# Microbial-ENzyme Decomposition (MEND) Model MANUAL

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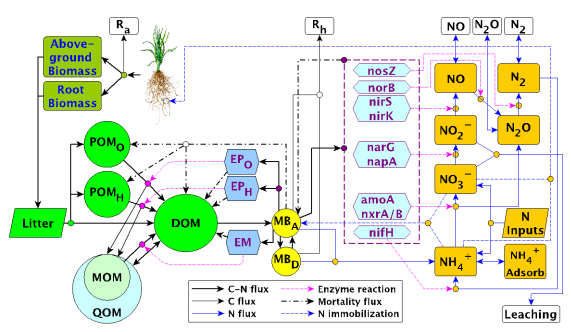


Figure 1 Diagram of the C-N coupled Microbial-ENzyme Decomposition (MEND) model

R<sub>a</sub> and R<sub>h</sub> are autotrophic and heterotrophic respiration, respectively. POM<sub>O</sub> and POM<sub>H</sub> are particulate organic matter (POM) decomposed by oxidative (EP<sub>O</sub>) and hydrolytic enzymes (EP<sub>H</sub>), respectively. MOM is mineral-associated OM, which is decomposed by a mixed enzyme group EM. Dissolved OM (DOM) interacts with the active layer of MOM (QOM) through sorption and desorption. Litter enters POM<sub>O</sub>, POM<sub>H</sub>, and DOM. Microbes consist of active (MB<sub>A</sub>) and dormant microbes (MB<sub>D</sub>). DOM can be assimilated by MB<sub>A</sub>. Mineral N deposition and fertilization enter NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup> that can be immobilized by microbes and taken up by plant roots. NH<sub>4</sub><sup>+</sup> adsorption is also considered. N fixation, nitrification and denitrification are mediated by nitrogenase (nifH), ammonia oxidases (amoA, nxrA/B) and N-reductases (narG/napA, nirS/nirK, norB, nosZ), respectively. Mineral N loss pathways include leaching (NO<sub>3</sub><sup>-</sup> and NO<sub>2</sub><sup>-</sup>) and gas emission (NO, N<sub>2</sub>O, and N<sub>2</sub>) from soil to atmosphere.

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# Table of Contents

1	Mod	Model Repositories4				
2	Dire	ctories	4			
3	Majo	or Subroutine Calls	5			
3.	.1	MEND & MENDcn	5			
3.	.2	MEND_mult				
4	Sour	ce Code Files & Major Functions & Subroutines	7			
5	Cont	rol & Output files	8			
6	Deriv	ved Rates: *_RATE_hour.out	9			
7	SOIL	_INI.dat: Model Initialization	10			
8	Inpu	t Data	11			
9	Unce	ertainty Quantification (UQ) by Critical Objective Function Index (COFI): Inputs &				
Out	puts		12			
10	sINI9	%iScenario	13			
11	Scen	ario Design	13			
12	sINI9	%iKenetics	14			
13	sINI9	%iHR	14			
14	sINI9	%iTmp_Func	14			
15		· – %iError				
16	MEN	ID State Variables (C & N Pools)	16			
17		ID Parameters				
1	7.1	MEND Parameters				
1	7.2	MEND Parameters that may be pre-determined				
18	Resp	onse Variables for Model Calibration/Optimization	22			
18	8.1	Calibration Variables in MEND_namelist.nml	22			
	8.2	Add a new Calibration Variable				
19		meter Optimization Algorithm				
20		%VARopt_int				
21	CN R	atios in 3 Litter Pools	24			
22	MEN	ID (non-MPI) Configuration/Compiling on OSX & Cygwin/Unix	25			
2	2.1	Dependency of source code				
	2.2	gcc/gfortran Installation				
	2.3 2.4	gdb Installation on Mac OS Mojave  NetBeans Configuration on OSX & Windows				
	2.4 2.5	Compiling on Unix				
	2.6	Text Editor				
2	2.7	Eclipse Configuration for non-MPI MEND	32			
23	Tips	on unix and git commands	34			
2	3.1	iTerm2:	34			
2	3.2	git	34			

# Model Repositories

Model Version	Notes
MEND	Carbon-Nitrogen coupled version: <b>sINI%Carbon only = .FALSE.</b>
	https://github.com/wanggangsheng/MEND.git
MEND mult	Multiple-Case Version of MEND: run multiple cases in one-run
_	https://github.com/wanggangsheng/MEND mult

# Directories

ID	Directory	Notes
0	model dir	Control Files: MEND_namelist.nml
1	src	Source code
2	Userio/inp	Input data
		MEND_mult: includes sub-folder for each case
3	Userio/out	Output data
		MEND_mult: includes sub-folder for each case

# Major Subroutine Calls

# 3.1 MEND & MENDcn

0	1	2	3	4
MEND main	MENDIN			
	SCE	fMEND_OBJ	Par: sINI%LCI0	
			Par: sINI%r0	
			subMEND_INI	subMEND_Files_Open
				subMEND_CPOOL_UPDATE1
				subMEND_NPOOL_UPDATE1
				subMEND CN UPDATE0
			subMEND_RUN	subMEND_PAR
				subMEND
				subMEND_output_rate
				subMEND_output
				sOUT_OPT_h
				sOUT_Day2Mon
				Extract data for
				comparison/calibration

#### 3.2 MEND\_mult

0	1	2	3	4
MEND main	MENDIN			
	SCEUA	fMEND_OBJ	Par: sINI%LCI0	
			Par: sINI%r0	
			DO iCase=1,nCase	
			MENDIN_CASE	
			subMEND_INI	
			subMEND_RUN	subMEND_PAR
				subMEND
				subMEND_output_rate
				subMEND_output
				sOUT_OPT_h
				sOUT_Day2Mon
			END DO	

- 'SIM\_obs.out' combines all '\*\_SIM\_obs.out' for all cases into 1 output file.
- After each MEND run, the total objective function (fMEND\_OBJ) for multiple cases is calculated by reading data in this file.
- MEND\_mult allows different calibration-variables used in various cases. e.g., case1 with 2 variables (CO2 & MBC), case2 with 2 variables (CO2 & DOC). Under this condition, the total OBJ (fMEND\_OBJ) will include 3 single objectives (CO2, MBC, & DOC).
- 'MEND\_namelist.nml' Cali\_OBJ\_Weight(·) defines OBJ weighting factors for these objectives combined from multiple cases; whereas the OBJ weighting factors for each case ('case.ini' Line 48-57) are NOT used for calibration/optimization.
- MEND\_mult runs much slower than MEND. Because MEND\_mult REPEATEDLY reads input
  data for each case (subroutine 'MENDIN\_CASE') during each model run. However, MEND
  only read input data ONCE for a single case.

# 4 Source Code Files & Major Functions & Subroutines

ID	F90 file	Notes	Major Functions & Subroutines
1	MEND main	Main program	,
2.1	MEND IN	Control file	○ MENDIN(sSCE,sINI)
2.2	MENDIN_CASE	Input data for each case MEND mult only	, ,
3	MOD MEND TYPE	Data structure for MEND	
4	MOD_MEND	MEND model; Depends on  MOD_MEND_TYPE  MOD_USRFS	<ul> <li>subMEND: MEND model</li> <li>subMEND_PAR: MEND parameters modified by temperature, moisture, etc.</li> <li>subMEND_RUN: run model continuously with multiple time-steps</li> <li>subMEND_INI: model initialization</li> <li>fMEND_OBJ: objective function for model evaluation &amp; optimization</li> <li>sINP_Read: read input, soil temp &amp; moisture</li> <li>subMEND_Files_Open: open output files</li> <li>subMEND_Files_close: close output files</li> <li>sOUT_OPT_VAR_Extract: extract HOURLY outputs for response variables used for optimization</li> <li>subMEND_output: HOURLY outputs for all state variables &amp; fluxes</li> <li>sOUT_tscale: convert HOURLY outputs (subMEND_output) to DAILY, MONTHLY &amp; YEARLY outputs</li> <li>sOUT_ALL_tscale: convert HOURLY outputs (subMEND_output) to DAILY, MONTHLY &amp; YEARLY outputs, called by sOUT_tscale</li> <li>sSOBOL_VAR_OBJ: calculate obj for Sobol sensitivity analysis</li> <li>sSOBOL_VAR_Read: Read _VAR_day.out to calculate Sobol obj</li> </ul>
5	MOD_OPT_TYPE	Data structure for model optimization	
6	MOD_OPT	Optimization algorithm	<ul><li>○ SCE</li><li>Depends on</li><li>■ MOD_OPT_TYPE</li><li>■ MOD_MEND</li></ul>
7	MOD_STRING	String utility	<ul><li>StrCompress()</li></ul>
8	MOD_USRFS	User Functions and Subroutines	
9	MOD_MCMC	MCMC algorithm	

# 5 Control & Output files

ID	FILE	Notes	
	CONTROL		
	FILE:		
1	MEND namelist.nml	(1) MEND CONTROL file, stored in the model root dir	
-		o iModel =	
		'0'-run MEND model with parameter sets 'Pinitial';	
		'1'-model calibration/optimization	
		'2'- uncertainty quantification (UQ) using COFI method, output COFIpar.out	
		'3'-generate objective function values for Sobol Sensitivity analysis	
		'4'-UQ using MCMC	
		'5'-UQ with COFI, output both COFIpar.out & COFIvar.out (see Table 7)	
		'6'-find unfinished parameter sets in previous run (iModel = 3)	
		o sSite: site name	
		(2) MEND_mult CONTROL file	
		Line 11-14: define multiple cases, case names are dir names in inp & out	
2	**.ini	CONTROL/INITIAL file for each case, in inp/casedir	
	<b>INITIAL FILE:</b>		
1	SOIL_INI.dat	Initialization of SOC pools, stored in 'userio/inp'	
	<b>OUTPUT FILES:</b>	MEND_mult: Outputs of optimization for all-case are saved in	
		out; Outputs for each case are saved in the out/casedir	
1	* SIM obs.out	Outputs for SIM vs. OBS for those specific days with OBS.	
		VAR: corresponds to VARid in 'MEND_namelist.nml'.	
		Note: The 'best' parameter set among those 'best' from multiple opt-runs is	
		shown at the last line of this file. You may copy this line to	
		'MEND namelist.nml' for future run.	
2	* SIM day.out	Continuous daily SIM results for those variables used for calibration	
3	* SIM mon.out	Continuous monthly SIM results for those variables used for calibration	
4	*_VAR_hour.out	All state variables: hourly	
5	* VAR_day.out	All state variables: daily	
6	* VAR mon.out	All state variables: monthly	
7	* VAR_year.out	All state variables: yearly	
8	* FLX hour.out	All fluxes: hourly	
9	* FLX day.out	All fluxes:daily	
10	* FLX mon.out	All fluxes: monthly	
11	* FLX year.out	All fluxes: yearly	
12	* ITW hour.dat	External input (litter), T, SWC, SWP: hourly	
13	* ITW day.dat	External input (litter), T, SWC, SWP: daily	
14	* ITW mon.dat	External input (litter), T, SWC, SWP: monthly	
15	* RATE hour.out	Derived rates: hourly, e.g., active fraction	
16	* RATE day.out	Derived rates: daily	
17 18	* RATE mon.out  * RATE year.out	Derived rates: wearly	
19	* PAR hour.out	Derived rates: yearly Parameters modified by T, SWP, pH, etc: hourly	
20	* PAR day.out	Parameters modified by T, SWP, pH, etc. hourly  Parameters modified by T, SWP, pH, etc. daily	
21	* PAR day.out	Parameters modified by T, SWP, pH, etc. daily  Parameters modified by T, SWP, pH, etc. monthly	
22	* PAR year.out	Parameters modified by T, SWP, pH, etc. monthly  Parameters modified by T, SWP, pH, etc. yearly	
23	* OPT end.out	'best' parameter sets from multiple independent optimizations	
24	* OPT ini.out	Initial parameters for optimization, log file for any MEND run	
25	* OPT all.out	All parameter sets during optimization, used for COFI UQ	
26	* Ndep hour.out	Mineral N input: hourly	
∠∪	nuep_nour.out	Ministra in input, hourry	

# 6 Derived Rates: \*\_RATE\_hour.out

Col	Name	Units	Notes
1	Hour		Hourly time-step
2	kPOC1	h-1	Equivalent 1st-order decomposition rate;
			k=VP1*EP1/(POC1 + KP1)
3	kPOC2	h-1	Equivalent 1st-order decomposition rate;
			k=VP2*EP2/(POC2+ KP2)
4	kMOC	h-1	Equivalent 1st-order decomposition rate;
			k=VM*EM/(MOC+ KM)
5	kDOC	h-1	Equivalent 1st-order turnover rate;
			k=[(Vg+Vmt)/Yg]*MBa/(DOC + KD)
6	kMBa	h-1	Equivalent 1st-order turnover rate;
			k=[(Vg+Vmt)*(1/Yg-1)]*phi + rMORT +
			pEP+ pEM) * Vmt
7	kMBa_in	h-1	Equivalent 1st-order microbial uptake rate,
			k=[(Vg+Vmt)/Yg]*phi +
			Resuscitation/MBa
8	kMBd	h-1	Output rate of dormant microbes,
			k=(Resuscitation+Maintenance)/MBCd
9	kMBd_in	h-1	Input rate for dormant microbes,
			k=Dormancy/MBCd
10	kMB	$h^{-1}$	Turnover rate of total MBC,
			k=(CO2_gm+ENZ_prod+Mortality)/MBC
11	kMB_in	h-1	Assimilation rate of total MBC,
			k=uptake/MBC
12	Phi	_	DOC saturation level, = DOC/(DOC +
			KD)
13	Active_Fraction	_	Fraction of active microbes
14	CUE	_	Apparent carbon use efficiency = [uptake -
			CO2_gmo]/uptake
15	NUE	_	Apparent nitrogen use efficiency = $[CN_{MB}]$
			$-CN_{MB\_min}$ ]/ $(CN_{MB\_max} - CN_{MB\_min})$
16	Balance_Error	mg C g <sup>-1</sup> soil or	Error for balance check, RE=(TOCend -
		mg C cm <sup>-3</sup> soil	TOCbeg) - (TOCinp - TOCout)*dt
17	TOCbeg	mg C g <sup>-1</sup> soil	Total organic carbon at the beginning of
		mg C cm <sup>-3</sup> soil	the time-step
18	TOCend	Same as above	Total organic carbon at the end of the time-
			step
19	TOCinp	Same as above	TOC input during the time-step
20	TOCout	Same as above	TOC output during the time-step
21	STP	°C	Soil temperature
22	SWC	cm <sup>3</sup> cm <sup>-3</sup>	Soil water content
23	SWP	MPa	Soil water potential
24	рН	-	Soil pH

# 7 SOIL\_INI.dat: Model Initialization

nrow= 26

Depth: cm; SOC/POC/MOC/MBC: mgC/cm<sup>3</sup>

1         Depth         10         Soil depth (cm)           2         Sand         0.259         Sand fraction           3         Clay         0.509         Clay fraction           4         CN_MB_mean         8         Mean CN ratio of microbial biomass           5         CN_MB_min         2         Min CN ratio of microbial biomass           6         CN_MB_max         14         Max CN ratio of microbial biomass           7         CN_EP1         3         CN ratio of oxidative enzymes           8         CN_EP2         3         CN ratio of hydrolytic enzymes           9         CN_EM         3         CN ratio of MOM enzymes           10         fQOM         0.01         Fraction of QOM in MOM           11         SOC         21.14         Actually NOT used for modeling           12         POC         5.67         POC1 + POC2           13         MOC         15.47         Mineral-associated Organic Carbon           14         DOC         0.31         Dissolved Organic Carbon           15         MBC         0.53         Default value = 2.5% SOC           16         EP1         0.0011         Default value = 0.005% SOC           18	ID	Property	Value	Notes
3   Clay   0.509   Clay fraction	1	Depth	10	Soil depth (cm)
4         CN_MB_mean         8         Mean CN ratio of microbial biomass           5         CN_MB_min         2         Min CN ratio of microbial biomass           6         CN_MB_max         14         Max CN ratio of microbial biomass           7         CN_EP1         3         CN ratio of oxidative enzymes           8         CN_EP2         3         CN ratio of hydrolytic enzymes           9         CN_EM         3         CN ratio of MOM enzymes           10         fQOM         0.01         Fraction of QOM in MOM           11         SOC         21.14         Actually NOT used for modeling           12         POC         5.67         POC1 + POC2           13         MOC         15.47         Mineral-associated Organic Carbon           14         DOC         0.31         Dissolved Organic Carbon           15         MBC         0.53         Default value = 2-5% SOC           16         EP1         0.0011         Default value = 0.005% SOC           17         EP2         0.0011         Default value = 0.005% SOC           18         EM         0.0014         Default value = 0.005% SOC           19         CN_FOM         26         CN ratio of Particulate Organic Matter (POM	2	Sand	0.259	Sand fraction
5         CN_MB_min         2         Min CN ratio of microbial biomass           6         CN_MB_max         14         Max CN ratio of microbial biomass           7         CN_EP1         3         CN ratio of oxidative enzymes           8         CN_EP2         3         CN ratio of hydrolytic enzymes           9         CN_EM         3         CN ratio of MOM enzymes           10         fQOM         0.01         Fraction of QOM in MOM           11         SOC         21.14         Actually NOT used for modeling           12         POC         5.67         POC1 + POC2           13         MOC         15.47         Mineral-associated Organic Carbon           14         DOC         0.31         Dissolved Organic Carbon           15         MBC         0.53         Default value = 2-5% SOC           16         EP1         0.0011         Default value = 0.005% SOC           17         EP2         0.0011         Default value = 0.005% SOC           18         EM         0.0014         Default value = 0.005% SOC           19         CN_SOM         13         CN ratio of SOM           20         CN_POM         26         CN ratio of Particulate Organic Matter (POM), will be calculated	3	Clay	0.509	Clay fraction
6         CN_MB_max         14         Max CN ratio of microbial biomass           7         CN_EP1         3         CN ratio of oxidative enzymes           8         CN_EP2         3         CN ratio of hydrolytic enzymes           9         CN_EM         3         CN ratio of MOM enzymes           10         fQOM         0.01         Fraction of QOM in MOM           11         SOC         21.14         Actually NOT used for modeling           12         POC         5.67         POC1 + POC2           13         MOC         15.47         Mineral-associated Organic Carbon           14         DOC         0.31         Dissolved Organic Carbon           15         MBC         0.53         Default value = 2.5% SOC           16         EP1         0.0011         Default value = 0.005% SOC           17         EP2         0.0011         Default value = 0.005% SOC           18         EM         0.0014         Default value = 0.005% SOC           19         CN_SOM         13         CN ratio of SOM           20         CN_POM         26         CN ratio of Particulate Organic Matter (POM), will be calculated by model           21         CN_MOM         10         CN ratio of DOM, default	4	CN_MB_mean	8	Mean CN ratio of microbial biomass
7         CN_EP1         3         CN ratio of oxidative enzymes           8         CN_EP2         3         CN ratio of hydrolytic enzymes           9         CN_EM         3         CN ratio of MOM enzymes           10         fQOM         0.01         Fraction of QOM in MOM           11         SOC         21.14         Actually NOT used for modeling           12         POC         5.67         POC1 + POC2           13         MOC         15.47         Mineral-associated Organic Carbon           14         DOC         0.31         Dissolved Organic Carbon           15         MBC         0.53         Default value = 2-5% SOC           16         EP1         0.0011         Default value = 0.005% SOC           17         EP2         0.0011         Default value = 0.005% SOC           18         EM         0.0014         Default value = 0.005% SOC           19         CN_SOM         13         CN ratio of SOM           20         CN_POM         26         CN ratio of Particulate Organic Matter (POM), will be calculated by model           21         CN_MOM         10         CN ratio of DOM, default = CN_MOM           22         CN_DOM         10         CN ratio of microbial bioma	5	CN_MB_min	2	Min CN ratio of microbial biomass
8         CN_EP2         3         CN ratio of hydrolytic enzymes           9         CN_EM         3         CN ratio of MOM enzymes           10         fQOM         0.01         Fraction of QOM in MOM           11         SOC         21.14         Actually NOT used for modeling           12         POC         5.67         POC1 + POC2           13         MOC         15.47         Mineral-associated Organic Carbon           14         DOC         0.31         Dissolved Organic Carbon           15         MBC         0.53         Default value = 2.5% SOC           16         EP1         0.0011         Default value = 0.005% SOC           17         EP2         0.0011         Default value = 0.005% SOC           18         EM         0.0014         Default value = 0.005% SOC           19         CN_SOM         13         CN ratio of SOM           20         CN_POM         26         CN ratio of Particulate Organic Matter (POM), will be calculated by model           21         CN_MOM         10         CN ratio of MOM, CN_MOM < CN_SOM	6	CN_MB_max	14	Max CN ratio of microbial biomass
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10         fQOM         0.01         Fraction of QOM in MOM           11         SOC         21.14         Actually NOT used for modeling           12         POC         5.67         POC1 + POC2           13         MOC         15.47         Mineral-associated Organic Carbon           14         DOC         0.31         Dissolved Organic Carbon           15         MBC         0.53         Default value = 2-5% SOC           16         EP1         0.0011         Default value = 0.005% SOC           17         EP2         0.0011         Default value = 0.005% SOC           18         EM         0.0014         Default value = 0.005% SOC           19         CN_SOM         13         CN ratio of SOM           20         CN_POM         26         CN ratio of Particulate Organic Matter (POM), will be calculated by model           21         CN_MOM         10         CN ratio of MOM, CN_SOM           22         CN_DOM         10         CN ratio of DOM, default = CN_MOM           23         CN_MB         5.5         CN ratio of microbial biomass           24         NH4         0.003           25         NO3         0.01	8	CN_EP2	3	CN ratio of hydrolytic enzymes
11         SOC         21.14         Actually NOT used for modeling           12         POC         5.67         POC1 + POC2           13         MOC         15.47         Mineral-associated Organic Carbon           14         DOC         0.31         Dissolved Organic Carbon           15         MBC         0.53         Default value = 2.5% SOC           16         EP1         0.0011         Default value = 0.005% SOC           17         EP2         0.0011         Default value = 0.005% SOC           18         EM         0.0014         Default value = 0.005% SOC           19         CN_SOM         13         CN ratio of SOM           20         CN_POM         26         CN ratio of Particulate Organic Matter (POM), will be calculated by model           21         CN_MOM         10         CN ratio of MOM, CN_MOM < CN_SOM	9	CN_EM	3	CN ratio of MOM enzymes
12         POC         5.67         POC1 + POC2           13         MOC         15.47         Mineral-associated Organic Carbon           14         DOC         0.31         Dissolved Organic Carbon           15         MBC         0.53         Default value = 2-5% SOC           16         EP1         0.0011         Default value = 0.005% SOC           17         EP2         0.0011         Default value = 0.005% SOC           18         EM         0.0014         Default value = 0.005% SOC           19         CN_SOM         13         CN ratio of SOM           20         CN_POM         26         CN ratio of Particulate Organic Matter (POM), will be calculated by model           21         CN_MOM         10         CN ratio of MOM, CN_SOM           22         CN_DOM         10         CN ratio of DOM, default = CN_MOM           23         CN_MB         5.5         CN ratio of microbial biomass           24         NH4         0.003           25         NO3         0.01	10	fQOM	0.01	Fraction of QOM in MOM
13         MOC         15.47         Mineral-associated Organic Carbon           14         DOC         0.31         Dissolved Organic Carbon           15         MBC         0.53         Default value = 2-5% SOC           16         EP1         0.0011         Default value = 0.005% SOC           17         EP2         0.0011         Default value = 0.005% SOC           18         EM         0.0014         Default value = 0.005% SOC           19         CN_SOM         13         CN ratio of SOM           20         CN_POM         26         CN ratio of Particulate Organic Matter (POM), will be calculated by model           21         CN_MOM         10         CN ratio of MOM, CN_MOM < CN_SOM	11	SOC	21.14	Actually NOT used for modeling
14         DOC         0.31         Dissolved Organic Carbon           15         MBC         0.53         Default value = 2-5% SOC           16         EP1         0.0011         Default value = 0.005% SOC           17         EP2         0.0011         Default value = 0.005% SOC           18         EM         0.0014         Default value = 0.005% SOC           19         CN_SOM         13         CN ratio of SOM           20         CN_POM         26         CN ratio of Particulate Organic Matter (POM), will be calculated by model           21         CN_MOM         10         CN ratio of MOM, CN_SOM           22         CN_DOM         10         CN ratio of DOM, default = CN_MOM           23         CN_MB         5.5         CN ratio of microbial biomass           24         NH4         0.003           25         NO3         0.01           26         rCN_LIC2LAR         2           See Table 19	12	POC	5.67	POC1 + POC2
15         MBC         0.53         Default value = 2-5% SOC           16         EP1         0.0011         Default value = 0.005% SOC           17         EP2         0.0011         Default value = 0.005% SOC           18         EM         0.0014         Default value = 0.005% SOC           19         CN_SOM         13         CN ratio of SOM           20         CN_POM         26         CN ratio of Particulate Organic Matter (POM), will be calculated by model           21         CN_MOM         10         CN ratio of MOM, CN_MOM < CN_SOM	13	MOC	15.47	Mineral-associated Organic Carbon
16         EP1         0.0011         Default value = 0.005% SOC           17         EP2         0.0011         Default value = 0.005% SOC           18         EM         0.0014         Default value = 0.005% SOC           19         CN_SOM         13         CN ratio of SOM           20         CN_POM         26         CN ratio of Particulate Organic Matter (POM), will be calculated by model           21         CN_MOM         10         CN ratio of MOM, CN_SOM           22         CN_DOM         10         CN ratio of DOM, default = CN_MOM           23         CN_MB         5.5         CN ratio of microbial biomass           24         NH4         0.003           25         NO3         0.01           26         CN_LIC2LAR         3           26         CN_LIC2LAR         3           See Table 19	14	DOC	0.31	Dissolved Organic Carbon
17         EP2         0.0011         Default value = 0.005% SOC           18         EM         0.0014         Default value = 0.005% SOC           19         CN_SOM         13         CN ratio of SOM           20         CN_POM         26         CN ratio of Particulate Organic Matter (POM), will be calculated by model           21         CN_MOM         10         CN ratio of MOM, CN_MOM < CN_SOM	15	MBC	0.53	Default value = 2-5% SOC
18 EM         0.0014 Default value = 0.005% SOC           19 CN_SOM         13 CN ratio of SOM           20 CN_POM         26 CN ratio of Particulate Organic Matter (POM), will be calculated by model           21 CN_MOM         10 CN ratio of MOM, CN_MB < CN_MOM < CN_SOM	16	EP1	0.0011	Default value = 0.005% SOC
19         CN_SOM         13         CN ratio of SOM           20         CN_POM         26         CN ratio of Particulate Organic Matter (POM), will be calculated by model           21         CN_MOM         10         CN ratio of MOM, CN_MOM < CN_SOM	17	EP2	0.0011	Default value = 0.005% SOC
20 CN_POM 26 CN ratio of Particulate Organic Matter (POM), will be calculated by model 21 CN_MOM 20 CN_MB < CN_MOM < CN_SOM 21 CN_DOM 22 CN_DOM 23 CN_MB 24 NH4 25 NO3 26 CN_CN_LIC2LAP 26 CN_Tatio of Particulate Organic Matter (POM), will be calculated by model CN ratio of MOM, CN_MB < CN_MOM < CN_SOM CN_Tatio of DOM, default = CN_MOM CN_Tatio of microbial biomass CN_TATIO ORGANICATION CN_TATIO OF Particulate Organic Matter (POM), will be calculated by model CN ratio of MOM, CN_MB < CN_MOM < CN_SOM CN_Tatio of DOM, default = CN_MOM CN_TATIO OF TATION	18	EM	0.0014	Default value = 0.005% SOC
20       CN_POM       26       (POM), will be calculated by model         21       CN_MOM       10       CN ratio of MOM, CN_MOM < CN_SOM	19	CN_SOM	13	CN ratio of SOM
21 CN_MOM   10   CN ratio of MOM,   CN_MB < CN_MOM < CN_SOM   22 CN_DOM   10   CN ratio of DOM,   default = CN_MOM   23 CN_MB   5.5   CN ratio of microbial biomass   24 NH4   0.003   25 NO3   0.01     See Table 19	20	CN POM	26	S
21         CN_MOM         10         CN_MB < CN_MOM < CN_SOM           22         CN_DOM         10         CN ratio of DOM, default = CN_MOM           23         CN_MB         5.5         CN ratio of microbial biomass           24         NH4         0.003           25         NO3         0.01           36         rCN_HC2LAP         3           See Table 19		div_i on		
22       CN_DOM       10       CN ratio of DOM, default = CN_MOM         23       CN_MB       5.5       CN ratio of microbial biomass         24       NH4       0.003         25       NO3       0.01         36       rCN_HC2LAP       3         See Table 19	21	CN_MOM	10	·
23 CN_MB       5.5 CN ratio of microbial biomass         24 NH4       0.003         25 NO3       0.01         36 PCN_HC2LAP       3 See Table 19	22	CN DOM	10	
24 NH4 0.003 25 NO3 0.01 26 PCN LIC2LAP 3 See Table 19	-	_		
25 NO3 0.01  26 rCN LIC2LAP 2 See Table 19	-			2.1.2.330 0
26 rCN LICZLAP 2 See Table 19				
				-

# 8 Input Data

Note: all input data will be converted to hourly rate; see input data format in dir: userio/inp (MEND\_mult: see casedir in inp)

Input Data	Notes/Sample data	Time scale
Litter input	1) Continuous input, e.g., litter fall,  'MEND_namelist.nml'  Type-1 Input  Soil depth (cm) in SOIL_INI.dat  will be used to convert  mg C cm <sup>-2</sup> hour <sup>-1</sup> to  mg C cm <sup>-3</sup> hour <sup>-1</sup>	Monthly, daily, or hourly units:  mg C cm <sup>-2</sup> d <sup>-1</sup> or  mg C cm <sup>-2</sup> month <sup>-1</sup> or  mg C cm <sup>-2</sup> hour <sup>-1</sup>
	2) Other constant input, e.g., only annual amount available,  'MEND_namelist.nml'  Type-2 Input	Convert annual amount to hourly rate
	3) Other constant input during a specific period, e.g., dead roots, 'MEND_namelist.nml'  Type-3 Input	Specify total amount & the period (beginning & ending dates)
Soil temperature	'MEND_namelist.nml' preferred units: °C STP.dat	Hourly, daily, or monthly
Soil moisture	'MEND_namelist.nml' preferred units: % will be converted to soil water potential (MPa) using the retention curve parameters given in soil retention curve See 'fSWC2SWP' in 'MOD_MEND.F90' SWC.dat	Hourly, daily, or monthly
Mineral N input	NH4dep.dat; NO3dep.dat	Monthly
C:N ratios	<ol> <li>Input C:N, see         'MEND_namelist.nml'</li> <li>SOM C:N, see SOIL_INI.dat</li> </ol>	

# 9 Uncertainty Quantification (UQ) by Critical Objective Function Index (COFI): Inputs & Outputs

# (1) MEND

iModel = 5

Category	File Name	Notes
Input File	COFIpar.dat	Copy 'out/OPT_all.out' to 'inp/COFIpar.dat', add
		"OBJ_critical= $J_{cr}$ $J_{min}$ " to 1 <sup>st</sup> line, where $J_{min}$ denote
		the optimial/minimum OBJ; and <i>J<sub>cr</sub></i> denotes the
		critical OBJ for UQ:
		$J_{cr} = J_{opt} \cdot \eta = J_{opt} \cdot \left(1 + \frac{p}{n-p} F_{\alpha, p, n-p}\right)$
		$J_{opt}$ is the optimum (min) OBJ, $n$ is the number of
		OBS, p is the number of PARs, and $F_{\alpha,p,n-p}$ is the
		value of the F-distribution for $\alpha$ , $p$ , and $n-p$ .
Output Files	COFIpar.out	Save parameters that result in $J_{\min} \le fObj \le J_{cr}$
	COFIvar.out	Save variables on those observational time steps
		predicted by the PARs in 'COFIpar.out'

#### (1) MEND\_mult

• enabled/corrected on 10/4/2020

MOD MEND.F90

REAL(8) function fMEND OBJ(xx, sPAR, sINI, sOUT)

```
~Line 363–366:
if (sINI%iModel.eq.3) then
write(format101, *) "(", k, "E15.3)"
write(sINI%iFout_UQvar, format101)dOBS_SIM(:,2)
```

#### ATTENTION:

end if

• MEND.ini

iModel = 1; iprint = 1 !! during optimization iModel = 3 !! for COFI

• input file: inp/COFIpar.dat

### 10 sINI%iScenario

Scenario design: data used to conduct scenario analysis beyond calibration period See 'MEND namelist.nml' siScenario

iScenario	Notes
1	Repeatedly use 1-yr mean hourly data derived from observed data
2	Repeatedly use multiple-year hourly (observed) data

### 11 Scenario Design

Parameters controlling the changes in Temperature, Water content & Litterfall Input; specifically, for the subtropical Dinghushan case study, may not be applied to other case studies

See 'MEND namelist.nml' Scenario design

Parameter	Note				
STP_delta	Gradually annual change in Temperature, °C/a				
SWC logis(3)	$\theta(t)$ p				
	$\frac{\theta(t)}{\theta(t_0)} = \frac{p}{1 - (1 - p) \cdot e^{-r(t - t_0)}}$				
	Soil Water Content (SWC, $\theta$ ) parameters in logistic equation; SWC				
	DECREASES with time				
	(1) $p \in (0,1)$ : lower bound for the ratio of $\frac{\theta(t \to \infty)}{\theta(t_0)}$				
	(2) $r > 0$ : steepness; $r = 0$ means no changes				
	(3) t <sub>0</sub> : reference year				
SIN_logis(4)	L(t) 1				
	$\overline{L_{max}} = \frac{1 + \exp[\beta_0 - \beta_1 \cdot (t - t_0)]}{1 + \exp[\beta_0 - \beta_1 \cdot (t - t_0)]}$				
	Litterfall input parameters in logistic equation; Litterfall				
	INCREASES with time				
	$(1) \beta_0 > 0$ : intercept				
	(2) $\beta_1 > 0$ : steepness; $\beta_1 = 0$ means no changes				
	(3) t <sub>0</sub> : reference year				
	(4) fDOC_delta (NOT included in the logistic equation): gradually				
	annual change of DOC fraction in SOC input				

### 12 sINI%iKenetics

### **Decomposition Kinetics for POC/MOC**

See 'MEND\_namelist.nml' siKinetics (MEND\_mult: Line22-23)

iKinetics	Mic-Enz	Kinetics	Equation	Sample
				MEND_namelist.nml
0	Both	<ul><li> Michaelis-Menten</li><li> MOM decomposition</li><li> No MOM-QOM interaction</li></ul>	$F_{dec} = \frac{V_M \cdot E \cdot S}{K_M + S}$	MEND_Enz-Mic.ini
1	No Enz	First Order	$F_{dec} = k_M \cdot S$	MEND_No-Enz.ini
2	Both	Second Order	$F_{dec} = k_M \cdot E \cdot S$	
10	Mic-Enz	<ul><li> Michaelis-Menten</li><li> QOM decomposition</li><li> MOM-QOM interaction</li></ul>	$F_{dec} = \frac{V_M \cdot E \cdot S}{K_M + S}$	
11	None	First Order	$F_{dec} = k_M \cdot S$	MEND_No-Enz-

#### 13 sINI%iHR

Calculation Method for Growth & Maintenance Respiration from Active Microbes

Culculation	i vicinda idi Gidwen & iviaintenance Respiration	THOM TICHTE WHICH OBES
iHR	Method	Notes
0	$HR_g = \max \left[ D, \left( \frac{1}{Y_G} - 1 \right) \frac{V_g \cdot BA \cdot D}{K_D + D} \right]$	$HR_g$ or $HR_m$ is constrained by DOC
	$HR_{m} = \max \left[ D, \left( \frac{1}{Y_{G}} - 1 \right) \frac{V_{m} \cdot BA \cdot D}{K_{D} + D} \right]$	
1	$HR_{gm} = \max \left[ D, \frac{1}{Y_G} \times \frac{\left(V_g + V_m\right) \cdot BA \cdot D}{K_D + D} \right] \times \left(1 - Y_g\right)$	Total microbial uptake is constrained by DOC, $HR_g$ or $HR_m$ is a
	$HR_g = HR_{gm} \times \frac{V_g}{V_g + V_m}, HR_m = HR_{gm} \times \frac{V_m}{V_g + V_m}$	fraction of total uptake

# 14 sINI%iTmp\_Func

**Temperature Response Function** 

p	11050011001	
iTmp_Func	fT	Notes
0	fTArh: Arrhenius Equation	See function in MOD_MEND
1	fTQ10: Q10 method	See function in MOD MEND

# 15 sINI%iError

Error code during model runs

sINI%iError	Description		
< 0	Mass balance error		
	function fMEND_OBJ(xx, sPAR, sINI, sOUT):		
	slNI%rOBJ = dabs(const_FillValue)		
	fMEND_OBJ = dabs(const_FillValue)		
	SUBROUTINE sSOBOL_VAR_OBJ():		
	if(itype.eq.0) then		
	dSIM_t0 = dabs(const_FillValue)		
	else		
	Sobol_obj = dabs(const_FillValue)		
	end if		
-11	Error in 1 <sup>st</sup> carbon balance check		
-12	Error in 1 <sup>st</sup> nitrogen balance check		
-21	Error in 2 <sup>nd</sup> carbon balance check		
-22	Error in 2 <sup>nd</sup> nitrogen balance check		

# 16 MEND State Variables (C & N Pools)

ID	Soil C and/or N pool	Pool Name	Variable name
1	Particulate organic matter (POM) decomposed by oxidative enzymes	POMo	C pool: <i>PO</i> ; N pool: <i>PON</i>
2	POM decomposed by hydrolytic enzymes	$POM_{H}$	PH; PHN
3	Mineral-associated organic matter	MOM	M;MN
4	Dissolved organic matter	DOM	D;DN
5	Active MOM interacting with DOM	QOM	Q;QN
6	Active microbial biomass	$MB_A$	BA; BAN
7	Dormant microbial biomass	$MB_D$	BD; BDN
8	Oxidative enzymes decomposing POM <sub>0</sub>	EPo	EPO; EPON
9	Hydrolytic enzymes decomposing $POM_H$	$EP_{H}$	EPH; EPHN
10	Enzymes decomposing MOM	EM	EM; EMN
11	Ammonium oxidase	ENH4	ENH4; ENH4N
12	Nitrate reductase	ENO3	ENO3; ENO3N
13	Nitrite reductase	ENO2	ENO2; ENO2N
14	Nitric oxide reductase	ENO	ENO; ENON
15	Nitrous oxide reductase	EN2O	EN2O; EN2ON
16	Nitrogenase	EN2	EN2; EN2N
17	Adsorbed ammonium	NH <sub>4</sub> <sup>+</sup> Adsorb	NH4ads
18	Ammonium	$\mathrm{NH_4}^+$	NH4
19	Nitrate	$NO_3^-$	NO3
20	Nitrite	$NO_2^-$	NO2
21	Nitric oxide	NO	NO
22	Nitrous oxide	$N_2O$	N2O
23	Dinitrogen	$N_2$	N2

### **17 MEND Parameters**

#### **17.1 MEND Parameters**

('MEND namelist.nml' INITIAL PARAMETERS) (MEND mult: Line 28-57)

		inenst.nim intital parameters) (NE			
ID	Parameter	Description	Range	Units	Eq#
1	$LF_0$	Initial fraction of $PO$ , $LF_0 = PO/(PO+PH)$	(0.1, 1.0)	_	
2	$r_0$	Initial active fraction of microbes, $r_0 = BA/(BA+BD)$	(0.01, 1)	_	
3	$fR_a$	Scaling factor for autotrophic respiration $(R_a)$	(0.1, 0.4)	_	11
4	fINP	Scaling factor for litter input rate	(0.1, 0.9)	_	1
5	Vd	Maximum specific decomposition rate $Vd_{PO} = Vd_{PH} = Vd_{M} = Vd$	(0.1, 100)	mg C mg <sup>-1</sup> C h <sup>-1</sup>	24–26
6	$K_{PO}$	Half-saturation constant (HSC) for PO decomposition	(40,100)	mg C cm <sup>-3</sup> soil	24
7	$fK_M$	$K_M = K_{PO} \times fK_M, K_{PH} = K_{PO}/fK_M$	(2, 20)	_	25–26
		$K_{PH}$ and $K_M$ are HSC for $PH$ and $M$ , respectively			
8	$Q_{ m max}$	Maximum sorption capacity	(0.5, 5.0)	mg C cm <sup>-3</sup> soil	27
9	$K_{ba}$	Binding affinity for DOM, sorption rate $k_{ads} = k_{des} \times K_{ba}$	(1, 16)	(mg C cm <sup>-3</sup> soil) <sup>-1</sup>	27
10	$k_{des}$	Desorption rate for DOM	(1e-4, 0.01)	mg C cm <sup>-3</sup> soil h <sup>-1</sup>	28
11*	$r_E$	Enzyme turnover rate	(1e-4, 0.01)	mg C mg <sup>-1</sup> C h <sup>-1</sup>	39, 41
12*	$p_{EP}$	$[V_m \times p_{EP}]$ is the production rate of $EP$ ( $EPO + EPH$ ), $V_m$ is the specific maintenance rate for $BA$	(1e-4, 0.05)	_	38
13*	fр <sub>ЕМ</sub>	$fp_{EM} = p_{EM}/p_{EP}$ , $[V_{mt} \times p_{EM}]$ is the production rate of EM	(0.1, 5.0)	_	38
14*	$f_D$	Fraction of decomposed PO and PH allocated to D	(0.05, 1)	_	3
15*	$g_D$	Fraction of dead BA allocated to D	(0.01, 1)	_	1
16	$g_{PO}$	$(1-g_D) \cdot g_{PO}$ is the fraction of dead BA entering PO	(0.05, 0.2)	_	2
17*	$V_g$	Maximum specific uptake rate of D for growth	(1e-3, 0.1)	mg C mg <sup>-1</sup> C h <sup>-1</sup>	29
18*	α	$= V_m/(V_g + V_m)$ , $V_m$ is max specific maintenance rate	(0.01, 0.5)	_	29
19*	$K_D$	HSC for microbial uptake of D	(1e-4, 0.5)	mg C cm <sup>-3</sup> soil	29
20*	$Y_g(T_{\rm ref})$	Intrinsic C use efficiency at reference temperature ( $T_{ref}$ )	(0.1, 0.6)	_	29
21	$k_{Yg}$	Slope for $Y_g$ dependence of temperature	(1e-3, 0.016)	1/°C	29
22*	$Q_{10}$	Q <sub>10</sub> for temperature response function	(1.2, 2.5)	_	
23	γ	Max microbial mortality rate = $V_m \times \gamma$	(0.1, 20)	_	32
24	β	Ratio of dormant maintenance rate to $V_m$	(5e-4, 0.05)	_	36
25	ΨA2D	Soil water potential (SWP) threshold for microbial	(-0.6, -0.2)	MPa	
		dormancy; both $\psi_{A2D}$ & $\psi_{D2A} < 0$			
26	τ	$\psi_{D2A} = \psi_{A2D} \times \tau$ , $\psi_{D2A}$ is the SWP threshold for microbial resuscitation	(0.1, 0.9)	_	
27	ω	Exponential in SWP function for microbial dormancy or resuscitation	(1, 6)	_	
28	$VN_{im,BA}$	Max specific microbial N immobilization rate	(1e-4, 0.1)	mg N mg <sup>-1</sup> C h <sup>-1</sup>	48, 49
-	· tiit,DA	$VN_{im,NH4} = VN_{im,BA} \cdot NH4/(NH4 + NO3)$		8 8	-, -
		$VN_{im,NO3} = VN_{im,BA} \cdot NO3/(NH4 + NO3)$			
29	$KSN_{BA1}$	HSC for microbial immobilization of NH <sub>4</sub> <sup>+</sup>	(1e-4, 0.01)	mg N cm <sup>-3</sup> soil	48–50
30	$KSN_{BA2}$	HSC for microbial immobilization of NO <sub>3</sub> <sup>-</sup>	(1e-4, 0.01)	mg N cm <sup>-3</sup> soil	48–50
31*	$VN_{nit}$	Max specific nitrification rate (VN <sub>1</sub> )	(0.1, 1000)	mg N mg <sup>-1</sup> C h <sup>-1</sup>	42
32*	$VN_{denit}$	Max specific denitrification rate, $VN_j = VN_{denit}$ , $j=2-5$	(1e-5, 0.1)	mg N mg <sup>-1</sup> C h <sup>-1</sup>	44
33*	$VN_{fix}$	Max specific N fixation rate $(VN_6)$	(1e-4, 0.1)	mg N mg <sup>-1</sup> C h <sup>-1</sup>	45
34	KSN <sub>1</sub>	HSC for nitrification	(1e-4, 0.1)	mg N cm <sup>-3</sup> soil	42
35	KSN <sub>2</sub>	HSC for denitrification of NO <sub>3</sub> <sup>-</sup> and NO <sub>2</sub> <sup>-</sup> (KSN <sub>3</sub> )	(1e-4, 0.1)	mg N cm <sup>-3</sup> soil	44
36	KSN <sub>4</sub>	HSC for denitrification of NO and N <sub>2</sub> O (KSN <sub>5</sub> )	(1e-4, 0.1)	mg N cm <sup>-3</sup> soil	44
37	KSN <sub>6</sub>	HSC for N fixation	(1e-4, 0.1)	mg N cm <sup>-3</sup> soil	45
38*	$VN_{VG}$	Max plant N uptake rate; $VN_{VG,NH4} = VN_{VG,NO3} = VN_{VG}$	(1e-6, 1e-3)	mg N cm <sup>-3</sup> h <sup>-1</sup>	51, 52
39	KSN <sub>VG1</sub>	HSC for plant uptake of NH <sub>4</sub> <sup>+</sup>	(1e-4, 0.01)	mg N cm <sup>-3</sup> soil	51
40	KSN <sub>VG2</sub>	HSC for plant uptake of NO <sub>3</sub> <sup>-</sup>	(1e-4, 0.01)	mg N cm <sup>-3</sup> soil	52
41	ωνG	Exponential for calculating $rGPP$ as a function of GPP	(0.01, 1)		52
42*	NH4 <sub>max</sub>	Maximum sorption capacity for NH <sub>4</sub> <sup>+</sup>	(1e-5, 0.01)	mg N cm <sup>-3</sup> soil	53
43	K <sub>ba,NH4</sub>	Binding affinity for NH <sub>4</sub> <sup>+</sup>	(1, 1e4)	(mg N cm <sup>-3</sup> soil) <sup>-1</sup>	53
44	rleach	Scaling factor for NO <sub>3</sub> <sup>-</sup> and NO <sub>2</sub> <sup>-</sup> leaching	(0.01, 1)	_	54
				•	

Note1: 'MEND\_namelist.nml' Cali\_Calibrate (MEND\_mult: Line 31-57), Column 'Calibrate': '1' means the parameter will be calibrated, '0' mean no-calibration.

Note2: 'MEND\_namelist.nml' Pintital (MEND\_mult: Line 61), initial parameter values, will override the values in Column 'Initial' at Line 78-104.

# 17.2 MEND Parameters that may be pre-determined

Parameter	Regression Equation	$\mathbb{R}^2$	p-value
$K_{\rm Pl}$ (mg C g <sup>-1</sup> soil): half-saturation	$K_{\rm P1} = 41.58 \times {\rm DOC} + 44.95$	0.97	0.01
constant for POC decomposition	$K_{\rm P1} = 0.62 \times (\% {\rm Sand}) + 45.56$	0.82	0.10
1	$K_{\rm Pl} = 1.07 \times {\rm POC} + 50.13$	0.64	0.20
$K_{\rm M}$ (mg C g <sup>-1</sup> soil): half-saturation	$K_{\rm M} = 455.11 \times {\rm DOC} + 400.42$	0.98	0.01
constant for MOC decomposition	$K_{\rm M} = 4.13 \times {\rm MOC} + 425.75$	0.73	0.15
$K_{\rm D}$ (mg C g <sup>-1</sup> soil): half-saturation	$K_{\rm D} = 0.33 \times {\rm MBC} + 0.21$	0.63	0.21
constant for microbial uptake of			
DOC			
$Q_{\text{max}}$ (mg C g <sup>-1</sup> soil): maximum	$Q_{\text{max}} = 0.012 \times \text{MOC} + 3.43$	0.98	0.01
sorption capacity	$Q_{\text{max}} = 0.011 \times \text{SOC} + 3.39$	0.91	0.04
$K_{\text{ba}}([\text{mg C g}^{-1} \text{ soil}]^{-1})$ : binding affinity	$K_{\text{ba}} = -0.082 \times (\% \text{Sand}) + 11.23$	0.84	0.08
$K_{\text{des}}$ (mg C g <sup>-1</sup> soil h <sup>-1</sup> ): desorption	$K_{\text{des}} = -0.0059 \times \text{DOC} + 0.0059$	0.95	0.02
rate	$R_{\text{des}} = 0.0039 \times \text{DOC} + 0.0039$	0.93	0.02
Tate			
$V_{death} = V_{mt} \times gamma$ : microbial	[Hansen, 1990, DAISY]		
death rate	$\begin{bmatrix} 0.001 - 0.01  d^{-1}, \end{bmatrix}$		
death rate	$4 \times 10^{-5} - 4 \times 10^{-4}  h^{-1}$		
	+^10		
Vd	We can also may shiv determine Vd		
	We can also roughly determine Vd		
	by a few model trials.		
	(1) Try to fix the value of Vd		
	and calibrate other		
	parameters to fit Rh		
	observations. If the		
	goodness-of-fit of Rh is		
	good (i.e., $R^2 > 0.4$ ) and the		
	PBIAS of Rh is good (i.e.,		
	$ PBIAS  = \left  \frac{\bar{Y}_{Sim} - \bar{Y}_{obs}}{\bar{Y}_{obs}} \right $		
	<10%), we would		
	conclude the simulated		
	enzyme activities ( $EA =$		
	$Vd \cdot EC$ ) or		
	decomposition rates		
	$(FR = (Vd \cdot EC)/(K +$		
	S = EA/(K+S) are		
	good.		
	(2) Check the simulated total		
	SOM-degrading enzyme		
	concentrations (EC:		
	variable name ENZD C in		
	xxx VAR xxx.out). Total		
	AAA_ V AIR_AAA.UUI J. TUIdi	<u> </u>	

calculate the fRa directly use $lm(Ra \sim GPP+0)$ , i.e., $fR_a = \frac{R_a}{GPP}$ fINP: scaling factor for litter input rate  we can roughly determine fINP, particularly for the control treatment, which is often assumed to be at quasi steady state, i.e., average input rate to $SOC = mean$ heterotrophic respiration rate. Thus $fINP = \frac{R_h}{GPP}$ Regarding the fINP for other treatments, we may use the same fINP as the control treatment or adjust it carefully.  Basically, fINP should NOT be calibrated simultaneously with other parameters. If we need to calibrate fINP, we have to set a narrow parameter range for fINP based on our knowledge.  fD, gD  I prefer NOT to calibrate fD and gD if we could fix them (e.g., fD=0.75, gD=0.50) to achieve good model performance.  However, we may need to calibrate fD and gD to obtain good results.  rE, pEP, fpEM  usually I will calibrate these three parameters: FE, pEP, fpEM.  Yg and kYg  avoid calibrating Yg(Tref) and kYg simultaneously. We need to fix one of them and calibrate the other one.		<u></u>	
		enzyme concentrations	
(3) If EC values are too high, increase the value of $Vd$ , do model calibration again, we would see a decrease in EC.  Re. Scaling factor for autotrophic respiration (Re.)  As $Ra = fRa * GPP$ , we could calculate the fRa directly use $\lim(Ra \sim GPP + 0)$ , i.e., $fR_a = \frac{\overline{R}_a}{\overline{GPP}}$ we can roughly determine flNP, particularly for the control treatment, which is often assumed to be at quasi steady state, i.e., average input rate to $SOC = \text{mean heterotrophic}$ respiration rate. Thus $fINP = \frac{\overline{R}_k}{\overline{GPP}}$ Regarding the flNP for other treatments, we may use the same flNP as the control treatment or adjust it carefully. Basically, flNP should NOT be calibrated simultaneously with other parameters. If we need to calibrate flNP, we have to set a narrow parameter range for flNP based on our knowledge.  As $Ra = fRa * GPP$ , we could calculate the flNP and the flNP, we can be a control treatment of the flNP as the control treatment or adjust it carefully. Basically, flNP should NOT be calibrate flNP, we have to set a narrow parameter range for flNP based on our knowledge.  As $Ra = fRa * GPP$ , we could calculate flD and gD if we could fix them (e.g., flD=0.75, gD=0.50) to achieve good model performance. However, we may need to calibrate flD and gD to obtain good results.  The fixed provides a flow of the flNP and kYg simultaneously. We need to fix one of them and calibrate the other one.		should have the magnitude	
increase the value of $\nabla d$ , do model calibration again, we would see a decrease in EC. $R_a$ : Scaling factor for autotrophic respiration ( $R_a$ )  As $Ra = FRa * GPP$ , we could calculate the fRa directly use $Im(Ra \sim GPP + 0)$ , i.e., $Im(Ra \sim GPP + 0$		of 0.1–1% of MBC.	
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$ \begin{array}{c} \text{model calibration again, we} \\ \text{would see a decrease in EC.} \\ \hline \textit{Re. Scaling factor for autotrophic respiration} \\ \textit{(R.)} \\ \hline \textit{Rs. Ra} = \textit{FRa} * \textit{GPP}, \text{ we could} \\ \text{calculate the fRa directly use} \\ \text{Im}(\text{Ra} \sim \text{GPP+0}), \text{i.e.,} \\ \hline \textit{fRa} = \frac{\overline{R_a}}{\overline{GPp}} \\ \hline \textit{fINP: scaling factor for litter input} \\ \text{rate} \\ \hline \\ \textit{mex. GPP+0}, \text{i.e.,} \\ \hline \textit{fRa} = \frac{\overline{R_a}}{\overline{GPp}} \\ \hline \textit{fint} \\ \text{mex. GPP+0}, \text{i.e.,} \\ \hline \textit{fint} \\ \text{particularly for the control} \\ \text{treatment, which is often} \\ \text{assumed to be at quasi steady} \\ \text{state, i.e., average input rate to} \\ \text{SOC} = \text{mean heterotrophic} \\ \text{respiration rate. Thus} \\ \hline \textit{fINP} = \frac{\overline{R_h}}{\overline{GPp}} \\ \hline \text{Regarding the fINP for other} \\ \text{treatments, we may use the same} \\ \text{fINP as the control treatment or} \\ \text{adjust it carefully.} \\ \hline \text{Basically, fINP should NOT be} \\ \text{calibrated simultaneously with} \\ \text{other parameters. If we need to} \\ \text{calibrate fINP, we have to set a} \\ \text{narrow parameter range for fINP} \\ \text{based on our knowledge.} \\ \hline \textit{fD, gD} \\ \hline \text{I prefer NOT to calibrate fD and} \\ \text{gD if we could fix them (e.g.,} \\ \text{fD=0.75, gD=0.50) to achieve} \\ \text{good model performance.} \\ \hline \text{However, we may need to calibrate} \\ \hline \text{fD and gD to obtain good results.} \\ \hline \textit{vertical parameters: fE, pEP, fpEM.} \\ \hline \textit{avoid calibrating Yg(Tref) and kYg} \\ \text{simultaneously. We need to fix one} \\ \text{of them and calibrate the other one.} \\ \hline \end{tabular}$			
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$R_{o} \cdot \text{Scaling factor for autotrophic respiration} \\ (R_{o}) \\ R_{o} \cdot \text{Scaling factor for autotrophic respiration} \\ (R_{o}) \\ R_{o} \cdot \text{Scaling factor for litter input} \\ \text{rate} \\ R_{a} = \frac{R_{a}}{\overline{GPP}} \\ R_{a} = \frac{R_{a}}{\overline{GPP}} \\ R_{o} \cdot \text{TiNP: scaling factor for litter input} \\ \text{rate} \\ R_{o} \cdot \text{Timput} \\ \text{rate} \\ $		_	
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fINP: scaling factor for litter input rate		$f_{P} = \frac{R_a}{R_a}$	
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If we want to fix kYg, try default		If we want to fix kYg, try default	

	value 0.01 first. If we want to fix Yg(Tref), try default value 0.3 first.	
Q10	check namelist file: siTmp_Func = 0: Arrhenius Equation, siTmp_Func = 1: Q10 Equation is used.	
γ	try to fix it by default value $\gamma = 0.1$ first.	

### Notes:

SOC: soil organic carbon; POC: particulate organic carbon; MOC: mineral-associated organic carbon; DOC: dissolved organic carbon; MBC: microbial organic carbon. Units:  ${\rm mg}\ {\rm C}\ {\rm g}^{-1}$  soil.

%Sand: percentage of sand in soil.

# 18 Response Variables for Model Calibration/Optimization

#### 18.1 Calibration Variables in MEND\_namelist.nml

Note: 'MEND\_namelist.nml' CALIBRATION DATA; see data format in dir: userio/inp MEND mult: Line 45-57 in '\*.ini' in 'userio/inp/casedir'

Column	Column Name	Notes		
1	VARid	Variable ID		
2	VAR	Variable Name		
3	Units	Units		
4	Calibrate	'1'-data available for calibration, '0'-unavailable		
5	tstep	Time-step: 0-hourly, 1-daily, 2-monthly,		
		5-mean value of last2years,		
		6-mean_value_of_last_80%data.		
		Usually the observations are regarded as data		
		measured during an hourly-scale, the simulation		
		results will be averaged during 1 day (24-h) to		
		compare with the observations		
6	Obs_file	File with observation data if available		
7	Obs_file_column	Column id for the data, currently not used		
8	OBJ	Type of objective function (minimization):		
		<ul> <li>NSEC: Nash-Sutcliffe Efficiency Coefficient (Coefficient</li> </ul>		
		of Determination), see 'f1NSE' in 'MOD_USRFS.F90'		
		MARE: Mean Absolute Relative Error, see 'fMARE' in		
		'MOD_USRFS.F90'		
		CORR: correlation coefficient, see 'f1CORR'		
		CORI: CORR with log10-tranformed data, see 'f1CORR'		
		MART: MARE with tolerance, see 'fMARE_tolerance'		
		NSEn: NSEC for normalized data, see 'f1NSE_norm'		
		MARn: MARE for normalized data, see 'fMARE_norm'		
0	ODI Waialet	AVGr:  ratio – AVGsim/AVGobs , see 'f1RAVG_ratio'      Weighting fortunation and ODI will be a green in a discount.		
9	OBJ_Weight	Weighting factor for each OBJ, will be normalized in		
		the code		

Microbial biomass carbon (MBC) should be included in model calibration. If we don't have MBC measurements. Assume MBC = 2-5% SOC, which is used for both initial MBC and reference value for simulated MBC. As for model calibration:

Cali\_tstep: 5 (mean MBC of last 2 years) or 6 (mean MBC of last 80% data)

Cali OBJ: 'MARt'

Cali\_OBJ\_Tolerance: 0.1 or 0.05 (mean MBC within 10% or 5% of the reference MBC)

#### 18.2 Add a new Calibration Variable

(1)	MEND_namelist.nml (	see Section 1	.6): add a new	variable, e.g.,	No.41 for rMBA	(active
	fraction of microbes)					
(	Cali varid(41:44)	= 41,	42,	43,	44	

Cali varid(41:44) = 41,42, 43,

```
Cali VAR(41:44)
                         = 'rMBA',
                                           'TBD1',
                                                          'TBD2',
                                                                        'TBD3'
                                                           'mgN-cm3-h',
  Cali Units(41:44)
                        = 'NONE',
                                          'mgN-cm3-h',
                                                                            'mgN-
cm3-h'
  Cali Calibrate(41:44)
                          = 0,
                                        0,
                                                     0,
                                                                 0
  Cali tstep(41:44)
                                                   2,
                                                               2
                        = 2,
                                       2,
                                           'EA NO2.obs',
  Cali obs file(41:44)
                         = 'rMBA.obs',
                                                             'EA NO.obs',
'EA N2O.obs'
  Cali obs file column(41:44) = 2,
                                            2,
                                                                     2
  Cali OBJ(41:44)
                         = 'MARE',
                                           'MARE',
                                                           'MARE',
'MARE'
  Cali OBJ Weight(41:44)
                                                                    2
                                            2,
                                                        2,
  Cali OBJ Tolerance(41:44) = 0,
                                            0,
                                                         0,
                                                                     0
```

(2) MOD\_MEND\_TYPE.f90

INTEGER, PARAMETER:: const nVARO = 44 !# of variables for calibration

(3) MOD\_MEND.F90

SUBROUTINE **sOUT\_OPT\_h**(nVAR,nHour,iHour,dSIM,sPAR,sOUT,VARopt\_int)

case (41) !!Active fraction of microbes

dSIM(iHour,j) = sOUT%CPOOL%MBA/max(sOUT%CPOOL%MB,1d-6)

or

#### SUBROUTINE

sOUT\_OPT\_VAR\_Extract(nVAR,nHour,iHour,dSIM,sPAR,sOUT,VARopt\_int)

(4) Prepare observed data file for rMBA: rMBA.obs

# 19 Parameter Optimization Algorithm

Note: 'MEND namelist.nml' SCE parameters, (MEND mult: Line 16-20)

only 3 parameter 'nrun', 'iniflg' and 'iprint' are required to edit

Parameter	Notes		
nrun	# of runs for optimization, these opt-runs are independent of each other.		
	There is a different random seed for each opt-run.		
	nrun <= 200		
iniflg	flag on whether to include the initial point (Line 108) in population = 0, not included		
	= 1, included		
iprint	Flag on whether to print all points generated during optimization		
	= 0, no printing		
	=1, print to *_OPT_all.out for COFI UQ analysis		

# 20 sINI%VARopt\_int

#### sINI%VARopt int(sINI%nVARopt, 3)

Column1: i	Column2: n	Column3: t: tstep
Index of calibrated	# of observations	0(hourly)
output VARiable		1(daily)
		2(monthly)
		3(seasonal): to_do
		4(yearly)
		5(mean, last 2 years)
		6(mean, last 80% period)

#### 21 CN Ratios in 3 Litter Pools

3 litter pools: Lignin, Cellulose, and Labile

Assume their fractions in Carbon mass = f Lig, f Cel, f Lab

$$f_Lig + f_Cel + f_Lab = 1$$

Assume CN Cel = 500

Assume CN Lig/CN Lab = rCN

Assume overall CN in litter pool = CN

N mass balance:
$$N = \frac{C}{CN} = \frac{C \cdot f_{Lig}}{CN_{Lab} \cdot rCN} + \frac{C \cdot f_{Cel}}{CN_{Cel}} + \frac{C \cdot f_{Lab}}{CN_{Lab}}$$
thus

$$CN_{Lab} = \frac{\left[\frac{f_{Lig}}{rCN} + f_{Lab}\right]}{\left[\frac{1}{CN} - \frac{f_{Cel}}{CN_{Cel}}\right]}$$

$$CN_{Lig} = CN_{Lab} \times rCN$$

### 22 MEND (non-MPI) Configuration/Compiling on OSX & Cygwin/Unix

#### 22.1 Dependency of source code

```
Edit 'nbproject/Configurations.xml' to define dependency of source code for compilation,
   e.g.,
MOD MEND depends on 2 modules (MOD MEND TYPE & MOD USRFS);
MOD OPT depends on 3 modules (MOD OPT TYPE, MOD MEND, &
MOD USRFS).
See './Configurations.xml' for example:
   <item path="src/MEND IN.F90" ex="false" tool="2" flavor2="0">
    <fortranCompilerTool>
     <additionalDep>${OBJECTDIR}/src/MOD STRING.o ${OBJECTDIR}/src/MOD USRFS.o
${OBJECTDIR}/src/MOD MEND TYPE.o ${OBJECTDIR}/src/MOD MEND.o
${OBJECTDIR}/src/MOD OPT TYPE.o</additionalDep>
    </fortranCompilerTool>
   </item>
   <item path="src/MEND main.F90" ex="false" tool="2" flavor2="0">
    <fortranCompilerTool>
     <additionalDep>${OBJECTDIR}/src/MOD USRFS.o ${OBJECTDIR}/src/MOD MEND TYPE.o
${OBJECTDIR}/src/MOD_MEND.o ${OBJECTDIR}/src/MOD_OPT_TYPE.o
${OBJECTDIR}/src/MOD_OPT.o ${OBJECTDIR}/src/MOD_MCMC.o</additionalDep>
    </fortranCompilerTool>
   </item>
   <item path="src/MOD MCMC.F90" ex="false" tool="2" flavor2="0">
    <fortranCompilerTool>
     <additionalDep>${OBJECTDIR}/src/MOD_USRFS.o ${OBJECTDIR}/src/MOD_MEND.o
${OBJECTDIR}/src/MOD OPT TYPE.o</additionalDep>
    </fortranCompilerTool>
   </item>
   <item path="src/MOD_MEND.F90" ex="false" tool="2" flavor2="0">
    <fortranCompilerTool>
     <additionalDep>${OBJECTDIR}/src/MOD_USRFS.o
${OBJECTDIR}/src/MOD MEND TYPE.o</additionalDep>
    </fortranCompilerTool>
   </item>
   <item path="src/MOD MEND TYPE.F90" ex="false" tool="2" flavor2="0">
   </item>
   <item path="src/MOD OPT.F90" ex="false" tool="2" flavor2="0">
    <fortranCompilerTool>
     <additionalDep>${OBJECTDIR}/src/MOD USRFS.o ${OBJECTDIR}/src/MOD MEND.o
${OBJECTDIR}/src/MOD OPT TYPE.o</additionalDep>
    </fortranCompilerTool>
            <item path="src/MOD OPT TYPE.F90" ex="false" tool="2" flavor2="0">
   </item>
   </item>
   <item path="src/MOD STRING.F90" ex="false" tool="2" flavor2="0">
   <item path="src/MOD_USRFS.F90" ex="false" tool="2" flavor2="0">
   </item>
```

#### 22.2 gcc/gfortran Installation

(1) OSX

#### https://github.com/fxcoudert/gfortran-for-macOS/releases

(2) Windows (install gcc/gfortran/make) Cygwin, <a href="https://www.cygwin.com/">https://www.cygwin.com/</a> Install package Devel

#### 22.3 gdb Installation on Mac OS Mojave

- (1) sudo port install gdb@8.0.1
  - Note: version after gdb8.0.1 does NOT compatible with High Sierra
- (2) In -sf /opt/local/bin/ggdb /usr/local/bin/gdb
- (3) create gdb certificate (e.g., gdbcert)
  - 1) Open Applications/Utilities/Keychain Access
  - 2) In menu, open Keychain Access > Certificate Assistant > Create a Certificate
  - Give it a name (e.g. gdbcert)
     Identity type: Self Signed Root
     Certificate type: Code Signing
     Check: Let Me Override Defaults
  - 4) Continue until "Specify a Location For"
  - 5) Set Keychain location to System. If this yields the following error: Certificate Error: Unknown Error =-2,147,414,007 Set Location to Login, Unlock System by click on the lock at the top left corner and drag and drop the certificate gdbcert to the System Keychain.
  - 6) Create certificate and close Certificate Assistant.
  - 7) Find the certificate in System keychain.
  - 8) Double click certificate.
  - 9) Expand Trust, set Code signing to Always Trust
- (4) Restart taskgated in terminal: sudo killall taskgated

Another method:

ps aux | grep taskgated

in the terminal to find the PID number of taskgated, and then killed it with sudo kill -9 <insert here your PID number>

#### (5) (Option) Enable root account:

- Open System Preferences
- Go to User & Groups > Unlock
- Login Options > "Join" (next to Network Account Server)
- Click "Open Directory Utility"
- Go up to Edit > Enable Root User

#### (6) Codesign gdb certificate:

Create a file gdb.xml:

```
</dict>
</plist>
```

sudo codesign --entitlements gdb.xml -fs gdb2 /usr/local/gdb-8.0.1/bin/gdb

if this command doesn't print anything, it means the binary is correctly signed.

### 5/3/2019: gdb works now@

- (7) (Option) Disable root account (see #10)
- (8) gdb does NOT work in NetBeans. However, it will work if we run NetBeans with sudo: sudo '/Applications/NetBeans/NetBeans 8.2.app/Contents/Resources/NetBeans/bin/netbeans'
- (9) gdb works in command line: sudo gdb ./mend

**Problem**: gdb stop at a line without breakpoint in MEND\_main.F90

**Solution**: in SourceTree, commit the code changes

#### 22.4 NetBeans Configuration on OSX & Windows

- (1) Install NetBeans with JDK
- NetBeans 11.0 (April 30, 2019) (Not Working sometimes) <a href="https://github.com/carljmosca/netbeans-macos-bundle">https://github.com/carljmosca/netbeans-macos-bundle</a>./install.sh
- NetBeans 8.2 with JDK 8u111 (April 30, 2019) (Working for both OSX and Windows) <a href="https://www.oracle.com/technetwork/java/javase/downloads/jdk-netbeans-jsp-3413139-esa.html">https://www.oracle.com/technetwork/java/javase/downloads/jdk-netbeans-jsp-3413139-esa.html</a>

Accept License Agreement Select file to download

➤ NetBeans 12.1 (Nov 6, 2020 Working with jdk-{Version <= 13})

Failed to install any plugin with jkd-15, e.g., c/c++

• Check jdk versions:

ls /Library/Java/JavaVirtualMachines

- copy and paste pack200, unpack200 and unpack.dil from bin of jdk-{Version <= 13} into the bin of jdk-{Version > 13}.
- C/C++ plugin for NetBeans 9.0
  - o **Tools > Plugins > Settings** tab **>** click the **Add** button.
  - o On the **Update Center Customizer** screen:
    - o Enter some value in the **Name** field (e.g. "My plugins"),
    - o in the **URL** field Enter

http://updates.netbeans.org/netbeans/updates/8.2/uc/final/distribution/catalog.xml.gz

- Click the **OK** button.
- This should create a new entry in the Configuration of Update Centers list in the Settings tab.
- Checking that new entry should instantly add plugins to the **Available Plugins** tab.
- Click the **Available Plugins** tab, then click the **Category** column to sort the entries by category.
- The **Name** of the entry at the top of the list should be **C/C++**. If so, you have successfully made the plugin available:

#### ➤ JDK 11

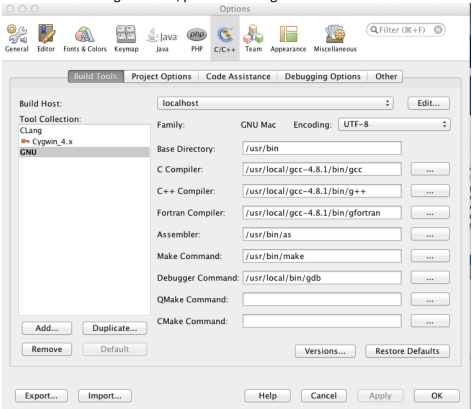
https://www.oracle.com/technetwork/java/javase/downloads/jdk11-downloads-5066655.html

#### (2) gcc/gfortran Compiler configuration

(i) preference->Options->C/C++->Build Tools

If something is missing (e.g., gfortran), we need to install it first, then click the button "..." to locate the package.

If we want to debug the code, please install "gdb" first.



- (4) A Project Folder will be generated, e.g., "/a1/a2/MEND"
- (5) Delete the Sample Source Files "bar.f". & "foo.f"
- (6) Copy the following folders & files into "/a1/a2/MEND": src, userio, MEND namelist.nml
- (7) Edit 'nbproject/Configurations.xml' to define dependency of source code for compiling (see Section 21.1)
- (8) Right click on project name "MEND" in left window, Project Properties->Build->Fortran Compiler->Tool

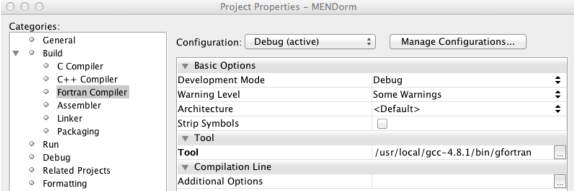
Specify the absolute path for "gfortran" Additional Options:

- -ffpe-trap=invalid,zero,overflow (check floating point exceptions)
- **-finit-local-zero** (initialize local INTEGER, REAL, and COMPLEX variables to zero, LOGICAL variables to false, and CHARACTER variables to a string of null bytes)

this will modify /Users/wgs/Dropbox\_job/Model/MEND/nbproject/Makefile-Debug.mk:

# Fortran Compiler Flags

FFLAGS=-ffpe-trap=invalid,zero,overflow -finit-local-zero



- (9) Compile a single FORTRAN file by right-clicking the file name and click "Compile File (F9)"
- (11)Run model in NetBeans: click icon (12)Run model in Terminal: cd MEND

./mend

(13) Solution to Error: Id: library not found for -ISystem

Solution 1: xcode-select --install

Solution 2 (11/4/2020: works for new laptop):

sudo softwareupdate -1

the Command Line Tools were listed as an update, so I installed them using sudo softwareupdate -i -a

https://developer.apple.com/download/more/

#### 22.5 Compiling on Unix

- (1) Copy the NetBeans configured MEND folder to another system (Windows-Cygwin or Unix)
- (2) Edit file "/nbproject/Makefile-Debug.mk": replace "FC=/\*\*/gfortran" with the full path of "gfortran" on the new machine system, e.g., FC=/usr/local/gcc-8.1/bin/gfortran-8.1

```
# Fortran Compiler Flags
FFLAGS=-ffpe-trap=invalid,zero,overflow -finit-local-zero
```

- (3) "cd" into the MEND model directory
- (4) make clean
- (5) **make -k**: NOT required anymore, as 'nbproject/**Configurations.xml**' defines dependency of source code for compilation
- (6) make
- (7) The "mend" or "mend.exe" file will be generated if there are no errors
- (8) If the path for "mend.exe" is "/dist/Debug/.../mend.exe", type "./dist/Debug/.../mend.exe" to run model
- (9) Another option to run model: copy "/dist/Debug/.../mend.exe" to root-dir of MEND model, type "./mend.exe"
- (10) A 3<sup>rd</sup> option to run model
  cd MEND
  ln -sf ./dist/Debug/.../mend.exe mend.exe
  ./mend.exe

```
##Example: Begin-------
## login into cluster 129.15.40.234
ssh your-username@129.15.40.234
## pass: *******
```

## make a new directory makdir test

## switch to your local machine ## copy a folder from local machine to remote machine ## enter the local directory containing MEND scp -r MEND your-username@129.15.40.234:~/test/MEND2

## switch to 129.15.40.234 which gfortran

/usr/bin/gfortran

vi nbproject/Makefile-Debug.mk ## edit the following line FC=/usr/bin/gfortran ## save and exit

make clean

```
## build the model, generate excutable file
   make
   Is -It dist/Debug/GNU-MacOSX/
   ## link the executable file
   In -sf dist/Debug/GNU-MacOSX/mend mend
   ## run executable file
   ./mend
##Example: End-----
22.6 Text Editor
   ➤ Sublime Text
https://www.sublimetext.com/
Tools -> Command Palette -> install package control
Tools -> Command Palette -> Package Control: Install Package -> SideBarEnhancements
Tools -> Command Palette -> Package Control: Install Package -> Fortran
Preferences -> Setting
Preferences.sublime-settings – User
       "color scheme": "Packages/Color Scheme - Default/Monokai.sublime-color-
scheme",
      "ignored_packages":
             "Vintage"
       "word wrap": false
}
```

#### 22.7 Eclipse Configuration for non-MPI MEND

- ➤ Installing Photran in Eclipse on a Machine With Internet Access To install Photran, start Eclipse, then...
  - 1. Click on Help > Install New Software...
  - 2. Click on the "Available Software Sites" hyperlink
  - 3. Look for <a href="http://download.eclipse.org/tools/ptp/updates/luna">http://download.eclipse.org/tools/ptp/updates/luna</a> in the list.
    - 1. If it is in the list, check the box next to it (if it is not already checked).
    - 2. If it is not in the list...
      - 1. Click on the "Add..." button
      - 2. In the Location field,
        - type http://download.eclipse.org/tools/ptp/updates/luna
      - 3. Click OK to close the Add Site dialog. This will return you to the Install dialog.
  - 4. Click OK to close the Preferences dialog (i.e., the list of available software sites). This will return you to the Install dialog.
  - 5. In the "Work with:" dropdown, choose <a href="http://download.eclipse.org/tools/ptp/updates/luna">http://download.eclipse.org/tools/ptp/updates/luna</a>
  - 6. Expand "Fortran Development Tools (Photran)" and check the box next to "Photran End-User Runtime"
  - 7. If you are running Linux and have the Intel Fortran Compiler installed, or if you are on a Macintosh and have the IBM XL Fortran compiler installed, expand "Fortran Compiler Support" and select the appropriate compiler.
  - 8. Click on the "Next" button
  - 9. If you get an error message, see below for troubleshooting information.
  - 10. Click the Finish button and agree to the license to complete the installation.
- New -> Fortran Project -> Executable (Gnu Fortran on MacOS X) First. Check "Generate Makefiles automatically"
- ➤ **ATTENTION**: Fortran-90 files must have extension of "f90", NOT "F90", otherwise resulting in failure to build project in Eclipse
- > ATTENTION: module: the Fortran file name must be the same as the module name
- Compile code: After "makefile" and "subdir.mk" are generated, uncheck "Generate Makefiles automatically". Now we can modify "makefile" and "subdir.mk", e.g.,

FC=/usr/local/gcc-8.1/bin/gfortran-8.1 Then use \$(FC)

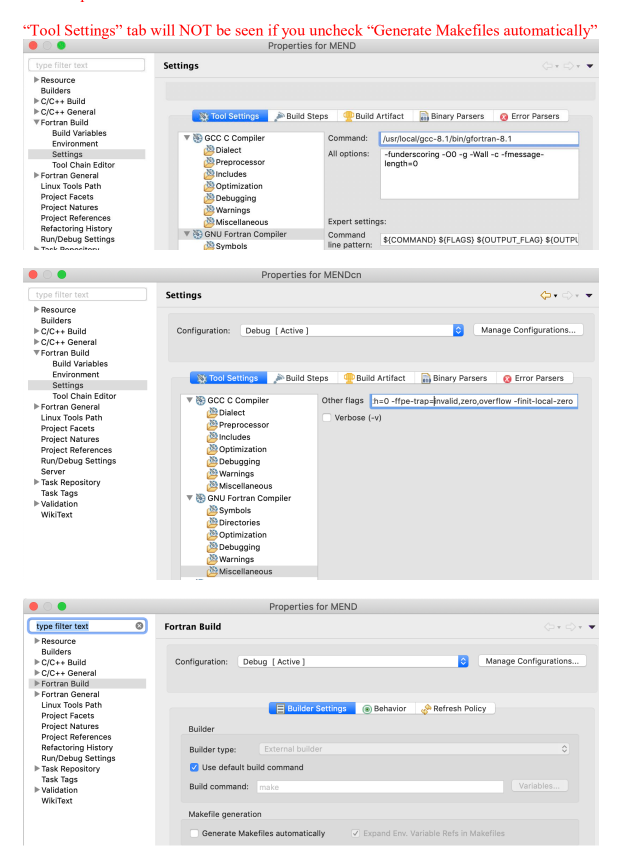
Clean:

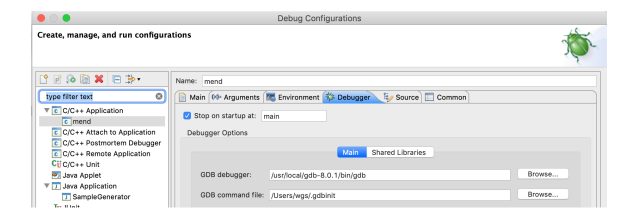
-\$(RM) ./\*.mod

> Debug with gdb

ATTENTION: use full path
/usr/local/gdb8.3.1/bin/gdb
/Users/wgs/.gdbinit

#### set startup-with-shell off





#### 23.1 iTerm2:

### https://iterm2.com/

- disable line wrapping: tput rmam
- enable line wrapping: tput smam

#### 23.2 git

gitignore NOT working correctly git rm -r --cached .
 git add .