1 Title: Facilitating feedbacks between field

² measurements and ecosystem models

- 3 Running Title: Feedbacks between measurements and models
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12 Abstract

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Ecological models help us understand how ecosystems function, predict responses to
   global change, and identify future research needs. However, widespread use of models is
   limited by the technical challenges of model-data synthesis and information management.
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   To address these challenges, we present a ecoinformatic workflow, the Predictive
   Ecosystem Analyzer (PEcAn), that facilitates model analysis. Herein we describe the
   PEcAn modules that synthesize plant trait data to estimate model parameters, propagate
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   parameter uncertainties through to model output, and evaluate the contribution of each
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   parameter to model uncertainty. We illustrate a comprehensive approach to the
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   estimation of parameter values, starting with a statement of prior knowledge that is
   refined by species level data using Bayesian meta-analysis; this is the first use of a
   rigorous meta-analysis to inform the parameters of a mechanistic ecosystem model.
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   Parameter uncertainty is propagated using ensemble methods to estimate model
   uncertainty. Variance decomposition allows us to quantify the contribution of each
   parameter to model uncertainty; this information can be used to prioritize subsequent
   data collection. By streamlining the use of models and focusing efforts to identify and
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   constrain the dominant sources of uncertainty in model output, the approach used by
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   PEcAn can speed scientific progress.
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   We demonstrate PEcAn's ability to incorporate data to reduce uncertainty in
   productivity of a perennial grass monoculture (Panicum virgatum L.) modeled by the
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   Ecosystem Demography model. Prior estimates were specified for fifteen model
   parameters, and species-level data were available for seven of these. Meta-analysis of
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   species-level data substantially reduced the contribution of three parameters (specific leaf
   area [SLA], maximum carboxylation rate, and stomatal slope) to overall model
   uncertainty. By contrast, root turnover rate, root respiration rate, and leaf width had
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- 37 little effect on model output, therefore trait data had little impact on model uncertainty.
- For fine root allocation the decrease in parameter uncertainty was offset by an increase in
- model sensitivity. Remaining model uncertainty is driven by growth respiration, fine root
- allocation, leaf turnover rater, and SLA. By establishing robust channels of feedback
- between data collection and ecosystem modeling, PEcAn provides a framework for more
- 42 efficient and integrative science.
- keywords: traits, ecoinformatics, ecophysiology, Ecosystem Demography, sensitivity
- analysis, variance decomposition, ecological forecast, Bayesian, meta-analysis, ecosystem
- 45 model

46 Introduction

- In the face of unprecedented global change there is growing demand for predictions of ecosystem responses that provide actionable information for policy and management (Clark et al., 2001). Currently, the response of the terrestrial biosphere remains one of the largest sources of uncertainty in projections of climate change (Denman et al., 2007). This uncertainty comes from a combination of the uncertainties about our conceptual understanding of ecological systems, as captured by the structure and assumptions of the models used to make ecological forecasts, the uncertainties in the parameters of these 53 models, and the uncertainties associated with the underlying data itself (McMahon et al., 2009). Reducing these uncertainties requires that we be able to synthesize existing information, efficiently identify the dominant sources of model uncertainty and target them with further field research. Despite the acknowledged importance of these activities, there is often a disconnect between model simulation and data collection. Both model-data synthesis and the investigation of uncertainty remain challenging, while the use of models to quantitatively inform data collection is extremely rare. Most modeling uses a single point estimate for each parameter, effectively treating each parameter value as completely certain. However, 62 such point estimates do not account for the degree to which we understand a parameter 63 based on observations. Furthermore, the rationale for a particular estimate is often unclear, as is the degree to which the estimate represents the process being observed or its representation in a model. In many cases, parameter values are chosen iteratively to "tune", or "calibrate" the model output to observations. A first step toward constraining 67 model uncertainty is to account for uncertainty in model parameters instead of relying on point estimates.
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More rigorous approaches to estimating parameter values include model optimization and

data assimilation (Medvigy et al., 2009; Reichstein et al., 2003), as well as Bayesian model-data fusion (Luo et al., 2011). However, these approaches have generally started with uninformative or vague prior estimates of model parameters. These vague priors ignore available data that could directly inform parameter values; the most commonly used vague prior distribution is a uniform. A uniform prior assigns equal probability to parameter values over its entire range, in many cases over many orders of magnitude. The 76 use of such vague priors often exacerbates problems with equifinality (Richardson and Hollinger, 2005; Williams et al., 2009; Luo et al., 2009) which can produce unidentifiable 78 parameters, as well as biologically unrealistic parameter sets that generate the right model 79 output for the wrong reasons (Beven and Freer, 2001; Beven, 2006; Williams et al., 2009). Another reason to use informed priors is to take advantage of one of the key strengths of the Bayesian paradigm: the ability to synthesize multiple sources of information in a 82 rigorous and consistent framework. For example, plant traits related to leaf stoichiometry and photosynthetic capacity are often well constrained by previous research (Skillman, 2008; Reich and Oleksyn, 2004; Wright et al., 2004; Wullschleger, 1993), while other traits, such as root respiration rate, are more difficult to measure and data are sparse. Informed priors allow existing information to be formally integrated into model parameterization, 87 even if there is no data for the particular species or plant functional type (PFT) being measured; the level of confidence in a parameter value is reflected in its variance. Models have rarely been used to quantify the value of data with respect to reducing uncertainty. Instead, data collection is often focused on answering specific questions in specific spatial, temporal, and taxonomic contexts. In these contexts, the value of a particular data set is based on the ability to answer a particular question. However, the same data set may have a very different value in the context of reducing model uncertainty. For example, a single data point used to inform a poorly understood but influential model parameter can reduce model uncertainty more than a large collection of

data on a trait that is relatively well studied. In a modeling context, the value of an additional data point depends both on how much it constrains parameter uncertainty and the sensitivity of model output to the parameter. Thus, the ability to comprehensively utilize available data in model parametrization can help to identify gaps in existing 100 knowledge, improve the ability of models to account for current understanding, and 101 inform data collection efforts by identifying the knowledge gaps most responsible for 102 uncertainty. 103 While the increasing sophistication of model-data fusion and uncertainty accounting is a 104 critical step in the right direction, the complexity of such analyses can make models even 105 less accessible. One of the reasons to make models more accessible, and to make them 106 better at synthesizing existing data, is that they are fundamentally a formal, quantitative 107 distillation of our current understanding of how a system works. As such, models can be 108 used to identify gaps in our understanding and target further research. This feedback 109 between models and data could be improved if models were routinely evaluated in a way 110 that quantifies the value of data with respect to reducing uncertainty. We fundamentally 111 believe that streamlining the informatics of modeling – the need to track, process, and 112 synthesize data and model output – will make the development and application of 113 ecological data and models more accessible, transparent, and relevant. In this paper we present the Predictive Ecosystem Analyzer (PEcAn) as a step toward 115 meeting these objectives. PEcAn is a scientific workflow that manages the flows of data 116 used and produced by ecological models, and that assists with model parametrization, 117 error propagation, and error analysis. PEcAn accomplishes two goals: first, it synthesizes 118 data and propagates uncertainty through an ecosystem model; second, it places an 119 information value on subsequent data collection that enables data collection that 120 efficiently reduces uncertainty. In addition to quantifying the information content of any 121 prediction or assessment, these techniques also help identify the gaps in our knowledge of 122

ecological and biogeochemical processes (Saltelli et al., 2008). 123 PEcAn addresses the challenge of synthesizing plant trait data from the literature in a 124 way that accounts for the different scales and sources of uncertainty. Available data is 125 synthesized using a Bayesian meta-analysis, and the meta-analysis posterior estimates of 126 plant traits are used as parameters in an ecosystem model. 127 A model ensemble is a set of model runs with parameter values drawn from the 128 meta-analysis posteriors estimate of plant traits. Output from a model ensemble 120 represents the posterior predictive distributions of ecosystem responses that account for 130 trait parameter uncertainty (hereafter "model posterior" refers to the "model ensemble 131 output"). Sensitivity analysis and variance decomposition help to determine which traits 132 (model parameters) drive uncertainty in ecosystem response (model posterior) (Saltelli 133 et al., 2008; Larocque et al., 2008). These analyses help target parameters for further 134 constraint with trait data, forming a critical feedback loop that drives further field 135 research and provides an informative starting point for data assimilation. Here we 136 illustrate an application of PEcAn to the assessment of aboveground yield in a perennial 137 grass monoculture. 138 In the sections below, we provide an overview of the components of PEcAn's integrated 139 framework for data synthesis and ecological prediction. We start with a description of the methods implemented in the workflow (Implementation). This includes descriptions of 141 the database, Bayesian meta-analysis, ensemble analysis, sensitivity analysis, and 142 variance decomposition. Finally we present an example of the application of the system 143 (Application) to analyze the aboveground biomass of switchgrass (Panicum virgatum L.), 144

by the Ecosystem Demography model, version 2.1 (ED2) (Medvigy et al., 2009;

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Moorcroft et al., 2001).

147 Implementation

$_{ ext{ iny 48}}$ PEcAn workflow

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The Predictive Ecosystem Analyzer (PEcAn) manages the flow of information into and
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    out of ecosystem models.PEcAn is not a model itself, it is a scientific workflow consisting
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    of discrete steps, or modules. Individual modules are building blocks of the workflow,
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    represented by the rectangles in Figure 1, while flows of information are represented by
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    arrows. This makes PEcAn an encapsulated, semi-automated system for model
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    parametrization, error propagation, and analysis.
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    A central objective of the PEcAn workflow is to make the entire modeling process
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    transparent, reproducible, and adaptable to new questions (sensu Stodden et al., 2010;
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    Ellison, 2010). To achieve this objective, PEcAn's adheres to "best practice" guidelines
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    for ecological data management and provenance tracking (Jones et al., 2006; Michener
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    and Jones, 2012).
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    PEcAn uses a database to track data provenance and a settings file to control workflow
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    analyses and model runs. The database records the site, date, management, species, and
    treatment information for each trait observation used in the meta-analysis. Settings
    related to the experimental design and computation are set and recorded in a separate file
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    for each analysis.
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    The PEcAn source code, as well as the inputs and output used in the analysis described
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    below (see Application) are provided as an appendix. However, new users are encouraged
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    to utilize the latest release available on the project web site (www.pecanproject.org).
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    This site also provides a virtual machine and a web-interface that minimize the effort
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    required to run PEcAn and begin using an ecosystem model. The PEcAn "virtual
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    machine" provides all of the required software dependencies in a pre-configured desktop
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    environment that can be run on any standard operating system using a freely available
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virtualization software such as VirtualBox (www.virtualbox.org/) or VmWare Player 172 (www.vmware.com). The virtual machine minimizes the installation time and 173 pre-requisite knowledge required to perform analyses, and can be used to support 174 investigation, development, and education. The web interface is even easier to use, but 175 does not provide access to all of PEcAn's functionality. 176 The PEcAn software is primarily written in R and developed in a Linux environment. It 177 also relies on a MySQL database, bash, JAGS, and specialized R packages. PEcAn has a 178 family of model-specific functions that manage the details of launching of model runs and 179 reading model output. 180 Although PEcAn does not depend on any specific model, it was developed to support 181 ecosystem models that run in high-performance computing environments; for this reason, 182 it is capable of running models locally, remotely, or through queuing systems regardless of 183 whether PEcAn is compiled locally or run as a virtual machine. The PEcAn 1.1 release 184 described herein runs with the Ecosystem Demography model, in addition, the current 185 release also supports SIPNET (Moore et al., 2008) and near term support for IBIS 186 (Kucharik et al., 2000), DayCent (Parton et al., 1998), and BioCro (Miguez et al., 2009) 187 is under development. 188

189 Trait Database

Model parameters are associated with corresponding prior distributions, and in many
cases, with species-level data. Both prior distributions and data are stored in a relational
database (Appendix B). PEcAn directly accesses the database, which contains additional
meta-data for each data set, including site descriptions, measurement conditions,
experimental details, and citations.

195 Trait Priors

A fundamental component of the Bayesian approach to parameter estimation is the use of 196 priors. Priors formally incorporate knowledge of a parameter based on previous studies 197 into a new analysis. In the current study, we leverage previously collected data from 198 non-target species to place biologically informed constraint on the distribution of a plant 190 trait parameter. When additional data for a specific species or plant functional type is 200 available, priors are further constrained before being used as model parameters. When no 201 additional data are available, these priors are used directly to parameterize the model. 202 For the P. virgatum example described below, priors were set using data from all plant 203 species, from only grass species, or from just C4 grass species depending on availabile 204 data. Sources of this prior information included data from previous and ad-hoc syntheses, 205 expert knowledge, and biophysical constraints (Table 1). 206 Prior distributions used in the meta-analysis were fit to one of four types of information: 207 1) data from multiple species, 2) the posterior predictive distribution for a new species 208 from a meta-analysis of data (when error estimates were available), 3) a central tendency 200 informed by data with expert constraint on the confidence interval, or 4) expert 210 constraints on both the central tendency and confidence intervals. In case number 2, the 211 across-species meta-analysis "posterior" informs the prior for the species-level 212 meta-analysis. In all cases, maximum likelihood estimation was used to fit a prior 213 distribution. When more than one candidate distribution was considered, Aikake's 214 Information Criterion (AIC) was used to select the best fit distribution. The choice of 215 prior was confirmed by visually inspecting the prior density functions overlain by data or expert constraints (Figure 2). 217

Meta-analysis 218

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A Hierarchical Bayes meta-analytical model (Figure 3) formally synthesizes available trait 219 data from multiple studies while accounting for various sources of uncertainty. This 220 Hierarchical Bayes approach integrates prior information and provides a flexible approach 221 to variance partitioning and parameter estimation. 222 The meta-analytical framework is useful for summarizing data sets that include summary 223 statistics. The trait data queried by PEcAn consist of a trait name, sample mean, sample 224 size, and a sample error statistic. PEcAn transforms error statistics to exact or 225 conservative (i.e., erring toward inflating the variance) estimates of precision ($\tau = 1/SE^2$) 226 (Appendix C). 227 The sample mean is drawn from a normal distribution:

$$Y_k \sim N(\Theta_k, \tau_k) \tag{1}$$

Where Y_k is the sample mean of the k^{th} unique site by treatment combination (sample unit), Θ_k is the unobserved 'true' value of the trait for the k^{th} sample unit. 230 The meta-analysis partitions trait variability into among site, among treatment, and 231 within-unit variance. The unobserved 'true' trait mean Θ_k is a linear function of the 232 global trait mean, β_0 plus random effects for study site (β_{site_i}) and treatment $(\beta_{\text{tr}|\text{site}_i,j})$ 233 and a fixed effect for greenhouse (β_{gh}) :

$$\Theta_k = \beta_0 + \beta_{\text{site}i} + \beta_{\text{tr}|\text{site}ij} + \beta_{\text{gh}}I(i)$$
 (2)

Where i indexes study site, j indexes each treatment within a study, and I(i) is an 235 indicator variable set to 0 for field studies and 1 for studies conducted in a greenhouse, 236 growth chamber, or pot experiment. The parameter used in the ecosystem model is the 237 posterior estimate of the global mean trait value, β_0 . β_0 , has an informed prior functional 238

form and parameter specification that varies by trait and species or PFT. Methods used 239 to elicit priors for the present study are provided in the Application section under Priors. 240 The "site" random effects (β_{site}), accounts for the spatial (among-site) heterogeneity of a 241 parameter. The "treatment" random effect ($\beta_{\text{tr|site}}$) accommodates differences among 242 experimental treatments. These random effects of treatment and site are assumed to be 243 Normally distributed with zero mean and they have diffuse Gamma priors on precision 244 $\tau_{\rm site}$ and $\tau_{\rm tr}$. Control treatments and observational studies have $\beta_{\rm tr|site} = 0$. PEcAn 245 dynamically adjusts the meta-analysis model specification to include terms for each level 246 of site and treatment, or greenhouse studies as required by available data. To ensure that 247 the prior on precision remains sufficiently diffuse when the magnitude of a parameter is 248 small, the scale parameters in the gamma priors on random effect precision terms ($\tau_{\rm site}$ 249 and $\tau_{\text{tr|site}}$) are scaled to $(\bar{\beta_0}^2/1000)$ when the prior on β_0 has a mean $\bar{\beta_0} < \sqrt{10}$. 250 A "greenhouse" fixed effect β_{gh} accounts for potential biases associated with plants grown 251 in a greenhouse, growth chamber, pot, or other controlled environment. This "greenhouse" 252 effect, $\beta_{\rm gh}$, has a diffuse Normal prior with a mean of zero and a precision of 0.01. 253 The observation precision (precision = 1/variance) for the k^{th} sample mean, τ_k , is 254 determined based on the within-unit precision, τ_Y , and the sample size, n, as $\tau_k = n \times \tau_Y$ 255 (since $SE = SD/\sqrt{n}$). A common within sample unit precision, τ_Y , is assumed in order 256 to accommodate literature values with missing sample sizes or variance estimates. The 257 sample standard error, se_k , is drawn from a Gamma distribution with parameters 258 informed by the sample size, n, and within-site precision, τ_Y : 259

$$\frac{1}{n \times se_k^2} \sim \text{Gamma}(\frac{n}{2}, \frac{n}{2\tau_Y}) \tag{3}$$

 au_Y has a diffuse gamma prior. Unlike the mean and variance parameters, missing values of n cannot be estimated and are conservatively set either to 2 (when existence of a

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variance estimate indicates n \geq 2) or to 1 (if no variance estimate is given).
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    The random and fixed effects and the among study, among treatment, and within-unit
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    precisions are used to evaluate the importance of different sources of uncertainty.
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    The meta-analysis module in PEcAn is fit using JAGS software (version 2.2.0, (Plummer,
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    2010)) called from within R code that handles data manipulations and meta-analysis
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    model specification in JAGS. JAGS uses standard Markov Chain Monte Carlo (MCMC)
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    methods (Gelman and Rubin, 1992) to approximate the posterior distribution of the
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    terms in the meta-analysis. To overdisperse the n MCMC chains, initial values of \beta_0 are
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   set to the \frac{1}{n+1}, \dots, \frac{n}{n+1} quantiles of the prior on \beta_0; for the default n=4 chains, this
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    would be the {0.2, 0.4, 0.6, 0.8} quantiles. Following Gelman and Shirley (2011), PEcAn
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    discards the first half of each chain, thins each chain to 5000 samples and then combines
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    the chains into a single vector of samples for each term in the meta-analysis model. Trace
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    plots and the Gelman-Rubin convergence diagnostic (Gelman and Rubin, 1992) are used
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    to assess chain convergence. Density plots (Figure 4) are used to visually compare the \beta_0
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    chain to data and priors. The significance of the greenhouse effect is evaluated by
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   calculating a two-sided probability that \beta_{gh} \neq 0.
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    When species-level data are unavailable, the posterior distributions are equivalent to the
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    priors.
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    Each term in the meta-analysis is represented as a vector of MCMC samples from the
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    posterior distribution. Statistical summaries of the parameters can easily be calculated
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    from these chains, and chains can also be directly sampled for use in ecosystem model
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    parameterization. When the \beta_0 chains are sampled for the ecosystem model ensemble,
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the meta-analysis posteriors become the model ensemble priors.

285 Model Analysis

286 Ensemble Analysis

Typically, ecosystem models are run for a single model parameterization. For example, 287 the model could be evaluated at the median value of each parameter. However, this 288 approach only provides a point estimate with no accounting for parameter uncertainty. 289 To propagate parameter uncertainty through the ecosystem model, PEcAn uses standard 290 ensemble-based Monte Carlo approaches. An ensemble of model runs is a set (e.g. 500 or 1000) of model runs that are parameterized by sampling from the trait parameter 292 distributions. For each ensemble member, parameter sets are sampled from the full joint 293 parameter distribution of β_0 , the vector of all model parameters. As a result, the model 294 ensemble approximates the posterior distribution of the ecosystem model output. The 295 model ensemble produces a posterior distribution of the ecosystem model output that can be summarized with standard statistics (e.g. mean, standard error, and credible interval). 297

298 Sensitivity Analysis

Sensitivity analyses are used to understand how much a change in a model parameter affects model output; sensitivity is the derivative, $df/d\beta_{0t}$, of the model (f) with respect 300 to the estimate of β_0 for trait t. PEcAn approximates the sensitivities based on univariate 301 perturbations of model parameters. These approximations are necessary because 302 analytical solutions for sensitivity are not tractable for most ecosystem models, and 303 PEcAn is designed to be flexible and applicable to any such model. One disadvantage of 304 traditional perturbation-based sensitivity analyses is that the perturbations are usually 305 arbitrary, for example varying each parameter by a fixed percentage of its value 306 (Larocque et al., 2008) rather than over a meaningful range of the parameter. These 307 traditional approaches make interpretation of sensitivities difficult because they fail to 308

acknowledge the distribution or uncertainty of each parameter. In this regard, PEcAn 309 offers a distinct advantage over traditional sensitivity analyses because parameters are 310 varied based on the meta-analysis posterior parameter distributions. 311 Based on initial exploratory analyses, we found a local perturbation to be inadequate for 312 capturing the responses in most parameters so we instead estimate sensitivities using a 313 global univariate sensitivity analysis. By default, PEcAn evaluates each parameter at the 314 posterior median and at the six posterior quantiles equivalent to $\pm [1, 2, 3]\sigma$ in the 315 standard normal while holding all other variables constant at their posterior median. The 316 relationship between model output and each model parameter β_{0t} is then approximated 317 by a natural cubic spline $g_t(\beta_{0t})$ that interpolates through the evaluation points. The 318 model sensitivity to each parameter is approximated by the derivative of the spline 319 $(dg_t/d\beta_{0t})$ at the parameter mean. In addition to the sensitivity analysis, this set of 320 spline approximations is used in the variance decomposition, in partitioning residual 321 variance, and in evaluating the effect of ensemble size on the estimate of model variance. 322 To facilitate comparisons among the trait sensitivities, despite differences in the units on 323 different traits, we tabulate the coefficient of variation (normalized parameter variance) 324 and the elasticity (sensitivity with terms df and $d\beta_{0t}$ standardized by the mean model 325 output and parameter mean respectively).

27 Variance Decomposition

Variance decomposition aims to explain how much each input parameter contributes to
uncertainty in model output (Cariboni et al., 2007). Although the present analysis
focuses on model parameters, these methods can be extended to address uncertainty in
initial conditions or model drivers.

The Delta Method uses Taylor series expansion to approximate the probability
distribution of a continuous function of random variables (Oehlert, 1992; pp. 240–245 in

Casella and Berger, 2001). In this study, the model output $f(\beta_0)$ is a function of a vector of the full set of parameters. After approximating the distribution of $f(\beta_0)$, it is possible to estimate the variance of the model output. The first step is to derive the Taylor series approximation of the variance of a function (Casella and Berger, 2001, equation 5.5.9 in):

$$Var(f(\boldsymbol{\beta_0})) \approx \sum_{t=1}^{m} Var\left(f(\overline{\beta_{0t}}) + \frac{df}{d\beta_{0t}}(\beta_{0t} - \overline{\beta_{0t}}) + \ldots\right)$$
(4)

$$= \sum_{t=1}^{m} \left(\frac{df}{d\beta_{0t}}\right)^2 Var(\beta_{0t}) + \omega \tag{5}$$

where m is the number of parameters in the model, and the error term ω accounts for higher order terms in the Taylor series, and β_{0t} is the estimate of β_0 from the 330 meta-analysis (equation 2) for each trait, t. 340 With this approximation, it is straightforward to estimate the variance contributed by each parameter. The terms in this form of the variance decomposition can be estimated 342 directly from the preceding analyses: $Var(f(\beta_0))$ is the variance of the model ensemble; 343 $Var(\beta_{0t})$ is the posterior variance of trait β_{0t} from the meta-analysis (equation 2); and $df/d\beta_{0t}$ is the model sensitivity at the parameter mean $\overline{\beta_{0t}}$. The resulting assertion is that the variance of model output is equal to the sum over the variance of each trait times its sensitivity squared plus a closure term, ω . 347 We found that the traditional Taylor polynomial approach to variance decomposition 348 produced a poor closure of the total variance of the model output: for more sensitive 349 parameters, a linear approximation of $f(\beta_0)$ provided unrealistic estimates of the sensitivity function that overestimated variance. Increasing the order of the Taylor series 351 expansion actually exacerbated this problem (results not shown). One problem with the 352 polynomial approximation is that, unlike polynomials, most response variables in 353 ecosystems and ecosystem models tend to be asymptotic at both high and low values of a 354 trait. For example, when assessing aboveground biomass there is a lower bound of zero 355

biomass and most parameters become progressively less sensitive, if not genuinely
asymptotic, at their upper bound. This asymptotic behavior is poorly approximated by a
polynomial because polynomials are unbounded at extreme parameter values. Therefore,
we sought a better approximation for the variance decomposition.

First, we formulated a more generalized form of the variance decomposition (equation 4):

$$Var(f(\boldsymbol{\beta_0})) = \sum_{t=1}^{m} Var(g_t(\beta_{0t})) + \omega$$
 (6)

The spline $g_t(\beta_{0t})$ is a statistical emulator of the model response to trait t that transforms β_{0t} from the parameter domain to the model domain. The univariate 362 contribution of each parameter to variance of the model output is thus $Var(g_t(\beta_{0t}))$. 363 Equation 6) only requires β_{0t} from the preceding meta-analysis, $g_t(\beta_{0t})$ from the sensitivity analysis, and $Var(f(\beta_0))$ from the ensemble analysis. 365 The final term, ω , is the closure between the right hand side and the left hand side of the 366 variance decomposition; ω represents the effects of the higher order terms in the Taylor 367 approximation and the covariance terms between parameters. This closure term is 368 intended to represent parameter interactions that are excluded from the univariate variance decomposition (equation 6). Negative trade-offs among physiological traits 370 would result in ω less than zero. However, our estimate of ω also includes errors 371 associated with using finite sample sizes, the spline approximation in each $g_t(\beta_{0t})$, and 372 biological range restrictions on model output that are not reflected in the variance 373 decomposition (equation 6). 374 One approach to partition the error in the closure term is to use the univariate spline 375 functions from the sensitivity analysis to estimate what the model output would be for 376 each of the parameter sets used in the model ensemble; we call this estimate the "spline

378 ensemble":

$$\boldsymbol{g}_{\ell}(\boldsymbol{\beta}_{0}) = \boldsymbol{g}(\hat{\boldsymbol{\beta}}_{0}) + \sum_{t=1}^{m} \left(g_{t}(\beta_{0t\ell}) - g_{t}(\hat{\beta_{0t}}) \right)$$
 (7)

In this equation, $g_{\ell}(\beta_0)$ is the spline estimate of the model output for the ℓ^{th} ensemble member and $\hat{\beta}_{0t}$ is the posterior median parameter value. 380 Although the individual splines may respect range restrictions on output variables (e.g. 381 biomass values cannot fall below zero), combinations of the splines evaluated for a set of unfavorable traits can fall outside these ranges. For parameter sets that give a 383 biologically implausible estimate of negative biomass $(g_{\ell}(\beta_0) < 0)$, the estimate is set to 384 zero. The only difference between the variance of the spline ensemble (equation 7) and 385 the variance decomposition (equation 6) is that range restrictions are not corrected for in 386 the variance decomposition. Therefore, the spline ensemble allows us to estimate the effect of using combinations of spline estimates that do not respect the zero bound on biomass in the variance decomposition. The difference between the model ensemble and 389 the spline ensemble provides an estimate of parameter interactions in the model because 390 the spline ensemble does not include the parameter interactions that exist in the model. 391 The precision of the estimate of model ensemble variance is affected by the number of 392 runs in the ensemble. When the computational expense of the model itself limits the 393 ensemble size, there can be significant uncertainty in the estimate of ensemble variance. 394 The uncertainty in a sample variance is estimated as 395

$$Var(s^2) = \frac{1}{n} \left(\mu_4 - \frac{n-3}{n-1} \sigma^4 \right)$$
 (8)

Mood et al., 1974, , p 239) where μ_4 is the fourth central moment. $Var(s^2)$ scales inversely with sample size. The effect of the limited model ensemble size on uncertainty in the estimate of ensemble variance is measured in two ways. The first way is to calculate $Var(s^2)$ for the model ensemble (n = 500). The second way is to compare

- $Var(s^2)$ of the spline ensemble with 500 and 10,000 runs. The 95%CI for s^2 is calculated
- 401 as $s^2 \pm 1.96 s_{s^2}$ where $s_{s^2} = \sqrt{Var(s^2)}$.
- The errors introduced from using a spline approximation of the model response can not
- be estimated based on the existing output, but it is small in comparison to the other
- effects given the range restrictions imposed by the spline interpolation.
- The results of a model ensemble are posterior estimates of aboveground biomass.
- However, we also distinguish between ensembles depending on the nature of model
- parameters. First, we ran a "prior model ensemble" using an ensemble of parameter sets
- drawn from prior distributions, and then a "posterior model ensemble" drawn from
- 409 meta-analysis posteriors.

410 Application: Switchgrass Monoculture

- We demonstrate the application of PEcAn to estimate the aboveground yield of an
- experimental switchgrass (*Panicum virgatum*) monoculture. The first step to applying
- PEcAn was to construct an appropriate set of priors based on data syntheses and expert
- knowledge. These priors were conservative estimates of the plant trait parameters based
- on information other than species level data. Next, switchgrass trait data from both
- 416 previous studies and field measurements were summarized using meta-analysis to
- constrain the prior parameter estimates. The Ecosystem Demography model version 2.1,
- (Medvigy et al., 2009; Moorcroft et al., 2001) was used to simulate plant growth.
- The model ensemble and sensitivity analysis were performed using both the prior and
- posterior parameter estimates. By comparing the prior model ensemble to the posterior
- model ensemble, we are able to evaluate the ability of species level data to reduce model
- 422 uncertainty.
- To evaluate model performance, we compare the ensemble estimates of aboveground

biomass with observed yields (Heaton et al., 2008; Wang et al., 2010, Figure 5).

425 Site

Switchgrass (Panicum virgatum) is a perennial grass native to North America that has 426 received attention as a potential cellulosic biofuel crop (McLaughlin and Kszos, 2005; 427 Wang et al., 2010). We modeled the aboveground biomass production of a switchgrass monoculture and compared model estimates to a monoculture planted in 2002 at the 420 University of Illinois Agricultural Research and Education Center in Urbana, IL 430 (40.09 N, 88.2 W). The climate at this site is characterized by hot, humid summers and 431 cold winters with a 50 year (1959-2009) mean annual temperature of 11 °C and mean 432 annual precipitation of 1000 mm yr⁻¹ (Angel, 2010). Meteorological data used to drive 433 the model were downloaded from the North American Regional Reanalysis (Mesinger 434 et al., 2006). Soil is a silt loam from the Drummer-Flanagan soil series; texture data was 435 obtained through the USDA NRCS web soil survey website (websoilsurvey.nrcs.usda.gov). 436 The yield and other aspects of this ecosystem have previously been reported (Heaton et al., 2008). 438

439 Ecosystem Demography Model

We used the Ecosystem Demography Model, version 2 to model the productivity and soil
carbon pools in this switchgrass agro-ecosystem. ED2 is a terrestrial biosphere model
that couples age- and stage-structured plant community dynamics with ecophysiological
and biogeochemical models. The biophysical land-surface model in ED2 allows plant
uptake and growth to respond dynamically to changes in weather and soil hydrology
(Medvigy et al., 2009). ED2 has the ability to link short-term, physiological responses to
environmental conditions with realistic, long-term successional changes in ecosystem
structure and composition (Moorcroft et al., 2001). While other models have both

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succession and physiology, ED2 also has explicit spatial scaling, a sub-daily time-step,
    and the ability to couple with to a land surface model (Dietze and Latimer, 2011).
440
    ED2 incorporates a mechanistic description of plant growth that accounts for the fast
450
    temporal responses of plants to changes in environmental conditions. In this study, we
451
    vary fifteen model parameters based on observable plant traits that control carbon
452
    uptake, carbon allocation, turnover, and reproduction (Table 1, Figures 2, 4).
453
    ED2 calculates photosynthetic rates using the enzyme kinetic model developed for C3
454
    plants (Farguhar and Sharkey, 1982; Ball et al., 1987) and the modifications for C4
455
    (Collatz et al., 1992). V<sub>c,max</sub> sets the upper bound on the rate of Rubisco-limited
456
    photosynthesis, while light limited photosynthesis is constrained by the quantum
457
    efficiency parameter, and a threshold parameter controls the minimum temperature at
458
    which photosynthesis will occur. Stomatal conductance is calculated using the Leuning
459
    variant of the Ball-Berry model (Leuning, 1995) and is controlled by the stomatal slope
460
    parameter. Leaf boundary layer conductance depends on the leaf width parameter.
461
    Together, stomatal conductance and leaf boundary layer conductance affect carbon and
462
    moisture fluxes and the leaf energy balance. Specific leaf area (SLA) determines the
463
    amount of leaf area produced per unit leaf biomass investment.
464
    In addition to photosynthesis, ED2 also accounts for carbon allocation to growth,
465
    respiration, and for the turnover rate of carbon pools. These parameters include: one to
466
    partition between leaf and fine root growth; one for allocation to reproduction; two
467
    respiration parameters associated with growth respiration and root maintenance
468
    respiration; and two parameters to control the rates of leaf and root turnover.
469
    Finally, three demographic parameters control seed dispersal, seedling mortality, and
470
    adult mortality due to carbon limitation (Table 1).
471
```

472 Priors

473 Priors from data

Priors were estimated by finding the best fit distribution to raw data sets include SLA 474 and leaf turnover rate from the GLOPNET database (Wright et al. (2004), n = 125, 40475 respectively), root turnover rate (Gill and Jackson (2000), n = 66), and quantum yield 476 (Skillman (2008), n = 56). Candidate distributions for these priors were Gamma, 477 Weibull, log-Normal, and F because each of these traits is bound at zero. In all cases we are interested in using the full distribution of across-species data as our prior constraint 479 on what one individual species is capable of doing, as opposed to using the estimate of 480 the mean of this distribution as our prior. 481 Quantum yield data represent a survey of published values of quantum yield in C4 482 monocots (Skillman, 2008); original data were provided by the author and restricted to 483 measurements made under photorespiratory conditions (ambient CO_2 and O_2) (J. 484 Skillman, personal communication). Given the narrow range of data (CV = 11%), the 485 normal distribution was also considered but was not the best fit. 486

Priors from meta-analysis

We used meta-analysis to calculate a prior from data when summary statistics and
sample sizes were available. The meta-analysis model used to calculate prior distributions
is similar to the one used by PEcAn to summarize species-level data (equation 2), with
three differences. First, there are no site, treatment, or greenhouse effects. Second, data
from multiple species were used. Third, we generated a posterior predictive distribution
to predict the distribution of trait values for an unobserved C4 plant species, unlike the
species-level meta-analysis, which estimated the global mean parameter value. Thus, the
model included plant functional type (PFT) as a random effect:

$$\Theta_{\text{species}} = \beta_0 + \beta_{\text{PFT}} \tag{9}$$

Stomatal slope is the empirical slope coefficient in the (Leuning, 1995) model of stomatal conductance. Estimates of this parameter are sparse, so we collected new data for this 497 study. Stomatal slope was estimated using measurements of four leaves from each of five 498 field-grown energy crop species during the 2010 growing season (Appendix A). The five 499 species included two C4 grasses: Miscanthus (Miscanthus x giganteus) and Switchgrass 500 (P virgatum) planted in 2008 and three deciduous tree species: Red Maple 501 (Acer rubrum), Eastern Cottonwood (Populus deltoides, and Sherburne Willow 502 Salix x Sherburne) planted in 2010 as 2 year old saplings. All plants were grown at the 503 Energy Biosciences Institute Energy Farm (40°10'N, 88°03"W). We used the data from 504 the three tree species and Miscanthus to calculate the posterior predictive distribution of 505 an unobserved C4 grass species, and used this distribution as the prior estimate for 506 Switchgrass stomatal slope. 507 Maximal carboxylation rate (V_{cmax}) data consists of ninety-four C3 species (Wullschleger, 508 1993) plus three C4 species (Kubien and Sage, 2004; Massad et al., 2007; Wang et al., 509 2011). To express V_{cmax} at a common temperature of $25^{\circ}C$ for all species, we applied an 510 Arrhenius temperature correction (Appendix C). The Wullschleger (1993) data set 511 included a 95% CI and an asymptotic error calculated by the SAS nlin proceedure (Stan 512 Wullschleger, personal communication). We used the asymptotic error as an estimate of 513 SE, the 95% CI to estimate SD $\left(\text{SD} = \frac{\frac{1}{2}\text{CI}}{1.96}\right)$, and then estimated n as $\hat{n} = \left(\frac{\text{SE}}{\text{SD}}\right)^2$. Plant 514 species were classified into five functional types: C3 grass, C4 grass, forb, woody 515 non-gymnosperm, and gymnosperm based on species records in the USDA PLANTS 516 Database (USDA and NRCS, 2011). Ambiguous species (those with both forb and woody 517 growth forms, n = 15) were excluded.

Leaf width data represent 18 grass species grown in a common garden greenhouse 519 experiment (Oyarzabal et al., 2008). P. virgatum was among the 18 species, and was 520 excluded from the prior estimation but used as raw data in the meta-analysis. The 521 remaining seventeen species were divided into C3 and C4 functional types. Relative to 522 the small sample of C4 grasses, switchgrass leaf width was an outlier; inflating the 523 variance four-fold reduced the prior information content to account for this descrepency. 524 Root respiration rate values were measured on thirty-six plants representing five 525 functional types, including six C4 grass species (Tjoelker et al., 2005). As before, P. 526 virgatum data was excluded from the prior estimation and used as raw data in the 527 species-level meta-analysis.

Priors from limited data and expert knowledge

When available data were limited to a few observations, these were used to identify a 530 central tendency such as the mean, median, or mode, while expert knowledge was used to 531 estimate the range of a confidence interval. An optimization approach was used to fit a 532 probability distribution to this combination of data and expert constraint. 533 In order to estimate the fine root to leaf ratio for grasses, we assume fine roots account 534 for all belowground biomass (Jackson et al., 1997) and that leaves account for all above ground biomass. Roots account for approximately 2/3 of total biomass across temperate 536 grassland biomes (Saugier et al., 2001, Table 23.1), so we constrained a beta prior on the 537 root fraction to have a mean of 2/3 by setting $\alpha = \beta/2$ since the mean of a beta is 538 defined as $\frac{\alpha}{\alpha+\beta}$. To convert from proportion to ratio, we used the identity: if 539 $X \sim \text{Beta}(\frac{\alpha}{2}, \frac{\beta}{2}) \text{ then } \frac{X}{1-X} \sim F(\alpha, \beta) \times \frac{\alpha}{\beta}.$ We constrained the 95%CI = [1/3, 10/11],equivalent to a fine root to leaf ratio with a mean fixed at two and a 95%CI = $[^{1}/_{2}, 10]$. 541 We simulated the distribution of the fine root to leaf ratio by drawing 10000 samples 542 from a $F(2\alpha, \alpha)$ distribution and multiplying these samples by two.

Seed dispersal in ED2 represents the proportion of seed dispersed outside of a 7.5m radius plot, which we approximate as a beta distribution. Although no direct measurements of 545 seed dispersal were available, it was straightforward to parametrize a ballistic model of 546 seed dispersal (Ernst et al. (1992), from Creemer 1977): $D = \frac{V_w H}{V_t}$. This model relates dispersal distance D to terminal velocity V_t , wind speed V_w , and seed height H. Although more sophisticated treatments of dispersal exist and are important for estimating low 549 probability long distance dispersal events (Clark et al., 1999; Thompson and Katul, 550 2008), the Ernst et al. (1992) model is sufficient for relatively short dispersal distances 551 required in the present context. 552 Values of terminal velocity, V_t , of grass seeds were taken from two references, (Ernst 553 et al., 1992; Jongejans and Schippers, 1999) and these data were best described as 554 $V_t \sim \text{Gamma}(2.93, 1.61).$ 555 Next the heights from which the seeds are dropped was estimated from calibrated 556 photographs of reproductively mature switchgrass at a field site in Urbana, IL: 557 $H \sim N(2, 0.5)$. Finally, wind speed observed at the site were fit to a Weibull distribution 558 (Justus et al., 1978). $V_w \sim \text{Weibull}(2.4, 0.712)$ (Marcelo Zeri, unpublished wind and 559 height data). Given these distributions of V_w , H, and V_t , sets of 100 dispersal distances 560 were simulated 10000 times to calculate the fraction of seeds in each simulation dispersed 561 beyond 7.5m, 562

⁵⁶³ Priors informed by expert knowledge

When no data were available, expert knowledge was used to estimate the central tendency and confidence interval for a trait parameter. Again, optimization was used to fit a probability distribution to these constraints.

The minimum temperature of photosynthesis for C4 grasses was given a prior based on expert knowledge with a mean of 10°C and a 95%CI = [8, 12]°C that fits a normal

- $(\mu = 10, \sigma = 1.02)$ distribution (Don Ort, UIUC, personal communication, 2010).
- 570 The growth respiration factor is the proportion of daily carbon gain lost to growth
- respiration. Because it is a proportion, the beta distribution was fit with a mean set
- equal to the ED2 default parameter value, 0.33 and a 95%CI = [0.05, 0.60], conservatively
- based on the range of construction costs reviewed by Amthor (2000).
- 574 Seedling mortality factor represents the proportion of carbon allocated to reproduction
- that goes directly to the litter pool. Given the default ED2 parameter is 0.95, we stated a
- beta prior with a median at 0.95, and a 95%CI = $[^2/_3, 1]$.
- 577 The mortality factor in ED2 is the rate parameter in the negative exponential
- relationship between carbon balance and mortality (Medvigy et al., 2009). The default
- parameter for all plant functional types (PFT's) in ED2 is 20, and our weakly informed
- gamma prior sets this as the median and gives a 95%CI = [5, 80].
- Reproductive allocation represents the proportion of carbon in the storage pool allocated
- to reproduction. This parameter is a proportion and has a default value of 0.33 in ED.
- The Beta(2,4) distribution has a mean of $^{1}/_{3}$ and a 95%CI = [0.05, 0.72] representing
- relatively high uncertainty. This distribution implies that the plant may allocate any
- fraction of the carbon pool to reproduction between but not equal to 0 and 1 and has an
- 80% probability that less than half of the carbon pool will be allocated to reproduction.

587 Switchgrass Trait Meta-analysis

- 588 Switchgrass trait data used to constrain model parameters are stored in the Biofuel
- Ecophysiological Trait and Yield database (BETYdb, www.betydb.org), a database
- designed to support research on biofuel crops. BETYdb includes both previously
- published and primary data (Appendix A). Prior to entry in the database, data were
- 592 converted to standard units chosen for each variable (Table 1).
- Trait data available for *Panicum virgatum* include V_{cmax}, SLA, leaf width, fine root to

leaf ratio, root respiration, stomatal slope, and root turnover rate (Figure 4, Table 2).

Methods used to collect these data and site descriptions are available in the source

references (Appendix A). Each meta-analysis was run with four 50,000 step MCMC

chains.

598 Model Analysis

We executed a ten-year, 500 run ensemble of ED2 using parameter values drawn from the prior or posterior parameter distributions. The model was run for the years 1995-2006 to 600 simulate the field trials conducted by Heaton et al. (2008). The model output of interest 601 was the December mean aboveground biomass (AGB) during the years 2004–2006, 602 simulating the yields of the mature stand (Heaton et al., 2008). The ensemble estimate 603 was also compared to the larger set of all reported switchgrass yield data (Wang et al., 604 2010). 605 Runs resulting in yields less than 2 Mg/ha were considered non-viable parameter 606 combinations. To test if prior and posterior parameter sets resulted in different fractions of non-viable runs, we estimated the posterior probability of a non-viable run as a 608 binomial posterior from a beta-binomial model with a flat (Beta(1,1)) prior. Then, we 609 calculated the two-tailed probability that the difference between these binomial posteriors 610 was $\neq 0$. 611

Results

613 Trait Meta-analysis

Switchgrass data were collected from the literature and field for seven of the model parameters: specific leaf area (SLA) (n = 24), leaf width (39), V_{cmax} (4), fine root to leaf allocation ratio (4), stomatal slope (4), root respiration rate (1), and root turnover rate

(1). Table 2 summarizes the meta-analysis for each of these parameters, including the 617 posterior mean and 95% CI of the global mean, the fixed greenhouse effect, and each of 618 the variance components (reported as standard deviations). 619 SLA and leaf width data were from from multiple sites, but the meta-analysis provided 620 no evidence for among site variability in excess of within site variability (σ_Y and σ_{site} , 621 respectively, in Table 2). For the remaining traits, there was insufficient spatial sampling 622 to assess site to site variability. Greenhouse growing conditions had a positive effect on 623 both SLA (P = 0.027), and leaf width (P = 0.052). 624 Figure 4 compares parameters before and after incorporating data in the meta-analysis. 625 A reduction in parameter uncertainty is seen as the reduction in the spread of the posterior (black) compared to the prior (grey) parameter distributions. The influence of 627 the prior information on the posterior distribution increased when the prior was more 628 constrained or when less data were available for use in the meta-analysis. For example, 629 data substantially constrained the uncertainty in the V_{cmax} and SLA posteriors relative to 630 the priors. By contrast, there was little effect of additional data on the parameter 631 estimates for fine root to leaf allocation and root respiration rate; these parameters had 632 relatively well constrained priors and limited species-specific data. 633

Model Analysis

635 Ensemble

Within the model ensemble analysis (Figure 5), both the prior and posterior
parameterizations produced yield estimates that were consistent with yields observed at
the Urbana site for which the model was run (Heaton et al., 2008) and with 1902
previously reported yields of switchgrass (Wang et al., 2010). In both the prior and
posterior ensembles, the predicted aboveground biomass was clearly bimodal. These two

modes had little overlap and a distinct break at two Mg/ha. We infered that the first peak represents non-viable plants generated by unrealistic parameter sets so plants with 642 aboveground biomass less than two Mg/ha were considered "non-viable". When 643 summarizing the model output, we consider viable and non-viable ensemble members separately; all runs are considered in the senstivity analysis and variance decomposition. A greater percentage of runs in the prior ensemble fell below this threshold (53.4 vs 36.6, 646 $P \simeq 0$). 647 Compared to the prior ensemble prediction, the data-constrained posterior runs had lower 648 median yields and a more constrained 95% credible interval (16.5[7.2, 37] Mg/ha vs 25[7.7, 56] Mg/ha). This reflects the substantial shrinkage of the posterior relative to the prior SD estimates of model output uncertainty (from $\sigma = 19.7$ to $\sigma = 11.9$). In 651 particular, the upper tail of the modeled yield was reduced toward the observed yields. 652 Despite the reduction in ensemble uncertainty, the ensemble posterior yield was still 653 relatively imprecise and had much greater uncertainty than the field trial (Heaton et al., 2008, $\sigma = 4.1$) or the meta-analysis of all observations (Wang et al., 2010, ($\sigma = 5.4$)). 655 The spline ensemble viable plants had a median 18.8[2.9, 48] and $\sigma = 13$. 656

657 Sensitivity Analysis

Sensitivity analysis demonstrated that traits varied in their effect on on aboveground 658 biomass (Figure 6), and many of these relationships are clearly non-polynomial. For 659 example, parameters associated with photosynthesis and carbon allocation - including 660 V_{cmax} , SLA, growth respiration, and root allocation - were particularly sensitive. For 661 particularly sensitive parameters, the sensitivity functions had coverage of unrealistic 662 yields greater than 30 Mg/ha. Constraining SLA and V_{cmax} parameters with data 663 resulted in a more realistic range of yields. On the other hand, aboveground biomass was 664 largely insensitive to leaf width, seed dispersal, and mortality rate. 665

Variance Decomposition

The variance decomposition showed that data-constrained parameters substantially 667 reduced their contribution to overall model variance (Figure 7). Prior to including 668 species-specific trait data, SLA contributed the most to model uncertainty, followed by growth respiration, fine root allocation, V_{cmax}, seedling mortality, and stomatal slope 670 (right panel, grey bars Figure 7). Incorporating species level data substantially reduced 671 the contributions of SLA, V_{cmax}, seedling mortality, and stomatal slope to model 672 uncertainty. For example, SLA fell from first to fourth and stomatal slope fell from sixth 673 to fourteenth in rank contribution to ensemble variance. Although the addition of data 674 reduced parameter uncertainty for fine root to leaf allocation, aboveground biomass was 675 more sensitive to this parameter at the posterior median. These changes cancelled each 676 other out, and as a result the contribution of the fine root allocation parameter to 677 ensemble variance remained constant. The variance of the ensemble was less than the variance calculated in the variance 679 decomposition, and this difference is the closure term, ω . The closure term accounted for 680 approximately 28.455981108897% of the variance decomposition estimate (Table 3b). 681 There was no effect of increasing the sample size from 500 to 10000 on the variance 682 estimates (Table 3a). 683

Discussion

Switchgrass Trait Meta-analysis

When species-level data were available, the meta-analysis constrained estimates of the trait mean parameter (Figure 4) and provided insight into the sources of parameter uncertainty (Table 2). In the context of constraining model parameters, we were

interested in accounting for but not directly investigating the specific effects of site, treatment, or greenhouse effects. However, we can use the meta analysis results to 690 identify sources and scales of parameter variability. This insight into parameter 691 variability helps inform future sampling designs, development of the ecosystem model, 692 and improvement of methods used to parametrize the ecosystem model. 693 Where data from multiple sites were available, we could evaluate the relative importance 694 of within versus among-site variance for the range of sites represented in the data 695 (Table 2). Recent studies demonstrate important effects of intraspecific trait variability 696 on ecosystem functioning (Breza et al., 2012; Albert et al., 2011; Violle et al., 2012). 697 Therefore, for traits that do exhibit greater variability across than within sites, explicit 698 inclusion of spatial, environmental, and even genetic information into the meta-analytical 690 model would be justified. This approach would enable the estimation of site-specific 700 parameters for use in the ecosystem model and will be investigated in future development 701 of the meta-analysis module. 702 For the other parameters (V_{cmax} , fine root allocation, root respiration rate, and root 703 turnover rate) data came from one site, so it is not possible to estimate the across-site 704 variability. For these traits, obtaining data from additional sites would better constrain 705 both the global mean and the across-site variance. This additional data collection is 706 particularly justified for traits that contribute most to the uncertainty in the model 707 ensemble. 708

$_{\scriptscriptstyle{09}}$ Model Ensemble

Despite the large reduction in model uncertainty from the prior to the posterior model
ensemble, the uncertainty in projected yield is substantial (Figure 5) and further
constraint would increase the utility of this model output. However, the explicit
accounting of parameter uncertainty is an important first step toward producing more

informative model output. If model parameters had been treated as fixed constants, we 714 would have no estimate of model uncertainty; without an estimate of uncertainty, the 715 similarity between the modeled (16.5 Mg/ha) and observed (12.0 Mg/ha) median yields 716 would be difficult to interpret; a naive interpretation could create false confidence in the 717 model. Including the non-viable plants would have made the model mean more accurate 718 (Figure 5), but the 90%CI would have been less accurate, containing the possibility that 719 switchgrass would not grow in Champaign County, Illinois, even though extensive 720 research (Heaton et al., 2008; VanLoocke et al., 2012, personal observation) demonstrates 721 that it does grow very well in this area. 722 The reduction in median modeled yield in the posterior relative to the prior model 723 ensemble 5 is consistent with the reduced probability of high SLA and V_{cmax} values in the 724 posterior relative to the prior distributions. As expected, the use of switchgrass trait data 725 to inform model parameters succeeded in both reducing total uncertainty and bringing 726 modeled yield in line with observations of switchgrass yields both at this site (Heaton 727 et al., 2008) and globally (Wang et al., 2010). Reducing uncertainty in model outputs, in 728 this case yield, is key to increasing the value of ecological forecasts (Clark et al., 2001). 729 While reducing uncertainty does not necessarily increase model accuracy, an estimate of 730 model uncertainty is a first step toward generating meaningful statistical inference from the model itself. Without an estimate of model uncertainty, it is not possible to make 732 such a basic inference as the probability that the model predictions overlap with observed 733 data; this limits the capacity of models to inform research and applied problems (Clark 734 et al., 2001). Instead, comparisons of ecosystem models with observations have focused 735 on differences and correlations between model output and data (Bellocchi et al., 2010; 736 Schwalm et al., 2010; Dietze et al., 2011) without providing a confidence interval around 737 the model output itself. The ability to identify, with confidence, the conditions under 738 which a model produces valid output helps determine appropriate applications of the 730

model and it helps to identify targets for further model development (Williams et al., 740 2009). While parameter uncertainty is clearly just one of many sources of uncertainty in 741 models (McMahon et al., 2009), and constraining model parameters by no means 742 guarantees that a model will match reality, is difficult to assess the accuracy of a model if 743 it has low precision. In deterministic models, such as most ecosystem models, parameter uncertainty is a major driver of the precision of a model. 745 In this study, we can state with 90% Confidence that the mean Switchgrass yield during 746 the Heaton et al. (2008) study should have been between 7.2 and 37, and if we had made 747 this prediction in advance, we could have said that we were correct because the mean did fall within this range. We can also see that the model uncertainty contains the 90% CI 749 for observed switchgrass yields globally (Wang et al., 2010), even though we know that 750 important drivers of variability in the global meta-analysis (e.g., climate, soil) are 751 different from the source of uncertainty in our model predictions (e.g., parameters). The 752 model ensemble left open the possibility that the yields could have been much more or 753 much less than was actually observed, and we conclude that much of this variability could 754 be constrained with additional trait level data or data assimilation. Wang et al 201x (in 755 review, Ecological Applications #12-0854) provides an example of combining the PEcAn 756 meta-analysis and variance decomposition with data assimilation of biomass to constrain uncertainty in parameter estimates and improve the accuracy and precision of model 758 output. Once the model can make more precise predictions, it will be possible to begin 759 investigation of other sources of uncertainty, such as model structure and state variables 760 (e.g. climate, soil). 761 Although the present analysis focuses on modeled aboveground biomass, PEcAn can 762 analyze any model output, including pools and fluxes of water, energy, and carbon. 763 Indeed, PEcAn's approach to the synthesis of data and mechanistic models is 764 independent of the system being modeled, and thus has potential applications far beyond 765

the scope of its current development to support ecosystem modeling.

7 Variance Decomposition

Variance decomposition quantified the contribution of each parameter to model 768 uncertainty, helping to identify a subset of parameters for further constraint. SLA, V_{cmax}, 769 fine root to leaf ratio, and leaf turnover rate dominated uncertainty in yield prior to incorporating species level data. Therefore, SLA, which can be measured quickly and at 771 low cost, would make a good first target for reducing uncertainty when a new species is 772 evaluated. SLA also correlates strongly with other important model parameters, such as 773 photosynthetic rate, leaf lifespan, and nitrogen content (Wright et al., 2004). The ranking of parameters based on variance contribution depends on the response variable of choice 775 (in this case, aboveground biomass) as well as the conditions of the run (duration, soil, 776 climate), and the species or PFT being evaluated. In general, for a given species and 777 model output, overall patterns of parameter importance are consistent across a broad 778 range of climates (Wang et al., 201x, in review, Ecological Applications #12-0854). Variance decomposition (equation 6) demonstrates that it is not parameter uncertainty or 780 model sensitivity alone, but the combination of the two, that determines the importance 781 of a variable. For example, despite the high uncertainty in seed dispersal, switchgrass 782 yield is insensitive to this parameter (Figures 6, 7), therefore a better understanding of 783 switchgrass seed dispersal would not reduce model uncertainty. By contrast, although 784 uncertainty in the growth respiration is not particularly large, switchgrass yield was very 785 sensitive to growth respiration, and for this reason growth respiration is the greatest 786 contributor to model uncertainty. In addition, although no seedling mortality data were 787 available, model sensitivity to this parameter was much lower in the posterior compared 788 to prior runs. Using the sensitivity analysis or parameter uncertainties alone would thus 789 lead to incorrect conclusions about what parameters are most important and an 790

791 inefficient approach to reducing predictive uncertainties.

This analysis only represents the first step toward more comprehensive accounting of 792 known sources of uncertainty. The next step in reducing uncertainty would be to use the 793 results of the variance decomposition to target the most influential model parameters for 794 further constraint through data collection. We have demonstrated how the use of 795 species-level data to constrain parameter uncertainty reduced ensemble variance, resulting 796 in a new set of targets for additional field observations and refined literature surveys. 797 Traits that are difficult to measure, such as belowground carbon cycling, can be indirectly 798 constrained with ecosystem-level observations using data assimilation (Luo et al., 2009, 799 2011). Integrating data assimilation into PEcAn will allow ecosystem-level observations 800 to further constrain parameters for which trait level observations are difficult or 801 impossible to obtain. To date most Bayesian data assimilation approaches applied by 802 ecologists have employed flat, uninformative priors (assigning equal probability to values 803 over many orders of magnitude), which has lead to the problems of parameter 804 identifiably and the criticism that model parameters are allowed to take on biologically 805 unrealistic values. The use of the meta-analysis posteriors as priors in the data 806 assimilation step ensures that any parameter estimates are consistent with what is known 807 about different plant traits. In this way Bayesian methods are, in effect, updating the 808 literature-derived estimates with new data and providing a coherent and rigorous 809 framework for combining multiple different types of data. 810 In addition, by effectively restricting parameter space based on observed values, the use 811 of informed priors in data assimilation reduces problems of equifinality and identifiability. 812 This is consistent with the argument by Beven and Freer (2001) that only the feasible 813 parameter range should be sampled. 814 To a first order the spline interpolations of the univariate relationships between 815 parameters and aboveground biomass (Figure 6) provide a good estimate of the total 816

model variance. The closure term (Table 3b) accounted for approximately 5.2Mg ha⁻¹ or 28% of the variance decomposition estimate (18.1 Mg ha⁻¹, Table 3a), suggesting that 818 while parameter interactions are important, univariate parameter uncertainty drives 819 overall model variance. One key concern of parameter interactions is that the 820 combination of the variance decomposition terms would result in the prediction of 821 negative yields, which is clearly biologically impossible. By comparing the spline 822 ensemble, where this term is truncated, to the spline-based variance decomposition we 823 can conclude that this truncation effect accounts for 4.1 or 80% of the closure term in the 824 variance decomposition. 825 By contrast, evaluating the spline ensemble for different ensemble sizes shows that 826 ensemble size had negligible effect on the mean variance estimate although it does 827 improve the precision of this estimate (Table 3a). Finally, the difference between the 828 model and spline ensembles (Table 3b) suggests that the absence of parameter 829 interactions in the variance decomposition account for the remaining 20% of the closure 830 term (< 6% of the variance decomposition calculation), which could be improved by a 831 multivariate meta-analysis and sensitivity analysis, both of which are planned for future 832 development of PEcAn. Overall, the closure term is relatively well constrained even when 833 the parameter interactions are assumed to be linear.

35 Model-field work feedback

Variance decomposition can be used to inform data collection by identifying candidate parameters for further refinement based on their contributions to model variance. Recall that this variance contribution is a function of parameter sensitivity and the parameters' probability density (equation 6, Figure 7). Sensitivity is a function of the model and so there is no direct way to reduce sensitivity. However, because $Var(f) \propto Var(\beta_0)$, it is possible to reduce the model uncertainty by reducing parameter variances.

Through simple power analyses one can explicitly estimate the relationship between an increase in sample size and the reduction in posterior variance. Not only can we calculate 843 the reduction in parameter uncertainty that would be expected for a given sample size, 844 but using equation 6 we can also express this in terms of reductions in the variance of the 845 model output. This then allows us to directly compare the value of different data types in 846 a common currency. 847 Quantitatively evaluating the relationship between data and model uncertainty provides a 848 path of communication between field research and modeling, opening the door for a new 849 framework in which modeling and field work can be mutually informative. Given the current data and model uncertainties, it is possible to identify effective data acquisition 851 strategies based on this analysis. For example, data could be ranked by the ratio of 852 reduction in model uncertainty to the cost of acquiring each sample in terms of dollars 853 and/or man hours. In this way, data collection could be optimized in terms of the efficiency at which new information is gained. 855 These approaches close the model-data loop by enabling models to inform data collection, 856 and data to inform models. Such a shift has the potential to put field ecologists and 857 modelers in closer connection. It also gives us the tools to answer the long standing 858 question among many field ecologists about what exactly modelers need to know. Indeed, 859 this shift highlights a need for greater accessibility to models by the general research 860 community so that field ecologists can drive this loop directly. This is exactly the 861 objective of PEcAn – to encapsulate these tasks in a way that makes the analysis of 862 models transparent, repeatable, and accessible. 863 In addition to informing sample size, the parameter meta-analysis can inform 864 experimental design by providing valuable information on the scales of variability. For 865 example, when data from multiple sites is available, the meta-analysis partitions among 866 site and within site variance. This information can be used to construct optimal sampling 867

environmental covariates that should be measured in order to explain parameter 869 variability. 870 Based on our switchgrass example, variance partitioning also highlights broad data needs 871 and the discrepancy between the relative ease of parameterizing aboveground processes 872 compared to below ground processes. Indeed, one of the greatest challenges in ecosystem 873 ecology is the ability to constrain below ground processes such as root allocation, 874 respiration, and turnover. These parameters are uncertain precisely because measurement 875 is difficult, often indirect, and data may reflect the diverse methods used to indirectly estimate a pool or flux. Many parameters in the ED2 model correspond to processes that 877 are not directly observable. For example, the root respiration parameter in ED2 is not 878 total root respiration but just maintenance respiration, while measurements typically can 879 not separate growth, maintenance, and rhizosphere respiration. Whole-plant growth 880 respiration, which is currently the most important model parameter, is also difficult to 881 estimate directly from measurements (Amthor, 2000). In this case, data assimilation is 882 likely the most efficient route to constrain this parameter; data assimilation would 883 effectively use mass balance of ecosystem carbon exchange to estimate this respiration 884 parameter once other parameters are more directly constrained by data.

designs which balance intensive vs extensive sampling, and may help identify

886 Future Directions

The analyses presented here represent the first phase in the development of the PEcAn project. In the near term we intend to expand the existing analyses to include a multivariate meta-analysis and sensitivity analysis to reduce model uncertainty by accounting for parameter covariances. In addition, we plan to implement the power analyses discussed above to more quantitatively inform data collection. A data assimilation module is in progress for PEcAn that will allow the use of ecosystem level

data including plot-level inventory data, eddy covariance fluxes, and remote sensing imagery to enter the analysis and provide additional constraint on uncertainty in both 894 parameters and output. The basic concept of variance decomposition will also be 895 expanded to investigate other sources of variability, such as uncertainty in initial 896 conditions or in driver data. We are implementing ecosystem models other than ED2 897 within the PEcAn workflow. This will provide opportunities for multi-model ensemble 898 forecasting and assessing data requirements across models. 890 Integrating modeling into a workflow system has distinct advantages not just for model 900 analysis but also for managing the flows of information coming in and out of the model. 901 In this sense we also envision PEcAn as an informatics and data management tool. 902 Finally, it is our hope that other researchers will find PEcAn useful and contribute 903 modules that extend the functionality of the system in creative and exciting ways. 904

905 Conclusion

In this paper, we demonstrate an approach to the parametrization of a terrestrial biosphere model that synthesizes available data while providing a robust accounting of parameter uncertainty. We also present a scientific workflow that enables more efficient constraint of this uncertainty by identifying the key areas requiring data collection and model refinement. By quantifying the effect that each parameter has on model output uncertainty, this analysis identifies additional data that, once obtained, would improve model precision. In addition, the analysis calculates probabilities of alternate potential outcomes, resulting in more useful assessments.

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1187	1	Prior Distributions Prior distributions used in meta-analysis and model	
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1192	2	Meta-analysis Results Results of meta-analysis of Switchgrass data for six	
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3 a) Variance Estimates Comparison of sample variances (s, on stan-1199 dard deviation scale) for the aboveground biomass estimated from data-1200 constrained parameters calculated from model ensemble, spline-emulated 1201 model ensembles, and variance decomposition. Values in parentheses are 1202 estimates of uncertainty in the sample estimate of variance. Sample size, n, 1203 refers to the size of the sample from the posterior parameter distribution. 1204 b) Components of closure term, ω The closure term ω (equation 6) is 1205 5.2, the difference between the variance decomposition and model ensemble 1206 estimates of σ . The closure due to parameter interactions is estimated as the 1207 difference between the spline ensemble and the model ensemble; the closure 1208 due to the absence of a lower bound of zero on the spline functions is esti-1209 mated as the difference between the variance decomposition and the spline 1210 ensemble estimates. * Analysis of the closure term is based on estimates 1211 with n = 10000 parameter sets, except in the case of the model ensemble 1212 because evaluation of the model ensemble at n = 10000 is computationally 1213 1214

Parameter Specific Leaf Area Leaf Turnover Rate	$\frac{\mathrm{Units}}{\mathrm{m}^2 \ \mathrm{kg}^{-1}}$	Clade Grass Grass	Distribution Gamma Weibull	2.06	b 19.00 0.63	N 125 40	b N mean LCL 9.00 125 17 3.2 0.63 40 4.6 0.91	3.2 0.91	UCL 36 11	Citation (Wright et al., 2004) (Wright et al., 2004)
Root Turnover Rate	1/3	Grass	Gamma	1.67	0.66	99	0.59	0.073	1.4	(Gill and Jackson, 2000)
Quantum Efficiency	percent	C4 grasses	Weibull	90.90	1580.00	99	0.058	0.046	0.07	(Skillman, 2008)
Stomatal Slope	ratio	C4 Grass	Gamma	3.63	3.81	4	3.4	1.4	5.5	*
Vcmax	umol CO2 $m^{-2} s^{-1}$	graminoid	Gamma	3.49	24.70	26	22	8.6	36	(Wullschleger, 1993)
Leaf Width	mm	C4 Grass	Weibull	26.10	5.94	18	4.4	2.9	6.2	(Oyarzabal et al., 2008)
Root Respiration Rate	umol $CO_2 \text{ kg}^{-1} \text{ s}^{-1}$	C4 Grass	Ţ	5.61	2.33	35	5.6		10	(Tjoelker et al., 2005)
Fine Root Allocation	ratio	Grass	Beta	0.80	0.81	0	3.1	0.46	11	(Chapin III et al., 2002)
Seed Dispersal	percent	Grass	log-Normal	20.10	74.90	30	0.21	0.14	0.3	(Jongejans and Schippers, 1
Photosynthesis min temp	Celsius	C4 Grass	Ţ	10.00	1.02	0	10	∞	12	*
Growth Respiration	percent	Grass	log-Normal	2.63	6.52	0		0.062	9.0	*
Seedling Mortality	percent	monocots	log-Normal	3.61	0.43	0		0.5		*
Mortality Coefficient	$1/\mathrm{yr}$	plants	Weibull	1.47	90.0	0		1.8	80	*
Reproductive Allocation	percent	Plants	log-Normal	2.00	4.00	0		0.053	0.72	*

Table 1

Variable	n	β_0	σ_Y	$\sigma_{ m site}$	$\sigma_{ m treatment site}$	$eta_{ m greenhouse}$
Specific Leaf Area	24	16(12,20)	2.8(2.5, 3.2)	3.2(1.6, 7.3)	2.4(1.1,6)	6.5(1,12)
Leaf Width	39	6(4.7, 6.6)	0.46(0.44, 0.48)	0.47(0.2, 2.1)	6.4(1.9, 130)	1.6(-0.033, 3.5)
Vcmax	4	24(18,30)	12(8.1, 17)		1.2(0.098, 47)	
Fine Root Allocation	4	1.3(0.5, 2.6)	2.2(1.2, 6.2)			
Root Respiration Rate	\vdash	5.1(3.7, 6.6)	1.2(0.39, 2.3)			
Root Turnover Rate	\vdash	0.67(0.2, 1.1)	0.45(0.14, 0.88)			
Stomatal Slope	4	4.1(3.9, 4.3)	0.33(0.23, 0.45)			

Table 2

	model	spline	variance
	ensemble	ensemble	decomposition
n	$Sf(oldsymbol{eta_0})$	$S_{m{g}}(m{eta_0})$	$\sum s_{g_i(\beta_{0_i})}$
500	13(14)	13.8(13)	18.2(6)
10000	*	14.1(2.8)	18.1(1.2)
		(a)	

	calculation	Mg/ha
$\omega_{ ext{total}}$	$\sum s_{g_i} - s_f$	5.2
$\omega_{ m covariance}$	$s_{\boldsymbol{g}} - s_f$	1.1
$\omega_{\mathrm{truncation}}$	$\sum s_{g_i} - s_{\boldsymbol{g}}$	4.1
	(b)	

Table 3

List of Figures

Overview of the PEcAn workflow. The synthesis of plant trait data begins by querying a database of plant trait data for data on a single species or a plant functional type, and then mapping these data to the model parameters that they inform. The database also provides probability distributions that describe our prior information about the range of values that a model parameter can take. Next, this information is synthesized in a Bayesian meta-analysis, resulting in a posterior trait distribution that summarizes the uncertainty in each parameter. The ensemble of model runs produces the posterior distribution of model outputs, representing a probabilistic assessment or forecast that accounts for input parameter uncertainty. The final steps in the workflow are the sensitivity analysis and variance decomposition; these steps gives insight into the relative contribution of each parameter to the uncertainty in the model output, and can be used to guide additional data collection that will most efficiently reduce model uncertainty.

Prior distributions PDFs of priors with data constraints. Parameter value is on the x-axis and probability density is on the y-axis, and the area under each curve equals one. Three points on each line, from left to right, indicate the 2.5th, 50th, and 97.5th quantiles. (From top left) Four priors fit to data (data points shown as rug plot) using maximum likelihood: specific leaf area and leaf turnover rate (Wright et al., 2004), root turnover rate (Gill and Jackson, 2000), and quantum yield (Skillman, 2008). Four priors fit to the posterior predictive distribution of an unobserved C4 grass species using Bayesian meta-analysis of data from multiple plant functional types (C4 data shown in black, other functional types in grey): stomatal slope (present study data provided in Appendix A), V_{cmax} of C3 plants (Wullschleger, 1993) and C4 grasses (Kubien and Sage, 2004; Massad et al., 2007; Wang et al., 2011), leaf width (Oyarzabal et al., 2008), and root respiration (Tjoelker et al., 2005). Priors fit to 95% CI (dashed grey line) and median (solid grey line) based on ED2 defaults and expert opinion as described in the text: fine root to leaf ratio (Chapin III et al., 2002), seed dispersal (Ernst et al. (1992) model parameterized with site level data), minimum temperature of photosynthesis (Don Ort, personal communication), growth respiration, seedling mortality factor, mortality factor, and reproductive allocation. . .

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Overview of the Hierarchical Bayesian meta-analysis model. For each trait, the posterior estimate of the global trait mean (β_0) is used as an input parameter in the sensitivity analysis and model ensemble (Figures 6 and 5). Results from the meta-analysis of specific leaf area are as an illustrative example; x-axes have units of m^2kg^{-1} and all plots are on the same scale. Each of the k sample means (Y_k) are taken from published articles and unpublished field measurements, and may be associated with a sample standard error and sample size. When sufficient data were available, site, treatment, and greenhouse effects were estimated. The within-unit standard deviation, σ_Y , is estimated from se and n. Site and treatment random effects, represented by β_{site} and $\beta_{\text{tr}|\text{site}}$, are estimated for each site and treatment within site with from normal distributions with mean zero and standard deviations σ_{site} and $\sigma_{\text{tr}|\text{site}}$, respectively. Greenhouse is a fixed effect. Table 2 summarizes the global mean, variance terms, and greenhouse effect for the seven model parameters informed by species-level data.

Ensemble average 2004-2006 post-senescence yield. Histogram of results from prior ensemble runs (dashed), posterior ensemble runs (solid line), and the spline ensemble (gray line). The gray box on the left represents non-viable ensemble members (\leq 2Mg/ha, see text). Horizontal bars provide a summary of yields, from top: a three year trial at the modeled site (Heaton et al., 2008), all 1902 observations included in a recent meta-analysis (Wang et al., 2010), viable runs from the ED2 ensemble based on prior and posterior parameterizations. Diamonds indicate the median; thick and thin lines indicating the 68% and the 95% CI, respectively. Histogram-style plots provide comparison of the distributions of observations and model runs. For clarity, non-viable and viable runs are plotted with different width bins. .

Univariate relationships between parameters and 2004-2006 average modeled yield. Parameter values are on the x-axis and biomass is on the y-axis while runs centered around the prior median are in gray and those centered around the posterior median are in black. The univariate responses were estimated using a cubic spline to fit model output at the median and $\pm [1, 2, 3]\sigma$ quantiles of each parameter while holding other parameters to the median value.

Partitioning of variance by parameter results from variance decomposition conducted before (grey) and after (black) updating parameter estimates with species-level data in the meta-analysis. From left to right, panels present: a) the uncertainty associated with each parameter (coefficient of variation, $CV = \sigma/\mu$). The degree to which some parameters have been constrained by data is indicated by the reduction in CV in the black compared to the grey bars; sample sizes are indicated in Table 2. b) the sensitivity of modeled aboveground biomass to each parameter presented as elasticity (elasticity is normalized sensitivity, and an elasticity of one indicates that model output will double when the parameter value doubles). Sensitivity is the slope of the line at the median in Figure 6). Parameters with larger bars have greater influence on model output. c) Partial variance is the contribution of each parameter to explained variance. This is a function of both the parameter variance and sensitivity. Parameters with both large CV and elasticity contribute the most to uncertainty in model output.

Figure 1

The Predictive Ecosystem Analyzer Data Figure 2 Data Collection $-\beta_{0t}$ Priors $\Theta_k = \beta_0 + \beta_{\text{site}i} + \beta_{\text{tr}|\text{site}ij} + \beta_{\text{gh}}I(i)$ Meta-analysis Meta-analysis Posteriors β_{0t} Ecosystem Model Sensitivity Analysis Model Ensemble Figure 5 Variance Decomposition Ensemble Posterior $f(\boldsymbol{\beta_0})$ $g_t(eta_{0t})$ $Var(f(\boldsymbol{\beta_0})) = \sum_{t=1}^{m} Var(g_t(\beta_{0t})) + \omega$ Figure 6

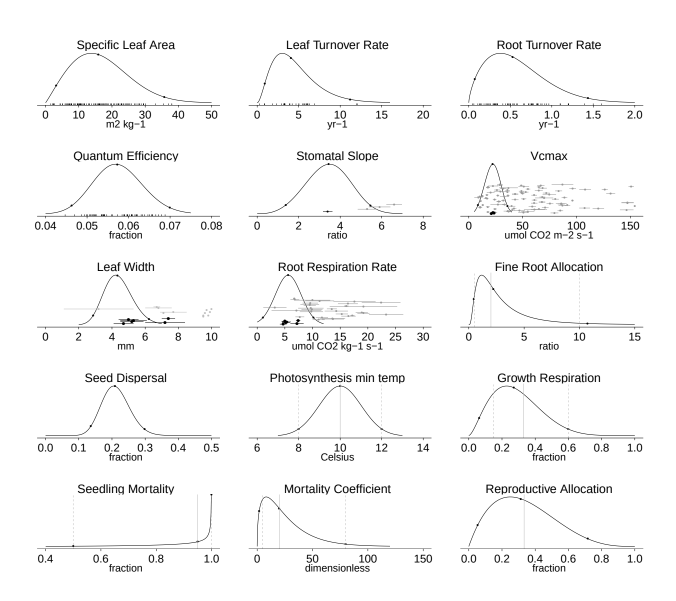


Figure 2

Figure 3

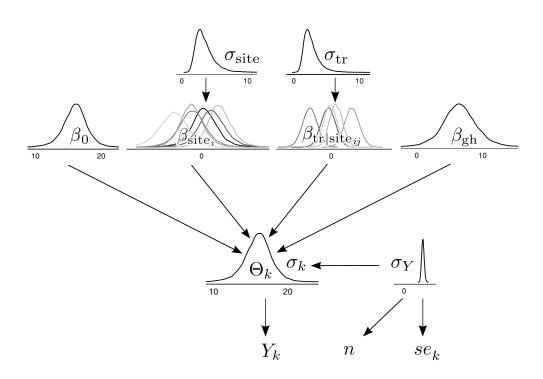
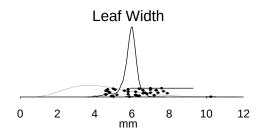
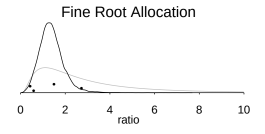
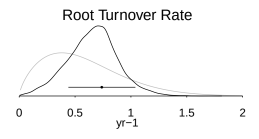
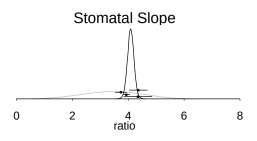


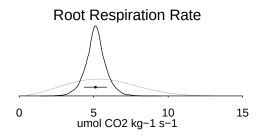
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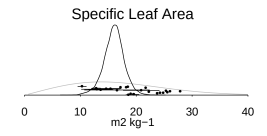












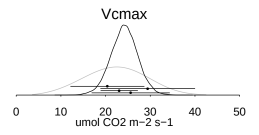
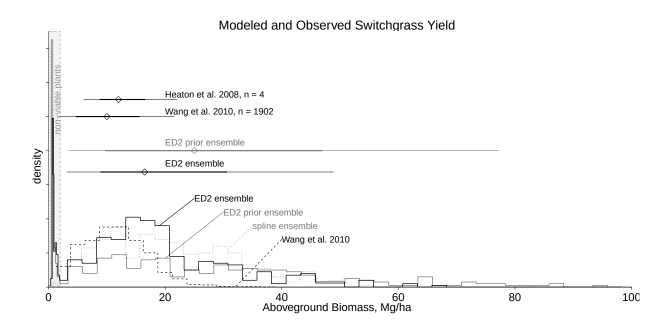


Figure 5



Sensitivity of Aboveground Biomass (Mg/ha) to Fifteen Plant Traits

Figure 6

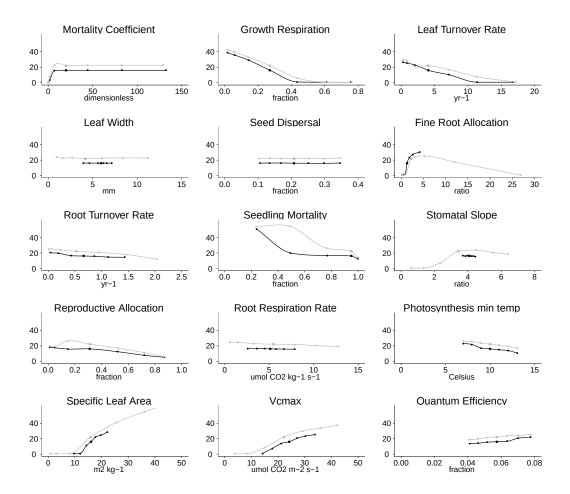


Figure 7

