

The International Land Model Benchmarking (ILAMB) System: Design, Theory, and Implementation

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Abstract

The increasing complexity of Earth system models (ESMs) has inspired efforts to quantitatively assess model fidelity through rigorous comparison with best-available measurements and observational data products. ESMs exhibit a high degree of spread in predictions of land biogeochemistry, biogeophysics, and hydrology, which are sensitive to forcing from other model components. Based on insights from prior land model evaluation studies and community workshops, the authors developed an open source model benchmarking software package that generates graphical diagnostics and scores model performance in support of the International Land Model Benchmarking (ILAMB) project. Employing a suite of *in situ*, remote sensing, and reanalysis datasets, the ILAMB package performs comprehensive model assessment across a wide range of land variables and generates a hierarchical set of webpages containing statistical analyses and figures designed to provide the user insights into strengths and weaknesses of multiple models or model versions. Described here is the benchmarking philosophy and mathematical methodology embodied in the most recent implementation of the ILAMB package. Comparison methods unique to a few specific datasets are presented, and guidelines for configuring an ILAMB analysis and interpreting resulting model performance scores are discussed. ILAMB is being adopted by modeling teams and centers during model development and for model intercomparison projects, and community engagement is sought for extending evaluation metrics and adding new observational datasets to the benchmarking framework.

1 Introduction

As Earth system models (ESMs) become increasingly complex and observational data volumes rapidly expand, there is a growing need for comprehensive and multi-faceted evaluation of model fidelity. Process-rich ESMs pose challenges to developers implementing new parameterizations or tuning process representations, and to the broader community seeking information about the skill of model predictions. Model developers and software engineers require a systematic means for evaluating changes in model results to ensure that developments improve the scientific performance of target process representations while not adversely affecting results in other, possibly less familiar, parts of the model. To advance understanding and predictability of terrestrial biogeochemical processes and their interactions with hydrology and climate under conditions of increasing atmospheric carbon dioxide, rigorous analysis methods, employing best-available observational data, are required to objectively assess and constrain model predictions, inform model development, and identify needed measurements and field experiments (*Hoffman et al.*, 2017).

Building upon past model evaluation work (*Randerson et al.*, 2009), we developed an extensible model benchmarking package in support of the goals of the International Land Model Benchmarking (ILAMB; <https://www.ilamb.org/>) activity. ILAMB's goals are to

1. develop internationally accepted benchmarks for land model performance by drawing upon international expertise and collaboration;
2. promote the use of these benchmarks by the international community for model intercomparison and development;
3. strengthen linkages among experimental, remote sensing, and climate modeling communities in the design of new model tests, benchmarks, and measurement programs; and
4. support the design and development of a new, open source, benchmarking software system for use by the international community.

Three ILAMB workshops have been held—in Exeter, United Kingdom, in 2009; Irvine, California, United States, in 2011 (Luo *et al.*, 2012); and Washington, DC, United States, in 2016 (Hoffman *et al.*, 2017)—to engage the modeling, measurements, and remote sensing communities in the identification of observational datasets and the design of model evaluation metrics. In this way, community consensus was sought for the curation of observational data and the methodology of model evaluation and scoring, which are described below.

Recognition that the capacities of the terrestrial and marine biosphere to store anthropogenic carbon will weaken under climate warming (Cox *et al.*, 2000; Friedlingstein *et al.*, 2001; Fung *et al.*, 2005; Denman *et al.*, 2007; Randerson *et al.*, 2015; Mahowald *et al.*, 2017; Moore *et al.*, 2018) and that uncertainties in carbon cycle feedbacks must be quantified and reduced to improve projections of future climate change (Friedlingstein *et al.*, 2006; Gregory *et al.*, 2009; Arora *et al.*, 2013; Ciais *et al.*, 2013; Friedlingstein *et al.*, 2014; Hoffman *et al.*, 2014) has inspired efforts to quantitatively evaluate model performance through comparison with *in situ* and remote sensing observations (Anav *et al.*, 2013; Eyring *et al.*, 2016). Multi-model simulation results from the third Coupled Model Intercomparison Project (CMIP3; Meehl *et al.*, 2007) and fifth Coupled Model Intercomparison Project (CMIP5; Taylor *et al.*, 2012), which informed the Intergovernmental Panel on Climate Change (IPCC) Fourth and Fifth Assessment Reports (AR4 and AR5), provided opportunities for developing and testing model evaluation diagnostics, formal metrics, and exploration of benchmarking concepts and techniques. Early work on coupled model evaluation and establishing formal metrics focused primarily on atmospheric variables (Reichler and Kim, 2008; Gleckler *et al.*, 2008). Following the first two ILAMB workshops, the land modeling community began exploring standardized and comprehensive benchmarking for terrestrial carbon cycle models (Cadule *et al.*, 2010; Blyth *et al.*, 2011; Abramowitz, 2012; Kelley *et al.*, 2013; Dalmonch and Zaehle, 2013; Piao *et al.*, 2013; Anav *et al.*, 2013; Bouskill *et al.*, 2014; Ghimire *et al.*, 2016). While some researchers define benchmarking as a series of model tests based on a pre-defined expected level of performance (Abramowitz, 2005; Best *et al.*, 2015), most of the systematic benchmarking strategies explored by the land modeling community to date do not depend upon the establishment of an expected level of performance.

The ILAMB software package, hereafter referred to as ILAMB, shares some of the same goals as existing model diagnostic and evaluation tools, such as the Protocol for the Analysis for Land Surface models (PALS; Abramowitz, 2012), the Program for Climate Model Diagnosis and Intercomparison (PCMDI) Metrics Package (PMP; Gleckler *et al.*, 2016), the Earth System Model Evaluation Tool (ESMValTool; Eyring *et al.*, 2016), the Land surface Verification Toolkit (LVT; Kumar *et al.*, 2012), and a wide variety of often custom-developed diagnostic packages in use at international modeling centers. Some of these tools provide model-to-model comparisons, a large collection of standalone graphical diagnostics, or workflow infrastructure that allows one to regenerate analysis results from previously published studies but with new model outputs. In contrast, ILAMB was designed to compare multiple models or model versions with observations simultaneously, assess functional relationships between prognostic variables and one or more forcing variables through variable-to-variable comparisons (e.g., gross primary production vs. precipitation), and score model performance across a suite of metrics, variables, and datasets. Model performance is evaluated for variables in categories of biogeochemistry (Table 2), hydrology (Table 3), radiation and energy (Table 4), and climate forcing (Table 5).

For every variable, ILAMB generates graphical diagnostics (spatial contour maps, time series line plots, and Taylor diagrams (Taylor, 2001)) and scores model performance for the period mean, bias, root mean squared error (RMSE), spatial distribution, interannual coefficient of variation, seasonal cycle, and long-term trend. Model

performance scores are calculated for each metric and variable and are scaled based on the degree of certainty of the observational dataset, the scale appropriateness, and the overall importance of the constraint or process to model predictions, following a customizable rubric described below (Table 1). Scores are aggregated across metrics and datasets, producing a single scalar score for each variable for every model or model version. As shown in Figure 1, these scalar scores are presented graphically. On the left side we use a stop-light color scheme to indicate aggregate performance for each model by variable. On the right, we show relative performance (i.e., Z-score), indicating which models or model versions perform better with respect to others contained in the overall analysis.

We do not view these aggregate absolute scores as a determinant of ‘good’ or ‘bad’ models. We envision the scores as a tool to more quickly identify relative differences among models and model versions which the scientist must then interpret. As in any evaluation methodology, many of our choices are subjective and must be considered as the scores are interpreted. Where possible, the ILAMB implementation allows for users to customize weights and diagnostics in order to incorporate aspects of model performance relevant to their scientific goals. ILAMB may be thought of as a framework which may be expanded to incorporate community ideas regarding model benchmarking. Thus while our choices are subjective, they are informed by the preferences of a larger community and can be considered as an initial suggestion.

The remainder of this paper describes the ILAMB methodology used to compute aggregate absolute scores. First we describe how we compare an individual observational dataset to model output (Section 2). Then we explain how scores are aggregated across datasets for each variable and present the datasets used in the land model evaluation (Section 3). In Section 4 we present some salient points about how the ILAMB software is designed. Finally, in Section 5 we discuss what ILAMB scores mean and how they should be used.

2 Methodology

In this section we describe the methodology used to assess how well a model captures information contained in a reference (e.g. observational) dataset. For the purposes of this section, we discuss the analysis of a generalized variable $v(t, \mathbf{x})$ which we assume represents a piecewise discontinuous function of constants in space and time. This means that the temporal domain, represented by the variable t , is defined by the beginning and ending of time intervals and the spatial domain, represented by the variable \mathbf{x} (bolded to emphasize it is a vector quantity), represents the areas created by cell boundaries or the areas associated with data sites. When necessary, we use the subscript ‘ref’ to reflect a variable whose source is a reference or observational dataset, and the subscript ‘mod’ for model datasets.

While many statistical quantities may be computed, the goal of our initial methodology is to examine the mean state and variability around the mean over monthly to decadal time scales and grid cell to global spatial scales. While we intend to uniformly apply this analysis procedure to all variables, we also implement a mechanism to skip certain aspects when deemed inappropriate. For example, if a reference dataset only contains average information across a span of years, the annual cycle is undefined and automatically skipped in our implementation. The implementation also allows users to skip aspects of the analysis that are deemed inappropriate even if it is possible to compute these metrics using the available data. For example, the interannual variability may be poorly characterized in a reference dataset even though the quantity could be computed.

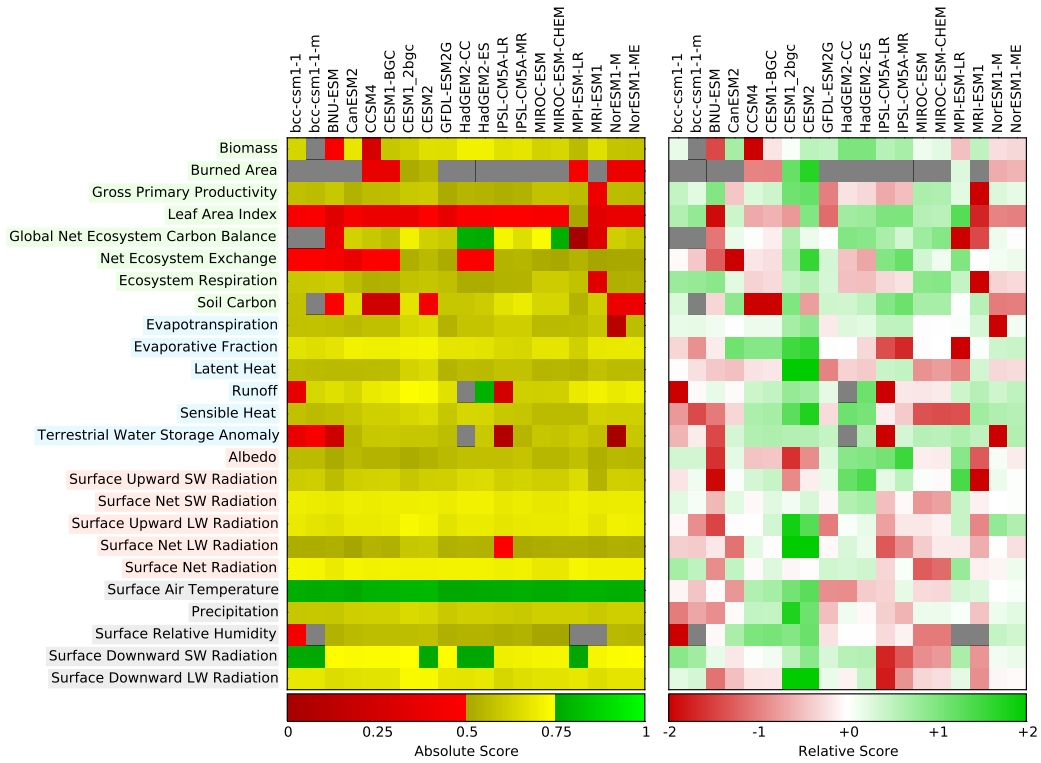


Figure 1: The ILAMB top-level graphic uses stop-light colors to show how different models or model versions (across the top) score with respect to each variable (down the left) in an absolute sense (left rectangle) and with respect to each other (right rectangle). Grey boxes reflect missing or unavailable data.

2.1 Preliminary Definitions

Before presenting the specifics of the ILAMB methodology, we first present some definitions used throughout the paper. While the following definitions are widely used in the community, there are many subtle choices in their implementation that affect the interpretation of the results. We present them here with precise meanings to emphasize where a choice has been made and our reasoning for making it.

2.1.1 Mean values over time

When calculating mean values over the time period of the benchmark dataset, denoted by a bar superscribing the variable, we use the midpoint quadrature rule to approximate the integral,

$$\begin{aligned}\bar{v}(\mathbf{x}) &= \frac{1}{t_f - t_0} \int_{t_0}^{t_f} v(t, \mathbf{x}) dt \\ &\approx \frac{1}{T(\mathbf{x})} \sum_{i=1}^n v(t_i, \mathbf{x}) \Delta t_i\end{aligned}\tag{1}$$

where n represents the number of time intervals on which v is defined between the initial time, t_0 , and the final time, t_f , and Δt_i is the size of the i^{th} time interval, modified to exclude time which falls outside of the integral limits,

$$\Delta t_i = \min(t_f, t_f^i) - \max(t_0, t_0^i)\tag{2}$$

where t_0^i and t_f^i are the initial and final time of each time interval. The average value is obtained by dividing through by the amount of time in the interval, $t_f - t_0$, replaced in our discrete approximation by the following function.

$$T(\mathbf{x}) = \sum_{i=1}^n \Delta t_i \text{ if } v(t_i, \mathbf{x}) \text{ is valid}\tag{3}$$

In words, Equation (3) addresses temporally discontinuous data by summing all the time step interval sizes only if the corresponding variable data is marked as valid. This means that if a function has some values masked or marked as invalid at some locations, we do not penalize the averaged value by including this as a time at which a value is expected. If an integral (or sum) is desired instead of an average, then we simply omit the division by $T(\mathbf{x})$ in Equation (1).

2.1.2 Mean values over space

When computing spatial means over various regions of interest, denoted by a double bar over a variable, we use the midpoint rule for integration to approximate the following weighted spatial integral,

$$\begin{aligned}\bar{\bar{v}}(t) &= \frac{1}{\int_{\Omega} w(\mathbf{x}) d\Omega} \int_{\Omega} v(t, \mathbf{x}) w(\mathbf{x}) d\Omega \\ &\approx \frac{1}{A(\Omega)} \sum_{i=1}^{n(\Omega)} v(t, \mathbf{x}_i) w(\mathbf{x}_i) a_i\end{aligned}\tag{4}$$

over a region Ω , also referred to as a area-weighted mean. Here the function $w(\mathbf{x})$ is an optional generic weighting function defined over space. The summation is over $n(\Omega)$, that is the integer number of spatial cells whose centroids fall into the region of interest. A function evaluation at a location \mathbf{x}_i refers to the constant value which corresponds to that spatial cell. The value of a_i is the area of the cell, which could be some fraction of the total cell area if integrating over land in coastal regions. We then divide through by the measure, the sum of the grid areas with the weights,

$$A(\Omega) = \sum_{i=1}^{n(\Omega)} w(\mathbf{x}_i) a_i \quad \text{if } v(t, \mathbf{x}_i) \text{ is valid} \quad (5)$$

Note that if no weighting is required, this is a normalization by the sum of the area over which we integrate. As with the temporal mean, if an integral only is required, we simply omit the division by $A(\Omega)$. In cases where a mean over a collection of sites is needed, the spatial integral reduces to an arithmetic mean across the sites.

If we are spatially integrating a variable from a single source, then its spatial grid is clearly defined and Equation (4) can be directly applied to compute the quantity of interest. However, if the integrand involves quantities from two different sources, as in computing the global bias or RMSE, then there is likely a disparity in both resolution and representation of land areas. We address resolution differences by interpolating both sources to a grid composed of the cell breaks, the location at which two neighboring cells meet, of both data sources. Consider two spatial grids whose cells are defined by the outer product of 1D vectors representing the cell breaks in spherical coordinates,

$$\mathcal{G}_1 := \theta_1 \otimes \varphi_1 \quad (6)$$

$$\mathcal{G}_2 := \theta_2 \otimes \varphi_2 \quad (7)$$

where θ refers to the latitude, φ to longitude, and \otimes a operator which creates a two-dimensional grid from one-dimensional vectors. We address differences in resolution by defining a composite grid which consists of the outer product of the union of these two grids' cell breaks,

$$\mathcal{G}_c := (\theta_1 \cup \theta_2) \otimes (\varphi_1 \cup \varphi_2). \quad (8)$$

Once constructed, quantities defined on both \mathcal{G}_1 and \mathcal{G}_2 may be interpolated to \mathcal{G}_c by nearest neighbor interpolation with zero interpolation error due to the nested nature of the grids. This can be seen visually by comparing the three plots shown in Figure 2(a). In each plot, the tick marks along the x-axis represent the cell breaks of the particular one-dimensional grid left coarse for illustration. The cyan curve represents a step function defined on the grid of a reference dataset \mathcal{G}_1 and the magenta curve on that of the model dataset \mathcal{G}_2 . Both are interpolated to the composed grid \mathcal{G}_c without loss of information, albeit on a new grid containing more cells of variable size. Once on a composite grid, the quantities may be compared directly. As the ILAMB methodology has been envisioned for comparisons with model output from CMIP5, we have made an implicit assumption that each source grid, \mathcal{G}_1 and \mathcal{G}_2 , is regular and can be represented by one-dimensional vectors. While the implementation does provide naive interpolation for non-regular grids, the user is encouraged to employ a conservative interpolation scheme of their choosing prior to applying the ILAMB methodology.

In addition to resolution differences, we observe that data sources vary in the underlying representation of the distinction between land and water. We illustrate

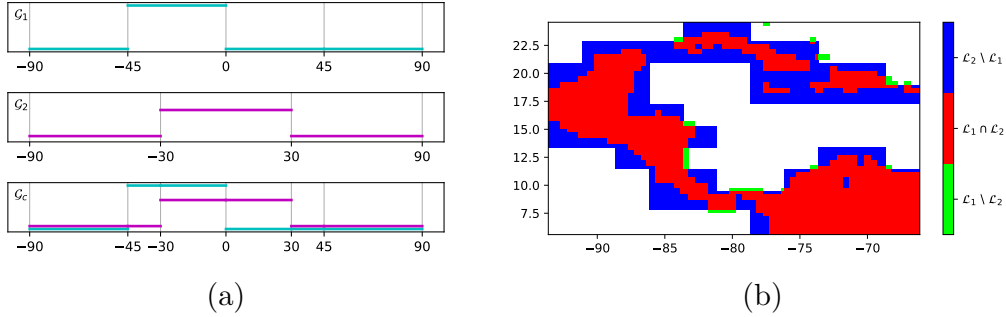


Figure 2: When comparing two spatial variables of varying resolution, we interpolate both to a common grid composed of the cell breaks of both variables over the intersection of what both variables agree is land. (a) Interpolation of sample step functions defined on grids \mathcal{G}_1 and \mathcal{G}_2 both interpolated to a composite grid \mathcal{G}_c using nearest neighbor interpolation with zero interpolation error. The vertical grid lines reflect the cell boundaries in each grid., (b) Differences in the representation of land from a reference and model dataset zoomed into Central America for emphasis. The red region represents where both sources are in agreement, the blue is land for the model but not the reference and the green is land for the reference but not the model.

this concept in Figure 2(b) where we compare a fine scale representation of land \mathcal{L}_1 to a relatively coarse representation \mathcal{L}_2 . This is a typical situation encountered when comparing high resolution observational data to lower resolution model output. The red region represents the intersection of land areas $\mathcal{L}_1 \cap \mathcal{L}_2$, that is, where both sources report the presence of land. However, there are missed land areas from both sources, represented by the blue and green colors. As much of the disagreement over what is considered land occurs around islands in tropical regions (for example Central America and Equatorial Asia), these non-represented areas can constitute a nontrivial percentage of the total represented variable v .

For transparency, the ILAMB implementation is built with the capability of reporting integrals over each of these three land areas. Unless specifically stated otherwise, when spatially integrating a quantity from a single source, we use the original grid and land areas given by that source. This is to remain as true to the original intent of the provider as we can. However, when comparing two data sources of varying resolution and land representation, we perform this integration over what both report to be land, $\mathcal{L}_1 \cap \mathcal{L}_2$ (the red area in Figure 2(b)).

2.1.3 Computing normalized scores from errors

In the following sections 2.2 and 2.3, we detail how we compute errors and transform them into normalized scores on the unit interval. This approach is intended to synthesize model performance across a range of dimensions with respect to a given dataset. We achieve this by taking a measure of the relative error, generically represented here as ε , and passing it through the exponential function,

$$s = e^{-\alpha\varepsilon} \quad (9)$$

where s is a score on the interval $[0, 1]$ and α is a parameter which can be used to tune the mapping of error to score. The classic expression of relative error is prone to numerical instabilities for denominator values near or which cross zero. Furthermore

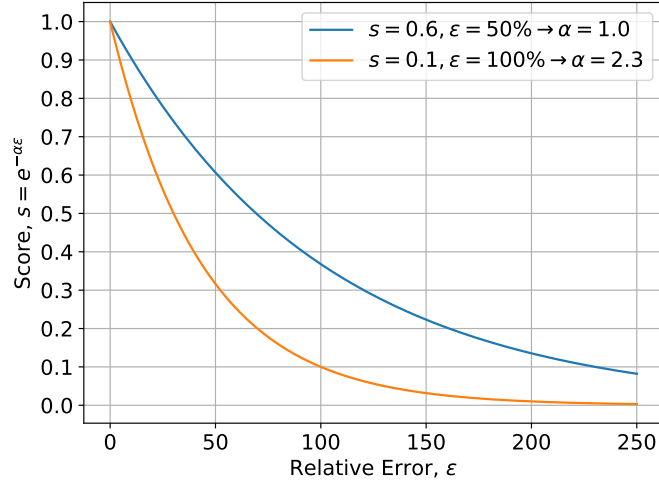


Figure 3: Mapping function of relative error ϵ to a score s on the unit interval. Two choices of α are shown: $\alpha = 1$, shown in blue, which equates a score of 0.6 to a relative error of 50%, and $\alpha = 2.3$, shown in orange, which equates a score of 0.1 to a relative error of 100%.

the magnitude of the error can depend on the units selected. For this reason we depart from the standard definition of relative error and develop specialized expressions in Equations (13, 18, 26).

While the choice of the exponential function is arbitrary, it was chosen because it maps zero error to a score of one and smoothly reduces the score as the error grows, never reaching exactly zero. This is important as we want to improve the score when the error improves, no matter how large of error we observe. If the user wants a relative error of $\hat{\epsilon}$ to equate to a score of \hat{s} , then

$$\alpha = -\frac{\ln(\hat{s})}{\hat{\epsilon}} \quad (10)$$

In Figure 3 we plot this function with two choices for α , which illustrates how the relative error may be controlled. Unless stated otherwise, we use an implicit $\alpha = 1$ throughout the manuscript.

2.2 Mean State Analysis

In this section, we describe the various metrics and plots that our methodology generates. While presented in terms of the abstract variable v , we also include sample plots of a comparison of the GBAF (*Jung et al.*, 2010) gross primary productivity (GPP) with CLM4.5 (*Oleson et al.*, 2013) for the purpose of illustration. In practice, ILAMB produces thousands of such plots and scalars, which are browsable in a website designed to aid modelers in understanding the benchmarking results.

2.2.1 Bias

We find the mean value in time, $\overline{v_{\text{ref}}}(\mathbf{x})$, over the time period of the reference, as well as that of the model, $\overline{v_{\text{mod}}}(\mathbf{x})$, over the same time period. These are spatial

variables that are included in the standard output as plots, as shown in Figure 4(a-b). We also compute the bias,

$$bias(\mathbf{x}) = \overline{v_{\text{mod}}}(\mathbf{x}) - \overline{v_{\text{ref}}}(\mathbf{x}) \quad (11)$$

as well as its mean over a given region, $\overline{bias(\mathbf{x})}$. To score the bias, we need to non-dimensionalize it as a relative error. We have chosen to do this by using the centralized root mean square of the reference data,

$$crms(\mathbf{x}) = \sqrt{\frac{1}{t_f - t_0} \int_{t_0}^{t_f} (v_{\text{ref}}(t, \mathbf{x}) - \overline{v_{\text{ref}}}(\mathbf{x}))^2 dt}, \quad (12)$$

which makes the relative error in bias given as,

$$\varepsilon_{bias}(\mathbf{x}) = |bias(\mathbf{x})| / crms(\mathbf{x}) \quad (13)$$

where the $|\cdot|$ operator represents the absolute value. The bias score as a function of space is,

$$s_{bias}(\mathbf{x}) = e^{-\varepsilon_{bias}(\mathbf{x})} \quad (14)$$

and the scalar score

$$S_{bias} = \overline{s_{bias}}(\mathbf{x}), \quad (15)$$

that is, the spatially integrated bias score. The motivation behind Equation (13) is to normalize the bias by the variability at any given spatial location. However, this also leads to the consequence that in areas where the given variable v has a small magnitude, simple noise can lead to large relative errors. For example, in Figure 4(d) we observe a poor score in the dry regions of Australia where GPP is small. Given the small contribution, it is undesirable that these errors induce a large negative contribution to the overall score. To address this issue, we introduce the concept of *mass weighting*. That is, when performing the spatial integral to obtain a scalar score (Equation (15)), we weight the integral by the period mean value of the reference variable using Equation (4) with $w = \overline{v_{\text{ref}}}$. In some instances the variable is truly a mass, but other times a flux or rate. The main motivation is to weight in areas where the variable is active. So while in our conceptual example, there is large relative error in GPP over deserts, these values will not negatively contribute to the overall score as the value of GPP is low in this area.

We apply mass weighting when the variable v represents a mass or flux of carbon or water as in GPP or precipitation. For variables representing energy states or quantities, such as temperature and radiation, we omit the weighting and perform a spatial integral only. We report plots of the bias and its score as well as the scalar integrated mean values.

2.2.2 Root mean squared error

For reference datasets with seasonal and interannual variability, we compute the root mean squared error over the time period of the reference dataset,

$$rmse(\mathbf{x}) = \sqrt{\frac{1}{t_f - t_0} \int_{t_0}^{t_f} (v_{\text{mod}}(t, \mathbf{x}) - v_{\text{ref}}(t, \mathbf{x}))^2 dt} \quad (16)$$

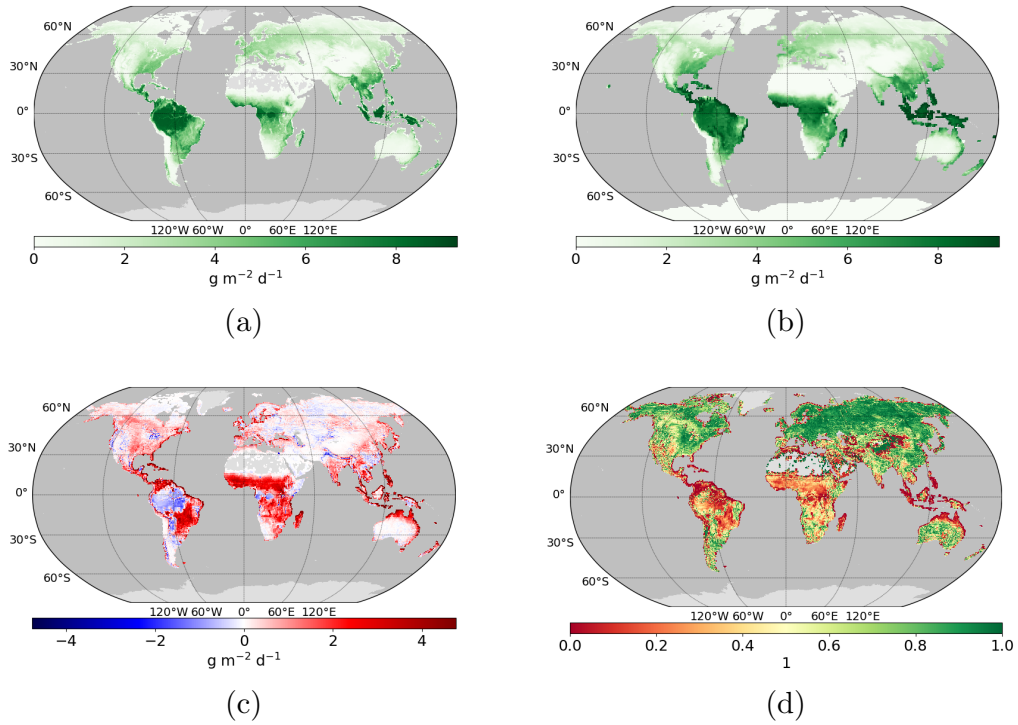


Figure 4: Comparisons of GPP between the reference (GBAF) and the model (CLM4.5) dataset. Each period mean is plotted over the original grid of the dataset. We highlight here that the reference (a) is not defined over Antarctica, Greenland, and part of the Sahara desert whereas the model (b) is defined over all land areas. Yet when the bias (c) and its score (d) is reported, the area represented is what both the reference and model agree on as land. (a) Reference period mean, $\overline{v_{\text{ref}}}(\mathbf{x})$, (b) Model period mean, $\overline{v_{\text{mod}}}(\mathbf{x})$, (c) Bias, $\text{bias}(\mathbf{x})$, (d) Bias Score, $s_{\text{bias}}(\mathbf{x})$

and include plots and the scalar $\overline{rmse(\mathbf{x})}$ in the standard output (Figure 5(a)). To score the root mean square error, we normalize the centralized root mean square error,

$$crmse(\mathbf{x}) = \sqrt{\frac{1}{t_f - t_0} \int_{t_0}^{t_f} ((v_{\text{mod}}(t, \mathbf{x}) - \overline{v_{\text{mod}}(\mathbf{x})}) - (v_{\text{ref}}(t, \mathbf{x}) - \overline{v_{\text{ref}}(\mathbf{x})}))^2 dt} \quad (17)$$

by the centralized root mean square of the reference dataset, Equation (12). This leads to a relative error of

$$\varepsilon_{rmse}(\mathbf{x}) = crmse(\mathbf{x})/crms(\mathbf{x}) \quad (18)$$

and a spatial RMSE score

$$s_{rmse}(\mathbf{x}) = e^{-\varepsilon_{rmse}(\mathbf{x})}. \quad (19)$$

The scalar score is obtained by

$$S_{rmse} = \overline{s_{rmse}}(\mathbf{x}), \quad (20)$$

where we again employ mass weighting when necessary. We score the centralized root mean squared error to decouple the bias score from the RMSE score. Computing the RMSE score by normalizing the RMSE would lead to a double counting of errors. That is, a large error in bias also leads to a large error in RMSE. By scoring the centralized RMSE, we remove the bias from the RMSE, allowing the RMSE score to focus on an orthogonal aspect of model performance.

2.2.3 Phase Shift

We evaluate the phase shift of the annual cycle of many datasets that have monthly variability by comparing the timing of the maximum of the annual cycle of the variable, $c(v)$ at each spatial cell across the time period of the reference dataset. We then approximate the phase shift of the reference and model datasets by subtracting these two values,

$$\theta(\mathbf{x}) = \arg \max_t (c_{\text{mod}}(t, \mathbf{x})) - \arg \max_t (c_{\text{ref}}(t, \mathbf{x})) \quad (21)$$

expressed in days. As the units for phase shift are consistent across all variables, no normalization is needed and we can remap the shift to the unit interval by

$$s_{\text{phase}}(\mathbf{x}) = \frac{1}{2} \left(1 + \cos \left(\frac{2\pi\theta(\mathbf{x})}{365} \right) \right) \quad (22)$$

and then spatially integrate the score over the appropriate region to find the scalar score,

$$S_{\text{phase}} = \overline{s_{\text{phase}}}(\mathbf{x}), \quad (23)$$

where again mass weighting is employed when appropriate. We include plots of the phase shift and its score in the standard output and represent them here in Figure 5(c-d). In addition to plots which show the time averaged variables as a map, we include

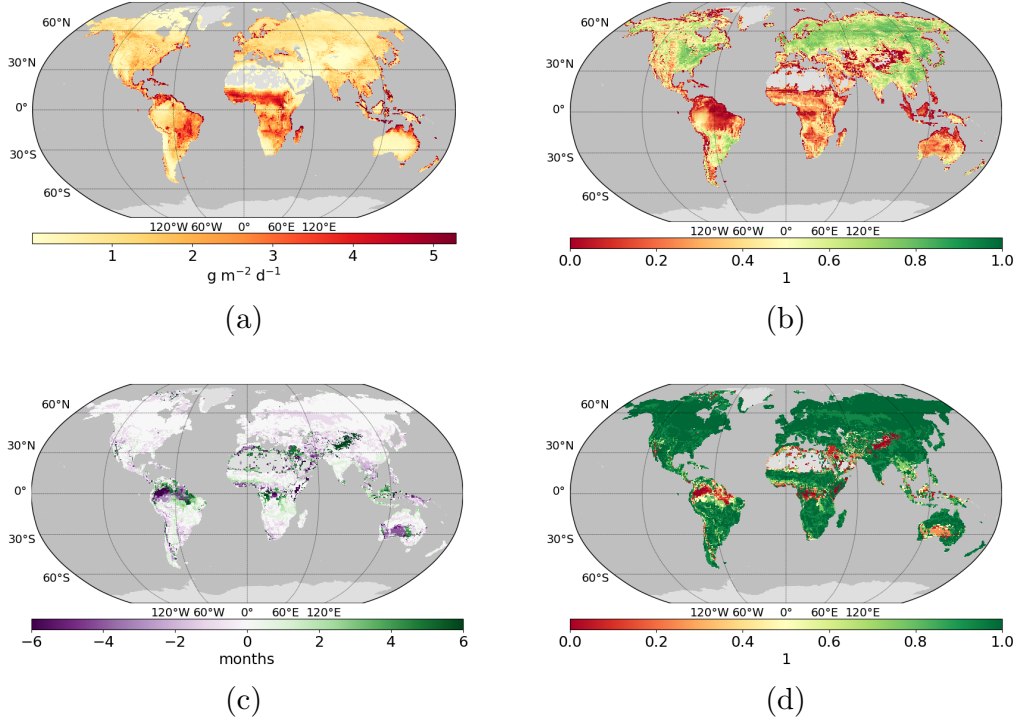


Figure 5: Comparisons of the RMSE and phase of GPP between the reference (GBAF) and the model (CLM4.5) dataset. (a) RMSE, $rmse(\mathbf{x})$, (b) RMSE score, $s_{rmse}(\mathbf{x})$, (c) Phase shift, $\theta(\mathbf{x})$, (d) Phase shift score, $s_{cycle}(\mathbf{x})$

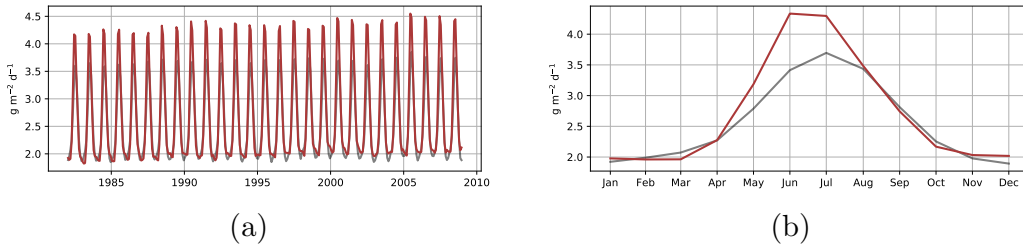


Figure 6: Spatial means of GPP of the reference (GBAF) shown in grey and the model (CLM4.5) in maroon. (a) Spatially integrated mean, $\overline{v_{ref}}(t)$ and $\overline{v_{mod}}(t)$, (b) Mean annual cycle, $\overline{v_{ref}}(t)$ and $\overline{v_{mod}}(t)$

line plots of the mean annual cycle and the spatially averaged variables, $\overline{v_{ref}}(t)$ and $\overline{v_{mod}}(t)$ shown in Figure 6.

2.2.4 Interannual Variability

A score for the interannual variability is computed by removing the annual cycle from both the reference and the model,

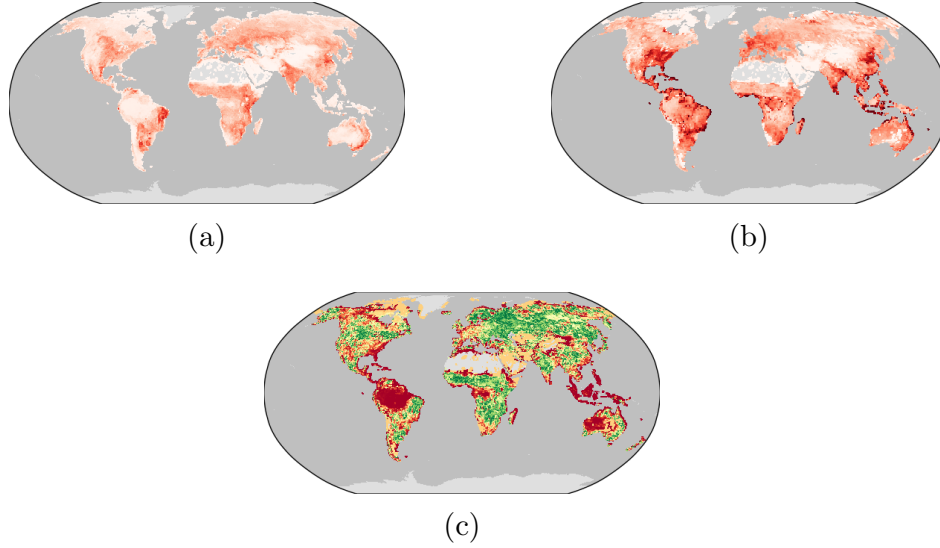


Figure 7: Comparisons of the interannual variability of GPP between the reference (GBAF) and the model (CLM4.5) dataset. (a) Reference interannual variability, $iav_{\text{ref}}(\mathbf{x})$, (b) Model interannual variability, $iav_{\text{mod}}(\mathbf{x})$, (c) Interannual variability score, $s_{iav}(\mathbf{x})$

$$iav_{\text{ref}}(\mathbf{x}) = \sqrt{\frac{1}{t_f - t_0} \int_{t_0}^{t_f} (v_{\text{ref}}(t, \mathbf{x}) - c_{\text{ref}}(t, \mathbf{x}))^2 dt} \quad (24)$$

$$iav_{\text{mod}}(\mathbf{x}) = \sqrt{\frac{1}{t_f - t_0} \int_{t_0}^{t_f} (v_{\text{mod}}(t, \mathbf{x}) - c_{\text{mod}}(t, \mathbf{x}))^2 dt} \quad (25)$$

$$\varepsilon_{iav}(\mathbf{x}) = (iav_{\text{mod}}(\mathbf{x}) - iav_{\text{ref}}(\mathbf{x})) / iav_{\text{ref}}(\mathbf{x}) \quad (26)$$

and then computing a score as a function of space,

$$s_{iav}(\mathbf{x}) = e^{-\varepsilon_{iav}(\mathbf{x})}. \quad (27)$$

The scalar score is then obtained by

$$S_{iav} = \overline{s_{iav}}(\mathbf{x}), \quad (28)$$

where mass weighting is used when necessary. We include plots of the variability and the score in the standard output and show them here in Figure 7. Note that while here we have shown the interannual variability of the GBAF product for illustration, in the default ILAMB configuration, the interannual variability is currently omitted for the GBAF products because its representativeness is considered to be poor (see Figure 10 of (Kumar *et al.*, 2016)).

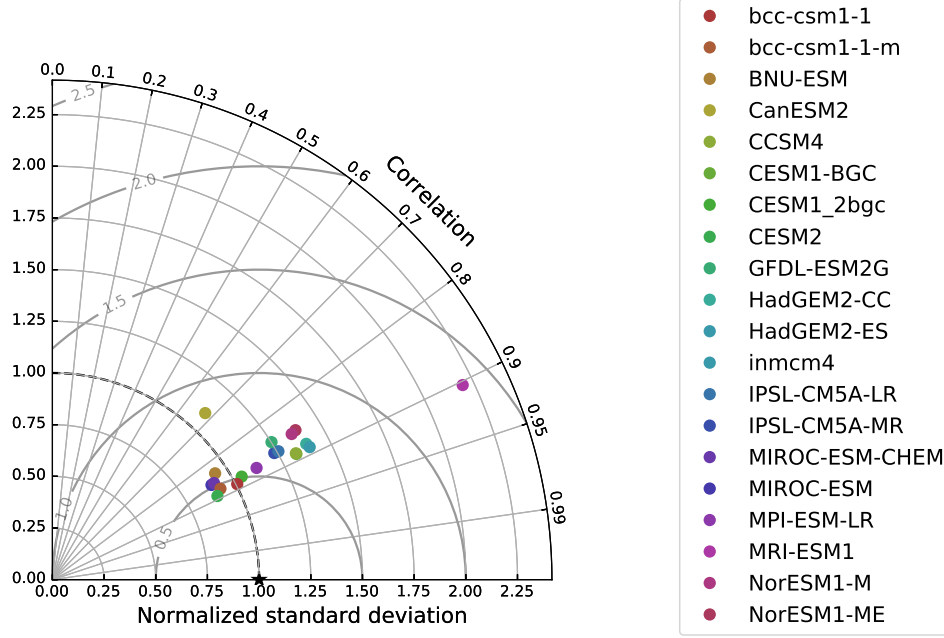


Figure 8: Taylor diagram comparing the spatial distribution of GPP of the reference (GBAF) shown as a black star to the CMIP5 models shown in colors.

2.2.5 Spatial Distribution

We score the spatial distribution of the time averaged variable by generating a Taylor (*Taylor, 2001*) diagram. We do this by computing the normalized standard deviation,

$$\sigma = \frac{\text{stdev}(\overline{v_{\text{mod}}}(\mathbf{x}))}{\text{stdev}(\overline{v_{\text{ref}}}(\mathbf{x}))} \quad (29)$$

and the spatial correlation R of the period mean values $\overline{v_{\text{ref}}}(\mathbf{x})$ and $\overline{v_{\text{mod}}}(\mathbf{x})$, and then assigning a score by the following relationship

$$S_{\text{dist}} = \frac{2(1 + R)}{(\sigma + \frac{1}{\sigma})^2}, \quad (30)$$

where the main idea is that we penalize the score when R and σ deviate from a value of 1. We include the Taylor plot in the standard output and represent it here in Figure 8.

2.2.6 Overall Score

The overall score for a given variable and data product is a composite of the suite of metrics defined above. We use a weighted sum,

$$S_{\text{overall}} = \frac{S_{\text{bias}} + 2S_{\text{rmse}} + S_{\text{phase}} + S_{\text{iav}} + S_{\text{dist}}}{1 + 2 + 1 + 1 + 1}, \quad (31)$$

where the RMSE score is doubly weighted to emphasize its importance.

2.3 Relationship Analysis

As models are frequently calibrated using the mean state scalar measures described in Section 2.2, a higher score does not necessarily reflect a more process-oriented model. In order to assess the representation of mechanistic processes in models, we also evaluate variable-to-variable relationships. For example, we look at how well models represent the relationship that GPP has with precipitation, evapotranspiration, and temperature. For the purposes of this section, we represent a generic dependent variable as v , as before, and score its relationship with an independent variable u . We then quantify the variable-to-variable relationship of the time period mean, $\bar{u}(\mathbf{x})$ on $\bar{v}(\mathbf{x})$, derived from the combination of reference datasets to the relationship diagnosed in models. We use the mean values over the reference time period to establish relationships as they represent a logical starting point. In the future, we plan to extend the relationship analysis to include seasonal and interannual variability.

2.3.1 Functional Response

We estimate a functional response by a 1D histogram, binned in terms of the independent variable $\bar{u}(\mathbf{x})$ with a number of bins, initially set to $n_{\text{bins}} = 25$. Then in each bin, we compute the mean value of the corresponding dependent variable, $\bar{v}(\mathbf{x})$ to approximate the functional dependence of u on v . We represent this binning with the operator \mathcal{F} that operates on the dependent and independent variables. We use it to compute functions from both the reference and model datasets.

$$f_{\text{ref}}(u) = \mathcal{F}(\bar{v}_{\text{ref}}(\mathbf{x}), \bar{u}_{\text{ref}}(\mathbf{x})) \quad (32)$$

$$f_{\text{mod}}(u) = \mathcal{F}(\bar{v}_{\text{mod}}(\mathbf{x}), \bar{u}_{\text{mod}}(\mathbf{x})), \quad (33)$$

where both curves are plotted in Figure 9(a) for the case of GPP compared to surface air temperature. These response curves are then scored by computing a relative error based on the RMSE,

$$\varepsilon_{\text{func}}^u = \sqrt{\frac{\int (f_{\text{ref}}(u) - f_{\text{mod}}(u))^2 du}{\int f_{\text{ref}}(u)^2 du}}, \quad (34)$$

where the integrals are approximated by the midpoint rule over the bins of the independent variable $\bar{u}(\mathbf{x})$. Then we use Equation (9) to map this relative error to a score by,

$$S_{\text{func}}^u = e^{-\varepsilon_{\text{func}}^u}. \quad (35)$$

The superscript u reinforces that this score represents functional performance with respect to a given independent variable u . The ILAMB implementation allows for any number of independent variables to be studied. In terms of our sample, ILAMB scores the functional relationship of GPP with respect to each independent variable separately (precipitation, evapotranspiration, temperature, *etc.*) and then computes the mean of these scores for the overall relationship score.

2.3.2 Hellinger Distance

In addition to the one-dimensional histograms, we also build normalized two-dimensional histograms ($n_{\text{bins}} = 25$ in both dimensions) from the time averaged data $\bar{v}(\mathbf{x})$ and $\bar{u}(\mathbf{x})$, represented here by the operator \mathcal{D} . We represent these distributions by,

$$d_{\text{ref}}(u) = \mathcal{D}(\overline{v_{\text{ref}}}(\mathbf{x}), \overline{u_{\text{ref}}}(\mathbf{x})), \quad (36)$$

$$d_{\text{mod}}(u) = \mathcal{D}(\overline{v_{\text{mod}}}(\mathbf{x}), \overline{u_{\text{mod}}}(\mathbf{x})), \quad (37)$$

as depicted in Figure 9(b–c). If we represent individual elements from these distributions $d_{\text{ref}}(u) = (p_1, \dots, p_{n_{\text{bins}}^2})$ and $d_{\text{mod}}(u) = (q_1, \dots, q_{n_{\text{bins}}^2})$, we can compute the so-called Hellinger distance (*Law et al.*, 2015)

$$S_{\text{dist}}^u = \frac{1}{\sqrt{2}} \sqrt{\sum_{i=1}^{n_{\text{bins}}^2} (\sqrt{p_i} - \sqrt{q_i})^2}$$

as a measure of how similar two distributions are to each other. While there are other choices, such as the Kullback-Leibler divergence, which are more commonly employed (*Dirmeyer et al.*, 2014), the Hellinger distance comes with the added benefit of being already normalized $[0, 1]$ and thus, further normalization is not necessary to use this directly as a score.

However, we only report the Hellinger distance as a scalar and do not include it in the scoring of the relationships. This is due to the fact that a bias in an independent variable can cause a density shift in the 2D distribution that would cause the score to unreasonably decrease. In terms of our example, a bias in precipitation (e.g. arising from a coupled model) could result in a poor relationship score with GPP, even if there is no underlying deficiency in the land-model simulated precipitation versus GPP relationship.

3 Datasets

In this section we explain how we utilize the methodology presented in Section 2 to evaluate model performance with respect to a collection of datasets (Tables 2–5) assembled by the ILAMB community. Errors in measurements, lack of measured or reported uncertainties, and inconsistencies in measurement methodology or instrumentation leading to ambiguous confidence in derived or synthesized data products all represent challenges in using observational data for benchmarking. In addition, the spatial and temporal coverage of different data products can vary substantially.

To account for the lack of quantitative uncertainties and scale mismatches between observations and models, and to bring a quantitative objectivity to model–data comparison, we developed a three-element rubric for weighting datasets as represented in Table 1. The first weight is based on a qualitative estimate of the certainty we have in a particular dataset. This weight encompasses both our certainty in the process used to obtain the observational information as well as the presence of quantitative uncertainty in the measurements themselves. A second weight for each dataset reflects its spatial and temporal coverage. The datasets employed in ILAMB are diverse and include site-level data, reanalysis data products, and remotely-sensed data. As our aim is to provide insight in land model performance on global and decadal scales, we give more weight to global products which are time series that extend for several years. The weights are combined multiplicatively to assign a total weight for each dataset. Then we normalize the weight by the sum of the weights of all the datasets for a given variable. For example, from Table 2 we see that there are two datasets used to benchmark GPP: Fluxnet and GBAF. For the Fluxnet product, we assign a certainty weight of 3 because while the collection is discussed in the published literature, there is no quantitative uncertainty provided. We assign a scale weight of 3 because the collection of sites covers multiple years of a substantial region of the globe yet has sparse

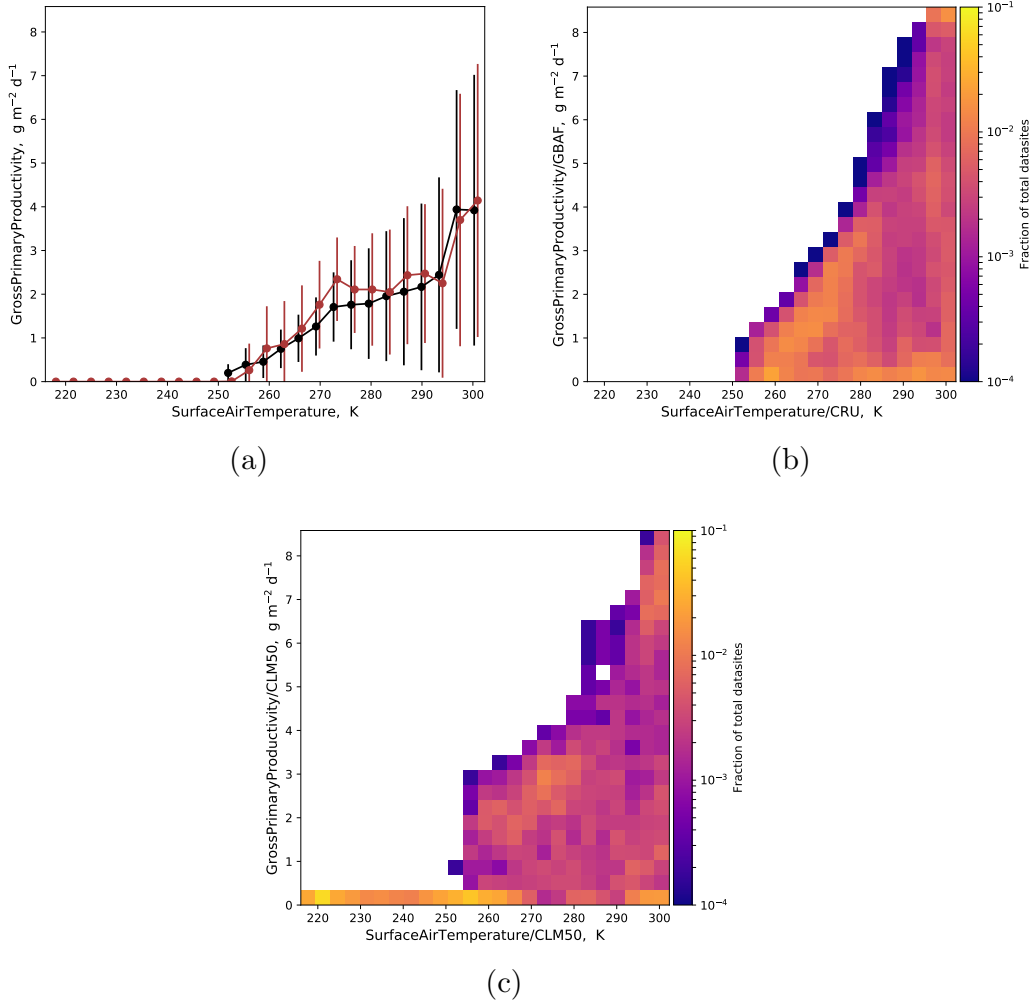


Figure 9: Variable-to-variable relationship plots which are a part of the standard output from the ILAMB methodology. (a) Functional responses, the reference $f_{\text{ref}}(u)$ in black, and the model $f_{\text{mod}}(u)$ in maroon. Data points reflect the mean for each independent value and the error bars reflect the standard deviation range., (b) Reference distribution, $d_{\text{ref}}(u)$, (c) Model distribution, $d_{\text{mod}}(u)$

coverage over important regions such as the tropics. The GBAF product is assigned a certainty weight of 3 for the same reason and a scale weight of 5 as it provides global coverage spanning multiple years. Then the total weight for the GPP variable which the GBAF dataset carries is

$$w_{\text{GBAF}}^{\text{GPP}} = \frac{3 \cdot 5}{3 \cdot 3 + 3 \cdot 5} \approx 63\%.$$

We use these weights to blend the overall score (Equation (31)) from each dataset for each variable. In this way ILAMB remains flexible to adding datasets as they are developed, allowing more weight to be given to those that the community believes are more credible and that are more comparable in scale to models.

A third weight reflects how useful the measured variable is in the focus of a model intercomparison project. Here, as an example, we show weighting for an analysis of model performance in representing the carbon cycle. We use these weights to blend the overall scores from each variable into a complete score across all variables for a given model. This allows ILAMB to include comparisons that are important for a complete understanding of the carbon cycle without necessarily allowing them to heavily influence the overall score. For example, the radiation and energy cycle datasets in Table 4 are all weighted comparatively low because, while they help one understand the carbon cycle, they are not as influential in the overall behavior.

We emphasize that this rubric is particular to our overarching goal of understanding the carbon cycle on global and decadal scales. However, the implementation is flexible and allows for an arbitrary weighting scheme to be developed that suits the needs of the user, community, or model intercomparison project that it serves.

The references and weights for each dataset that we have selected may be found in Tables 2–5. Each table represents a different aspect of the model: the ecosystem and carbon cycle in Table 2, the hydrological cycle in Table 3, the radiation and energy cycle in Table 4, and the forcings in Table 5. For the majority of these datasets, we make a direct comparison of the observed quantity to model outputs, or algebraic combinations of model outputs using the methodology described in section 2. However, there are a few special cases which require specific handling which we describe in the next section.

3.1 Special cases

In general, a consistent methodology is applied to compare model output with each dataset. This consistency across variables and datasets is a strength of the ILAMB methodology. However, this is not always possible, and here we enumerate a few exceptions and how they are handled.

3.1.1 Evaporative Fraction

To test the partitioning of surface energy, we compare the evaporative fraction derived from the GBAF (*Jung et al.*, 2010) data product to that of the models. The evaporative fraction is an algebraic expression in terms of the latent heat $L_e(t, \mathbf{x})$ and the sensible heat $S_h(t, \mathbf{x})$, given as

$$ef(t, \mathbf{x}) = \frac{L_e(t, \mathbf{x})}{L_e(t, \mathbf{x}) + S_h(t, \mathbf{x})}. \quad (38)$$

The expression can cause nonsensical results because in winter, the sensible heat flux can be negative, leading to a change of sign in the evaporative fraction. The expression can also lead to large evaporative fraction values since the magnitudes of both the

Table 1: The ILAMB rubric used to assign relative weights of a dataset. A score for each dataset is assigned in each of three areas. These scores are then combined multiplicatively and used to determine relative importance for a dataset with respect to a given variable.

Score	Certainty	Scale	Process
1	No given uncertainty, significant methodological issues affecting quality	Site level observations with limited space/time coverage	Observations that have limited influence on the targeted Earth system dynamics
2	No given uncertainty, some methodological issues affecting quality	Partial regional coverage, up to 1 year	Observations have direct influence on the targeted Earth system dynamics
3	No given uncertainty, methodology has some peer review	Regional coverage, at least 1 year	Observations useful to constrain processes that contribute to the targeted Earth system dynamics
4	Qualitative uncertainty, methodology accepted	Important regional coverage, at least 1 year	Observations well-suited to constrain important processes
5	Well-defined and relatively low uncertainty	Global scale spanning multiple years	Observations well-suited for discriminating critical processes among models

Table 2: References and weighting of datasets used to measure the ecosystem and carbon cycle. Weights are chosen using the rubric in Table 1 and reflect a focus on understanding the carbon cycle.

Variable/Dataset	Certainty	Scale	Process
Biomass			5
Tropical (<i>Saatchi et al., 2011</i>)	4	4	
NBCD2000 (<i>Kellndorfer et al., 2013</i>)	4	2	
USForest (<i>Blackard et al., 2008</i>)	4	2	
BurnedArea			4
GFED4S (<i>Giglio et al., 2010</i>)	4	5	
GrossPrimaryProductivity			5
Fluxnet (<i>Lasslop et al., 2010</i>)	3	3	
GBAF (<i>Jung et al., 2010</i>)	3	5	
LeafAreaIndex			3
AVHRR (<i>Myneni et al., 1997</i>)	3	5	
MODIS (<i>De Kauwe et al., 2011</i>)	3	5	
GlobalNetEcosystemCarbonBalance			5
GCP (<i>Le Quéré et al., 2016</i>)	4	5	
Hoffman (<i>Hoffman et al., 2014</i>)	4	5	
NetEcosystemExchange			5
Fluxnet (<i>Lasslop et al., 2010</i>)	3	3	
GBAF (<i>Jung et al., 2010</i>)	2	2	
EcosystemRespiration			4
Fluxnet (<i>Lasslop et al., 2010</i>)	2	3	
GBAF (<i>Jung et al., 2010</i>)	2	2	
SoilCarbon			5
HWSD (<i>Todd-Brown et al., 2013</i>)	3	5	
NCSCDV22 (<i>Hugelius et al., 2013</i>)	3	4	

Table 3: References and weighting of datasets used to measure the hydrology cycle. Weights are chosen using the rubric in Table 1 and reflect a focus on understanding the carbon cycle.

Variable/Dataset	Certainty	Scale	Process
Evapotranspiration			5
GLEAM (<i>Miralles et al., 2011</i>)	3	5	
MODIS (<i>De Kauwe et al., 2011</i>)	3	5	
EvaporativeFraction			5
GBAF (<i>Jung et al., 2010</i>)	3	3	
LatentHeat			5
Fluxnet (<i>Lasslop et al., 2010</i>)	3	1	
GBAF (<i>Jung et al., 2010</i>)	3	3	
Runoff			5
Dai (<i>Dai and Trenberth, 2002</i>)	3	5	
SensibleHeat			2
Fluxnet (<i>Lasslop et al., 2010</i>)	3	3	
GBAF (<i>Jung et al., 2010</i>)	3	5	
TerrestrialWaterStorageAnomaly			5
GRACE (<i>Swenson and Wahr, 2006</i>)	5	5	

latent and sensible heat can become small. For this reason, we apply a mask to ef , L_e , and S_h only considering values for which $S_h > 0$, $L_e > 0$, and $S_h + L_e > \phi$, where $\phi = 20 [Wm^{-2}]$ is a surface energy threshold.

Equation (38) is used to study how models partition the surface energy throughout the relevant season. Thus we use that expression when computing the RMSE or seasonal cycle. However, when comparing period mean values and the bias, Equation (38) leads to a combination of averaging methods. For this reason, when computing the mean evaporative fraction over time and the bias, we use a ratio of means in place of the mean of the ratio,

$$\overline{ef}(\mathbf{x}) = \frac{\overline{L_e}(\mathbf{x})}{\overline{L_e}(\mathbf{x}) + \overline{S_h}(\mathbf{x})}. \quad (39)$$

Beyond this change, the evaporative fraction is evaluated using the methodology defined in Section 2.

3.1.2 Albedo

We compare the albedo derived from observational data products (*Kato et al., 2013; Stackhouse Jr. et al., 2011; König-Langlo et al., 2013*) to that of models using the following expression,

$$al(t, \mathbf{x}) = \frac{R_u(t, \mathbf{x})}{R_d(t, \mathbf{x})}. \quad (40)$$

where R_u and R_d is the upward and downward shortwave radiation, respectively. As with the evaporative fraction in Section 3.1.1, the albedo expression can become numerically unstable when R_d approaches 0. Thus we again apply a mask, ignoring

Table 4: References and weighting of datasets used to measure the radiation and energy cycle. Weights are chosen using the rubric in Table 1 and reflect a focus on understanding the carbon cycle.

Variable/Dataset	Certainty	Scale	Process
Albedo			1
CERES (<i>Kato et al., 2013</i>)	4	5	
GEWEX.SRB (<i>Stackhouse Jr. et al., 2011</i>)	4	5	
MODIS (<i>De Kauwe et al., 2011</i>)	4	5	
SurfaceUpwardSWRadiation			1
CERES (<i>Kato et al., 2013</i>)	4	4	
GEWEX.SRB (<i>Stackhouse Jr. et al., 2011</i>)	4	5	
WRMC.BSRN (<i>König-Langlo et al., 2013</i>)	4	3	
SurfaceNetSWRadiation			1
CERES (<i>Kato et al., 2013</i>)	4	5	
GEWEX.SRB (<i>Stackhouse Jr. et al., 2011</i>)	4	5	
WRMC.BSRN (<i>König-Langlo et al., 2013</i>)	4	3	
SurfaceUpwardLWRadiation			1
CERES (<i>Kato et al., 2013</i>)	4	5	
GEWEX.SRB (<i>Stackhouse Jr. et al., 2011</i>)	4	5	
WRMC.BSRN (<i>König-Langlo et al., 2013</i>)	4	3	
SurfaceNetLWRadiation			1
CERES (<i>Kato et al., 2013</i>)	4	5	
GEWEX.SRB (<i>Stackhouse Jr. et al., 2011</i>)	4	5	
WRMC.BSRN (<i>König-Langlo et al., 2013</i>)	4	3	
SurfaceNetRadiation			2
CERES (<i>Kato et al., 2013</i>)	4	5	
Fluxnet (<i>Lasslop et al., 2010</i>)	4	3	
GEWEX.SRB (<i>Stackhouse Jr. et al., 2011</i>)	4	5	
WRMC.BSRN (<i>König-Langlo et al., 2013</i>)	4	3	

Table 5: References and weighting of datasets used to measure the forcings. Weights are chosen using the rubric in Table 1 and reflect a focus on understanding the carbon cycle.

Variable/Dataset	Certainty	Scale	Process
SurfaceAirTemperature			2
CRU (<i>Harris et al., 2014</i>)	5	5	
Fluxnet (<i>Lasslop et al., 2010</i>)	3	3	
Precipitation			2
CMAP (<i>Xie and Arkin, 1997</i>)	4	5	
Fluxnet (<i>Lasslop et al., 2010</i>)	3	3	
GPCC (<i>Schneider et al., 2014</i>)	4	5	
GPCP2 (<i>Adler et al., 2012</i>)	4	5	
SurfaceRelativeHumidity			3
ERA (<i>Dee et al., 2011</i>)	2	5	
SurfaceDownwardSWRadiation			2
CERES (<i>Kato et al., 2013</i>)	4	5	
Fluxnet (<i>Lasslop et al., 2010</i>)	4	3	
GEWEX.SRB (<i>Stackhouse Jr. et al., 2011</i>)	4	5	
WRMC.BSRN (<i>König-Langlo et al., 2013</i>)	4	3	
SurfaceDownwardLWRadiation			1
CERES (<i>Kato et al., 2013</i>)	4	5	
GEWEX.SRB (<i>Stackhouse Jr. et al., 2011</i>)	4	5	
WRMC.BSRN (<i>König-Langlo et al., 2013</i>)	4	3	

regions where no significant incoming radiation is observed, $R_d < \delta$. Equation (40) is used when comparing the RMSE and seasonal cycle. When the period mean and bias are computed, we compute the period mean average albedo based on the ratio of averages,

$$\overline{al}(\mathbf{x}) = \frac{\overline{R_u}(\mathbf{x})}{\overline{R_d}(\mathbf{x})}. \quad (41)$$

3.1.3 Global Net Ecosystem Carbon Balance

The observational datasets for the global net ecosystem carbon balance (*Le Quéré et al., 2016; Hoffman et al., 2014*) represent global totals, yet models return this value as fluxes defined over space. To create a model quantity commensurate with the observational data, ILAMB must integrate over the globe using Equation (4). As the observational dataset is a time series, much of our scoring methodology does not apply. For this discussion we will represent the global rate of carbon as nbp [$PgC\ yr^{-1}$]. We compute the accumulation of nbp

$$anbp(t) = \int_{t_0}^t nbp(t) dt \quad (42)$$

and score the difference in accumulated total at the end of the time period. The precise method differs slightly in each observational dataset.

The Global Carbon Project (GCP) dataset is derived by taking the land sink (uncertainty of ± 0.8 [$PgC\ yr^{-1}$]) and subtracting the emissions from land-use change (uncertainty of ± 0.5 [$PgC\ yr^{-1}$]). This means that the total uncertainty of the accumulated nbp at the end of 2010 is $\sqrt{0.5^2 + 0.8^2} \cdot (2010 - 1959) = 48.1$ [PgC]. We use this uncertainty to normalize the difference in accumulation at the end of the time period as a measure of relative error,

$$\varepsilon_{GCP} = \left| \frac{anbp_{\text{mod}}(2010) - anbp_{\text{ref}}(2010)}{48.1} \right| \quad (43)$$

and then again Equation (9) to compute a score of the difference

$$S_{GCP}^{\text{diff}} = e^{-\alpha_{nbp}\varepsilon_{GCP}}, \quad (44)$$

where $\alpha_{nbp} = 0.287$ and is chosen such that if a model falls within the certainty bounds of the accumulated amount through 2010, the corresponding score is at minimum 0.75. We see this as an important first step in incorporating uncertainty into the comparison methodology. We use the uncertainty to tune the scoring methodology, giving a good score to models that fall inside this uncertainty bound. We also compare the global rates of carbon across the time period in the form of a Taylor score of the time series, S_{GCP}^{dist} Equation (30) where the correlation and standard deviation are taken across the temporal dimension. Then the overall score is

$$S_{GCP}^{\text{nbp}} = \frac{1}{2} (S_{GCP}^{\text{diff}} + S_{GCP}^{\text{dist}}) \quad (45)$$

In the *Hoffman et al.* (2014) dataset, we only score the accumulated amount at the end of the observed period. We omit providing a Taylor scoring of the rates because there appears to be some smoothing of the rate data inherent in the process of producing this dataset. However, this dataset explicitly provides a lower and upper bound on uncertainty as a function of time throughout the dataset. So we determine the integrated uncertainty at the end of 2010 by accumulating the upper (52.4 [PgC]) and lower (-32.1 [PgC]) limit of uncertainty, computing the difference, and then halving the value resulting in an uncertainty of 42.3 [PgC]. We then use the same approach to score the difference,

$$\varepsilon_{\text{Hoffman}} = \left| \frac{a_{\text{mod}}(2010) - a_{\text{ref}}(2010)}{42.3} \right| \quad (46)$$

$$S_{\text{Hoffman}}^{\text{nbp}} = e^{-\alpha_{nbp}\varepsilon_{\text{Hoffman}}} \quad (47)$$

3.1.4 Runoff

We use the *Dai and Trenberth* (2002) river discharge dataset to assess model performance of runoff for the world's 50 largest river basins. First, we compute the mean annual runoff from the model over the time period of the observational dataset. Then we take the river discharge data and distribute it over the area of the river basins and compare this to the mean runoff over the same basin. This simple approach was taken to allow us to compare runoff across models even if they do not have a river routing model.

We include plots of the mean runoff of the reference and model over river basins and the bias, represented in Figure 10. We also include regional mean runoff plots for each of the river basins included, but only show that of the Amazon river basin in Figure 10(d). The model performance is then scored using the bias (Section 2.2.1), the interannual variability (Section 2.2.4), and the spatial distribution (Section 2.2.5) metrics.

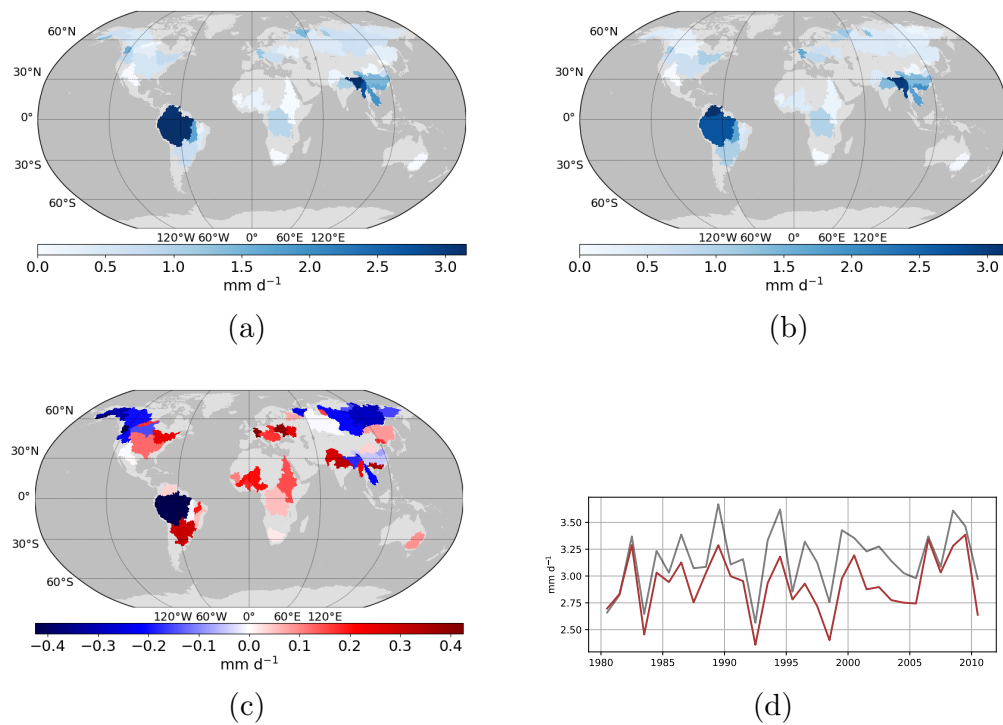


Figure 10: Comparisons of runoff between the reference (*Dai and Trenberth, 2002*) and the model (CLM4.5) dataset. (a) Reference mean runoff, (b) Model mean runoff, (c) Mean runoff bias, (d) Annual mean runoff for the Amazon river basin where the reference is shown in grey and the model in maroon.

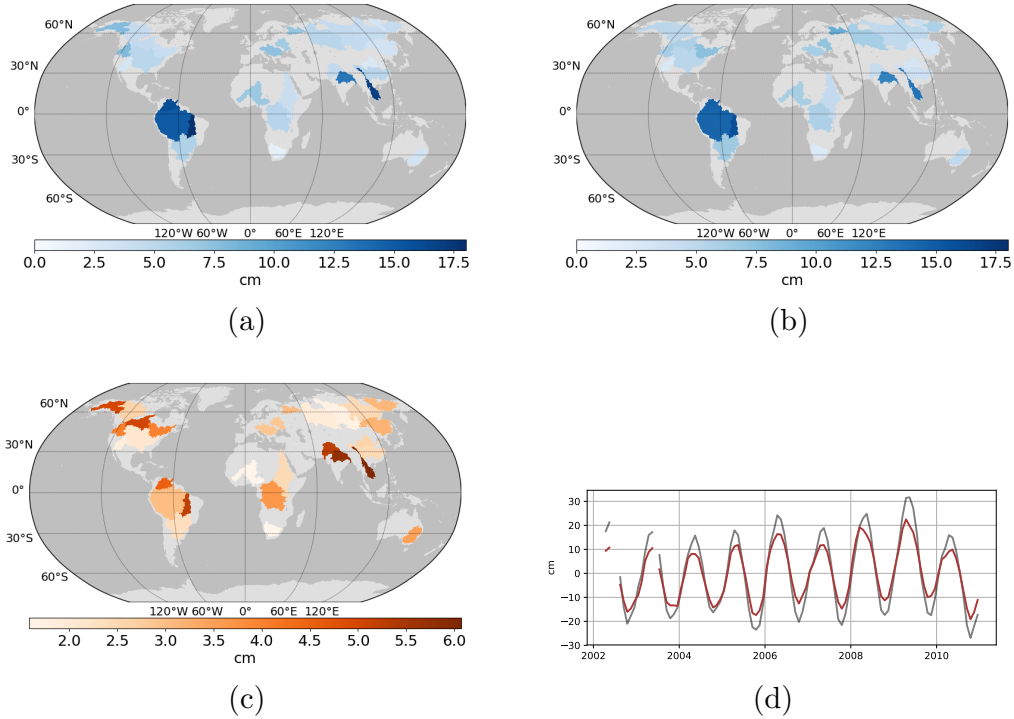


Figure 11: Comparisons of the terrestrial water storage anomaly between the reference (GRACE) and the model (CLM4.5) dataset. (a) Reference mean anomaly magnitude, (b) Model mean anomaly magnitude, (c) Mean anomaly RMSE, (d) Annual mean anomaly for the Amazon river basin where the reference is shown in grey and the model in maroon.

3.1.5 Terrestrial Water Storage Anomaly

We use the Gravity Recovery and Climate Experiment (GRACE) (*Swenson and Wahr, 2006*) dataset to assess the terrestrial water storage anomaly (twsa) in models. However, there are a few challenges in producing a fair comparison. The first of those is that models report only the storage and so the anomaly must be computed. The more serious challenge is that the resolution of this data is quite coarse (300–400 [km]) and thus, pointwise comparisons are not appropriate (*Swenson, 2013*). Instead we compare mean anomaly values over 30 of the world’s largest river basins. In this way the comparison is more fair as it is over large areas and automatically omits dry areas which are not of interest.

We include plots of the magnitude of the mean anomaly of the reference and model over river basins and the RMSE, represented in Figure 11. We also include regional mean anomaly plots for each of the river basins, but show only that of the Amazon river basin in Figure 11(d). The model performance is then scored using the RMSE (Section 2.2.2) and the interannual variability (Section 2.2.4) metrics.

4 Software

We have implemented the methodology described in Sections 2 and 3 into a software package that is freely available to the community. We previously developed a prototype implementation (*Mu et al., 2015*) based on the NCAR Command Language (NCL). We then moved the algorithm into an open-source, openly-developed python

package (Collier *et al.*, 2016) in an effort to produce a product to which the community can more easily make contributions. The referenced digital object identifier (DOI) will lead to the software repository, where the source code and documentation can be found. The documentation includes the public interface as well as tutorials that span topics such as installation, basic usage, adding models or benchmark datasets, and formatting benchmark datasets.

The ILAMB package is designed to ingest datasets which follow the Climate and Forecast (CF) convention (Eaton *et al.*, 2017). The CF website explains that the “conventions define metadata that provide a definitive description of what the data in each variable represents, and the spatial and temporal properties of the data. This enables users of data from different sources to decide which quantities are comparable, and facilitates building applications with powerful extraction, regridding, and display capabilities.” We have built the ILAMB package to embody this philosophy, making it directly useful to those who adhere to this standard. While model intercomparison efforts, such as CMIP5, have encouraged the use of these conventions among modelers, the observational community has not yet widely put them into practice. Much of the work in adding datasets to the collection is in encoding them to follow this convention.

For the purpose of communicating how the ILAMB package works, consider the configure file shown in Figure 12, which defines a set of observational datasets that will be used to confront models. The ‘h1’ bracket is a heading used to categorize variables, represented by the ‘h2’ heading. This comparison involves the surface upward short-wave radiation and the albedo, both of which are variables belonging to the radiation and energy cycle. Inside each ‘h2’ heading, we specify the variable name that will be compared (‘rsus’ is the netCDF variable name for surface upward shortwave radiation). However, we provide a mechanism for variable synonyms in this case by specifying alternate variable names. If the ILAMB system cannot find the main variable, it will try to find any alternates that the user specifies. This allows the software to encourage the use of standard variable names, but accounts for modeling groups wanting to use ILAMB without pre-processing. Also note the ‘derived’ keyword in the albedo section. While the components of albedo are part of standard model output, the albedo is not. The ILAMB package allows for users to specify algebraic relationships in the configure file process. This makes the process automatic and transparent to those who may read this configure file.

The ILAMB package will ingest this configure file and try to build commensurate quantities from model outputs. While observational datasets come in different forms (globally gridded remote sensing products, tower data collections, etc.), the ILAMB system reads the spatial and temporal information found in the file and uses it to trim, subsample, and/or coarsen the model data as appropriate.

5 Discussion

The ILAMB framework is designed to be both powerful and flexible. While we have made choices in the default configuration, described above, focused on global analysis for decadal to centennial scale ESMs, ILAMB allows the user to customize selection of variables, weighting of datasets, and spatial subsetting that make it useful for assessing results from mesoscale weather forecasting or other models. We envision developing a library of sample configuration files, targeting various well-known models and model applications.

As much of the usefulness of ILAMB depends on the quality of the underlying observational data, we recommend that data providers include explicit representations of the underlying spatial grids including the areas over which quantities have been averaged. Observational datasets frequently report mean values in a cell taken over an

```

[h1: Radiation and Energy Cycle]

[h2: Surface Upward SW Radiation]
variable = "rsus"
alternate_vars = "FSNS"

[CERES]
source = "DATA/rsus/CERES/rsus_0.5x0.5.nc"

[h2: Albedo]
variable = "albedo"
derived = "rsus/rsds"

[CERES]
source = "DATA/albedo/CERES/albedo_0.5x0.5.nc"

```

Figure 12: Sample ILAMB configure file defining comparisons to the surface upward shortwave radiation and albedo variables from the CERES (*Kato et al., 2013*) product.

area which may include land but also portions of lakes, rivers, and oceans. This leads to ambiguity with regard to the contribution of land cover types to the measurement itself and subsequently adds to the uncertainty when comparing values to model output.

5.1 Interpreting the Overall Score

The thrust of this paper is to detail a methodology for computing a single overall score that captures a model’s skill in reproducing patterns found in the observed record. However, we do not view the absolute value of the score as particularly meaningful beyond the precise definition described in this paper. In general, no model can achieve a perfect score for any given variable for several reasons.

First, there is measurement error and uncertainty in the observational data that makes a perfect score unlikely or undesirable against even a single dataset. This is what motivates some in the community (*Abramowitz, 2005; Best et al., 2015*) to pose that benchmarking requires an expectation of performance which is admittedly lacking in our approach. Second, despite that every attempt is made to employ multiple independent datasets of high quality for confrontation with models, these datasets are inconsistent with each other, making a perfect score across all datasets impossible. We do this as comparisons with multiple observational and synthesized datasets for a single variable offer the user more information about the robustness of model predictions within the limits of observational uncertainty at varying spatial and temporal scales. Third, a lower score with respect to a given variable is not necessarily a sign of a poor model. It may rather highlight the need for better measurement campaigns or improved metrics (i.e., sometimes we learn that our measurements are incomplete or do not acknowledge important uncertainties, or our metrics are inappropriate for a given dataset).

The overall score is meant to aid the scientist in discovering when meaningful changes have occurred in the model or across models. The holistic nature of the ILAMB suite of datasets and metrics helps provide a synthesis of model performance that directs the attention of the user to relevant aspects. While we present Figure 1 as the

main result of the ILAMB methodology, it is intended to merely indicate variables of particular interest for further consideration. ILAMB output is presented as a hierarchy of interactive webpages that employ JavaScript features to present information to users in a logical and intuitive fashion. From the graphical overview, the user can select individual variables and datasets from the “Results Table” tab to be led to pages which detail the contributing factors to the model’s overall score. On this new page, pre-defined spatial regions can be individually selected, causing the tabular data and diagnostics to be updated automatically to reflect information relevant only to that region. Although all the tabular information, scores, and graphical diagnostics are pre-computed and generated when ILAMB is run, the web-based interface is designed to facilitate discovery and understanding of model results. The overall score does not replace the scientist, it guides her/him to the relevant plots and diagnostics.

5.2 How is ILAMB Used?

The ILAMB package is particularly useful for verification, i.e., during model development to confirm that new model code improves performance in a targeted area without degrading performance in another area, and for validation, i.e., when comparing performance of one model or model version to that of other models or model versions.

In developing and applying the ILAMB package, we have incorporated a wide variety of representative observational datasets (see Tables 2, 3, 4, and 5), and we have favored data that have the most open data policies. In many cases, these data have been averaged or remapped to be more directly comparable with model output. As this collection of datasets grows, maintaining and distributing the latest versions will be challenging and require community collaboration. For tracking the evolving performance of models over the long term, it may be necessary to maintain access to older versions of data as well as the latest version since corrections to observational datasets can significantly impact model performance scores. Various technologies could fill this role, and the Observations for Climate Model Intercomparisons (obs4MIPs; <https://www.earthsystemcog.org/projects/obs4mips/>) activity shows promise as a potential solution to this challenge. The preferred solution would ideally support versioning and allow for long-lived versions associated with ILAMB releases. In the interim, we have implemented a simple scheme for sharing summarized and remapped datasets through a webserver.

The ILAMB package is currently being used by individual model developers and international modeling centers. ILAMB offers developers a quick and easy method for checking the impacts of new model development before committing code changes. For modeling centers, ILAMB provides a systematic assessment of historical simulation experiments and enables tracking of performance of model revisions. ILAMB will also be useful for model intercomparison projects (MIPs) as a starting point for evaluating model variability and uncertainty. As a part of such MIPs, investigators may wish to develop custom metrics or incorporate datasets specific to their purposes. ILAMB could be executed automatically as model results are uploaded to a system like the Earth System Grid Federation (ESGF; <https://esgf.llnl.gov/>) to give users a “first look” at variation in results and to determine if output should be downloaded for a particular study. ILAMB diagnostics can also be useful for parameter sensitivity studies or for optimization experiments in combination with an automated modeling framework like the Predictive Ecosystem Analyzer (PEcAn; <http://pecanproject.org/>; *LeBauer et al.*, 2013; *Dietze et al.*, 2014). For the assessments community, the results of a multi-model ILAMB evaluation could be useful for understanding which model results would be appropriate for use in studying impacts and which models may poorly capture processes relevant to the impacts under consideration.

5.3 Future Work

Development of the ILAMB package is ongoing, and the terrestrial modeling and observational communities are being engaged to identify *in situ* and remote sensing datasets, to define additional evaluation metrics, and to use the package for a wide variety of MIPs (Hoffman *et al.*, 2017). While most effort has been invested in global- and regional-scale model evaluation, new work is focused on improved benchmarking for site-level time series, spatial transects, and seasonal and diurnal variability. Future development will include incorporation of experiment-specific model evaluation metrics derived from prior studies, including Free-Air CO₂ Enrichment (FACE) (Zahle *et al.*, 2014; Walker *et al.*, 2014, 2015), nutrient addition, rainfall exclusion, and warming experiments (Bouskill *et al.*, 2014; Zhu *et al.*, 2016). Partner activities, like NASA’s Permafrost Benchmarking System project and the Arctic-Boreal Vulnerability Experiment (ABOVE), are integrating additional datasets and building metrics for specific regions, study areas, or processes of interest. We are applying the ILAMB methodology and code base to develop a marine biogeochemical model benchmarking tool, called the International Ocean Model Benchmarking (IOMB) package.

Based on previous prototypes and community discussion, we developed the ILAMB model benchmarking package for evaluating the fidelity of land carbon cycle models. The package generates graphical diagnostics and computes a comprehensive set of statistics through model–data comparisons, and scores model performance for a wide variety of variables for a suite of observational datasets. Rigorously defined model evaluation metrics and strategies for handling multiple resolutions and land masks are documented above. The ILAMB package is open source and is becoming widely adopted by modeling centers and for informing model intercomparison studies. We are actively seeking community involvement in adding more evaluation metrics and new observational datasets.

Acknowledgments

This manuscript has been authored by UT-Battelle, LLC under Contract No. DE-AC05-00OR22725 with the U.S. Department of Energy. The United States Government retains and the publisher, by accepting the article for publication, acknowledges that the United States Government retains a non-exclusive, paid-up, irrevocable, world-wide license to publish or reproduce the published form of this manuscript, or allow others to do so, for United States Government purposes. The Department of Energy will provide public access to these results of federally sponsored research in accordance with the DOE Public Access Plan (<http://energy.gov/downloads/doe-public-access-plan>).

This research was supported through the Reducing Uncertainties in Biogeochemical Interactions through Synthesis and Computation Scientific Focus Area (RUBISCO SFA), which is sponsored by the Regional and Global Climate Modeling (RGCM) Program in the Climate and Environmental Sciences Division (CESD) of the Office of Biological and Environmental Research (BER) in the U.S. Department of Energy Office of Science. Oak Ridge National Laboratory (ORNL) is managed by UT-Battelle, LLC for the U.S. Department of Energy under Contract No. DE-AC05-00OR22725. The National Center for Atmospheric Research (NCAR) is managed by the University Corporation for Atmospheric Research (UCAR) on behalf of the National Science Foundation (NSF). Lawrence Berkeley National Laboratory (LBNL) is managed and operated by the Regents of the University of California under Contract No. DE-AC02-05CH11231.

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