# A Quick Introduction to TMM

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#### Abstract

This document goes through how to use the Transport Matrix Model (TMM) using computational resources at the University of Minnesota's Minnesota Supercomputing Institution (MSI). All the source codes and documents (including this tutorial) that Tanioka made to TMM is available at my github website (https://github.com/tanio003/tmm/tree/TT\_Release).

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## 1 Setting up

### 1.1 Flow Chart

Before you do anything, read "README.txt" by Samar Khatiwala at the following website: <a href="https://github.com/samarkhatiwala/tmm">https://github.com/samarkhatiwala/tmm</a>. I will go over each of these steps specifically aimed at audiences using the computational cluster *Mesabi*.

- 1. Installing and configuring PETSc
- 2. Downloading all the scripts and transport matrices into your own local directory
- 3. Compiling the model (we use the BGC model MOPS2 for this example)
- 4. Running the model
- 5. Processing the model outputs
- 6. Displaying the model outputs

## 1.2 Steps

#### 1.2.1 Step 0: Logging into MSI and Mesabi

Open the terminal (assuming that you have a MAC or Linux environment) on your computer and log in to MSI with your x500 account:

```
$ ssh -Yt youremail@umn.edu
```

Log in to Mesabi:

```
$ ssh -X mesabi
```

Make a new directory called TMM2 in your home directory and enter into this directory. Everything related to TMM will go into this directory.

```
$ mkdir TMM2
$ cd TMM2
```

## 1.2.2 Step 1: Installing and configuring PETSc

Download the latest version of PETSc and save it in your TMM2 directory and unzip this package. If opened properly, you should see the new directory petsc-3.13.5.

```
$ wget http://ftp.mcs.anl.gov/pub/petsc/release-snapshots/petsc-lite-3.13.5.tar.gz
$ tar -xvf petsc-lite-3.13.5.tar.gz
$ ls
petsc-3.13.5
```

Import the required modules: (1) impi, (2) impi/intel, and (3) cmake. Also make sure that you are using python 3, not python 2 (= default for MSI).

```
$ module purge
$ module load intel
$ module load impi/intel
$ module load cmake
$ module load python3
$ module list
Currently Loaded Modulefiles:
1) intel/2018.release(default) 4) cmake/3.10.2(default)
2) intel/2018/release 5) python3/3.7.1_anaconda
3) impi/intel(default)
```

Set up \$PETSC\_DIR to your petsc-3.13.5 directory:

```
$ export PETSC_DIR=$HOME/TMM2/petsc-3.13.5
$ echo $PETSC_DIR
/.../TMM2/petsc-3.13.5
```

Configure PETSc. Although this part is quite tricky you can copy and use my config file "reconfigure-arch-linux-c-opt.py". If the config file does not work properly, let me know and I can show you a way to compile without using this .py file.

```
$ cd petsc-3.13.5
$ cp ~/../tanio003/TMM2/petsc-3.13.5/config/reconfigure-arch-linux-c-opt.py config/
$ config/reconfigure-arch-linux-c-opt.py
```

Don't worry about some warning signs. It takes few minutes to compile. If it's compiled properly you should see the notice "Conifgure stage complete." Then build PETSc library:

```
$ make all
```

Building process takes about 15-30 minutes. If you're very lucky it will go through in a single shot. But in most cases, it fails during the middle of the process. Don't worry if it fails the first time. Simply type "\$ make all" again and hopefully it will finish building from where it left off. If built properly, you should see the message "Now to check if the libraries are working do:...". Then type,

```
$ make check
...
Completed test examples
```

If you get this far, you've managed to build the PETSc successfully and you're ready to go to the next step. If it failed, read the error messages, debug, and try again. **Building PETSc is harder than it looks** so you need to be patient.

As more of a technical note, the procedure above uses the Intel compilers and Intel MPI library. By loading the cmake, the PETSc build system can learn more about the host machine. In addition to taking advantage of compiler optimizations and vectorization, the procedure above builds PETSc against the Intel Math Kernel Library (MKL) for BLAS, LAPACK and ScaLAPACK which gives a performance gain over the reference implementations. For the FORTRAN compiler, we specifically need to use mpiifort, and not mpif90 (the default compiler), because TMM codes are written in both F77 and F90. Also, since we don't require C++ for TMM we put the flag in the config file, --with-cxx=0. The reason we need to use MPI compilers, not regular gcc compilers, is because we want to run PETSc in a parallel mode (i.e., by using the command mpiexec in the runscript). For more details about building PETSc please check out https://www.mcs.anl.gov/petsc/documentation/installation.html.

#### 1.2.3 Step 2: Downloading all the scripts and transport matrices

 First download Matlab scripts from http://kelvin.earth.ox.ac.uk/spk/Research/TMM/tmm\_matlab\_code.tar.gz and put into the first level of your TMM2 folder Path. You could also get my copy.

```
$ cp -r ~/../tanio003/TMM2/tmm_matlab_code $HOME/TMM2/
```

2. Download transport matrices and related data for the model of your choice: <a href="http://kelvin.earth.ox.ac.uk/spk/Research/TMM/TransportMatrixConfigs/">http://kelvin.earth.ox.ac.uk/spk/Research/TMM/TransportMatrixConfigs/</a> and put into he first level of your TMM2 folder Path. You can download all 6 configurations but I warn you that MITgcm\_ECCO\_v4 and UVicKielIncrIsopycDiffTransient take a very long time. You could also grab my copy (e.g., to copy MITgcm\_ECCO).

```
$ cp -r ~/../tanio003/TMM2/MITgcm_ECCO $HOME/TMM2/
```

3. Download miscellaneous data called OceanCarbon from <a href="http://kelvin.earth.ox.ac.uk/spk/Research/TMM/MiscData/">http://kelvin.earth.ox.ac.uk/spk/Research/TMM/MiscData/</a>. You can get my copy by:

```
$ cp -r ~/../tanio003/TMM2/OceanCarbon $HOME/TMM2/
```

4. Download source codes for TMM and models:

```
$ git clone https://github.com/tanio003/tmm
```

This directory (/TMM2/tmm) contains the source codes from Khatiwala's master branch ("master") and my public release branch ("TT\_Release"). For our exercise, we will be using some of my new codes so switch from the master branch to my branch in the newly created tmm directory:

```
$ cd tmm
(master) $ ls
driver HOWTO.txt LICENSE.txt models README.txt
(master) $ git checkout TT_Release
(TT_Release) $ ls
driver HOWTO.txt LICENSE.txt models README.txt Tutorial_MSI
```

Notice that in the branch TT\_Release, there is a new directory Tutorial\_MSI, which was not present in the master branch.

(Optional) If you want to make start changing your own changes, I would suggest making a new branch in your local computer (e.g., yournewrepo), and leave master and TT\_Release untouched.

```
(TT_Release) $ git checkout -b yournewrepo
(yournewrepo) $ git branch --show-current
yournewrepo
```

5. Set the environment variable TMMROOT to point to the top level of the TMM directory.

```
(TT_Release) $ export TMMROOT=$HOME/TMM2/tmm
(TT_Release) $ echo $TMMROOT
/home/.../TMM2/tmm
```

#### 1.2.4 Step 3: Compiling the model

Here, let's try compiling biogeochemical MOPS.

For each model there is model-specific source codes
 (\$TMMROOT/models/current/mops2/src/); Matlab script(s) in
 \$TMMROOT/models/current/mops2/matlab/ to generate input data and read model output; and run scripts and other runtime data such as namelists in runscripts/.

First, we create a new "Run directory". I make a new base directory Runs and in that directory, I make subdirectories for specific experiments. I call it Runs/MOPS/Test\_spinup and copy here all the files needed.

```
$ cd ~/TMM2
$ mkdir -p Runs/MOPS/TMM_spinup
$ cd Runs/MOPS/TMM_spinup
$ cp -p $TMMROOT/models/current/mops2.0/src/Makefile .
$ cp -p -R $TIMMROOT/models/current/mops2.0/matlab/* .
$ cp -p $TMMROOT/models/current/mops2.0/runscripts/* .
```

2. Compile mops. Make sure that all the modules are loaded.

```
$ module load intel
$ module load impi/intel
$ module load cmake
$ make clean all
$ make mops
```

If compiled properly, you'd find a new executable "mops" created along with a bunch of objective .o files.

```
BGC_INI.o
BGC_MODEL.o
CAR_CHEM.o
                                            n7fluxes28.m
                                            n7physics.m
                                            n7tracers28.m
CAR_INI.o
external_forcing_mops_biogeochem.o
                                           n7tracersavg28.m
perry1996-runoff-noarctic_noname.txt
insolation.o
                                            perry1996-runoff_noname.txt
load_output.m
                                            petsc_matvec_utils.o
load_output_time_avg.m
load_pco2.m
                                            petsc_signal_utils.o
process_output.m
                                            runscript
make_input_files_for_mops_model.m
make_rivers.m
misfit_mops_biogeochem.o
                                            tmm_external_bc.o
                                            tmm_forcing_utils.o
tmm_forward_step.o
mops_biogeochem_copy_data.o
                                            tmm_main.o
mops_biogeochem_diagnostics.o
                                            tmm_monitor.o
mops_biogeochem_ini.o
                                            tmm_profile_utils.o
mops_biogeochem_misfit.o
                                            tmm_timer.o
{\tt mops\_biogeochem\_model.o}
                                            tmm_write.o
mops_biogeochem_set_params.o
```

3. Edit the file make\_input\_files\_for\_mops\_model.m. First thing to do is to make sure that variable base\_path point to the right directory for the TMM configuration.

```
% make_input_files_for_mops_model.m
% Set toplevel path to GCMs configuration
% base_path = '/data2/spk/TransportMatrixConfigs/MITgcm_2.8 deg';
% base_path = '/data2/spk/TransportMatrixConfigs/MITgcm_ECCO';
% base_path = '/data2/spk/TransportMatrixConfigs/MITgcm_ECCO_v4';
base_path = '/data2/spk/TransportMatrixConfigs/MITgcm_ECCO_v4';
base_path = '~/TMM2/MITgcm_2.8 deg';
addpath(genpath('~/TMM2/tmm_matlab_code'));% add tmm_matlab_code to the search path oceanCarbonBasePath = '~/TMM2/OceanCarbon'; % add OceanCarbon to the search path atmosDataPath = fullfile (oceanCarbonBasePath, 'AtmosphericCarbonData');
```

In the same matlab file, there are different switches with 0's and 1's. For this spin-up exercise, we **couple MOPS to a simple OCMIP-like carbon model** and a **fix atmospheric** pCO<sub>2</sub> **at 280 ppm**. So set the switches as following:

```
% make_input_files_for_mops_model.m
periodicForcing=1
periodic Matrix =1
dt=43200; % time step to use (43200s for ECCO and MIT2.8; 28800s for any other TMMs)
rearrangeProfiles=1
bigMat=0
writeFiles=1
writeTMs=1
useCoarseGrainedMatrix = 0\\
writePCFiles=0
READ_SWRAD=0
                                   % Read short-wave radiation?
useCarbon=1
                                   % Use simple inorganic carbon model?
useAtmModel=0
                                   % Use prognostic 1-box atmosphere?
pCO2atm_ini=280.0
                                   % Initial pco2?
useTimeVaryingPrescribedCO2=0
                                  % Use prescribed pco2 pathway?
% Use DIC and Alk to calculate E-P?
% Scaling factor for E-P (default = 1)
useVirtualFlux=1
empScaleFactor=1.0
% Modified by Tatsuro Tanioka 200907 to allow for Atmospheric CO2 option
% For a prescribed pCO2 run, useTimeVaryingPrescribedCO2=1 and choose a scenario
% Available options: 'historical', 'RCP3PD', 'RCP45', 'RCP6' and 'RCP85'
co2Scenario='RCP85';
```

Then open MATLAB and run make\_input\_files\_for\_mops\_model.m

This creates a bunch of periodic forcing files (xxx\_01, xxx\_02,...), initial tracer concentrations (po4ini.petsc, no3ini.petsc,...), and binary files (.bin and .petsc) related to model geometry and forcing.

#### 1.2.5 Step 4: Running the model

MSI systems use job queues to efficiently and fairly manage when computations are executed. The queuing system at MSI is called PBS (Portable Batch System) and to submit a job to a PBS queue users create PBS job scripts. PBS script contains information on the resources requested for calculation, as well as the commands for executing the calculation.

Below is the custom PBS script for submitting a new job to run MOPS2 using Mesabi. It's called runscript\_msi and should be in the current directory already. Here is the first 11 lines:

```
1 #!/bin/bash -1
2 #PBS -1 walltime = 01:00:00, nodes = 1:ppn = 24, pmem = 2580mb
3 #PBS -m abe
4 #PBS -j oe
5 #PBS -M tatsurobkkuk@gmail.com
6
7 cd $PBS_O_WORKDIR
8
9 module load intel
10 module load impi/intel
11 module load cmake
```

The first line defines which type of shell the script will be read. Here we will use the bash. The second line contains the PBS resource request. The current job wull require 1 hour, 1 nodes each with 24 processor cores (ppn), and 2580 megabytes of memory per core (pmem).

The two lines containing #PBS -m abe, and #PBS -M tatsurobkkuk@gmail.com. are both commands having to do with sending message emails to the user. The first of these lines instructs the PBS system to send a message email when the job aborts, begins, or ends. The second command specifies the email address to be used. Using the message emails is recommended because the reason for a job failure can often be determined using information in the emails. The seventh line sets the directory at which commands are executed and the lines 8-11 loads the necessary software modules.

The follwing lines  $(12\sim)$  contain the commands to execute and start a specific program. Different flags need to be change accordingly depending on the nature of experiments.

```
# 360 days per year with a time step of 2 steps per day:
14 mpiexec -np 24 -hostfile $PBS_NODEFILE ./mops \ # number of cores, models
15 -numtracers 9 \ # number of tracers (i.e. state variables)
16 — i po4ini.petsc,dopini.petsc,oxyini.petsc,phyini.petsc,zooini.petsc,detini.petsc,no3ini.petsc,dicini.petsc,alkini.petsc \ # files for initialization of BGC state variables
        -me Ae \ # the name of the explicit transport matrix
-mi Ai \ # the name of the implicit transport matrix
-t0 0.0 -iter0 0 \ # starting time[years] starting time[timesteps]: for initial run
-deltat_clock 0.001388888888888 \ # ocean timestep length[years]: 2 timesteps/day
17
         -max_steps 2160000 \ # total number of timesteps to be evaluated (here 3000 yrs) -write_time_steps 72000 \ # output frequency(in timesteps, here every 100 yrs)
23 -o po4out.petsc,dopout.petsc,oxyout.petsc,phyout.petsc,zooout.petsc,detout.petsc,no3out.petsc,dicout.petsc,alkout.petsc \ # files for output of 9 BGC state variables
         -external_forcing \ # calculate BGC explicitly
24
        -use_profiles \
-nzeuph 2 \ # number of layers in euphotic zone (2 for MIT2.8, 6 for ECCO)
         -biogeochem_deltat 43200.0 -days_per_year 360.0 \ # ocean timestep[seconds]
        -burial_sum_steps 720 \ # sum burial over a period of 720 timesteps = 1 yr
-pco2atm 280.0 \ # use fixed pCO2 of 280 ppm
-use_virtual_flux \ # use the global surface mean DIC and Alk to calculate E-P
-periodic_matrix \ # use of periodic transport matrix
28
29
30
31
         -matrix_cycle_period 1.0 -matrix_num_per_period 12 \ # the unit of time is year and
# circulation has a periodicity of 1 year; monthly mean tranport matrix (12 TMs/year)
33 -periodic_biogeochem_forcing \ # periodic biogeochemical forcing
34 -periodic_biogeochem_cycle_period 1.0 -periodic_biogeochem_num_per_period 12 \
         -num_biogeochem_steps_per_ocean_step 8 \ # the number of BGC timestep per ocean step -separate_biogeochem_time_stepping \ # timestep BGC model separately from ocean step -time_avg -avg_start_time_step 2159281 -avg_time_steps 60 \
35
36
# initial timestep for avg concentrations (for final year); average over 60 timesteps =1 m
         -avg\_files \ po4avg.petsc\ , dopavg.petsc\ , oxyavg.petsc\ , phyavg.petsc\ , zooavg.petsc\ , \\
detavg.petsc,no3avg.petsc,dicavg.petsc,alkavg.petsc \ # avg file names 39 -calc_diagnostics -diag_start_time_step 2159281 -diag_time_steps 60 \
# initial timestep for diagnostic fluxes (for final year); average over 60 timesteps = 1 mo
40  -diag_files fbgc1.petsc,fbgc2.petsc,fbgc3.petsc,fbgc4.petsc,fbgc5.petsc,fbgc6.petsc,
                                                           files names
fbgc7.petsc \ # diagnsotic flux
             > log
                          # outputting to logfile
```

Currently BGC model writes the following 7 diagnostics into the different files:

- 1. fbgc1.petsc primary production in each box [mmol P/m3/oceantimestep]
- 2. fbgc2.petsc zooplankton grazing in each box [mmol P/m3/oceantimestep]

- 3. fbgc3.petsc detritus sedimentation through upper boundary of each box [mmol P/m2/oceantimestep]
- $4. \ \ \textbf{fbgc4.petsc} \ remineralization \ of \ detrius \ and \ DIP \ in each \ box \ [mmol\ P/m3/ocean time step]$
- 5. fbgc5.petsc river runoff [mmol P/m3/oceantimestep]
- 6. fbgc6.petsc nitrogen fixation [mmol N/m3/oceantimestep]
- 7. fbgc7.petsc denitrification [mmol P/m3/oceantimestep]