echo 'libkpp
@echo 'libnc
                     Builds GEOS-Chem objs & libs only in NcdfUtil/'
@echo 'libutil
                     Builds GEOS-Chem objs & libs only in GeosUtil/'
@echo 'ncdfcheck
                     Determines if the netCDF library installation works'
@echo 'exe
                     Creates GEOS-Chem executable'
@echo 'clean
                     Removes *.o, *.mod files in source code subdirs only'
@echo 'realclean
                     Removes all *.o, *mod, *.lib *.a, *.tex, *ps, *pdf files everywhere'
                     Synonym for "make realclean",
@echo 'distclean
@echo 'doc
                     Builds GEOS-Chem documentation (*.ps, *.pdf) in doc/'
@echo 'docclean
                     Removes *.tex, *.pdf, *,ps from doc/'
                     Displays this help screen'
@echo 'help
```

@echo '

```
@echo ''
@echo 'Special targets for mercury simulation:'
@echo 'allhg
                      Default target for Hg simulation (synonym for "libhg exehg")'
@echo 'libhg
                      Builds GEOS-Chem code for Hg simulation'
@echo 'libgtmm
                      Builds GEOS-Chem + GTMM code for Hg simulation'
                      Creates GEOS-Chem executable for Hg simulation'
@echo 'exehg
@echo ''
@echo 'Special targets for TOMAS aerosol microphysics:'
                      Builds GEOS-Chem + TOMAS (synonym for "libtomas exetomas")'
@echo 'tomas
@echo 'libtomas
                      Builds GEOS-Chem + TOMAS objs & libs in GeosTomas/'
@echo 'exetomas
                      Creates GEOS-Chem + TOMAS executable'
                      Removes *.o *.mod files only in GeosTomas/'
@echo 'cleantomas
@echo ''
@echo 'Special targets for APM aerosol microphysics:'
                      Builds GEOS-Chem + APM (synonym for libapm exeapm)'
@echo 'apm
                      Builds GEOS-Chem + APM objs & libs in GeosApm/ subdir'
@echo 'libapm
                      Creates GEOS-Chem + APM executable in GeosApm/'
@echo 'exeapm
@echo 'cleanapm
                      Removes *.o *.mod files only in GeosApm/'
@echo ''
@echo 'OPTIONAL-FLAGS may be:'
                      Options: ifort pgi (default is ifort)'
@echo 'COMPILER=___
                      Disable precise floating point math optimization (for speed),
@echo 'PRECISE=no
@echo 'DEBUG=yes
                      Builds GEOS-Chem for a debugger (with -g -00)'
@echo 'BOUNDS=yes
                      Turns on subscript-array checking (for debug)'
                      Turns OpenMP parallelization on/off (default is yes)'
@echo 'OMP=[yes|no]
@echo 'IPO=yes
                      Turns on optmization options -ipo -static (ifort only)'
                     Turns on -traceback option (ifort only)'
@echo 'TRACEBACK=yes
@echo 'NONUMA=yes
                      Turns on -mp=nonuma option (pgi only)'
                      Specifies which simulation is done. Options: standard SOA'
@echo 'CHEM=___
@echo 'NTRAC=[43|54]
                      Specifies # of tracers for KPP chemical solver. Should use CHEM flag ins
@echo 'KPPSOLVER=___
                      Specifies the integrator used w/ KPP:'
                      Options: lsodes radau5 rosenbrock runge_kutta (default is rosenbrock)'
@echo '
```

(NOTE: This is set by default if you use "make tomas")