Description and Evaluation of the MIT Earth System Model (MESM)

MIT Joint Program working paper

October 6, 2017 (subject to future updates)

Andrei Sokolov¹, David Kicklighter², Adam Schlosser¹, Chien Wang¹, Erwan Monier¹, Benjamin Brown-Steiner¹, Ron Prinn¹, Chris Forest³, Xiang Gao¹, Alex Libardoni³ and Sebastian Eastham⁴

¹ Joint Program on the Science and Policy of Global Change, Massachusetts Institute of Technology, Cambridge, MA, USA
² The Ecosystems Center, Marine Biological Laboratory, Woods Hole, MA, USA
³ Department of Meteorology & Earth and Environmental Systems Institute, Pennsylvania State University, University Park, PA, USA
⁴ Laboratory for Aviation and the Environment, Department of Aeronautics and Astronautics, Massachusetts Institute of Technology, Cambridge, MA, USA

Correspondence to: sokolov@mit.edu

Abstract

The MIT Integrated Global System Model (IGSM) is designed for analyzing the global environmental changes that may result from anthropogenic causes, quantifying the uncertainties associated with the projected changes, and assessing the costs and environmental effectiveness of proposed policies to mitigate climate risk. The IGSM consists of the MIT Earth System Model of intermediate complexity (MESM) and the Economic Projections and Policy Analysis (EPPA) model. This paper documents version of the MESM, which includes a 2-dimensional (zonally averaged) atmospheric model with interactive chemistry coupled to the Global Land System model and an anomaly-diffusing ocean model.

1 Introduction

Projections of climate change over the next century have been hampered by our limited understanding and ability to model the complex interdependencies of the human-climate system. Among factors contributing to this difficulty are significant uncertainties in climate system properties that determine its response to transient forcing, such as climate sensitivity, the rate at
which the deep ocean absorbs heat and the strength of the carbon cycle. There are additional uncertainties in the forcing itself, especially in the forcing associated with aerosol-cloud interaction.

Unfortunately, the available directly-measured ocean, land and atmospheric data can only place limited constraints on these key quantities (e.g. Andronova & Schlesinger, 2001; Gregory et al., 2002; Forest et al., 2008, Libardoni and Forest, 2011). Existing general circulation models (AOGCMs) that couple atmosphere-ocean-land components also differ significantly in both climate sensitivity and rate of heat uptake (e.g. Andrews et al., 2013; Forster et al., 2013; IPCC, 2013). In spite of significant efforts, these uncertainties have not been reduced over the last few decades and are unlikely to be substantially reduced within the next decade or more, when important policy choices must nevertheless be made. Uncertainties in the strength of the carbon cycle and carbon-climate feedback also remain significant (Friedlingstein et al, 2006 and 2014), leading to a large spread in the future atmospheric CO$_2$ concentration and radiative forcing projected by different Earth System Models (ESM) for a given emission scenario.

These uncertainties, in turn, result in a rather wide uncertainty in projected future climate change. To account for these uncertainties, the latest report of Intergovernmental Panel on Climate Change (IPCC, 2013) provides mean values and probability intervals for projected changes of future climate, based on a multi-model ensemble (MME). There are, however, well known problems with MMEs, such as small sample size and the fact that different modes are neither independent nor likely plausible (IPCC, 2013 and literature referenced there). In addition, there are no guaranties that the existing AOGCMs sample the full ranges of uncertainties in different climate characteristics and moreover sample these ranges homogeneously. An alternative approach is to estimate the probability distributions of climate parameters based on available data for past climate and then carry out large (few hundred members) ensembles of future climate simulations by sampling parameter values from these distributions.

Even with much greater computational power than is available today, however, it will not be possible to carry out such an exercise using a fully complex state-of-the-art AOGCM or ESM. Therefore, such studies are usually carried out with Earth System Models of Intermediate Complexity (EMICs) (Knutti et al., 2003; Rogelj et al., 2012; Sokolov et al, 2009; Webster et al., 2003 and 2012). It has been shown that, with an appropriate choice of parameter values EMICs can reproduce global mean changes simulated by different AOGCMs under different forcing
scenarios (Raper et al., 2001; Sokolov et al., 2003; Meinshausen et al. 2011). Model intercomparisons also have shown that in many cases changes in climate predicted by models of intermediate complexity are very similar to those obtained in the simulations with AOGCMs (Gregory et al., 2005; Stouffer et al., 2005).

The MIT Earth System Model (MESM) was designed to provide the flexibility and computational speed required to handle uncertainty analysis while representing to the best degree possible the physics, chemistry and biology of the more computationally intensive AOGCMs. Also, within the MIT Integrated System Model (IGSM), the MESM is linked to a model of human interactions. The MESM can be run in both concentration-driven and emissions-driven modes. As a result, it can be used for the quantification of uncertainties in future climate. Specifically, Forest et al. (2002 and 2008), Libardoni and Forest (2011 and 2013) and Libardoni (2017) used the MESM to produce probability distributions for the climate sensitivity, the rate of heat uptake by the deep oceans, and the net forcing due to aerosol-radiation interaction by comparing observed temperature changes over the 20th century with results of the historical (concentration-driven) simulations in which model parameter values were varied over wide ranges. Then constructed distributions were used to carry out ensembles of future climate emissions-driven simulation and produce probability distributions for changes in different climate variables. Uncertainty in climate system response can be combined with uncertainty in anthropogenic emissions (Webster et al., 2002 and 2008) to estimate overall uncertainty in possible anthropogenic climate change (Sokolov et al., 2009; Webster et al., 2003 and 2012).  

The first version of the MESM was developed in the mid-1990s (Sokolov and Stone 1995 and 1997; Prinn et al., 1999) and has since been continually modified and extended (Sokolov et al., 2005). Different versions of the MESM were used in a number of model intercomparison projects (e.g., Gregory et al., 2005; Petoukhov et al., 2005; Stouffer et al., 2005; Brovkin et al., 2006; Plattner et al., 2008; Eby et al., 2013; Olsen et al. 2013; Zickfeld et al., 2013; Brasseur et al., 2016). The MESM shows generally comparable results to those of more complex models. For example, the study of the impact of aviation emissions on atmospheric chemical composition and climate showed that MESM performs within the envelope of the more complex 3-D chemistry-climate models (Brasseur et al., 2016; Olsen et al., 2013).

1 In previous publications acronym “IGSM” was used for both Integrated System Model as a whole and its climate component. Here we use acronym “MESM” for the latter.
In this paper, we describe the current version of the MESM, as of the middle of 2017. Description of the model components is given in Section 2. Section 3 provides a comparison of simulated present-day climate and historical climate change with results produced by CMIP5 models and available observations.

2 Model Description

The major model components of the MESM (Figure 1) include:

- An atmospheric dynamics, physics and chemistry model, which includes a sub-model of urban chemistry,
- An ocean model with carbon cycle and sea-ice sub-models,
- A linked set of coupled process-based land models, the Terrestrial Ecosystem Model (TEM), a fully integrated Natural Emissions Model (NEM), and the Community Land Model (CLM), that simulate terrestrial water, energy, carbon and nitrogen budgets including carbon dioxide (CO$_2$) and trace gas emissions of methane (CH$_4$) and nitrous oxide (N$_2$O).

The earth system depicted in Figure 1 represents a fully coupled system that allows simulation of critical feedbacks among its components. Time-steps used in the various sub-models range from 10 minutes for atmospheric dynamics, to 1 month for TEM, reflecting differences in the characteristic time-scales of different processes simulated by the MESM. The major model components of the MESM and linkages are summarized below.

2.1 Atmospheric Dynamics and Physics

The MIT two-dimensional (2D) atmospheric dynamics and physics model (Sokolov & Stone, 1998) is a zonally-averaged statistical dynamical model that explicitly solves the primitive equations for the zonal mean state of the atmosphere and includes parameterizations of heat, moisture, and momentum transports by large-scale eddies based on baroclinic wave theory (Stone & Yao, 1987, 1990). The model’s numerics and parameterizations of physical processes, including clouds, convection, precipitation, radiation, boundary layer processes, and surface fluxes, build upon those of the Goddard Institute for Space Studies (GISS) GCM (Hansen et al., 1983). The radiation code includes all significant greenhouse gases (H$_2$O, CO$_2$, CH$_4$, N$_2$O, CFCs and O$_3$) and eleven types of aerosols. The model’s horizontal and vertical resolutions are
variable, but in the standard version of MESM it has 4° resolution in latitude and eleven levels in the vertical.

The MIT 2D atmospheric dynamics and physics model allows up to four different types of surface in the same grid cell (ice free ocean, sea-ice, land, and land-ice). The surface characteristics (e.g., temperature, soil moisture, albedo) as well as turbulent and radiative fluxes are calculated separately for each kind of surface while the atmosphere above is assumed to be well mixed horizontally in each latitudinal band. The area weighted fluxes from different surface types are used to calculate the change of temperature, humidity, and wind speed in the atmosphere. Fluxes of sensible heat and latent heat are calculated in the atmospheric model by bulk formulas with turbulent exchange coefficients dependent on the Richardson number. The atmosphere’s turbulence parameterization is also used in the calculation of the flux derivatives with respect to surface temperature. To account for partial adjustment of near surface air temperature to changes in fluxes the derivatives are calculated under the assumption that the exchange coefficients are fixed. A more detailed discussion of technical issues involved in the calculations of these fluxes and their derivatives is given in Kamenkovich et al. (2002).

The moist convection parameterization, which was originally designed for the GISS Model I (Hansen et al. 1983), requires knowledge of sub-grid scale temperature variance. Zonal temperature variance associated with transient eddies is calculated using a parameterization proposed by Branscome (see Yao and Stone 1987). The variance associated with stationary eddies is represented by adding a fixed variance that follows more closely the climatological pattern (see Figure 7.8b of Peixoto and Oort 1992). In addition, the threshold values of relative humidity for the formation of large-scale cloud and precipitation varies with latitude to account for the dependence of the zonal variability of relative humidity on latitude. Zonal precipitations simulated by the atmospheric model are partitioned into land and ocean components using present day climatology. These changes led to an improvement in the zonal pattern of the annual cycle of land precipitation and evapotranspiration (Schlosser at al. 2007).

The atmospheric model’s climate sensitivity can be changed by varying the cloud feedback. Namely, the cloud fraction used in radiation calculation ($C_{\text{rad}}$) is adjusted as follows:

$$C_{\text{rad}} = C^0 (1.0 + \kappa \Delta T_{\text{surf}}),$$  \hspace{1cm} (1)
where \( C^o \) is a cloud fraction simulated by the model, \( \Delta T_{\text{surf}} \) is a difference of the global-mean surface air temperature from is values in the control climate simulation and \( \kappa \) is a parameter used to vary climate sensitivity.

This method was proposed by Hansen et al. (1993) and was extensively tested in simulations with the MIT climate model (Sokolov & Stone, 1998). The choice of cloud feedback seems very natural because differences in climate sensitivity between different AOGCMs are to large extent caused by large differences in this feedback (Cess et al. 1990; Colman 2003; Bony et al. 2006; Webb et al. 2006; Williams et al. 2006). The method was later modified by using \( \kappa \) of different signs for high and low clouds (Sokolov, 2006), accounting for the fact that the feedback associated with changes in cloud cover has different signs for high and low clouds. Therefore, using different signs in Equation 1 depending on cloud heights minimizes the value of \( \kappa \) required to obtain a specific value of climate sensitivity. In addition, use of modified method improves an agreement in simulated changes is surface fluxes between MIT climate model and different AOGCMs.

This approach to changing climate sensitivity was also tested in simulations with CAM3 (Sokolov and Monier, 2012), by comparison with perturbed physics approach.

### 2.2 Urban and Global Atmospheric Chemistry

To calculate atmospheric composition, the model of atmospheric chemistry includes the climate-relevant chemistry of gases and aerosols at two domains: the urban scale and the global scale. The urban model is a sub-grid scale chemistry model whose emissions and pollutants are exported (along with emissions from non-urban areas) into the 2D global zonal-mean model of atmospheric chemistry, which is linked to the atmospheric dynamics and physics model described above. This atmospheric model provides wind speeds, temperatures, solar radiation fluxes, and precipitation to both the urban and global scale chemistry models. The details of the sub-grid scale urban chemistry model and the 2D zonal-mean atmospheric chemistry model, and their coupling, are described below.

#### 2.2.1 Urban Atmospheric Chemistry

Urban emissions and air pollution have a significant impact on global methane, ozone, and aerosol chemistry, and thus on the global climate. However, the nonlinearities in the chemistry
cause urban emissions to undergo different net transformations than rural emissions, and thus urban chemistry is treated separately from non-urban emissions within the MESM. Accuracy in describing these transformations is necessary because the atmospheric lifecycles of exported air pollutants such as CO, O₃, NOₓ and VOCs, and the climatically important species CH₄ and sulfate aerosols, are linked through the fast photochemistry of the hydroxyl free radical (OH) as we will emphasize in the results discussed later in section 5. Urban air-shed conditions need to be resolved at varying levels of pollution. The urban air chemistry model must also provide detailed information about particulates and their precursors important to air chemistry and human health, and about the effects of local topography and structure of urban development on the level of containment and thus the intensity of air pollution events. This is an important consideration because air pollutant levels are dependent on projected emissions per unit area, not just total urban emissions.

The urban atmospheric chemistry model has been introduced as an additional component to the original global model (Prinn et al. 1999) in MESM (Calbo et al. 1998; Mayer et al. 2000; Prinn et al. 2007). It was derived by fitting multiple runs of the detailed 3D California Institute of Technology (CIT) Urban Airshed Model, adopting the probabilistic collocation method to express outputs from the CIT model in terms of model inputs using polynomial chaos expansions (Tatang et al. 1997). This procedure results in a reduced format model to represent about 200 gaseous and aqueous pollutants and associated reactions over urban areas that is computationally efficient enough to be embedded in the global model. The urban module is formulated to take meteorological parameters including wind speed, temperature, cloud cover, and precipitation as well as urban emissions as inputs. Calculated with a daily time step, it exports fluxes along with concentrations (peak and mean) of selected pollutants to the global model.

2.2.2 Global Atmospheric Chemistry

The 2D zonal mean model that is used to calculate atmospheric composition is a finite difference model in latitude-pressure coordinates, and the continuity equations for the trace constituents are solved in mass conservative, of flux, form (see Wang et al., 1998 for a more complete description). The model includes 22 chemical species with 41 gas-phase and 12 aqueous-phase reactions (Wang et al., 1998). For the longer-lived species (CFCl₃, CF₂CL₂, N₂O, CO, CO₂, NO, NO₂, N₂O₅, HNO₃, CH₄, CH₂O, SO₂, H₂SO₄, HFC, PFC, SF₆, and black carbon
and organic carbon aerosols), the chemistry model includes convergence due to transport, parameterized north-south eddy transport, convective transport, and local true production and loss due to surface emissions, deposition, and chemical reaction. For the very-reactive atoms (e.g. O), free radicals (e.g. OH), and molecules (e.g. H$_2$O$_2$), their concentrations are unaffected by transport due to their very short lifetimes, and only their chemical production and loss (in both the gaseous and aqueous phase) is considered. The scavenging of carbonaceous and sulfate aerosol species by precipitation is also included using a method based on a detailed 3D climate-aerosol-chemistry model (Wang, 2004). Water vapor and air (N$_2$ and O$_2$) mass densities are computed using full continuity equations as part of the atmospheric dynamics and physics model (described above) to which the chemical model is coupled.

2.2.3 Coupling of Urban and Global Atmospheric Chemistry

The urban chemistry module was derived based on an ensemble of 24-hour long CIT model runs and thus is processed in the IGSM with a daily time step, while the global chemistry module is run in a real time step with the dynamics and physics model, 20 minutes for advection and scavenging, 3 hours for tropospheric reactions. The two modules in the IGSM are processed separately at the beginning of each model day, supplied by emissions of non-urban and urban regions, respectively. At the end of each model day, the predicted concentrations of chemical species by the urban and global chemistry modules are then remapped based on the urban to non-urban volume ratio at each model grid. Beyond this step, the resultant concentrations at each model grid will be used as the background concentration for the next urban module prediction and also as initial values for the global chemistry module (Mayer et al. 2000).

2.3 Ocean Component

The ocean model used in the version of MESM described in this paper, consists of a mixed-layer model with a horizontal resolution of 4° in latitude and 5° in longitude and a 3000-m deep anomaly-diffusing ocean model beneath. Mixed-layer depth is prescribed based on observations as a function of time and location (Hansen et al., 1983). In addition to the temperature of the mixed layer, the model also calculates the average temperature of the seasonal thermocline and the temperature at the annual maximum mixed layer depth (Russell et al. 1985). Heat mixing into the deep ocean is parametrized by the diffusion of the difference of the temperature at the bottom
of the seasonal thermocline from its value in a pre-industrial climate simulation (Hansen et al. 1984; Sokolov and Stone 1998). Since this diffusion represents a cumulative effect of heat mixing by all physical processes, the values of the diffusion coefficients are significantly larger than those used in the sub-grid scale diffusion parameterizations in OGCMs. The spatial distribution of the diffusion coefficients used in the diffusive model is based on observations of tritium mixing into the deep ocean (Hansen et al. 1988). The rate of oceanic heat uptake is varied by multiplying diffusion coefficients by the same factor in all locations.

The coupling between the atmospheric and oceanic models takes place every hour. The heat flux \( FH \) at the longitude-latitude point \((i, j)\) is calculated as:

\[
FH(i,j) = FHZ(j) + \frac{\partial FHZ}{\partial T}(j) \times (Ts(i,j) - Tsz(j)),
\]

where \( FHZ(j) \) and \( \frac{\partial FHZ}{\partial T}(j) \) are zonally averaged heat flux and its derivative with respect to surface temperature, and \( Ts(i,j) \) and \( Tsz(j) \) are surface temperature and its zonal mean.

The mixed-layer model also includes specified vertically-integrated horizontal heat transport by the deep oceans, a so-called “Q-flux”. This flux has been calculated from a simulation in which sea surface temperature and sea-ice distribution were relaxed toward their present-day climatology with relaxation a coefficient of 300 Wm\(^{-2}\)/K, corresponding to an e-folding time-scale of about 15 days for the 100 m deep mixed-layer. Relaxing SST and sea ice on such a short time scale, while being virtually identical to specifying them, avoids problems with calculating the Q-flux near the sea ice edge.

A thermodynamic ice model is used for representing sea ice. The model has two layers and computes ice concentration (the percentage of area covered by ice) and ice thickness.

Different version of MESM was developed by replacing simplified ocean model with the MIT 3D OGCM (Dutkiewicz et al., 2005). A detailed comparison of the results of simulations with the two versions of the MESM was carried out to evaluate the performance of the anomaly diffusing ocean model (ADOM) on different time scales Sokolov et al. (2007). This comparison lead to significant modification of the ocean carbon model originally used in AODM (Holian et al. 2001).

In the current version of MESM formulation of carbonate chemistry and parameterization of air-sea carbon fluxes are similar to the ones used in the MIT 3D OGCM (Dutkiewicz et al., 2005).
Vertical and horizontal transports of the total dissolved inorganic carbon, though, are parameterized by diffusive processes. The values of the horizontal diffusion coefficients are taken from Stocker et al. (1994), and the coefficient of vertical diffusion of carbon ($K_{vc}$) depends on the coefficient of vertical diffusion of heat anomalies ($K_v$). Originally, $K_{vc}$ was assumed to be proportional to $K_v$ (Prinn et al. 1999; Sokolov et al. 1998). This assumption, however, does not take into account the vertical transport of carbon due to the biological pump. In the present version of MESM $K_{vc}$ is, therefore, defined as:

$$K_{vc} = K_{vco} + rK_v$$

(3)

Where $K_{vco}$ represents mixing due to the biological pump and $rK_v$ due to physical processes. Since $K_{vco}$ is a constant, the vertical diffusion coefficients for carbon have the same latitudinal distribution as the coefficients for heat anomalies. For simulations with different rates of oceanic uptake, the diffusion coefficients are scaled by the same factor in all locations. Therefore, rates of both heat and carbon uptake by the ocean are defined by the global mean value of the diffusion coefficient for heat. In the rest of the paper the symbol $K_v$ is used to designate the global mean value.

Comparisons with 3D ocean simulations have shown that the assumption that changes in ocean carbon can be simulated by the diffusive model with fixed diffusion coefficient works only for about 150 years. On longer timescales, the simplified carbon model overestimates the ocean carbon uptake. However, if $K_{vc}$ is assumed to be time dependent, the MESM reproduces changes in ocean carbon as simulated by the 3D OGCM on multi century scales (Sokolov et al. 2007). Thus, for the runs discussed here, the coefficient for vertical diffusion of carbon was calculated as:

$$K_{vc}(t) = (K_{vco} + rK_v) f(t)$$

(4)

Where $f(t)$ is a time dependent function constructed based on the analyses of the depths of carbon mixing in simulations with the 3D OGCM.

Overall results presented by Sokolov et al. (2007) show that in spite of its inability to depict feedbacks associated with the changes in the ocean circulation and a very simple parameterization of the ocean carbon cycle, the version of the MESM with the ADOM is able to reproduce the important aspects of the climate response simulated by the version with the OCGM through the 20th and 21st century and can be used to produce probabilistic projections of
changes in many of the important climate variables, such as surface air temperature and sea level, through the end of 21st century.

The MESM also has been shown, with an appropriate choice of the model’s cloud feedback and effective diffusion coefficient, to reproduce the transient surface warming and sea level rise due to thermal expansion of the ocean as calculated in various coupled AOGCMs for 120-150 year time-scales (Sokolov & Stone, 1998; Sokolov et al., 2003).

2.4 Land Component

The land component of the MESM estimates how fluxes of heat, water, carbon and nitrogen, both within land ecosystems and between land and the atmosphere, vary across the globe over time. In addition, the land component estimates how soil moisture and the storage of carbon and nitrogen in vegetation and soils vary across the globe over time. The land flux and storage estimates are determined driven by near-surface atmospheric states (e.g. air temperature, humidity, pressure, wind speed), fluxes (radiation, precipitation) as well as atmospheric chemistry (carbon dioxide, ozone) variables estimated by the atmospheric component of the MESM along with external data sets that prescribe the distribution of land cover and soil texture across the globe. The land component estimates of albedo, sensible heat, latent heat, evapotranspiration, and fluxes of carbon dioxide (CO₂), methane (CH₄), and nitrous oxide (N₂O) are then used by the atmospheric component of the MESM to determine changes in atmospheric physics and chemistry. In order to assess the value of global land resources, estimates of net primary production (NPP) are used by the EPPA model in the MIT IGSM to indicate how this ecosystem service influences economic activity.

Global processes in the land component are represented with a dynamically linked set of terrestrial biogeophysical (i.e., water and energy budgets) and biogeochemical (i.e., carbon and nitrogen budgets including carbon dioxide, methane, and nitrous oxide fluxes) sub-models. These biogeophysical and biogeochemical calculations are organized into a single, self-consistent framework for the global terrestrial environment and hereafter referred to as the Global Land Systems (GLS) framework (Schlosser et al. 2007). The GLS framework, employs three coupled sub-models to represent the terrestrial water, energy, and ecosystem processes:

• The Community Land Model (CLM) described by Bonan et al. (2002) calculates the global, terrestrial water and energy balances.
The Terrestrial Ecosystems Model (TEM) of the Marine Biological Laboratory (Melillo et al., 1993, 2009; Xiao et al., 1997, 1998; Tian et al. 1999; Felzer et al., 2004) simulates carbon and nitrogen fluxes and the storage of carbon and nitrogen in vegetation and soils including the uptake and release of CO$_2$ associated with net primary production, decomposition and carbon sequestration or loss.

The Natural Emissions Model (NEM) described by Liu (1996) and Prinn et al. (1999) simulates CH$_4$ and N$_2$O fluxes.

Water, energy and carbon are conserved among these sub-models. GLS is also designed to be flexibly run either with the meridional resolution (4°) of the zonally-averaged atmospheric model within the MESM, or over a latitude-longitude grid for targeted studies (e.g., 2° by 2° as in Gao et al. 2013, and 0.5°x0.5° as in Melillo et al., 2009).

Herein, we describe the coupled configuration of the GLS used within the MESM. In this GLS configuration, a vegetation mosaic scheme has been used to represent the distribution of vegetation within a latitudinal zonal band at the same spatial resolution for all submodels. Each latitudinal band is represented with a mosaic of 35 land cover or IGSMVEG types (Schloesser et al., 2007) based on the presence or absence of a dominant tree, shrub or grass cover, ecological region (i.e., boreal, temperate, tropical), moisture status (upland, floodplain or wetland) and land management (crop, pasture). In most applications of the GLS within the MESM, the land cover has been assumed to be potential vegetation (i.e., natural vegetation in the absence of human activity) such that the potential impacts of land-use change on the ability of land to store carbon and nitrogen have not been considered. However, a scheme for incorporating the influence of land-use change on land carbon dynamics in the GLS has been developed and applied in Eby et al. (2013) and Zickfeld et al. (2013).

Below we highlight the key features of each of the land sub-models and modifications made to these sub-models for their implementation in the MESM.

2.4.1 The Community Land Model (CLM)

As in previous implementations of land biogeophysical and hydrologic processes within the IGSM framework, we have drawn from the Community Land Model (CLM, Lawrence et al., 2011). The CLM has been developed from a multi-institutional collaboration of land models, and serves as the core land system model for energy, water, carbon, and nutrient cycle studies within
the Community Earth System Model (Oleson et al., 2010). CLM has also been widely used and documented in land data assimilation research (e.g., Zhang et al., 2012), hydrologic studies at the global (e.g. Pu et al., 2012), regional (e.g. Swenson et al., 2012; Zampieri et al., 2012), and river-basin (e.g. Vano et al., 2012) scales, as well as coupled climate prediction studies (e.g., Tseng et al., 2012, Xin et al., 2012). CLM is also benchmarked within the iLAMB framework (e.g. Randerson et al., 2009).

Within this presented implementation of MESM, we have employed CLM Version 3.5 that largely follows the detailed documentation provided by Oleson et al. (2010) as well as many of the features highlighted by Lawrence et al. (2011). We however have made some modifications to CLM’s configuration used within the MESM. Within CLM’s surface soil hydrologic formulation, infiltration of rainfall in the uppermost layer of the soil required further refinement for its implementation in MESM. In the initial testing of CLM within the MESM’s zonal configuration, it was found that CLM produced excessive infiltration into the soil column. This resulted in an appreciably low bias in runoff and subsequently an excessive amount of evapotranspiration as compared to our previous versions of CLM implemented in the model and also against a multi-model consensus of estimates (Schlosser et al., 2007 – see Fig. 15). The algorithm that describes the infiltration rate, \( q_{\text{inmax}} \), into the uppermost soil layer can be summarized as:

\[
q_{\text{inmax}} \sim \left(1 - \frac{2b\psi_{\text{suscat}}}{dz}\right) \times F_{\text{dry}} \times K_{\text{sat}}
\]

where \( b \) is the Clapp-Hornberger parameter, \( \psi_{\text{suscat}} \) is the soil suction from the top layer of soil, \( dz \) is the thickness of the top soil layer, \( F_{\text{dry}} \) is a dryness factor of the upper soil layer, and \( K_{\text{sat}} \) is the saturated hydraulic conductivity. This infiltration formulation closely follows that of the classic Green-Ampt formulation (1911) for “enhanced” (i.e. values greater than saturated hydraulic conductivity) infiltration rate for dry soils—and will sustain this condition for dry soils (i.e. sub-saturated) in the uppermost soil layer. The inherent assumption with this formulation is that saturated and unsaturated conditions in the uppermost soil layer will occur sporadically over a large heterogeneous landscape of intermittent precipitation. However, within our zonally resolved implementation of CLM, we have removed this enhancement effect. While the MESM does resolve the temporal episodic nature of precipitation provided to the GLS (see Schlosser et al., 2007) - the spatially heterogeneous nature of these conditions is not comprehensively resolved. Therefore, we simply set the maximum infiltration rate equal to saturated hydraulic
conductivity. As a result of this modification, we find that our estimates of runoff and subsequent evapotranspiration are more aligned with present-day estimates from the more detailed models – judged commensurately on a zonally averaged basis (discussed in Section 3.3).

The CLM, as well as atmospheric physics, is run at an hourly time scales in order to resolve diurnal variations in the surface energy budget and associated radiative and turbulent heat exchange as well as momentum flux between the land and atmosphere. All inputs to CLM that require a temporal sampling resolution at the time-step are provided by the atmospheric model (as shown in Figure 1); CLM then calculates surface heat, water and momentum fluxes that are passed back to the atmospheric model. The calculations of soil/vegetation water and energy states and fluxes (and corresponding storages and temperatures) are averaged and accumulated as necessary given the time-steps of TEM and NEM. CLM provides estimates of soil moisture and temperature profiles, as well as evapotranspiration rates that are required inputs for the TEM and NEM components and used to estimate fluxes of CO₂, N₂O and CH₄ between terrestrial ecosystems and the atmosphere.

2.4.2 The Terrestrial Ecosystems Model (TEM)

The Terrestrial Ecosystem Model (TEM) is a process-based biogeochemistry model that uses spatially referenced information on atmospheric chemistry, climate, elevation, soil texture, and land cover to estimate monthly fluxes and pool sizes of carbon, nitrogen and water among vegetation, soils and the atmosphere. The TEM is well documented and has been used to examine patterns of land carbon dynamics across the globe including how they are influenced by multiple factors such as CO₂ fertilization, ozone pollution, climate change and variability, and land-use change, (Felzer et al., 2004, 2005; Reilly et al., 2007, 2012; Sokolov et al., 2008; Melillo et al., 2009, 2016; Galford et al. 2010, 2011; Kicklighter et al. 2012, 2014).

In TEM, the uptake of atmospheric carbon dioxide by vegetation, also known as gross primary production or GPP is dependent upon photosynthetically active radiation (PAR), leaf phenology, air temperature, evapotranspiration rates, atmospheric concentrations of carbon dioxide and ozone, the availability of inorganic nitrogen in the soil, and the ratio of carbon to nitrogen (C:N) of new plant biomass (Raich et al. 1991; McGuire et al. 1997; Tian et al. 1999; Felzer et al. 2004). Carbon dioxide is released back to the atmosphere from terrestrial ecosystems as a result of the autotrophic respiration (Rₐ) of plants and the heterotrophic respiration (Rₜ)
associated with the decomposition of soil organic matter. Net primary production (NPP), which is an important source of food and fiber for humans and other organisms on earth, is the net uptake of atmospheric carbon dioxide by plants and is calculated as the difference between GPP and $R_A$.

Heterotrophic respiration depends upon the amount of soil organic matter, the C:N ratio of the soil organic matter, air temperature and soil moisture (Raich et al. 1991; McGuire et al. 1997; Tian et al. 1999). Within an ecosystem, carbon may be stored either in vegetation biomass or in detritus (i.e. litter, standing dead and soil organic matter). In TEM, the carbon in vegetation biomass and detritus are each represented by a single pool. The transfer of carbon between these two pools is represented by litterfall carbon ($L_C$), which is calculated as a proportion of vegetation carbon. Changes in vegetation carbon ($\Delta VEGC$, also known as biomass increment), detritus ($\Delta SOILC$) and terrestrial carbon ($\Delta TOTALC$) are then determined as a linear combination of these fluxes:

$$\Delta VEGC = GPP - R_A - L_C$$

$$\Delta SOILC = L_C - R_H$$

$$\Delta TOTALC = \Delta VEGC + \Delta SOILC = NPP - R_H = GPP - R_A - R_H$$

Carbon sequestration in terrestrial ecosystems can be estimated by the GLS either as the sum of the estimated changes in carbon in vegetation and detritus or by the difference between NPP and $R_H$ (Eq. 8) which is also known as net ecosystem production or NEP.

An important feature of TEM is that the model simulates the influence of terrestrial nitrogen dynamics on terrestrial carbon dynamics. First, the uptake of carbon dioxide by plants is assumed by TEM to be limited by nitrogen availability in most land ecosystems on earth. Tropical forests are the only exceptions, where nitrogen availability is not assumed to limit GPP under contemporary conditions. The effect of nitrogen limitation on GPP is determined by first calculating $GPP_C$ assuming no nitrogen limitation:

$$GPP_C = f(CO_2, ET) f(PAR) f(CANOPY) f(LEAF) f(T) f(O_3, ET)$$

where $CO_2$ is atmospheric CO$_2$ concentration, $ET$ is evapotranspiration, PAR is photosynthetically active radiation, CANOPY is the relative state of a vegetation canopy recovering from disturbance as compared to the canopy state of a mature stand, LEAF is the monthly leaf area relative to the maximum leaf area of a stand, $T$ is air temperature, and $O_3$ is atmospheric ozone concentration. The influence of atmospheric CO$_2$ and $O_3$ concentrations on GPP depends on the status of vegetation stomates, which close during drier conditions (i.e. low
ET) to reduce the uptake of CO$_2$ or O$_3$ and open during wetter conditions (i.e. high ET). Details of Equation 9 have been described elsewhere (e.g Raich et al. 1991; McGuire et al. 1992, 1995, 1997; Pan et al. 1998; Tian et al. 1999; Felzer et al. 2004).

GPP$_N$ is then calculated based on the effects of nitrogen supply on net primary production (NPP$_N$):

$$\text{NPP}_N = P_{CN} \times (\text{NUPTAKE} + \text{NMOBIL})$$  \hspace{1cm} (10)

$$\text{GPP}_N = \text{NPP}_N + \text{RA}$$  \hspace{1cm} (11)

where $P_{CN}$ is the carbon to nitrogen ratio (C:N) of newly produced plant tissue, NUPTAKE is the amount of inorganic nitrogen acquired by plants from the soil and NMOBIL is the amount of vegetation labile nitrogen mobilized during a particular month (McGuire et al. 1997; Pan et al. 1998; Tian et al. 1999). Similar to GPP$_C$, NUPTAKE also depends on local environmental conditions (Raich et al. 1991; McGuire et al. 1992; Felzer et al. 2004):

$$\text{NUPTAKE} = f(N_{AVL}, H_2O) \times f(\text{CANOPY}) \times f(T) \times f(O_3, ET)$$

where $N_{AVL}$ is soil available nitrogen and H$_2$O is soil moisture. Monthly GPP is then determined as follows:

$$\text{GPP} = \min (\text{GPP}_C, \text{GPP}_N)$$  \hspace{1cm} (12)

In TEM, the uptake of atmospheric CO$_2$ by plants is assumed to follow Michaelis-Menten kinetics, according to which the effect of atmospheric CO$_2$ at time $t$ as modified by ET on the assimilation of CO$_2$ by plants is parameterized as follows:

$$f(\text{CO}_2(t)) = \frac{(C_{\text{max}} \times \text{CO}_2(t))}{(kc + \text{CO}_2(t))}$$  \hspace{1cm} (13)

where $C_{\text{max}}$ is the maximum rate of C assimilation, and $kc$ is the CO$_2$ concentration at which C assimilation proceeds at one-half of its maximum rate (i.e., $C_{\text{max}}$). Because the response of carbon uptake by plants to atmospheric CO$_2$ concentrations is uncertain (Sokolov et al. 2009), we examine the influence of this uncertainty on carbon by adjusting the value of $kc$ in our uncertainty analyses.

The Natural Emissions Model (NEM) has been embedded within the TEM infrastructure as described in Schlosser et al. (2007). As such, the CH$_4$ and N$_2$O flux estimates by NEM are directly based on monthly TEM estimates of soil organic carbon. Although the simulated carbon and nitrogen dynamics of NEM are still mostly separated from those in TEM, the embedded NEM provides a platform for improving the linkages between the two biogeochemistry-models. For example, NEM estimates of CH$_4$ emissions diminish the stock of soil organic carbon.
estimated by TEM. The embedded NEM allows simulation of daily CH$_4$ and N$_2$O emissions based on monthly estimates of soil organic carbon by TEM combined with the CLM estimates of daily soil temperatures, daily and hourly soil moistures, hourly rainfall intensities and storm durations. CLM also provides all data on soil properties required by TEM/NEM. Fluxes of CO$_2$, CH$_4$ and N$_2$O are passed to atmospheric model and are used in calculations of corresponding gases by atmospheric chemistry sub-model.

3 Model Evaluation

As mentioned above, MESM can be run either in concentration-driven or emissions-driven mode. In historical simulations MESM is forced by the prescribed changes in by greenhouse gases and ozone concentrations, aerosols, and solar irradiance. Greenhouse gas concentrations and stratospheric aerosols from volcanic eruptions are taken from the NASA GISS modeling group forcing suite. Miller et al. (2014) describe the methods for updating the greenhouse gas concentrations from Hansen et al. (2007) and the volcanic aerosol optical depths from Sato et al. (1993). Sulfate aerosol loadings are from Smith et al. (2011) extended by Klimont et al. (2013), solar irradiance is from the Kopp and Lean (2011) dataset, and ozone concentrations are from the SPARC dataset used in the CMIP5 experiments (Cionni et al., 2011). In future climate simulations MESM is driven by anthropogenic emissions of different gases produced by the MIT Economic Projection and Policy Analysis (EPPA) model (Paltsev et al., 2005).

3.1 Distribution of climate parameters and characteristic describing model response to external forcing.

To determine climate model parameters that produce changes in surface air and ocean temperatures consistent with available observations, 1800 historical simulations from 1861 to 2010 were carried out changing climate sensitivity, the rate of ocean heat uptake and the strength of aerosol forcing over wide ranges. Probability distributions for climate parameters were constructed using the methodology described in Forest et al., (2002 and 2008) and Libardoni and Forest (2011 and 2013). Detailed descriptions of the distributions obtained using different observational data sets, different estimates of natural variability and other assumptions are given in Libardoni et al (2017). Below we described our final distributions and present the results of simulations with the set of climate parameters sampled from it. This distribution was derived
using multiple datasets for changes in surface temperature from 1905 to 2010 and in ocean heat content from 1990 to 2010. Individual distributions were merged across all surface datasets to produce the final distribution. Estimates of natural variability from all CMIP5 models were used to estimate the noise-covariance matrix.

<table>
<thead>
<tr>
<th>Climate sensitivity K</th>
<th>Square root of diffusion coefficient cm/s$^{1/2}$</th>
<th>Radiative forcing due to aerosol radiation interaction W/m$^2$</th>
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<tr>
<td>50% 3.2</td>
<td>1.8</td>
<td>-0.24</td>
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<tr>
<td>95% 4.6</td>
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Table 1. Percentiles for climate parameters.

Table 1 shows medians and 90% probability intervals for model climate parameters from distribution used in this paper. The median values for climate sensitivity (see also Figure 2a) is rather close to the median climate sensitivity of CMIP5 AOGCMs (3.2K), while the 90% probability interval is shifted toward higher values compared with the CMIP5 models (1.9 to 4.5K).

To assess uncertainty in the transient climate response (TCR, i.e., temperature change in the time of CO$_2$ doubling), we carried out a set of climate simulations with a 1% per year increase of CO$_2$ concentration using 400 samples of climate parameters. To estimate uncertainty in the carbon cycle we calculated carbon uptake by the ocean and terrestrial ecosystems in these simulations. As discussed above, the vertical diffusion coefficient for carbon depends on the vertical diffusion coefficient for heat anomalies. As a result, uncertainty in oceanic carbon uptake is defined by the uncertainty in the heat uptake. In all simulations described in this paper we used $K_{vco} = 1$ cm/s and $r=0.52$ (Eq. 3). Uncertainty in the terrestrial carbon uptake was taken into account by varying the half-saturation constant, $k_{c}$ (Eq. 13). The results of CO$_2$-enrichment studies suggest that plant growth could increase by 24% - 54% in response to doubled CO$_2$ given adequate nutrients and water (Raich et al. 1991; McGuire et al. 1992; Gunderson and Wullschleger 1994; Curtis and Wang 1998; Norby et al. 1999, 2005). In the stand-alone TEM, a value of 400 ppmv CO$_2$ is chosen for the half-saturation constant, $k_{c}$, so that $f$(CO$_2$(t)) increases by 37% for a doubling of atmospheric CO$_2$ (e.g. McGuire et al. 1997; Pan et al. 1998). A 24% and 54% response to doubled CO$_2$ would correspond to $k_{c}$ values of 215 and 800 ppmv CO$_2$,
respectively. The sensitivity of plant uptake on CO₂ increase is defined not by the absolute value of \( f(CO_2(t)) \), which decreases with \( kc \), but by the ratio of \( f(CO_2(t)) \) to \( f(CO_2(0)) \) which increases with \( kc \).

Based on the comparison with other terrestrial carbon models (Sokolov et al., 2008) and results from stand-alone TEM simulations, in the TEM version implemented in the MESM different values of \( kc \) are used for nitrogen-limited and non-limited ecosystems. In simulations discussed below value of \( kc \) was varied from 200 to 800 ppmv CO₂ for nitrogen-limited ecosystems and from 75 to 300 ppmv CO₂ for ecosystems with no nitrogen limitations.

Total carbon uptake can be estimated from available data on carbon emissions and observed CO₂ concentrations and is less uncertain than carbon uptake by the ocean and carbon uptake by terrestrial ecosystem separately. To take this into account low values of \( K_v \) (small carbon uptake by the ocean) are coupled with high values of \( kc \) (large terrestrial carbon uptake).

Figure 2b shows frequency distributions for TCR from 400 simulations. The median value for TCR (1.7 K) is close to that for CMIP5 model (1.8K), while the 90% probability interval (1.4 – 2.0K) is significantly narrower than estimates based on CMIP5 models (1.2 – 2.4K). The relatively small range of TCR in our simulations is explained in part by the correlation between climate sensitivity and the rate of oceanic heat uptake imposed by observations (Figure 3). Another characteristic often used to describe transient model response to forcing is a “realized warming” defined as a ratio of equilibrium climate sensitivity to TCR. In our simulations, this characteristic ranges from 0.35 to 0.66 (90% interval) with median value of 0.54. The fact that these values are smaller than corresponding values for CMIP5 model (0.46-0.72 and 0.58) indicates that the rate of oceanic heat uptake in CMIP5 models is most likely smaller than in our simulations.

The transient climate response to cumulative carbon emission (TCRE), is defined as the ratio of surface warming to cumulative implied carbon emissions at the time of CO₂ doubling from simulations with a 1% per year increase in CO₂ concentrations (Matthews et al. 2009). Values of TCRE in MESM simulations vary (90% range) from 1.2 to 1.9 K/ EgC (Figure 2c). According to Gillett et al., (2013), a similar range for CMIP5 models is 0.8-2.4K/EgC. At the same time observationally constrained range, obtained using CMIP5 simulations and observed temperature to 2010 is 0.7–2.0K/ EgC (Gillett et al., 2013).
To evaluate uncertainty in long-term climate system response implied by the distribution of climate parameters described above, we carried out an ensemble of simulations using RCP8.5 GHGs concentrations. Comparison between the results from this ensemble and those from the multi-model CMIP5 ensemble are presented in Table 2. The MESM simulates less surface warming during the 21st century than the CMIP5 ensemble which, in part, may be explained by the fact that most of CMIP5 models overestimate warming in the first decade of 21st century. At the same time, MESM simulates stronger temperature increase during 22nd and 23rd centuries than CMIP5 models. It should be kept in mind that from 39 CMIP5 models that ran the RCP8.5 simulation only 12 were run beyond 2100. As can be seen on Figure 12.5 of IPCC 2013, in 2100 multi-model mean surface warming is smaller for these 12 models than for all 39 models. Use of different number of CMIP5 models in different simulations somewhat complicates comparison between CMIP5 and MESM results. Shown above estimates for equilibrium climate sensitivity and TCR are from simulation with 23 and 30 CMIP5 models, respectively (Table 9.5 in IPCC 2013). TCRE estimates are based on the results of 15 CMIP5 ESMs (Gillett et al., 2013).

<table>
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<th>MESM</th>
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<td>1.7 (1.3,2.0)</td>
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<td>6.5 (3.3, 9.8)</td>
<td>7.0 (5.7,8.9)</td>
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<td>2281–2300</td>
<td>7.8 (3.0, 12.6)</td>
<td>8.9 (6.9,11.0)</td>
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</tbody>
</table>

Table 2. Median values and 90% probability intervals for surface air temperature anomaly relative to 1986-2005 mean in simulations with RCP8.5 scenario. CMIP5 results are from Table 12.2 of IPCC (2013).

3.2 Historical Climate Change

To assess the quality of the probability distributions for climate model parameters, we carried out 400 historical simulations from 1861 to 2010. The model reproduces the observed changes in surface air temperature very well (Figure 4a). Temperature averaged over the first decade of the 21st century increases in our simulations relative to the 1861-1880 mean between 0.67 and 1K (90% probability interval) with a mean value of 0.835K. Similar to the TCR, range of temperature change simulated by MESM is significantly narrower than one produced by CMIP5 models. The MESM ensemble mean agrees better with observation than the CMIP5 mean, especially after year 2000, when CMIP5 models overestimate observed warming. Changes in
global mean precipitation simulated by MESM (Figure 4b) also agree well with observations, especially the increase in the last 50 years, and improves upon the simulations by the CMIP5 models. However, inter-annual variability simulated by MESM is much smaller than observed. The MESM simulates a larger increase in the ocean heat content both in the top 700m and 2000m (Figure 5) compared with observations and CMIP5 models (Figure 9.17 in IPCC 2013). At the same time, sea level increases due to thermal expansion at the rate of 0.85 (0.625-1.4) mm/yr between 1971 and 2010 and at the rate of 1.2 (0.96-1.6) mm/yr between 1993 and 2010. These trends are not too different from the estimates given by IPCC 2013: 0.8(0.5 to1.1) mm/yr and 1.1(0.8 to 1.4) mm/yr, respectively (Table 3.1 in IPCC 2013).

As indicated below in the emissions-driven simulations CO₂ emissions associated with agricultural activity are provided by the EPPA model, therefore the terrestrial ecosystem model calculates carbon fluxes using potential land cover distribution. Nonetheless, terrestrial carbon uptake generally falls within the range of the Global Carbon Project multi-model analysis (Le Quere et al., 2016), while being slightly smaller than the Global Carbon Project’s estimate obtained as the residual from the global carbon budget (figure 6a). The median value of the terrestrial carbon uptake averaged over 2000-2009 is about 0.25 GtC/y smaller than the best estimate provided by IPCC 2013 (Figure 7a). This can be partially attributed to the fact that nitrogen deposition is not taken into account. Simulated uncertainties in terrestrial uptake are also smaller than those suggested by IPCC 2013. Ensemble mean carbon uptake by the ocean (Figure 6b) agrees very well with data from the Global Carbon Project (Le Quere et al., 2016). However, the range on oceanic carbon obtained in our simulations is slightly shifted towards high values compare to IPCC estimate (Figure 7b).

3.3 Present-day Climate
3.3.1 Meteorological Variables

In this section, we compare annual mean data from out simulations averaged over 1991-2010 period with available observations and results from CMIP5 (Taylor et al., 2011) simulations. While simulating really well changes in the global mean temperature and precipitation (Figure 4), MESM simulations have some difficulties matching observed zonal distributions for present-day climate. MESM simulates somewhat lower temperature in the southern hemisphere and noticeably higher temperature in the mid-latitudes in the northern hemisphere (Figure 8a).
MESM also overestimates precipitation in the descending branch of mean meridional circulation in the northern hemisphere and underestimate precipitation in the mid-latitude storm track regions (Figure 8b), which shows the limitations of using a zonal-mean atmosphere model. The MESM realistically simulates the general characteristics of both surface latent and sensible heat fluxes (Figure 8c,d) and their latitudinal distributions fit within the range of the CMIP5 state-of-the-art climate except over a few latitude bands, specifically 20S-Equator and 20N-35N for latent heat flux and 90S-80S and 65-75N for sensible heat flux. Latitude-height cross sections of annual mean temperature, specific humidity and relative humidity are shown in Figure 9. The MESM is able to reproduce the overall latitudinal and vertical distributions of temperature and humidity generally well despite a cold bias in the polar regions, the maximum specific humidity in the equatorial regions not extending into the upper troposphere and high relative humidity values, especially in the mid-latitudes and polar regions.

3.3.2 Terrestrial Water Cycles

Global precipitation over land constitutes a substantial segment of the terrestrial water cycle and strongly influences the carbon and nutrient cycles tracked by TEM. As such, we compared MESM to the observationally-based estimate from the Global Precipitation Climatology Project (GPCP, Adler et al. 2003 and Adler et al., 2012). We find that the MESM estimate of land-only precipitation depicts the key seasonal and zonal attributes to a level that is very comparable with the CMIP5 models (Figure 10). In a global basis (Table 3), MESM is closely aligned with the GPCP estimate and also is centrally placed across the estimates of the CMIP5 models (equally biased high and low across each of these climate models). Some notable discrepancies are that land-only precipitation is biased low during the NH summertime midlatitude regions. This deficiency in summertime precipitation contributes correspondingly to slightly lower evapotranspiration rates compared to most of the CMIP5 models (Figure 11 and Table 3), although MESM is still within the range of CMIP models. For total runoff, MESM produces more runoff on the global scale than the CMIP5 models (Table 3), although the predominant seasonal and latitudinal features are preserved (Figure 12). As previously discussed in Section 2.4, the modification of the soil-infiltration scheme (i.e. removal of enhanced hydraulic conductivity under excessive dry soil conditions) was of considerable benefit to the performance of evapotranspiration and runoff rates shown here.
3.3.3 Ecosystem Productivity and Natural Emissions of Trace Gases

**Net Ecosystem Productivity:** The results from the key water and energy fluxes of the land system provide a first-order expectation to the climatological behavior of the terrestrial carbon cycle within MESM. As in previous evaluations of the land systems implemented within this Earth-system model framework (Schlosser et al., 2007) we focus on the net exchange of carbon between the land and the atmosphere, which represents a key coupling. Further, the TEM model is commonly used in a “standalone” configuration to simulate historical conditions and thus driven by observed atmospheric conditions (e.g. Zhu et al., 2011), and previous evaluations have used this as baseline for TEM reduced-form configuration with the MESM framework (e.g. Schlosser et al., 2007). We extend this approach by considering the net ecosystem productivity (NEP) of TEM on a month versus latitude projection (Figure 13). We find that the implementation of TEM within the MESM structure preserves all of the salient seasonal and latitudinal attributes as seen in previous evaluations (Figure 19 of Schlosser et al., 2007), and the characteristics of these patterns are consistent and corroborated by recent and detailed simulations with TEM in standalone configuration (e.g. Chen et al., 2011; Zhu and Zhuang, 2013; and Lu et al., 2015). Overall, terrestrial ecosystems represented a net uptake of carbon both globally as well as across all latitudes through our historical period of evaluation (1981-2005). In addition, the strongest (annual) sinks of carbon are found to be across the boreal latitude bands are comprised of extensive forest cover. An additional relative peak is also seen across the southern sub-tropical latitudes and is comparable in magnitude to the sink produced by northern midlatitude ecosystems. Although these areas are strong sinks – they carry a distinct seasonality and serve as considerable carbon sources during the late Fall through early Spring months.

**Methane Emissions:** In keeping with our overall approach to evaluate the performance of the land biogeochemical fluxes, we have evaluated the methane emissions against previous evaluations with models that contain similar core parameterization recipes but also a recent multi-model assessment (the WETCHIMP study of Melton et al., 2013) with distinct structural differences in overall design as well as implementation of the participating models. In order to provide the most consistent comparative evaluation in this regard, we focus on the historical period of 1993-2004. We find that for nearly all latitudes, the annual emission of methane from
MESM falls within the range of the multi-model assessment (Figure 14). The most notable exception is found at boreal latitudes with the MESM estimate well above the upper bound of the WETCHIMP range. However, one notable feature of this multi-model assessment is that it did not contain a model participant with TEM as its core ecosystem model that MESM employs. Looking at model-based studies that have used TEM as the core ecosystem model (e.g. Zhuang et al., 2004) as well as an observation-based artificial neural-network method (Zhu et al., 2013) to estimate total methane emissions in boreal latitudes (north of 45˚N), they indicate values on the order of 44 to 54 Tg CH₄/yr. The methane emission rates from the MESM historical simulation provide a more consistent latitude profile of emission across the boreal zone in this regard. Given that the higher boreal emission rates are more closely aligned to the observation-based result, we are confident that the MESM estimate is providing not only a result that is representative of the core ecosystem model behavior (i.e. TEM) but also a value that is empirically defensible.

**Nitrous Oxide:** In a similar vein to our evaluation of NEP, we find that the MESM historical simulation and a standalone version of CLM coupled to a N₂O emissions model (Saikawa et al., 2009) share common features but also exhibit distinct differences. It is important, however, to note that this model-to-model comparison cannot provide any judgement on the fidelity or veracity of either model. Rather, this is an evaluation of the sensitivity to the configuration and application of the soil N₂O emissions module (a variant of the DNDC model) within and Earth-system model framework. As such, both of the models exhibit consistent latitudinal locations of relative maximum emission rates (Figures 15 and 16) – which occur along the equator as well as during the summer season in both the northern and southern sub-tropical bands. Consistent across all these simulations is that the tropics provide the strongest annual emission source. While both models provide an additional source of emissions across the mid and higher northern latitude bands, a distinct difference is seen in the seasonality of this feature. The CLMCN-N2O model exhibits a distinct summertime peak in emissions that is coincident and widespread across the middle and into higher latitudes. In the MESM zonal configuration – the extent and summer timing of the peak is aligned with the CLMCN-N2O estimate between 45-75˚N, however for latitudes 25-45˚N there is an earlier onset and terminus of this feature. Through the course of a number of variants and sensitivity simulations with the IGSM, we have identified that the most likely culprit to this behavior lies within the MESM simulation of land precipitation. When
compared to an observationally-based estimate (GPCP), we find that the MESM estimate of precipitation is biased low during the summertime across the corresponding latitude bands (Figure 10). In conjunction with low evapotranspiration rates during the spring (not shown) – soil moisture stores become elevated at these latitudes and trigger the soil anaerobic conditions necessary for denitrification, and thus leads to the earlier emissions peak. However, by the beginning of the summer – elevated evapotranspiration levels combined with a precipitation deficit support dry soils and the emissions processes in MESM become dormant. Nevertheless, we have made salient progress from our earlier implementation (Schlosser et al., 2007) in providing a more consistent depiction of soil N₂O emissions with a reduced form of the model compared to its more detailed counterpart. The scientific community has recently recognized (Butterbach-Bahl et al., 2013) key areas for improvement and will continue to make necessary strides in the detailed modeling of nitrous oxide emissions and verification studies. As such, this strategy will continue to be a critical element of our model development, particularly with regard to the N₂O emissions component.

**Table 3.** Summary statistics for simulated fluxes in the land water/energy cycles. Shown are correlations and bias (in mm/day) of precipitation, evapotranspiration, and runoff. The correlations are performed on the month vs. latitude projections shown and discussed. For precipitation, all model simulations are judged against the Global Precipitation Climatology Project (GPCP) observations. For evaporation and runoff, these metrics are performed for the CMIP5 models against the MESM result in order to convey the degree of consistency between the more complex CMIP models to MESM’s intermediate complexity. Note that for the ACCESS1-3 outputs of total runoff were not made publicly available (N/A). Sign convention for bias is positive when CMIP5 value exceeds MESM or model exceeds observation.

<table>
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### 3.4 Emissions-driven Projections of Future Climate

In the simulations discussed below the MESM was forced by anthropogenic emissions calculated by the EPPA model, including carbon emissions associated with land use. MESM also takes into account natural emissions of CH$_4$ and N$_2$O calculated by NEM.

Wang et al., (1998) describe a tropospheric chemistry and transport scheme for chemical tracer used in the MESM. In the framework of the Aviation Climate Change Research Initiative (ACCRI, Brasseur et al., 2016), MESM was used in a tracer transport comparison exercise. In these simulations, the ACCRI 2006 fuel burn inventory (Barret et al., 2010) above 8km were used as a proxy for tracer emissions. Simulations were carried out for six months starting in January and July with tracers being released in the atmosphere during the first month only. Below we show comparisons between results obtained in simulations with the MESM and GEOS-Chem models for three different types of tracers: an inert tracer (no losses), a tropospheric ozone-like tracer with a prescribed a 21-day e-folding lifetime and a tracer with a dry deposition removal process. Results are shown only for simulations started in January because results of simulations started in July look very similar. The total mass of the tracer with dry deposition decreases slightly faster in simulations with MESM than with GEOS-Chem, while the changes for the other two tracers are practically identical in both model simulations (Figure 17). Figure 18 shows latitude- pressure distributions for the ozone-like tracer for the first, third and sixth months of the simulations. Because the total mass of the tracer decreases exponentially, data for the third and sixth months were multiplied by factors of $10$ and $10^4$, respectively. In general, the study of the impact of aviation emissions on atmospheric chemical composition and

<table>
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<td>0.95</td>
<td>0.20</td>
<td>0.87</td>
<td>0.74</td>
<td>0.70</td>
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</table>
climate showed that MESM results lie well within the envelope of the more complex 3-D chemistry-climate models (Olsen et al., 2013).

Our simulations with interactive chemistry start in year 2006 as a continuation of an historical simulation and future projections are then usually given to 2100. Here, however, we will concentrate on short-term simulations (2006-2015) when model mixing ratios using model emissions can be compared with observed mixing ratios.

As can be seen, the MESM simulates global concentrations of the 3 major long-lived greenhouse gases rather well (figure 19). Figure 20 plots tropospheric mixing ratios for some climate-relevant species (O₃, CH₄, CO, SO₂, NO and NO₂) as well as the very short-lived OH species from the MESM output as well as the zonal average for the ACCMIP archived version of the CESM CAM-Chem, with CAM version 3.5 (available at http://browse.ceda.ac.uk/browse/badc/accmip). The CESM output has 26 vertical levels, which have been regridded to match the 11-level MESM output in Figure 20. In general, the MESM results are comparable to the CESM data, with the vertical and latitudinal distributions of O₃, CH₄, and OH in fairly close agreement. The MESM does not have the hemispheric asymmetries for CO and NO₂ that are shown in the CESM data, and the MESM vertical distribution of SO₂ peaks at the third level, as a result of SO₂ emissions being distributed evenly in the bottom two layers. In addition, in contrast to most chemistry models (including the CESM-CAM-Chem), MESM does not use prescribed surface concentrations as low boundary conditions.

Radiative forcing and simulated surface air temperature (not shown) are almost identical in historical (concentration-driven) and emissions-driven simulations. In general, simulated climates are very similar during overlapping period of two simulations (2006-2010).
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Figure 1. MIT Earth System Model of Intermediate Complexity (MESM)
Figure 2. Frequency distribution of (a) ECS, (b) TCR and (b) TCRE. Vertical line shows median value and horizontal bar shows 90% probability interval. Red line CMIP5 estimate, from Table 9.5 of IPCC (2013) for TCR and from Gillett et al., (2013) for TCRE.
Figure 3. Distribution of climate sensitivity and the rate of ocean heat uptake (square root of vertical diffusion coefficient). Red dots show values of CS and SQRT(Kv) for 400 samples. The contour lines are for the 5, 10, 25, 50, 75, 90 and 95% percentiles.
Figure 4. Time series of global mean surface air temperature and precipitation relative to 1861-1880 mean. The simulations with the MESM are shown in blue (ensemble mean and one standard deviation in shading) and the simulations from the CMIP5 multi-model ensemble are shown in red (multi-model ensemble mean and one standard deviation in shading). After 2006, simulations under the RCP8.5 are used. Observations are shown in black lines, namely the HadCRUT4 (Morice et al., 2012) and global reconstructed precipitation (REC) data (Smith et al., 2012) for temperature and precipitation respectively.
Figure 5. Changes in ocean heat content relative to 1971 in the top 700m (a) and top 2000 m (b). Red lines are observations from Levitus et al., (2012). Blue lines are ensemble means.
Figure 6. (a) Terrestrial and (b) oceanic carbon uptake. Data for comparison are from Le Quere et al (2016).
Figure 7. Frequency distribution of (a) terrestrial and (b) oceanic carbon uptake averaged over years 2000-2009. The black line represents the 90% interval from the MESM ensemble. The red line represents the AR5 estimate (Table 6.1 IPCC 2013).
Figure 8. Zonal distribution of (a) surface air temperature (in °C), (b) precipitation (in mm/day), (c) surface latent heat flux (in W/m²), and (d) surface sensible heat flux (W/m²) averaged over the 1991-2010 period. The MESM simulation with the median values of climate parameters (climate sensitivity, ocean heat uptake rate and net aerosol forcing) is shown in blue. Simulations from the CMIP5 multi-model ensemble are shown in red (multi-model ensemble mean and full range in shading). After 2006, simulations under the RCP8.5 are used. Observations are shown in black lines, namely the HadCRUT4 (Morice et al., 2012), GPCP v2.3 (Adler et al. 2003) and MERRA2 (Gelaro et al., 2017) for temperature, precipitation and the heat fluxes respectively.
Figure 9 Latitude-height cross section of (a) temperature (in °C), (b) specific humidity (in kg/kg), and (c) relative humidity (in %) averaged over the 1991-2010 period. Observations from MERRA2 (Gelaro et al., 2017) are shown on the left panels, the CMIP5 multi-model ensemble mean is shown in the middle panels and the MESM simulation with the median values of climate parameters (climate sensitivity, ocean heat uptake rate and net aerosol forcing) is shown in the right panels. After 2006, CMIP5 simulations under the RCP8.5 are used.
Figure 10. Month vs. latitude profiles of precipitation (mm/day) given by observations from the Global Climatology Precipitation Project (GPCP, upper left); MESM (upper right panel), as well as two simulations from the Coupled Model Intercomparison Project Phase 5 (CMIP5, lower panels). In each of the model-result panels, the correlation of its month vs. latitude profile to that of GPCP is given. The CMIP5 results show the pattern with the highest (lower left) and lowest (lower right) pattern correlation with the GPCP observations.
Figure 11. Month vs. latitude profiles of evapotranspiration (mm/day) given by observations from the MESM (upper right panel), as well as two simulations from the Coupled Model Intercomparison Project Phase 5 (CMIP5, lower panels). The CMIP5 results show the pattern with the highest (lower left) and lowest (lower right) pattern correlation with MESM. Also shown is the simulation from CCSM4 (upper left) panel.
Figure 12. Month vs. latitude profiles of runoff (mm/day) given by observations from the MESM (upper right panel), as well as two simulations from the Coupled Model Intercomparison Project Phase 5 (CMIP5, lower panels). The CMIP5 results show the pattern with the highest (lower left) and lowest (lower right) pattern correlation with MESM. Also shown is the simulation from CCSM4 (upper left panel).
Figure 13. Simulations of emissions of net ecosystem productivity (NEP) by the MIT Earth-System Model (MESM) within the IGSM framework. Top panel displays the month versus latitude results averaged over 1981-2005, and the bottom panel shows the corresponding annual fluxes by latitude. Units are in 10^9 kg-C/month and 10^9 kg-C/year, respectively. Shown also in the bottom panel with the shaded red area is the multi-model range from five of the CMIP5 Earth-system models that provided data from historical simulations that cover the evaluation period.
Figure 14. Simulations of emissions of methane by the MIT Earth-System Model (MESM) within the IGSM framework. Top panel displays the month versus latitude results (Units in Tg-CH4/month) averaged over 1993-2004, and the bottom panel shows the corresponding annual fluxes by latitude (units in Tg-CH4/year). Shown also in the bottom panel with the shaded red area is the multi-model range from the results of WETCHIMP with. More recent studies (not shown in figure) indicate that the excess in emissions from MESM at high northern latitudes is credible (see corresponding text for details).
Figure 15. Simulations of soil emissions of nitrous oxide by the CLMCN-N2O model (Saikawa et al., 2013) forced by two different meteorological datasets (CAS and GMFD). Shown are month versus latitude results averaged over 1981-2005. Units are in 108 kg-N2O/month.
Figure 16. Simulations of soil emissions of nitrous oxide by the MIT Earth-System Model (MESM) within the IGSM framework. The top panel shows month versus latitude results averaged over 1981-2005 with units in 10^8 kg-N2O/month. The bottom panel provides the annual emission rates (10^8 kg-N2O/year) for the corresponding latitudes of the top panel. Shown also in the bottom panel with the shaded red area is a multi-estimate range based on the results from Saikawa et al. (2011) and Hashimoto (2012).
Figure 17. Time evolutions of total mass ($10^9$ kg) for inert ozone-like and tracer with dry deposition in simulations with January emissions with MIT MESM (solid lines) and GEOS-Chem (dashed lines)
Figure 18. Latitude-pressure distributions of ozone-like tracer concentration (ppbm) in simulations with January emissions for first (top), third (middle) and sixth (bottom) months of simulation. Left column - results obtained with the MIT MESM. Right column – result from GEOS-Chem.
Figure 19. Observed (red) and projected (black) concentrations of CO$_2$ (left), CH$_4$ (middle) and N$_2$O (right).
Figure 20. Present-day simulated zonal output from the MESM (left) and zonally averaged output from the CESM CAM-Chem, CAM version 3.5 archived data from the ACCMIP archive (right) for O$_3$, CH$_4$, CO, SO$_2$, OH, and NO$_2$. The CAM data has been regridded to match the MESM levels, and only data below the tropopause (here defined as 150 ppbv O3) is shown.