



Ministério da
Ciência, Tecnologia
e Inovação



JULES-CCATT-BRAMS

Requisites Necessary

Guide to compile and run

Test Case



Guide to: JULES-CCATT-BRAMS1.0

Author: Demerval Soares Moreira (demerval.moreira@cptec.inpe.br)

CPTEC/INPE/Bazil

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1 INTRODUCTION:

This guide shows the requisites necessary to compile and run JULES-CCATT-BRAMS1.0 model and pre/post processing. Also have a description of a Test Case. Further details can be obtained in BRAMS homepage: <http://brams.cptec.inpe.br>

JULES-CCATT-BRAMS1.0 was originated of coupling between jules-v3.0 and CCATT_BRAMS4.3.3.

2 REQUISITES:

2.1 Necessary:

- a) A good machine (preference for scalar machine with over 100 processors)
- b) mpich2 (<http://phase.hpcc.jp/mirrors/mpi/mpich2/>)
- c) LINUX operating system (or UNIX, but not tested in this system)
- d) FORTAN-90 compiler (preference to PGI-FORTRAN)
- e) Netcdf library (<http://www.unidata.ucar.edu/downloads/netcdf/index.jsp>)
- f) HDF4 library (<http://www.hdfgroup.org/products/hdf4/>)
- g) zlib library (<http://www.zlib.net/>)
- h) jpeg library (<http://www.iijg.org/>)

2.2 Recommended versions (were used by me)

- a) CRAY Cluster (2.1 GHz AMD Opteron processors) using 360 processors
- b) xt-mpich2_5.5.4 (Cray MPICH2 Message Passing Interface)
- c) SUSE Linux - 2.6.27.48-0.12-default - x86_64
- d) PGI_VERSION 11.3 (pgf90 11.3-0 64-bit)
- e) netcdf-3.6.2
- f) hdf-4.2.5
- g) zlib-1.2.7
- h) jpeg-8d

3 TO COMPILE THE LIBRAIES (Netcdf, HDF4, zlib and jpeg):

➤ Libraries and JULES-CCATT-BRAMS must be compiled with the some compiler.

- a) Shell command: "export CC=<C compiler>" (ex: export CC=pgcc)
- b) Shell command: "export F77=<F77 compiler>" (ex: export F77=pgf77)
- c) Shell command: "export FC=<F90 compiler>" (ex: export FC=pgf90)
- d) Go to: ./source/LIBS/netcdf-3.6.2 or ./source/LIBS/hdf-4.2.5 or ./source/LIBS/zlib-1.2.7 or ./source/LIBS/jpeg-8d
- e) Execute: DIR=\$(pwd); ./configure --prefix=\$DIR/installed
- f) Execute: make
- g) Execute: make install

h) Repeat items d-g until to compile the four libraries

4 TO COMPILE JULES-CCATT-BRAMS:

- a) Open in a ascii editor the file: `./source/JULES-CCATT-BRAMS1.0/src/brams/jules/LIB/Makefile`
- b) On line 56 inform the netcdf PATH
- c) On lines 108 and 118 inform the compiler name
- d) If pgf is not your compiler, create a file to your compiler with name: `./source/JULES-CCATT-BRAMS1.0/build/bin/include.mk.<COMPILES>`, similar to: `./source/JULES-CCATT-BRAMS1.0/build/bin/include.mk.pgf`
- e) Open in a ascii editor the file: `./source/JULES-CCATT-BRAMS1.0/build/bin/include.mk.<COMPILES>`
- f) In lines 25 and 27, inform the compiler name
- g) In lines 31, 34 and 36 you can change the options to your compilation.
- h) Execute: `./source/JULES-CCATT-BRAMS1.0/build/bin/comp.bash`

➤ If everything was alright, was generated the executable: `./source/JULES-CCATT-BRAMS1.0/build/ccatt-brams-4.3-<COMPILES>-CO2_JULES`

5 TO EXECUTE JULES-CCATT-BRAMS:

5.1 Requisites:

- a) JULES-CCATT-BRAMS Executable
 - ⇒ obtained above (`ccatt-brams-4.3-<COMPILES>-CO2_JULES`)
- b) CCATT-BRAMS namelist (RAMSIN)
 - ⇒ There is one model in: `./source/JULES-CCATT-BRAMS1.0/`
 - ⇒ Documentation BRAMS variables:
 - <http://downloads.cptec.inpe.br/~rdown/upload/brams40-namelist-final.pdf>
 - ⇒ Documentation CCATT variables:
 - http://downloads.cptec.inpe.br/~rdown/upload/BRAMS4.0_CATT_First_Time_User_Guide_Ver1.pdf
- c) JULES namelist (jules.in)
 - ⇒ There is one model in: `./source/JULES-CCATT-BRAMS1.0/`
 - ⇒ Documentation JULES variables: `./source/JULES-CCATT-BRAMS1.0/src/brams/jules/LIB/DOCS/jules_v3.0_users_guide.pdf`
- d) Soil Carbon map (soil_carbon.txt), only if `cs!=-1` in jules.in (line 554)
 - ⇒ Is a ascii file with three cols (longitude, latitude and soil carbon (C/m²)) and a lot of lines (depends of region domain and resolution)
- e) Atmospheric and CO₂ initialization and boundary condition (ex: `dpCO2-2010-03-25-0000.vfm`)
 - ⇒ Use geraDP program (<http://brams.cptec.inpe.br/geraDP.shtml>) to convert atmospheric variable (wind, temperature, geopotential and relative humid) in BRAMS format (ex: `dp2010-03-25-0000`)

- ⇒ Use `./source/Utils/join-dp_co2.f90` to join file above (dp) with CO2 boundary condition (Ex. from TM5 model)
- f) Source emissions (ex: `Queima_source-T-2010-03-25-000000-g1.vfm`)
 - ⇒ Use: `./source/PREP-CHEM-SRC-1.2`
 (http://brams.cptec.inpe.br/in_data_catt_burn_map.shtml)
- g) NDVI (http://brams.cptec.inpe.br/in_data_ndvi_modis.shtml)
- h) Soil Moisture (http://brams.cptec.inpe.br/in_data_soil_moisture.shtml)
- i) SST (http://brams.cptec.inpe.br/in_data_gl_weekly_sst.shtml)
- j) Soil Textural Class (http://brams.cptec.inpe.br/in_data_soil_textural.shtml)
- k) Landuse (http://brams.cptec.inpe.br/in_data_landuse.shtml)
- l) Topography (http://brams.cptec.inpe.br/in_data_topography.shtml)

5.2 Hint to execute the model

- a) Create a folder named "run" (`mkdir run`)
- b) Enter in "run" (`cd run`)
- c) Create a link to executable (`ln -s ../source/JULES-CCATT-BRAMS1.0/build/ccatt-brams-4.3-pgf-CO2_JULES`)
- d) Create a link to tables folder (`ln -s ../source/JULES-CCATT-BRAMS1.0/tables`)
- e) Copy RAMSIN_model (`cp ../source/JULES-CCATT-BRAMS1.0/RAMSIN_model RAMSIN`)
- f) Copy jules.in_model (`cp ../source/JULES-CCATT-BRAMS1.0/jules.in_model jules.in`)
- g) Open RAMSIN in ascii editor (`gedit`, `nedit`, `vi`, ...) to make the necessary changes
- h) Create the outputs folders: IVAR, HIS, ANL, sfc and tmp (`mkdir IVAR HIS ANL sfc tmp`)
- i) Execute the model for phases: MAKESFC, MAKEVFILE and INITIAL (see: http://downloads.cptec.inpe.br/~rdown/upload/Brams_First_Time_User_Guide_Ver4.pdf)

6 POST PROCESSING (RAMSPOST)

- Documentation: http://downloads.cptec.inpe.br/~rdown/upload/RAMSPOST_User_guide_ver_01.pdf
- a) To Compile:
 - Open in a ascii editor the file: `./source/Ramspost-5.1/LIB/include.mk`
 - Change to the options of your compiler.
 - Execute: `./source/Ramspost-5.1/comp.bash`
 - b) Configure the namelist (`ramspost.inp`), see: http://downloads.cptec.inpe.br/~rdown/upload/RAMSPOST_User_guide_ver_01.pdf
 - c) Run executable: `ramspost_51`
 - d) Open the forecasts p. ex. in GrADS software (<http://www.iges.org/grads/>)

7 TEST CASE

7.1 Configuration:

- Grid with horizontal resolution of 20 km
- Amazon region (Grid center: 59.0W ; 3.2S)
- X, Y, Z point = 310, 210, 48
- Six hour of integration (01/Mar/2010 00Z until 01/Mar/2010 06Z)
- Forecast output in each hour

7.2 Requisites necessary to this Test Case:

- Download the tarballs: source.tgz and test_case.tgz
- Execute item 3 to compile all libraries in your machine
- Execute item 4 to compile the model in your machine
- Execute item 6 to compile post processing in your machine

7.3 Not necessary to this Test Case:

- Download surface data (topography, NDVI, ...)
- Download Initialization and boundary condition (Atmospheric and CO₂)
- Prepare source emissions
- Change RAMSIN
- Change jules.in

7.4 To execute Test Case:

- Execute item 7.2 and expand all tarballs
 - Execute: `“./TEST_CASE/test_case.bash <nproc>”`, where <nproc> is the number of processor that you want to use.
- If everything was alright, was generated the files:
- `./TEST_CASE/grads/test_g1.gra` (binary with variables: Temperature and wind)
 - `./TEST_CASE/grads/test_g1.ctl` (descriptor to GrADS software)
 - `./TEST_CASE/grads/test_field.png` (figure generated by GrADS)
 - `./TEST_CASE/grads/test_serie.png` (figure generated by GrADS)