Bold black font: Topical editor and reviewer comments Black font: Author response Blue font: Verbatim copy and paste from the revised manuscript

Topical Editor Decision: Publish subject to minor revisions (review by editor) (08 Apr 2019) by Christoph Müller

Comments to the Author: Dear Dr. Elshall,

the original reviewers have seen your revised paper again and suggest further amendments to the paper. I think what they suggest is easy enough to implement.

I look forward to your revised paper.

Cheers Christoph

Thank you very much for handling the original and the revised submissions. We have responded to all the Editorial Support and reviewers' comments, and revised the manuscript accordingly.

Editorial Support

Besides adjustments requested by the Topical Editor or Referees, please check your manuscript carefully for typos, missing co-authors and their affiliations, terminology, updates of data in tables, or updates of variables in equations.

We updated the affiliation of the second author and the funding information. We improved the article keywords. We carefully checked the manuscript, and corrected several typos and grammatical errors, as shown in the *marked-up manuscript version*. We also slightly improved the writing style in several parts of the article and added more clarifications as shown in the *marked-up manuscript version*.

Anonymous Referee #1

I revisit some of my previous comments, considering response from the authors in the discussion forum.

1. Contribution: the authors have now better articulated their contribution. The paper is somewhat incremental, since the methodology is not new and the findings are similar to applications in other disciplines. Nevertheless it is still useful to have an explicit evaluation and comparison of the impacts of statistical assumptions in soil respiration models.

Thank you for taking the time to review the manuscript and for your valuable feedback.

2. Problems with residual autocorrelation: I agree that the authors do not need to solve this issue in this paper. However, the alternative suggested approach (Evin et al) could be easily tested (simply swap the order in which correlation and heteroscedasticity are applied in the likelihood function). In their extended response in the discussion forum the authors seem to indicate that they have tested this. Why then not add it to the paper? If not in the results then in the discussion part of the paper ("preliminary

results show..."). My suggestion is to include the entire extended response posted in the discussion forum (under comment 2 of RC1) in the paper itself, as it provides more information that is useful for readers.

We prefer to keep the manuscript in its current form since we are afraid that we cannot adequately address this autocorrelation problem in this manuscript. We prefer not to present incomplete or preliminary results in a published paper. However, we improved the extended response, and add it to the manuscript as suggested by the reviewer. The added part reads:

4.3 Discussion on handling residual correlation

Accounting for autocorrelation can lead to biased parameter estimation (Figure 5) and poor predictive performance (Figure 10). Auto-correlated residuals may be attributed to model discrepancy, as shown in Lu et al. (2013). The most obvious solution to handle the autocorrelation is to reduce the autocorrelation by improving the soil respiration model. If model improvement is difficult for practical reasons, we can improve the data model to better characterize the autocorrelation. Addressing autocorrelation in a data model is challenging since it involves several interlinked factors as follows:

- (1) Non-stationarity due to wet-dry periods could be a reason for this problem. By drawing on similarity from surface hydrology, the study of Ammann et al. (2018) suggests that auto-correlated residuals might be attributed to non-stationarity due to wet-dry periods with half-hourly data. Accounting for non-stationarity could better address the problem of auto-correlated residuals (Ammann et al., 2018; Smith et al., 2010b).
- (2) The way of implementing autocorrelation could have an impact. Autocorrelation could be applied to raw residuals directly (e.g., Li et al., 2015), to transformed residuals based on covariance matrix of residuals L(e) (e.g., Lu et al., 2013), or to normalized residuals L(a) (e.g., Schoups and Vrugt, 2010; Evin et al., 2013). Note that e is a vector of transformed residuals, while a denotes a vector of independent and identically distributed random errors with zero mean and unit standard deviation. The L(e) approach based on covariance matrix of residuals is generally limited to Gaussian data models (e.g. Lu et al., 2013), while the L(a) approach for normalized residuals can be readily adopted for non-Gaussian data models.
- (3) The autocorrelation model could have an impact. Using an autoregressive model is a popular technique to account for auto-correlated residuals. However, using an autoregressive model with either joint inversion approach (e.g., this study and Schoups and Vrugt, 2010) or sequential approaches (e.g., Evin et al., 2013, 2014; Lu et al., 2013) removes correlation errors through a filter approach, which can lead to a loss of information content. As this may cause overcorrection of prediction especially at surge events, Li et al. (2015) developed a restricted autoregressive model to overcome this adverse effect. Other autocorrelation models include moving average model and mixed autoregressive-moving averaging model (Chatfield, 2004).
- (4) Joint versus sequential inversion for autocorrelation could have an impact. Sequential inversion approaches include two-step procedures (e.g. Evin et al., 2013, 2014; Lu et al., 2013) or the multistep procedure (Li et al., 2016a). These sequential approach estimates the autoregressive parameters sequentially in a later step after estimating the physical model parameters and other data model parameters. Evin et al. (2013, 2014) used a sequential approach to avoid the interaction between the parameters of the heteroscedasticity model and the autocorrelation model. In addition, the autoregressive model parameters can be deterministically calculated as an internal variables of the data model similar to Lu et al. (2013), and not as calibration parameters (e.g. Schoups and Vrugt; Evin et al. 2013; 2014). While the first step in the sequential approach would avoid the biased parameter estimation (Figure 10a-d), the second step can still lead a poor predicative performance since we are essentially using a filter approach to remove

residual correlation. To address this problem, Li et al. (2016) multi-step procedure that is based on Gaussian data model uses restricted autoregressive model. Generally, Ammann et al. (2018) states that the joint inversion is still preferred, and understanding the conditions where accounting for auto-correlation can be achieved remain poorly understood.

In addition, the text about autocorrelation in the conclusions section was accordingly shortened. The revised manuscript reads "While the reasons remain poorly understood (Ammann et al., 2018), it might be attributed to non-stationarity due to wet-dry periods with half-hourly data (Ammann et al., 2018) or to the method of handling autocorrelation (e.g., Schoups and Vrugt, 2010, Evin et al., 2013; 2014; Lu et al., 2013; Li et al., 2015, 2016a; Ammann et al. 2018). Further investigation to address autocorrelation in soil respiration modeling is warranted in a future study."

3. Grammatical/spelling errors: the authors state that they have "corrected several other grammatical errors", but it's not clear what was corrected exactly.

Sorry for not listing the grammatical errors and typos the we corrected in the previous submission, which are as follows:

- Line 123: was used to select and the best model -> was used to select the best model
- Line 192: Laplace distribution used by (van Wijk et al., 2008) and (Ricciuto et al., 2011) -> Laplace distribution (van Wijk et al., 2008; Ricciuto et al., 2011).
- Line 207: to split a dataset of CO2 -> by splitting the dataset of CO2
- Line 329: model parameters are obtained jointly with -> model parameters jointly estimated with
- Line 420: each criteria -> each criterion
- Line 658: Accounting for ^ error term e -> Accounting for the error term e
- Line 688: the rest ^ six data models -> the rest of the six data models
- Line 722: the residuals, ^ thus resulting -> the residuals, and thus resulting
- Line 743: We tested eight data modeling -> We tested eight data models
- Line 807: The ^ conclusions above -> The above conclusions

Additional comment:

- the abstract states that "not accounting for heteroscedasticity ... will definitely underestimate uncertainty". It is not clear why this would be the case. Perhaps you mean to say that not accounting for any residual error beyond parameter uncertainty will underestimate predictive uncertainty (as in section 4.2)? Although that is not really a significant finding.

Comparing Figures 10a-b (with the SLS data model that does not account for residual error beyond parameter uncertainty) with Figures 10c-d (with the SEP data model that accounts for residual error beyond parameter uncertainty) shows that using an SEP data model with two additional parameters did not significantly impact the uncertainty. This is not the case for Figures 10e-h for data models WLS and WSEP that account for heteroscedasticity. Visual comparison of Figures 10a-d with Figures 10e-h and examination of the sharpness and predictive coverage metrics indicate that not accounting for heteroscedasticity will underestimate uncertainty. Accordingly, accounting for heteroscedasticity with WLS (Figures 10e-f) or WSELP (Figures 10g-h) makes the predictions more sensitive to peak carbon effluxes. We clarified this point, and the revised manuscript reads "Not accounting for heteroscedasticity will underestimate the predication uncertainty (Figure 10b and Figure 10d). This is mainly because the variance of the efflux residuals increases with the magnitude of the carbon effluxes (Figure 3a), and thus

assuming constant variance is not representative. Accordingly, accounting for heteroscedasticity using WLS (Figure 10e) or WSEP (Figure 10h) will make the predictions more sensitive to peck carbon effluxes. This and will generally improve the predictive coverage on the expense of sharpness and the central mean tendency."

Anonymous Referee #2

The revised version submitted by Elshall et al., generally answers quite well all my previous comment.

Thank you for taking the time to review the original and the revise manuscripts.

Nevertheless I still disagree with using a CUE maximum value of 0.6 and I would like to see some kind of sensitivity analysis to better understand the impact of such assumption. If the effect of changing the max CUE value is limited then the paper could be published in its current version. If not some discussions should be added.

We probably did not make ourselves clear. For the inverse modeling with MCMC sampling we did not assume CUE maximum value of 0.6. Thus, for parameter estimation and predictive performance, we did not impose this constraint. We merely obtained this CUE maximum value of 0.6 from literature, which is based on thermodynamic calculation (Fernández-Martínez et al., 2014; Li et al., 2014; Sinsabaugh, et al., 2013), to evaluate whether the posterior parameter distributions of CUE under different data models and different soil respiration models are within this physically reasonable range of $0 \sim 0.6$ or beyond. Thus, this assumption has no impact on the results. We clarified this in the revised manuscript as follows "Note that, for inverse modeling with MCMC sampling, we did not assume CUE maximum value of 0.6. In other words, for parameter estimation and predictive performance we did not impose the constraint that CUE is less than 0.6. We merely use this CUE maximum value of 0.6 to evaluate whether the posterior CUE parameter samples obtained using different data models and different soil respiration models are within the physically reasonable range of 0 are constraint that CUE is less than 0.6. We merely use this CUE maximum value of 0.6 to evaluate whether the posterior CUE parameter samples obtained using different data models and different soil respiration models are within the physically reasonable range of $0 \sim 0.6$."

Minor corrections: L376 you mean soil moisture I guess? L424 you mean Yi and not Xi, right?

Thank you for these two typos and we corrected both of them.

| 1 2 | Bayesian Inference and Predictive Performance of Soil Respiration Models in the Presence of Model Discrepancy |
|----------|---------------------------------------------------------------------------------------------------------------------------------|
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| | 47 48 49 | Key Points | | | |
|---|----------------|------------|-----------------------------------------------------------------------------------------------|--|--|
| | 49 50 | (1) | Bayesian inference and prediction are useful to evaluate multiple soil respiration models | | |
| | 51 | | with different levels of model complexity. | | |
| | 52 | (2) | Data models used in Bayesian inference have substantial impacts on model parameter | | |
| | 53 | | distributions and subsequently model predictions. | | |
| | 54 | (3) | Using exponential power distribution and considering heteroscedasticity in data models | | |
| | 55 | | improves Bayesian inference and prediction. | | |
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| | 68 | Keyw | ords: Soil respiration, modeling, Bayesian, likelihood function, data model, autocorrelation, | | |
| | 69 | hetero | scedasticity, skew exponential power distribution, cross-validation, relative model | | |
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76 Abstract

Bayesian inference of microbial soil respiration models is often based on the assumptions that the 77 residuals are independent (i.e. no temporal or spatial correlation), identically distributed (i.e. 78 Gaussian noise) and with constant variance (i.e. homoscedastic). In the presence of model 79 discrepancy, since no model is perfect, this study shows that these assumptions are generally 80 invalid in soil respiration modeling such that residuals have high temporal correlation, an 81 increasing variance with increasing magnitude of CO₂ efflux, and non-Gaussian distribution. 82 Relaxing these three assumptions stepwise results in eight data models. Data models are the basis 83 of formulating likelihood functions of Bayesian inference. This study presents a systematic and 84 comprehensive investigation of the impacts of data model selection on Bayesian inference and 85 predictive performance. We use three mechanistic soil respiration models with different levels of 86 model fidelity (i.e. model discrepancy) with respect to number of carbon pools and explicit 87 representations of soil moisture controls on carbon degradation, and accordingly have different 88 levels of model complexity with respect to the number of model parameters. The study shows data 89 models have substantial impacts on Bayesian inference and predictive performance of the soil 90 respiration models such that: (i) the level of complexity of the best model is generally justified by 91 the cross-validation results for different data models; (ii) not accounting for heteroscedasticity and 92 autocorrelation might not necessarily result in biased parameter estimates or predictions, but will 93 94 definitely underestimate uncertainty; (iii) using a non-Gaussian data model improves the parameter

estimates and the predictive performance; and (iv) separate accounting for autocorrelation or joint
inversion of correlation and heteroscedasticity can be problematic and requires special treatment.
Although the conclusions of this study are empirical, the analysis may provide insights for
selecting appropriate data models for soil respiration modeling.

99 1 Introduction

Developing accurate soil respiration models is important for realistic projection of global 100 carbon [C] cycle, as global soils store 2,300Pg carbon, an amount more than 3 times that of the 101 atmosphere (Schmidt et al., 2011) and release 60–75 Pg C/yr, about 7 times more CO_2 to the 102 atmosphere than all human-caused emissions (Le Quéré et al., 2014). The major work on soil 103 respiration modeling has been focused on advancing knowledge about model inputs and 104 calibration data (e.g. Janssens et al., 2003; Peters et al., 2007; Scott et al., 2009; Barron-Gafford et 105 106 al., 2011; Hilton et al., 2014) and on developing more advanced models for better representing soil microbial processes (e.g. Schimel and Weintraub, 2003; Allison et al., 2010; Davidson et al., 107 2011; Wieder et al., 2013, 2015; Xu et al., 2014; Zhang et al., 2014). Integration of data and 108 109 models is indispensable for improving predictability of the terrestrial carbon cycle, and statistical modeling is a vital tool for the model-data integration (Luo et al., 2011, 2014; Wieder et al., 2015). 110 In addition, use of state-of-the-art statistical methods is necessary to accurately quantify 111 uncertainty in parameters and structures of soil respiration models for improvement and practical 112 uses of the models (Katz et al., 2013). A data model that is also known as a residuals model or an 113 error model is used to characterize residuals (i.e., the difference between data and corresponding 114 model simulations). While a large number of data models have been used (e.g. Elshall et al., 2018; 115 Scholz et al., 2018) to our knowledge comprehensive and systematic evaluation of data models for 116 117 soil respiration modeling has not been reported in literature.

118 The objectives goal of this study are is to evaluate the impacts of data models on Bayesian 119 inference and predictive performance of three mechanistic soil respiration models, and to use these evaluation results findings to make broader recommendations. The three models were developed 120 by Zhang et al. (2014) to simulate the Birch effect (the peak soil microbial respiration pulses in 121 122 response to episodic rainfall pulses) at a site scale and a short temporal scale; understanding the, which are Birch effect is important for gaining mechanistic understanding of CO₂ efflux production 123 (Högberg and Read, 2006; Vargas et al., 2011). The models of Zhang et al. (2014) developed a 124 total five models, includingare based on an existing four-carbon pool model, but have and four 125 new models with additional carbon pools and/or explicit representations of soil moisture controls 126 on carbon degradation and microbial uptake rates. The models Zhang et al. (2014) were calibrated, 127 and Bayesian model selection was used to select the best model (Zhang et al., 2014). However, 128 129 this effort was based on a single data model. It is unknown whether the best model still remains the best (in terms of reproducing the both calibration data and the cross-validation data) if a 130 different data model is used. In addition, since predictive performance of the models was not 131 evaluated in Zhang et al. (2014), it is unknown whether the best model will give the best 132 predictions. These two questions are addressed in this study by considering eight data models and 133 by evaluating predictive performance in a manner of cross-validation. The top two models (also 134 135 the two most_high fidelity models) ranked by Zhang et al. (2014) are considered in this study, and the worst model (also the low fidelity model) is also considered in this study for comparison. We 136 use the terms model fidelity and model discrepancy interchangeably. Model fidelity refers to the 137 degree of realism of representing our scientific knowledge with respect to the real world system. 138 139 That is a high fidelity model has less discrepancy. Conducting Bayesian inference and Eevaluating predictive performance for the three models with different degrees of fidelity provides moreinsights than for a single model.

Bayesian inference in general uses the Bayes' theorem to update the prior distributions of 142 143 model parameters to posterior parameter distributions given a likelihood function of data. The mathematical formulation of the (formal and informal) likelihood function requires a probabilistic 144 data model that however is intrinsically unknown due to unknown errors in all model components 145 146 such as observation data, model structures, parameters, and driving forces. Bayesian inference of soil respiration models often adopts the assumption of independent, normally distributed and 147 homoscedastic residuals (e.g. Ahrens et al., 2014; Bagnara et al., 2015, 2018; Barr et al., 2013; 148 Barron-gafford et al., 2014; Braakhekke et al., 2014; Braswell et al., 2015; Correia et al., 2012; Du 149 et al., 2015, 2017; Hararuk et al., 2014; Hashimoto et al., 2011; He et al., 2018; Klemedtsson et 150 151 al., 2008; Menichetti et al., 2016; Raich et al., 2002; Ren et al., 2013; Richardson and Hollinger, 2005; Steinacher and Joos, 2016; Tucker et al., 2014; Tuomi et al., 2008; Xu et al., 2006; Yeluripati 152 et al., 2009; Yuan et al., 2012, 2016; Zhang et al., 2014; Zhou et al., 2010). These assumptions are 153 154 conveniently adopted to satisfysince the requirement of using an unknown probability model in Bayesian statistics, which is called "a basic dilemma" by (Box and Tiao, 1992). Box and Tiao 155 (1992). 156

Postulating the data models is always based on assumptions about residual statistics, and the most widely used assumptions are paired as follows: (i) independent vs. correlated residuals, (ii) homoscedastic vs. heteroscedastic residuals, and (iii) Gaussian vs. non-Gaussian residuals. For soil respiration modeling few studies have relaxed the non-correlation assumption(e.g. Cable et al., 2008, 2011; Li et al., 2016b), the homoscedasticity assumption_(e.g. Berryman et al., 2018; Elshall et al., 2018; Ogle et al., 2016; Tucker et al., 2013), and the non-Gaussian and homoscedasticity 163 assumptions (e.g. Elshall et al., 2018; Ishikura et al., 2017; Kim et al., 2014). TheA recent study 164 of (Scholz et al., (2018) relaxed these three assumptions using the generalized likelihood function developed by (Schoups and Vrugt, (2010). However, few studies have focused on investigating 165 appropriateness and impact of these assumptions for soil respiration modeling, by relaxing the 166 independent residuals assumption (Ricciuto et al., 2011) and the Gaussian residuals assumption 167 (Ricciuto et al., 2011; van Wijk et al., 2008). By relaxing these three assumptions stepwise 168 resulting in eight data models, to our knowledge this is the first study that systematically evaluates 169 the impact of data model selection on Bayesian inference and predictive performance of soil 170 171 respiration modeling. In addition, to our knowledge this is the first soil respiration modeling study that investigates the impact of data models in relation to model fidelity. 172

Relaxing these three assumption results in eight data models, which are shown in details in 173 174 Section 2. For example, combining the assumptions of independent, homoscedastic, and Gaussian residuals leads to the standard least squares data model. This model is the simplest one among the 175 eight data models, since it requires only one parameter, i.e., the constant variance of the Gaussian 176 177 distribution. Note that there is a difference between the soil respiration model parameters and the data model parameters. They technically can be jointly estimated together, but one arises from 178 assumptions about soil respiration processes, and the other from assumptions about the residuals. 179 Relaxing the homoscedastic assumption to heteroscedastic gives the weighted least squares data 180 model. It is more complex because it has extra parameters to account for multiple variances for 181 multiple data. Whenever one or combinations of the three assumptions (independence, 182 homoscedasticity, and normality) are relaxed, the resulting data models become more complex and 183 require more parameters. Such systematic evaluation of data models (McInerney et al., 2017; 184

185 Smith et al. 2010b, 2015) is necessary to evaluate appropriateness of residuals assumptions and186 their impacts on Bayesian inference.

The assumptions of heteroscedastic, correlated, and non-Gaussian residuals are accounted for 187 188 by using the method of Schoups and Vrugt (2010) in the following procedure: (i) the correlation is removed from the residuals by using an autoregressive model; (ii) the resulting residuals are 189 normalized by a linear model of variance; and (iii) the normalized residuals are characterized by 190 using the skew exponential power distribution. The data model parameters (i.e., coefficients of the 191 autoregressive model, the linear variance model, and the skew exponential power distribution) are 192 193 not specified by users, but estimated together with soil respiration model parameters during the Bayesian inference. The skew exponential power distribution is general in that by adjusting the 194 values of its kurtosis and skewness parameters the distribution can produce other distributions such 195 196 as the Laplace distribution (van Wijk et al., 2008; Ricciuto et al., 2011) and other distributions through using an exponential model with different kurtosis parameters (Tang and Zhuang, 2009). 197 It is worth pointing out that there exist other methods to account for the three assumptions. Evin 198 199 et al. (2013) suggested accounting for residual heteroscedasticity before accounting for residual autocorrelation. Lu et al. (2013) developed an iterative two-stage procedure to separately estimate 200 physical model parameters and data model parameters. Evin et al. (2014) developed a similar 201 procedure to first estimate model parameters and then estimate heteroscedasticity and 202 autocorrelation parameters. While this study uses the method of Schoups and Vrugt (2010), 203 exploring other methods is warranted in future studies. 204

After investigating the impacts of the data models on Bayesian inference, this study evaluates the impacts of the data models on predictive performance of the three soil respiration models. Using random samples generated during the Bayesian inference, a prediction ensemble is produced

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208 for each soil respiration model. The ensemble is used to evaluate predictive performance of the 209 models in a stochastic sense by estimating to what extent the models can predict future events. The evaluation in this study is done in a cross-validation manner by splitting the dataset of CO₂ efflux 210 into two parts for Bayesian inference and cross-validation, respectively. The evaluation of 211 predictive performance is important because different data models may give different parameter 212 distributions and accordingly different predictive performance. For example, the study of van Wijk 213 et al. (2008) concluded that the choice of the residual function is crucial to achieve accurate model 214 prediction and parameter estimation. Shi et al. (2014) showed that the posterior parameter 215 distributions and predictive performance given by two data models (weighted least square and 216 skew exponential power distribution after removing heteroscedasticity and autocorrelation) are 217 dramatically different, and a definitive conclusion was drawn that one data model is better than 218 219 the other. The evaluation of predictive analysis is conducted for the following two cases: (1) the prediction ensemble is generated by random samples of the soil respiration models only (i.e. 220 credible interval), and (2) the prediction ensemble is generated by random samples of not only the 221 soil respiration models but also the data models (i.e. predictive interval). The two cases lead to 222 different conclusions about the predictive performance. It is expected that the evaluation of 223 predictive performance conducted in this study can help select the most appropriate data model to 224 achieve optimal model predictions. 225

The remainder of the paper is organized as follows. Section 2 starts with a description of the evolving data models and their corresponding likelihood functions used in Bayesian inference, followed by a brief summary of the three soil respiration models. The results of Bayesian inference are discussed in Section 3 and Section 4, addressing the data model implications on parameter estimation and predictive performance, respectively. Section 5 summarizes the key findings andlimitations of this study, and provides recommendations for approaching data model selection.

232 **2** Methodology

This section starts with a description of the eight data models that account for the three pairs of assumptions about residuals in a stepwise manner in Section 2.1. The data models are used to build the likelihood functions used in Section 2.2 for Bayesian inference. The three soil respiration models and observations of CO_2 efflux are described in Sections 2.3 and 2.4, respectively. Metrics for evaluating predictive performance are presented in Section 2.5.

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239 2.1 Data models

This study considers eight evolving data models starting from a data model that assumes independent, homoscedastic, and Gaussian residuals to a data model that relaxes all the three assumptions. The eight data models are based on the generic normalized residual,

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$$a_t = \frac{\varepsilon_t}{\sigma_t}$$
 $a_t \sim X$, (1)

where $\varepsilon_t = d_t - Y_t$ is the residual (the difference between data d_t and its corresponding model simulation Y_t) at time or location t; σ_t is the standard deviation of the residual; and X is the probability density function (PDF) of a_t . The eight data models are formulated with different forms of ε_t , σ_t , and X. The standard least square (SLS) data model is

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$$a_t = \frac{\varepsilon_t}{\sigma_0} \qquad a_t \sim N(0,1) , \qquad (2)$$

249 where $\sigma_t = \sigma_0$ is a constant for all the data (i.e., homoscedasticity), and X is the standard normal 250 distribution, N(0,1). The unknown parameter σ_0 is estimated jointly with unknown physical model parameters. If σ_t is not a constant (i.e., heteroscedastic), SLS becomes the weighted least squared (WLS) data model. While heteroscedasticity can be accounted for through residuals transformation (e.g. Thiemann et al., 200; Smith et al., 2010b) or other similar approaches (Gragne et al., 2015) a linear heteroscedastic model $\sigma_t = \sigma_0 + \sigma_1 Y_t$ is assumed here by following other the studies of (Thyer et al.; (2009); Schoups and Vrugt; (2010), and; Evin et al.; (2013, 2014). With the linear model, there is no need to estimate σ_t for each data. Instead, σ_t is calculated by estimating only two parameters, σ_0 and σ_1 . The WSL data model is written as

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$$a_t = \frac{\varepsilon_t}{\sigma_0 + \sigma_1 Y_t} \qquad a_t \sim N(0, 1).$$
(3)

The two unknown parameters σ_0 and σ_1 are estimated jointly with unknown physical model parameters. The linear model assigns smaller weight to the data with larger simulation, Y_t . If the simulation is small and $\sigma_0 \gg \sigma_1 Y_t$, the weight becomes constant for all data. Both SLS and WLS assume that a_t is independently and identically distributed.

It is not uncommon that residuals are correlated in space and time, due to propagation of measurement errors (Tiedeman and Green, 2013) and model structure errors (Evin et al., 2014; Kavetski et al., 2013; Lu et al., 2013). The temporal correlation that occurs in the numerical example of this study can be accounted for <u>by</u> using a *p*-order autoregressive model. This leads to the data model of standard least square with autocorrelation (SLS-AC),

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$$a_t = \frac{\varepsilon_t - \sum_{i=1}^p \phi_i \varepsilon_{t-i}}{\sigma_0} \qquad a_t \sim N(0, 1)$$
(4)

where *p* is the order of autocorrelation, and ϕ_i is an autocorrelation coefficient. The unknown ϕ_i and σ_0 are estimated together with unknown model parameters. By extending the concept of correlated residuals to WLS leads to the weight least square with autocorrelation (WLS-AC),

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$$a_t = \frac{\varepsilon_t - \sum_{i=1}^p \phi_i \varepsilon_{t-1}}{\sigma_0 + \sigma_1 Y_t} \qquad a_t \sim N(0, 1)$$
(5)

273 The unknown parameters of σ_0 , σ_1 , and ϕ_i are estimated jointly with physical model 274 parameters. Equations (2) – (5) assume that the residuals are Gaussian.

The next four data models are similar to the previous four models except that the standard normal distribution of a_t is replaced by the skew exponential power distribution, $SEP(0,1,\xi,\beta)$, with zero mean and unit standard deviation (Schoups and Vrugt, 2010)

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$$p(a_t | \xi, \beta) = \frac{2\sigma_{\xi}}{\xi + \xi^{-1}} \omega_{\beta} \exp\left[-c_{\beta} |a_{\xi,t}|^{2/(1+\beta)}\right],$$
 (6)

279 where zero is mean, one is standard deviation, ξ is skewness, β is kurtosis,

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$$a_{\xi,t} = (\mu_{\xi} + \sigma_{\xi}a_t) / \xi^{\text{sign}(\mu_{\xi} + \sigma_{\xi}a_t)}$$
, $\mu_{\xi} = M(\xi - \xi^{-1})$, $\omega_{\beta} = \frac{\Gamma^{1/2}[3(1+\beta)/2]}{(1+\beta)\Gamma^{3/2}[(1+\beta)/2]}$,

281
$$\sigma_{\xi} = \sqrt{(1 - M^2)(\zeta^2 + \zeta^{-2}) + 2M^2 - 1}$$
, $M = \frac{\Gamma[1 + \beta]}{\Gamma^{1/2}[3(1 + \beta)/2]\Gamma^{1/2}[(1 + \beta)/2]}$, and

282
$$c_{\beta} = \left(\frac{\Gamma[3(1+\beta)/2]}{\Gamma[(1+\beta)/2]}\right)^{1/(1+\beta)}$$
 are derived variables of β and ξ , and $\Gamma[.]$ is the gamma function. The

kurtosis parameter { $\beta \in \mathbb{R}: -1 \le \beta \le 1$ } determines the peakness of the pdf such that the β values of -1, 0, and 1 give uniform, Gaussian and Laplace distributions, respectively. The skewness parameter { $\xi \in \mathbb{R}: 0.1 \le \xi \le 10$ } determines the skewness of the pdf such that the ξ values of 0.1, 1, and 10 give positively skewed, symmetric, and negatively skewed distributions, respectively. Setting $\beta = 0$ and $\xi = 1$ leads to $\mu_{\xi} = 0$, $\sigma_{\xi} = 1$, $\omega_{\beta} = 1/\sqrt{2\pi}$, $c_{\beta} = 1/2$ and $a_{\xi,t} = a_t$, and the skew exponential power distribution $SEP(0, 1, \xi = 1, \beta = 0)$ becomes the standard normal distribution,

290
$$p(a_t | \xi = 1, \beta = 0) = \frac{1}{\sqrt{2\pi}} \exp\left[-\frac{1}{2}(a_t)^2\right].$$
 (7)

which is the data model of SLS in equation (2).

292 Replacing $a_t \sim N(0,1)$ with $a_t \sim SEP(0,1,\xi,\beta)$ in equations (2) – (5) leads to the data models 293 SEP, WSEP, SEP-AC, and WSEP-AC as follows,

294
$$a_t = \frac{\varepsilon_t}{\sigma_0}$$
 $a_t \sim SEP(0, 1, \xi, \beta)$ (8)

295
$$a_t = \frac{\varepsilon_t}{\sigma_0 + \sigma_1 Y_t} \qquad a_t \sim SEP(0, 1, \xi, \beta) .$$
(9)

296
$$a_{t} = \frac{\varepsilon_{t} - \sum_{i=1}^{p} \phi_{i} \varepsilon_{t-1}}{\sigma_{0}} \qquad a_{t} \sim SEP(0, 1, \xi, \beta)$$
(10)

297
$$a_{t} = \frac{\varepsilon_{t} - \sum_{i=1}^{p} \phi_{i} \varepsilon_{t-1}}{\sigma_{0} + \sigma_{1} Y_{t}} \qquad a_{t} \sim SEP(0, 1, \xi, \beta)$$

$$(11)$$

In comparison with the Gaussian data models, the SEP-based data models have two more parameters (ξ and β) to be estimated jointly with physical model parameters. WSEP-AC dData model WSEP-AC, which is known as the generalized likelihood function, is the most commonly used SEP-based data model (e.g. Vrugt and Ter Braak, 2011; Hublart et al., 2016; Scholz et al., 2018). A summary table of the eight data models with corresponding parameters is provided in the supplementary materials.

304 2.2 Bayesian inference and likelihood functions

Consider a Bayesian inference problem for a nonlinear model, f, used to simulate state variables (e.g., CO₂ efflux), $d = Y(\theta) + e\underline{e}$, where d is a vector of data, θ is a vector of model parameters, and $e\underline{e}$ is a vector of residuals that may include errors in data, model parameters, and model structures. The goal of Bayesian inference is to estimate the posterior distributions, $p(\theta|d)$, of model parameters, θ , given data, d, using Bayes' theorem (Box and Tiao, 1992)

310
$$p(\theta | d) = \frac{p(d | \theta) p(\theta)}{\int p(d | \theta) p(\theta) d\theta}$$
(12)

where $p(\theta)$ is the prior distribution, and $p(d|\theta)$ is the likelihood function to measure goodness-offit between model simulations, $Y(\theta)$, and data, *d*. The prior distribution can be obtained from data of previous studies (e.g. Elshall and Tsai, 2014) or expert judgment. When prior information is lacking, a common practice is to assume uniform distributions with relatively large parameter ranges so that the prior distributions do not affect the estimation of posterior distributions.

The data models above can be used to construct the likelihood functions. For the Gaussian data models given in equations (2) - (5), the corresponding Gaussian likelihood functions are straightforward, and an example is equation (7). For the SEP data models, the corresponding likelihood that is called generalized likelihood function is (Schoups and Vrugt, 2010)

320
$$p(\boldsymbol{d} | \boldsymbol{\theta}) = p(\boldsymbol{\varepsilon}_{t} | \boldsymbol{\theta}) = \prod_{t=1}^{n} \sigma_{t}^{-1} \frac{2\sigma_{\xi}}{\xi + \xi^{-1}} \omega_{\beta} \exp\left(-c_{\beta} \left|a_{\xi,t}\right|^{2/(1+\beta)}\right).$$
(13)

where *n* is the dimension of *d*. The Gaussian likelihood functions are special case of the generalized likelihood functions. For example, by setting $\beta = 0$, $\xi = 1$, $\phi_i = 0$, $\sigma_t = \sigma_0$, $\sigma_{\xi} = 1$, $\mu_{\xi} = 0$,

323
$$\omega_{\beta} = 1/\sqrt{2\pi}$$
, $c_{\beta} = 1/2$, and $a_{\xi,t} = a_t$, equation (13) becomes the likelihood function corresponding

to the SLS data model. Replacing $\sigma_t = \sigma_0$ by $\sigma_t = \sigma_0 + \sigma_1 E_t$, equation (13) becomes the likelihood function of the WLS data model.

In this study, the posterior distributions of the data model parameters are jointly estimated with the soil respiration model parameters using the MT-DREAM_(ZS) code (Laloy and Vrugt, 2012). MT-DREAM_(ZS) implements a Markov chain Monte Carlo (MCMC) algorithm by running multiple Markov chains in parallel with adaptive proposal distribution, multiple-try sampling, -and sampling from an archive of past states. These state-of-the-art features assist in overcoming common challenges in the sampling landscape space such as multimodality, ill-conditioning, and high dimensionality, and thus allow for accurate exploration of the targeted distributions.

333

2.3 Soil respiration models

Zhang et al. (2014) studied the Birch effect (the peak soil microbial respiration pulses in 334 response to episodic rainfall pulses), and developed five models, evolving from an existing four-335 carbon pool model to models with additional carbon pools and/or explicit representations of soil 336 moisture controls on carbon degradation and microbial uptake rates. Three of the five models are 337 used in this study, and they are dented as 4C, 5C, and 6C. Note that model 4C is model 4C NOSM 338 of Zhang et al. (2014), not their model 4C. Figure 1 is the diagram of model 6C, the most complex 339 one among the five models. The simplest one, model 4C, has four carbon pools, i.e., soil organic 340 carbon (SOC), dissolved organic carbon (DOC), microbial biomass (MIC), and enzymes (ENZ), 341 and does not consider the soil moisture control on carbon degradation and microbial uptake rates. 342 343 Models 5C and 6C hasve an explicit representation of soil moisture controls on the rates. Based on the dual Arrhenius and Michaelis-Menten kinetics model, the original SOC degradation rate, 344 V_{decom} , is (Davidson et al., 2011; Davidson and Janssens, 2006) 345

$$346 \qquad V_{decom} = V_{max} C_{ENZ} \frac{C_{SOC}}{K_m + C_{SOC}} \tag{14}$$

where V_{max} [s⁻¹] is the maximum SOC degradation rate per unit enzyme when the substrates is not limiting, C_{ENZ} [gCm⁻³] is enzyme pool size, C_{SOC} [gCm⁻³] is SOC pool size, and K_m is the halfsaturation for SOC. The original microbial uptake rate, V_{uptake} , is (Davidson et al., 2011; Davidson and Janssens, 2006)

351
$$V_{uptake} = V_{\max_up} C_{MIC} \frac{C_{DOC}}{K_{m_up} + C_{DOC}} \frac{C_{O2}}{K_{m_upO2} + C_{O2}},$$
(15)

where V_{max_up} [s⁻¹] is the maximum DOC uptake rate when the substrates is not limiting, C_{MIC} [gCm⁻³] is the microbial biomass<u>MIC</u> pool size, C_{DOC} [gCm⁻³] is the DOC pool size, C_{O2} [m³m⁻³] ³] is the gas concentration of O₂ in the soil pore, and K_{m_up} [gCm⁻³] and K_{m_upO2} [m³m⁻³] are the corresponding half-saturation constants for DOC and O₂, respectively. With the explicit representation of soil moisture control, the two rates become (Zhang et al., 2014)

357
$$V_{decom} = V_{max} C_{ENZ} \frac{C_{SOC}}{K_m + C_{SOC}} \left(\frac{\theta}{\theta_s}\right)$$
(16)

358
$$V_{uptake} = V_{max_up} C_{MIC} \frac{C_{DOC}}{K_{m_up} + C_{DOC}} \frac{C_{O2}}{K_{m_upO2} + C_{O2}} \left(\frac{\theta}{\theta_s}\right)$$
 (17)

359 where θ [-] is the volumetric soil moisture, and θ_s [-] is the porosity.

In addition to using the new rate equations, models 5C and 6C have more carbon pools. In model 5C, DOC is split into two sub-pools for wet zone and dry zone of soil pores, and only the wet DOC is used by MIC, as shown in Figure 1. The moisture-controlled microbial uptake rate becomes

364
$$V_{uptake} = V_{max_up} C_{MIC} \frac{C_{DOC_w}}{K_{m_up} + C_{DOC_w}} \frac{C_{O2}}{K_{m_upO2} + C_{O2}} \left(\frac{\theta}{\theta_s}\right).$$
(18)

where C_{DOC_w} [gCm⁻³] is the DOC pool size in the wet soil pores. Model 6C is more complex in that ENZ is further split into two sub-pools for wet and dry pores, and both the wet and dry ENZ are subject to degradation, as shown in Figure 1. The moisture-controlled SOC degradation rate becomes

369
$$V_{decom} = V_{\max} C_{ENZ_{-}W} \frac{C_{SOC}}{K_m + C_{SOC}} \left(\frac{\theta}{\theta_s}\right)$$
(19)

370 for the wet ENZ and

371
$$V_{decom} = V_{\max} C_{ENZ_{-D}} \frac{C_{SOC}}{K_m + C_{SOC}} \left(1 - \frac{\theta}{\theta_s} \right) \varepsilon_D$$
(20)

for the dry ENZ, where C_{ENZ} w [gCm⁻³] is the wet soil pores enzyme pool size, C_{ENZ} [gCm⁻³] is 372 the enzyme pool size in the dry soil pores, and ε_D is the catalysis efficiency of the dry zone enzyme. 373 Due to considering the moisture control and adding more soil pools, model 5C is expected to 374 375 be significantly better than model 4C for simulating the Birch effect. Since the accumulated ENZ 376 in dry soil is secondary, model 6C is expected to be slightly better than model 5C. In terms of model structural error, model 4C has the largest model structure error, model 5C has significantly 377 378 less model structure error, and model 6C has the smallest model structural error. In other words, model 6C has the highest model fidelity (i.e. lowest model discrepancy) among the three models. 379 As shown below, the degree of model structural error is reflected in the process of Bayesian 380 381 inference and verified by the cross-validation.

382 2.4 Observations and parameter estimation

Figure 2 plots the time series of 17,016 observations of soil moistermoisture- and CO₂ efflux 383 used in this study. The observations were obtained during the entire year of 2007, covering a long 384 period of dry season prior to monsoon and episodic rainfall events during monsoon. The first two 385 third of this dataset is used for the Bayesian inference, and the last one third is used for cross-386 validation. The inference and cross-validation periods have both dry and wet periods, as shown in 387 Figure 2. The observation site is located within the Santa Rita Experimental Range (SRER, 388 31.8214°N, 110.8661°W, elevation 1,116 m) outside of Tucson, Arizona (Barron-Gafford et al., 389 2011; Scott et al., 2009). This savanna site was covered by 22% of perennial grass, forbs and 390 subshrubs and 35% of mesquite. The soils are uniformly Comoro loamy sand (77.6% sand, 11.0% 391 clay, and 11.4% silt). The half-hourly atmospheric forcing data were collected from measurements 392 through an eddy covariance tower (Scott et al., 2009). This includes downward shortwave, 393 longwave, precipitation, wind, air temperature, humidity, and pressure. Volumetric CO₂ 394 concentration was measured at a half-hourly interval through compact probes. The CO₂ efflux was 395 estimated from the gradient of CO₂ concentration measured at two depths of 2 cm and 10 cm 396 through Fick's first law of diffusion, and the estimates were validated against measurements from 397 a portable CO₂ gas analyzer. 398

The parameters estimated in this study include the parameters of the soil respiration models (4C - 6C) and the parameters of the data models described in Section 2.1. The estimated parameters of models 4C and 5C include the microbial carbon use efficiency (CUE) [g/g], enzyme production rate, k_e [g/m³s], microbial turnover rate, τ_m [1/s], and enzyme turnover rate τ_e [1/s]. Uniform distributions are used as the prior in the Bayesian inference, and the ranges of the four parameters are 0.2 - 1.00, $1 \times 10^{-12} - 1 \times 10^{-7}$, $1 \times 10^{-12} - 1 \times 10^{-5}$ and $1 \times 10^{-11} - 1 \times 10^{-6}$, respectively. The values of other parameters are fixed at the values used in Allison et al. (2010). Model 6C has two more parameters, and they are the catalysis efficiency ε_D [-] and the turnover rate of the dryzone enzymes τ_{en} [1/s]. The prior of the two parameters are uniform distributions with the ranges of 0.2 - 0.8 and $1 \times 10^{-12} - 1 \times 10^{-8}$, respectively.

The DREAM-based MCMC simulation is conducted for a total of 24 cases, the combinations of eight data models and three soil respiration models. For each case, the parameter distributions are obtained after drawing a total of 5×10^5 samples using five Markov chains. The Gelman and Rubin (1992) R-statistic is used for convergence diagnostic, and it approaches one in less than $413 \quad 40,000 \times 10^4$ samples. The initial 50% of the samples are discarded during the burn-in period.

414 2.5 Metrics for evaluating predictive performance

Three criteria are used to evaluate the predictive performance of the soil respiration models and data models, and they are central mean tendency, dispersion, and reliability. Each criterion is measured by a single metric. In addition, a newly defined metric <u>by (Elshall et al., 2018)</u> is also used for simultaneously measuring the three criteria.

The central mean tendency is measured in this study using the Nash-Sutcliffe model efficiency
(NSME) coefficient (Nash and Sutcliffe, 1970),

421
$$NSME = 1 - \sum_{i=1}^{n} (d_i - \overline{\mathbf{Y}}_i)^2 / \sum_{i=1}^{n} (d_i - \overline{\mathbf{d}})^2,$$
 (21)

where *n* is the number of cross-validation data, d_i is the *i*-th data, $\overline{\mathbf{d}}$ is the mean of the data, and $\overline{\mathbf{Y}_i}$ is the mean of the prediction ensemble, Y_i , for d_i . NSME ranges from $-\infty$ to 1, with *NSME* = 1 corresponding to a perfect match between data and mean prediction, i.e., the ensemble is centered on the data. *NSME* = 0 indicates that the model predictions are as only accurate as the mean of the

data, while an efficiency NSME < 1 indicates that the mean of data is a better prediction than the 426 mean prediction. 427

428

In addition to the central mean tendency, it is also desirable that the ensemble is precise with small dispersion and reliable to cover all the data. This study uses a nonparametric metric for 429 dispersion, and it is the sharpness of a prediction interval (e.g. Smith et al., 2010a) 430

431
$$Sharpness = 1/n \sum_{i=1}^{n} \left[Max(\mathbf{Y}_i) - Min(\mathbf{Y}_i) \right]$$
(22)

where \mathbf{Y}_i is the prediction ensemble within the 95% prediction interval₂ (the Bayesian credible 432 interval, not the confidence interval used in nonlinear regression (Lu et al., 2013). Smaller values 433 of sharpness indicate better prediction precision. Reliability is measured using predictive coverage. 434 (e.g. Hoeting et al., 1999), which is the percentages of data contained in the prediction interval. 435 Larger predictive coverage values are preferred. 436

To account for the trade-off between the three metrics, $\frac{1}{2}$ (2018b) -defined relative 437 model score (RMS) that simultaneously measure all the three criteria. Scoring rules are commonly 438 439 used in hydrology to assess predictive performance (e.g., Weijs et al., 2010; Westerberg et al., 2011). RMS is used in this study to measure the relative predictive performance of the 440 combinations of soil respiration models and data models. For combination M_i , RMS is defined as 441

442
$$RMS(M_j) = \sum_{i=1}^{n} \frac{p(d_i | \mathbf{Y}_{ij}, M_j)}{\sum_{j=1}^{m} p(d_i | \mathbf{Y}_{ij}, M_j)} \times 100$$
 (23)

where m is the number of combinations; and the ensemble prediction Y_{ij} is similar to Y_i above 443 where is with i index i over time and index j specific to the j-th combination. The density function 444 $p(d_i|\mathbf{Y}_{ij})$ can be evaluated by first obtaining the density function $p(\mathbf{Y}_{ij})$ of the ensemble prediction 445 Y_{ii} (e.g., by using the kernel density function) and then evaluating $p(d_i|Y_{ij})$ using interpolation 446

methods based on the intersection of Y_{ij} and d_i . More details of about evaluating *RMS* can be found in Elshall et al. (2018). This evaluation is based purely on the model predictions, and does not involve any assumptions on the models, their parameters, and likelihood functions. Larger RMS values indicate better overall predictive performance. A figure of our workflow scheme is presented in the supplementary materials.

452 **3** Results of Bayesian Inverse Modeling

This section analyzes the residuals of the best realization (with the highest likelihood value) of the MCMC simulation to understand whether the assumptions of the eight data models hold. The impacts of the data models on the posterior parameter distributions are also analyzed.

456

3.1 Residual characterization

Figure 3 shows residual plots for model 6C based on data models SLS and WSEP-AC. SLS is 457 458 the simplest data modelone with the assumptions of homoscedastic, independent, and Gaussian residuals, and the WSEP-AC is the most complex one without the assumptions. Model 6C is the 459 most complex model and also the best one as ranked by Zhang et al. (2014) using Bayesian model 460 selection. The variable a_t plotted in Figures 3a-3c and Figures 3d-3f is defined in equations (2) and 461 (11), respectively. Figures 3a - 3c show that all the three residual assumptions are violated when 462 463 SLS is used, because (i) the residual variance is not constant, but increases as a function of the simulated CO₂ efflux (Figure 3a); (ii) the autocorrelation function at most lags is beyond the 95% 464 confidence interval (Figure 3b); (iii) and the standard normal density function cannot adequately 465 466 characterize the residuals (Figure 3c). Figures 3d-f show that, after relaxing the three assumptions, the processed residuals, a_t , can be well characterized by WSEP-AC. Figure 3d shows that, after 467 normalizing ε_t with the linear variance ($\sigma_t = 0.034 + 0.099E_t$), the variation of the variance of 468 469 a_t becomes significantly smaller, although the variance is still not constant. Figure 3e shows that,

470 after removing a first-order autoregressive model from ε_t , a_t becomes less correlated, although the correlation is not fully removed. The two coefficients of the autoregressive model are $\phi_1 = 0.989$ 471 and $\phi_2 = 4.5 \times 10^{-6}$; the small value of ϕ_2 indicates that there is no need to attempt an autoregressive 472 model of higher order. Figure 3f shows that a_t follows the SEP distribution with the estimated 473 skewness coefficient of $\xi = 0.933$ and kurtosis coefficient of $\beta = 0.998$. As a summary, Figure 474 3 shows that it is important to examine the residuals and to determine whether the selecteda data 475 model is adequate for charactering the residuals. Although WSEP-AC still cannot perfectly 476 characterize ε_t , it is significantly better than SLS. 477

Although the Gaussian assumption used in SLS is violated for model 64C (Figure 3c), this is 478 479 not generally the case for other data models and soil respiration models. This is shown in Figure 4, which presents the quantile-quantile (Q-Q) plot for the eight data models and the three soil 480 respiration models. For SLS, WLS, SLS-AC, and WLS-AC, the theoretical quantiles are based on 481 the standard normal distribution, N(0,1); for SEP, WSEP, SEP-AC, and WSEP-AC, the theoretical 482 483 quantiles are based on the standard skew exponential power distribution, SEP(0,1,1,0). If the residuals follow the assumed standard distributions, the O-O plots fall on the 1:1 lines, which is 484 marked as the theoretical lines in Figure 4. If the residuals are Gaussian or SEP but not standard, 485 the Q-Q plots fall on a straight line but not the 1:1 line. Figures 4a and 4e show that, for all the soil 486 respiration models, the Q-Q plots of SLS and SEP deviate significantly from the theoretical lines 487 and exhibit fat-tail behaviors, which is an indication of outliers (Thyer et al., 2009). The deviation 488 is reduced after accounting for autocorrelation in SLS-AC and SEP-AC, as shown in Figures 4c 489 490 and 4g. It is interesting to observe from the two figures that the Q-Q plots of the three models are almost visually identical. The deviation is almost fully removed after accounting for 491 heteroscedasticity in WLS and WSEP in that their corresponding Q-Q plots fall on the 1:1 lines, 492

especially for models 5C and 6C, as shown in Figures 4b and 4f. However, the Q-Q plots start deviating from the 1:1 lines as shown in Figures 4d and 4h, after accounting for both heteroscedasticity and autocorrelation in WLS-AC and WSEP-AC. As a summary, Figure 4 shows that, for the numerical example of this study, either the Gaussian or the SEP distribution is valid if heteroscedasticity is accounted for in the data models. However, accounting for autocorrelation in the data models does not help improve the characterization of the residual distributions.

499

3.2 Posterior parameter distributions

While Figures 3 and 4 help understand validity of the three assumptions used in the data 500 501 models, the impacts of the data models on estimating model parameter distributions must be evaluated separately. This section discusses the impact of the data model selection on parameter 502 estimation with the objective of understanding whether if incorrect specification of the data model, 503 504 will necessarily leads to biased parameter estimates. Such assessment is not a trivial task for twohree main reasons. First, microbial soil respiration models aggregate complex natural processes 505 and spatial details into simpler conceptual representations. As a results several model parameters 506 507 are effective values of several complex natural processes that cannot be actually measured in the field as discussed by Vrugt et al. (2013). SecondIn addition, even for model parameter that can be 508 measured in the field, since the model structure is imperfect, calibrated it can be the case that 509 parameter values are sometimes can be accepted beyond their physically reasonable range, as 510 discussed by Pappenberger and Beven (2006). This is often undesirable, if we seek to make the 511 models more mechanistically descriptive. 512

513 We focus our discussion on carbon use efficiency (CUE) for microbial growth <u>due to two</u> 514 <u>reasons: (1) since CUE is a fundamental parameter in microbial soil respiration models, and (2) a</u> 515 <u>physically</u> reasonable <u>physical</u> range for CUE can estimated. The concept of microbial

516 CUE(Allison et al., 2010; Bradford et al., 2008; Manzoni et al., 2012; Wieder et al., 2013) has 517 been used to present fundamental microbial processes in recent microbial enzyme models (Allison et al., 2010; German et al., 2011; Schimel and Weintraub, 2003; Wang et al., 2013). The microbial 518 CUE, which is marked between MIC and CO2 in Figure 1, controls microbial growth, enzyme 519 production and microbial respiration. A physically reasonable range of CUE can be estimated from 520 the physical viewpoint (Tang and Riley, 2014). Sinsabaugh et al. (2013) study showeds that the 521 thermodynamic calculations support a maximum CUE of 0.60 and that methods used toprevious 522 studies that estimate CUE in terrestrial systems report a mean value of 0.55. Theoretically, there 523 is no lower limit for CUE as it can approach zero, and CUE< 0.1 has been are-reported for 524 terrestrial ecosystems (e.g., Fernández-Martínez et al., 2014) and used in modeling studies (Li et 525 al., 2014). Note that, for inverse modeling with MCMC sampling, we did not assume CUE 526 527 maximum value of 0.6. In other words, for parameter estimation and predictive performance we did not impose the constraint that CUE is less than 0.6. We merely use this CUE maximum value 528 529 of 0.6 to evaluate whether the posterior CUE parameter samples obtained using different data 530 models and different soil respiration models are within the physically reasonable range of $0 \sim 0.6$. Figure 5 plots the CUE posterior marginal density of the three soil respiration models obtained 531 using the eight data models. The physical range between zero and 0.6 is marked in yellow. Figure 532 5 shows that the CUE posterior parameter distribution offer Model 6C obtained using for all the 533 data models likelihood functions that does not account for autocorrelation are within thea 534 physically reasonable physical range. For models 4C and 5C, the posterior parameter samples are 535 outside the physical range for six data models. For model 4C, the posterior parameters are within 536 the physical range only for data models SEP and WSEP; for model 5C, the two data models are 537 538 WLS and WSEP. It is not surprising to find the posterior parameter distribution of models 4C and

539 5C, which have a certain degree of model structure error, to be out of the <u>physically</u> plausible 540 physical range. This can be attributed to two reasons. First, the model solution can be biased toward 541 the missing processes in the model structure such as the additional carbon pool in both 4C and 5C 542 or <u>missing</u> the explicit accounting for soil <u>moistermoisture</u> –in 4C. Second, biased parameter 543 estimation can compensate for model structure inadequacy and other sources of discrepancy in 544 both the physical model<u>s</u> and the <u>statistical data</u> model<u>s</u>.

In addition, it is important to understand how accounting for autocorrelation, heteroscedasticity 545 and non-Gaussian residuals can affect the parameter estimation. -First, it is observed in Figure 5e-546 547 h that we obtained biased parameter estimates that is are outside the reasonable physically reasonable range when autocorrelation is explicitly accounted for as shown in Figure 5e-h. This 548 may suggest again that accounting for heteroscedasticity is desirable but accounting for 549 550 autocorrelation is not. A possible reason is that filtering autocorrelation may reduce the residual space such that the transformed residual space cannot correspond to the parameter space of the 551 models. In other words, parameter information may be lost due to filtering out autocorrelation. 552 However, it is not fully understood why this does not occur for the model 6C under data model 553 SLS-AC (Figure 5e), and more research is warranted. Second, unlike accounting for auto-554 correlation, accounting only for heteroscedasticity (i.e., WLS and WSEP) since this will only 555 amplifiesy or reduces the variance without affecting the structure of the residual space. Figures 5c-556 d shows that account for heteroscedasticity (i.e. WLS and WSEP) tends to improve the parameter 557 estimation in comparison with homoscedastic data models (i.e., SLS and SEP) shown in Figure 558 559 5a-b. Finally, with respect to non-Gaussian residuals, Schoups and Vrugt (2010) proposes suggested that, compared to Gaussian pdf, the peaked pdf of the SEP with heavier a longer tails 560 561 compared to Gaussian pdf-is useful for making parameter inference robust against outliers. To a

562 certain degree, this can be substantiated by the results in Figure 5a-d, <u>insuch</u> that SEP and WSEP
 563 provide more favorable parameter estimates than SLS and WLS.

564 Finally, from Figure 5a we can also notice shows that the posterior parameter distributions of SLS (Figure 5a) is are very narrow for the three soil respiration models. Thise narrow posterior 565 parameter distributions of SLS compared to other likelihood functions can be attributed to several 566 reasons. Since SEP distribution can have heavier-longer tails than Gaussian distribution, this can 567 further increase the samples acceptance ratio from tails resulting in wider distribution (Figure 5b). 568 569 In addition, accounting for heteroscedasticity will result in wider the posterior parameter distribution (Figure 5c) due to accepting higher variances at peak effluxes. Moreover, filtering 570 correlation (Figure 5e-h) increases the entropy, and leads to wider distributions. 571

572

4.

Results of Predictive Performance

Based on the last one third of the CO_2 efflux observations, a cross-validation test was 573 conducted for all the 24 models, the combinations of three soil respiration models and eight data 574 575 models. Given For the cross-validation period, the predictive performance is examined using the four statistical metrics that are defined in Section 2.5. The metrics are also calculated for the 576 calibration period. This is not to perform Bayesian model selection given the calibration data, but 577 to better understand the impact of data models on predictive performance of the three soil 578 respiration models. For each calibration and each cross-validation data, a prediction ensemble is 579 generated from the two perspectives of parametric uncertainty only and total uncertainty, as 580 581 presented in Section 4.1 and 4.2, respectively.

582

583 4.1 Predictive performance with parametric uncertainty of soil respiration model

In this section the ensemble is generated by running the soil respiration models with the posterior samples (obtained from the Bayesian inference) of the physical model parameters. In other words, the ensemble addresses parametric uncertainty of the soil respiration models only. Considering the relative contribution of parametric uncertainty only will provide insights for modeling approaches that attempt to segregate various sources of uncertainty (e.g., Thyer et al., 2009 ; Tsai and Elshall, 2013).

The four statistics above (i.e. NSME, sharpness, coverage, and RMS) are calculated for the three soil respiration models and the eight data models. Taking data models SLS and WSEP-AC as an example, Figure 6 plots the data (for the calibration and cross-validation periods separately) along with the mean and 95% credible intervals of the prediction ensemble for the three models.

Figure 6 shows that the data models affect model simulations for all the models. The statistics, especially RMS, indicate that WSEP-AC has better predictive performance than SLS. This is most visually obvious for model 6C during the cross-validation period after 330 days, as the prediction ensemble of SLS (Figure 6k) cannot cover the observations, <u>whereasunlike</u> the prediction ensemble of WSEP-AC can (Figure 6l). This conclusion that WSEP-AC outperforms SLS agrees with that drawn from Figures 3 and 4.

Figure 7 plots the four statistics for all the soil respiration models and data models. Figures 7a and 7b show the predictive performance with respect to the central mean tendency <u>measured by</u> using NSME for both the calibration and cross-validation periods respectively. The results indicate that, <u>under all data models</u>, the low fidelity model 4C <u>under all data models will</u> over-fits the data <u>and</u> result<u>sing</u> in biased predictions, <u>such in</u> that the NSME values become significantly worse (<u>e.g.</u>, from 0.6 to -0.6) from the calibration to the cross-validation period. This is confirmed by the visual inspection of Figures 6a <u>and</u>, 6b, 6g for data model SLS, and <u>of Figures 6b and</u> 6h for data models SLS and WSEP-AC. For models 5C and 6C, their NSME values vary with the data models; and the central mean accuracy is the worst for SLS-AC that considers only autocorrelation (Figure 6b).

With respect to parametric uncertainty estimation, Figures 7c and 7d show <u>that</u> sharpness generally increases when the three assumptions in the data models are gradually relaxed from SLS to WSEP-AC. This is even more obvious during the validation period. Given that the prediction ensemble does not center on the data, the increasing sharpness is desirable as it improves reliability. This is confirmed by the reliability plots in Figures 7e and 7f. The exceptions are again for SLS-AC and SEP-AC that generally have the lowest coverage.

With respect to the overall predictive performance <u>measured by RMS</u>, the same variation pattern and exception are also observed in the RMS plots in Figures 7g and 7h. This is not surprising because RMS is the metric that can be used to measure all the three criteria (central mean tendency, sharpness, and reliability). Since the prediction ensemble is not centered on the data, the sharpness and reliability are the decisive factors for evaluating the predictive performance.

As a summary, while it is necessary to account for heteroscedasticity in a data model, caution is needed when accounting for autocorrelation in the manner described in Section 2.1. In addition, after comparing the RMS values of the residuals using the Gaussian and SEP distributions, the conclusion is that the SEP distribution outperforms the Gaussian distribution with respect to predictive performance. Finally, uncertainty underestimation <u>isas</u> evidencedt by the very small predictive coverage. The underestimation of uncertainty for all the physical models with all <u>likelihood functionsthe data model makes sense</u> not unexpected because only parametric 629 uncertainty is considered in this study. Considering the overall predictive uncertainty is the
630 subject of the next section.

631 4.2 Predictive performance with total uncertainty

The simulated output $\mathbf{Y}(\boldsymbol{\theta}_p)$ is will generally not be equal to the observed output **d**, and we 632 have a residual term **<u>e</u>** due to measurement, input and model structure errors such that 633 $\mathbf{d} = \mathbf{Y}(\boldsymbol{\theta}_p) + \boldsymbol{\varepsilon}$. Accounting for the error term_ $\mathbf{e}-\boldsymbol{\varepsilon}$ can be through separating various error terms. 634 For example, in section 4.1 we obtained uncertainty due to the physical model parameters. 635 Accounting for other sources of uncertainty can be done using a single model approach (e.g. Thyer 636 et al., 2009) or a multi-model approach (e.g. Tsai and Elshall, 2013). Alternatively, we can quantify 637 the uncertainty based on total residuals that separates out parametric uncertainty, so the residual 638 error includes errors ins measurements, model inputs, and model structures uncertainty (e.g. Thyer 639 et al., 2009; Schoups and Vrugt, 2010). This lumped approach is based on sampling the residuals 640 model $\varepsilon(\theta_{\varepsilon})$ with parameters θ_{ε} . SLS has one fixed parameter that is the constant variance, and 641 other data models have two to six parameters. Thus in this section the prediction ensemble 642 addresses parametric uncertainty of not only the soil respiration models but also the data models. 643 When generating the prediction ensemble in the procedure described by Schoups and Vrugt 644 (2010), an ensemble of residuals is first generated by running the data models with posterior 645 samples of the data model parameters for the positive carbon efflux domain; the residual ensemble 646 is then added to the prediction ensemble generated in Section 4.1. 647

We start by <u>athe</u> visual assessment of the predictive performance. Figure 8 is similar to Figure 649 6 with the exception that Figure 8 considers the overall <u>all</u>-predictive uncertainty (i.e. parametric 650 and output uncertainty), while Figure 6 considers the parametric uncertainty only. Figure 8 reveals 651 a practical observation about accounting for the overall uncertainty through the lumped approach of sampling the <u>dataresiduals</u> model<u>s</u>. For example, Figure 8b shows that, despite the wide prediction interval of model 4C, which has the model with significant model structure error, it could cannot capture the birch pulse around day 180. This clearly It indicates that proper using a data model for model residuals cannot compensate modeling of the residuals will not make up for of significant model structure error.

Figure 9 plots the four statistics (NSME, sharpness, predictive coverage, and RMS) of the three soil respiration models under the eight data models to assess the predictive performance. First wWith respect to central mean tendency, tThe NSME values in Figures 9a-9b are visually the same as those in Figures 7a-7b, indicating that the central mean accuracy under parametric uncertainty is the same as that under predictive uncertainty.

With respect to uncertainty, the values of sharpness and predictive coverage increase 662 substantially (Figures 9c - 9f). In particular, Figures 9e and 9f show that, except for SLS and SEP, 663 the predictive coverage of the rest of the six data models are close to 100% for all the three soil 664 respiration models, indicating that the prediction intervals cover almost all the data. This is 665 demonstrated in Figures 6 for WSEP-AC. Similar to Figures 7c and 7d, Figures 9c and 9d also 666 show a general pattern that the sharpness increases when the three assumptions in the data models 667 are gradually relaxed from SLS to WSEP-AC. The data models that account for autocorrelation 668 are still the exceptions. 669

With respect to the overall predictive performance, the RMS values are largely determined by <u>the mean accuracy and sharpness as the predictive coverage is similar for different data models.</u> Figures 9g and 9h of RMS show that the predictive performance of the four data models that account for autocorrelation is worse than that of the other four data models. This suggests again that one needs to be cautious when building autocorrelation into a data model. This is consistent

675 with the finding of Evin et al. (2013, 2014) that accounting for autocorrelation before accounting for heteroscedasticity or jointly accounting for autocorrelation and heteroscedasticity can result in 676 poor predictive performance. In summary, Figures 9g and 9h show for both the calibration and 677 678 prediction periods that accounting for heteroscedasticity (i.e. in WLS and WSEP) will-gives the best overall predictive performance, and accounting for autocorrelation without heteroscedasticity 679 (i.e.in SLS-AC and SEP-AC) will gives the worst overall predictive performance. Finally, for the 680 three soil respiration models, RMS shows that model 4C has the worst predictive performance for 681 both the calibration and cross-validation data. Generally speaking, the high fidelity model 6C 682 683 outperforms model 5C for both the calibration and cross-validation data, which justifies the complexity of model 6C. 684

To demonstrate the impacts of the data models on predictive performance of the soil respiration models, Figure 10 plots the model simulations and predictions given by model 6C during the calibration and cross-validation periods using all the eight data models.

In-Figure 10 is used to investigate we try to understand the predictive performance 688 characteristics of the different data models. bBy looking atexamining the predictive performance 689 of model 6C, \overline{S} specific predictive performance patterns can be identified. Figures 10-a – 10d 690 show that SLS and SEP have similar predictive performance with SEP generally having better 691 predictive performance especially during the validation period. Accounting for heteroscedasticity 692 using WLS as shown in Figures 10e and 10h Not accounting for heteroscedasticity will 693 underestimate the predication uncertainty (Figure 10b and Figure 10d). This is mainly because the 694 variance of the efflux residuals increases with the magnitude of the carbon effluxes (Figure 3a), 695 and thus assuming constant variance is not representative. Accordingly, accounting for 696 697 heteroscedasticity using WLS (Figure 10e) or WSEP (Figure 10h) will make the predictions more

698 sensitive to peck carbon effluxes. This and will generally improve the predictive coverage on the 699 expense of sharpness and the central mean tendency. While WLS and WSEP have similar predictive performance-, However, WSEP maintains has slightly better central mean tendency and 700 701 overall predictive performance than WLS. Figures 10i - 10l show that Aaccounting for autocorrelation using SLS-AC and SEP-AC as shown in Figures 10i and 10l reduces the 702 information content of the residuals, and thus resultings in wider uncertainty bands and 703 insensitivity to peak carbon effluxes as compared to SLS and SEP (Figures 10a-d), which may be 704 due to reduction of information content of the residuals. This resulteds in deteriorating the 705 706 sharpness, the central mean tendency and the capturing of peak carbon fluxes, especially during the validation period. Figures 10m - 10p show that Aaccounting for both heteroscedasticity and 707 autocorrelation using WLS-AC and WSEP-AC-will makes the inference robust against peaek 708 709 carbon effluxes. However, yet due to the loss of information content, the uncertainty bands are still wider, and uncertainty becomes overestimated especially during validation period as 710 compared to WLS and WSEP (Figures 10e - 10h). The results of Models 4C and 5C, which are 711 not shown here, also show the same prediction patterns with respect to non-Gaussian residuals, 712 heteroscedasticity, and autocorrelation. 713

Finally, we observe in Figure 10 that the data models that have good overall predictive performance as measured by RMS during the calibration period will maintain this good predictive performance during the validation period. For model 6C, RMS values for the calibration and validation periods are very well correlated with a correlation coefficient of 0.92. However, we note that for models 4C and 5C the overall predictive performances during the calibration and validation periods are not that well correlated as 6C, with correlation coefficients of 0.52 for model 4C and
0.61 for model 5C. This suggests that model 6C is more robust than 4C and 5C for forecasting andhindcasting.

722 4.3 Discussion on handling residual correlation

Accounting for autocorrelation can lead to biased parameter estimation (Figure 5) and poor 723 predictive performance (Figure 10). Auto-correlated residuals may be attributed to model 724 725 discrepancy, as shown in Lu et al. (2013). The most obvious solution to handle the autocorrelation is to reduce the autocorrelation by improving the soil respiration model. If model improvement is 726 difficult for practical reasons, we can improve the data model to better characterize the 727 728 autocorrelation. Addressing autocorrelation in a data model is challenging since it involves several interlinked factors as follows: 729 (1) Non-stationarity due to wet-dry periods could be a reason for this problem. By drawing on 730 similarity from surface hydrology, the study of Ammann et al. (2018) suggests that auto-731 correlated residuals might be attributed to non-stationarity due to wet-dry periods with half-732 733 hourly data. Accounting for non-stationarity could better address the problem of autocorrelated residuals (Ammann et al., 2018; Smith et al., 2010b). 734 (2) The way of implementing autocorrelation could have an impact. Autocorrelation could be 735 736 applied to raw residuals directly (e.g., Li et al., 2015), to transformed residuals based on covariance matrix of residuals L(e) (e.g., Lu et al., 2013), or to normalized residuals L(a) (e.g., 737 Schoups and Vrugt, 2010; Evin et al., 2013). Note that e is a vector of transformed residuals, 738 739 while a denotes a vector of independent and identically distributed random errors with zero mean and unit standard deviation. The L(e) approach based on covariance matrix of residuals 740 is generally limited to Gaussian data models (e.g. Lu et al., 2013), while the L(a) approach for 741 742 normalized residuals can be readily adopted for non-Gaussian data models.

| 743 | (3) The autocorrelation model could have an impact. Using an autoregressive model is a popular |
|-----|------------------------------------------------------------------------------------------------------|
| 744 | technique to account for auto-correlated residuals. However, using an autoregressive model |
| 745 | with either joint inversion approach (e.g., this study and Schoups and Vrugt, 2010) or |
| 746 | sequential approaches (e.g., Evin et al., 2013, 2014; Lu et al., 2013) removes correlation errors |
| 747 | through a filter approach, which can lead to a loss of information content. As this may cause |
| 748 | overcorrection of prediction especially at surge events, Li et al. (2015) developed a restricted |
| 749 | autoregressive model to overcome this adverse effect. Other autocorrelation models include |
| 750 | moving average model and mixed autoregressive-moving averaging model (Chatfield, 2004). |
| 751 | (4) Joint versus sequential inversion for autocorrelation could have an impact. Sequential inversion |
| 752 | approaches include two-step procedures (e.g. Evin et al., 2013, 2014; Lu et al., 2013) or the |
| 753 | multi-step procedure (Li et al., 2016a). These sequential approach estimates the autoregressive |
| 754 | parameters sequentially in a later step after estimating the physical model parameters and other |
| 755 | data model parameters. Evin et al. (2013, 2014) used a sequential approach to avoid the |
| 756 | interaction between the parameters of the heteroscedasticity model and the autocorrelation |
| 757 | model. In addition, the autoregressive model parameters can be deterministically calculated as |
| 758 | an internal variables of the data model similar to Lu et al. (2013), and not as calibration |
| 759 | parameters (e.g. Schoups and Vrugt; Evin et al. 2013; 2014). While the first step in the |
| 760 | sequential approach would avoid the biased parameter estimation (Figure 10a-d), the second |
| 761 | step can still lead a poor predicative performance since we are essentially using a filter |
| 762 | approach to remove residual correlation. To address this problem, Li et al. (2016) multi-step |
| 763 | procedure that is based on Gaussian data model uses restricted autoregressive model. |
| 764 | Generally, Ammann et al. (2018) states that the joint inversion is still preferred, and |
| I | |

765 <u>understanding the conditions where accounting for auto-correlation can be achieved remain</u>
 766 poorly understood.

767 **5.** Conclusions

768 In parameter estimation and prediction of soil carbon fluxes to the atmosphere, weone often assumes that residuals, which include errors in observations, model inputs, parameter estimates, 769 and model structureserrors, are normally distributed, homoscedastic and uncorrelated. We 770 studiedy these assumptions by calibrating three microbial enzymesoil inspiration models, which 771 have varying degrees of model structure errors. We further explore tested eight data models that 772 characterize the residuals statistically by starting with the standard least squares (SLS) and skew 773 exponential power (SEP) data models that assume homoscedastic and non-correlated residuals. 774 Given For these two distributions, we evaluated six other data models that account for 775 776 heteroscedasticity (WLS and WSEP), autocorrelation (SLS-AC and SEP-AC), and joint inversion of heteroscedasticity and autocorrelation (WLS-AC and WSEP-AC). To our knowledge this is the 777 778 first study that provides such detailed analysis for soil reparation inverse modeling. We also used 779 three solid respiration models with different degrees of model fidelity (i.e., model realismdiscrepancy) and model complexity (i.e. number of model parameters), to understand the 780 impact of model discrepancy on the calibration results under different data models. We analyzed 781 the <u>calibration</u> results with respect to $(\frac{1}{1})$ residual characterization, $(\frac{2i}{1})$ parameter estimation, 782 (3iii) predictive performance, and (4iv) impacts of model discrepancy. The main findings of this 783 study can be are summarized as follows: 784

(<u>1</u>i) With respect to residual characterization, residual analysis results suggest that the common assumption of not accounting for heteroscedasticity and <u>residual</u> autocorrelation of residuals
 (<u>i.e.in the data models</u> SLS and SEP) results in poor characterization of residuals. Explicit

accounting for heteroscedasticity <u>in(i.e.</u> WLS and WSEP) <u>can</u>-result<u>s</u> in <u>goodsignificantly</u>
<u>improved</u> characterization of the residuals, <u>and the improvement is larger than that obtained</u>
<u>by</u>, <u>and is followed by joint the inversion of accounting for both</u> heteroscedasticity and
autocorrelation (<u>i.e.in</u> WSL-AC and WSEP-AC). Accounting for autocorrelation only (<u>i.e. in</u>
SLS-AC and SEP-AC) <u>may notdoes not significantly</u> improve-<u>much</u> the characterization of
the residuals.

794 (2ii) With respect to parameter estimation, the impacts of the data models are evaluated by we focuseding -on carbon use efficiency (CUE), which is a central parameter in soil respiration 795 modeling. We found the Using SLS yields with relatively reasonable posterior parameter 796 distributions for CUE, yet very narrow posterior. The Ddata models consider autocorrelation 797 (i.e. SLS-AC, SEP-AC, WLS-AC and WSEP-AC that consider autocorrelation) tend to 798 799 generally yield CUE estimates that are physically non-unreasonable. We speculate that 800 filtering residual correlation can affect the mapping of the model physics (as implicitly 801 included in the residuals) into the likelihood parameter space, which might result in biased parameter estimates that are physically unreasonable. 802

(3iii) With respect to predictive performance, it is measured by four statistical criteria: we assessed 803 the central mean tendency, sharpness, coverage, and relative model scoreuncertainty bands and 804 805 the overall predictive performance for both the calibration and the cross-validation periods. Results show that accounting for autocorrelation (i.e. in SLS-AC, SEP-AC, WLS-AC, and 806 WSEP-AC) deteriorates the predicative performance, such that the predictive performance is 807 inferior to that of SLS in terms of the central mean tendency and overall predictive performance 808 (measured by the relative model score), especially during the cross-validation period. Results 809 810 also indicates that using thea SEP distribution can potentially improve the predictive

performance. The same is true for accounting for heteroscedasticity. Using SEP distribution
and accounting for heteroscedasticity (i.e. WSEP) can potentially improve the predictive
performance.

(4iv) With respect to the impact of model discrepancy, the high fidelity complex model (6C) gives
the best results with respect to parameter estimation and predictive performance. Model 6C
generally maintainsed its superior performance under different data models. This justifies the
complexity of model 6C relative to model 5C that has one less carbon pool. Model 4C that has
awith the lowest fidelity maintains its poor performance for different data models, because the
model model with has only four carbon pools and lacks the explicit representation of soil

820 moisture control, maintains its poor performance for different data models.

821 <u>Based on From</u> the empirical findings <u>above of this research</u>, we conclude the following:

(i1) Not accounting for heteroscedasticity and autocorrelation using a Gaussian or non-Gaussian
 data model might not necessarily result in biased parameter estimates or biased predictions
 with respect to central mean tendency, but will definitely underestimate uncertainty resulting
 in lower overall predictive performance.

- (2ii) Using a non-Gaussian residual errordata model can improve the parameter estimates ion, and
 theand predictive performance with respect to central mean tendency and uncertainty
 estimation quantification.
- 829 (<u>3</u>iii) Accounting for heteroscedasticity will definitely improves the uncertainty estimation with
 830 respect to reliability at the cost of having a wider predictive interval.

(4iv) This study confirms the other empirical findings and theoretical analysies (Evin et al., 2013;
2014; Li et al., 2015, Ammann et al. 2018) that separately accounting for autocorrelation or
jointly accounting for autocorrelation and heteroscedasticity can be problematic. While the

reasons remain poorly understood (Ammann et al., 2018), it might be attributed to non-834 stationarity due to wet-dry periods with half-hourly data (Ammann et al., 2018) or to the 835 method of handling autocorrelation (e.g., Schoups and Vrugt, 2010, Evin et al., 2013; 2014; 836 837 Lu et al., 2013; Li et al., 2015, 2016a; Ammann et al. 2018). Further investigation to address autocorrelation in soil respiration modeling is warranted in a future study. Accounting for non-838 stationarity (Smith et al., 2010b, Ammann et al. 2018) could address this problem. Relatively 839 poor performance with respect to autocorrelation can be also attributed to the implementation 840 scheme. The inference scheme such as joint inference as in this study, post-processing 841 inference approach for autocorrelation (Evin et al., 2013; 2014), residuals transformation 842 approach (e.g. Lu et al., 2013) or other strategies (Li et al., 2015, 2016a) could have an impact. 843 Yet Ammann et al., (2018) study states that the joint inversion is still preferred, and 844 understanding the conditions where accounting for auto-correlation can be achieved remain 845 poorly understood. Further investigation of this point is warranted in a future study. 846

The above conclusions are subject to several limitations. First, the conclusions are specific to 847 the soil respiration models developed and validated for semi-arid savannah. Performance 848 variations across different soil respiration models with different levels of complexities is possible. 849 Second, the conclusions are conditioned on the data that were obtained at the half-hour interval 850 over a one-year period. Different conclusions are possible if the data are thinned to daily or weekly 851 852 scales or data of longer observation periods are used. Third, ourthe study investigates effects of the residual assumptions of formal likelihood functions through direct conditioning of the residuals 853 model parameters, yet this can also be done through other approaches such as residuals 854 transformation (Thiemann et al., 2001), autorgressive bias model (Del Giudice et al., 2013), 855 856 approximate Bayesian computation (Sadegh and Vrugt, 2013), and data assimilation (Spaaks and

Bouten, 2013). Comparing different methods for accounting the residual assumptions are beyond
the scope of this work. Fourth, this study focuses on formal Bayesian computation using formal
likelihood functions, and comparison with other inference functions such as informal likelihood
functions or approximate Bayesian computation is warranted in a future study.

-Based on the aforesaid conclusions and limitations, we recommend to start calibrating soil 861 respiration models with simple SLS or SEP likelihood function. If the residuals characterization is 862 adequate (e.g., Scharnagl et al., 2011), then the underlying assumptions are met. Otherwise, 863 increase complexity of the data model until satisfactory results are obtained in terms of residuals 864 characterization, posterior parameter estimation, and predictive performance. This is similar to the 865 procedure given in Smith et al. (2015). Although the empirical findings of this study provide 866 general guidelines for data model selection of microbial for soil respiration modelings, more 867 comparative studies are needed to validate and refute the findings of this study. 868

869 Acronyms

| 870 | 4C | Four carbon pool model |
|-----|---------|-------------------------------------------------------------------|
| 871 | 5C | Five carbon pool model |
| 872 | 6C | Six carbon pool model |
| 873 | CUE | Microbial carbon use efficiency |
| 874 | DOC | Dissolved organic carbon |
| 875 | ENZ | Enzymes |
| 876 | MCMC | Markov chain Monte Carlo |
| 877 | MIC | Microbial biomass |
| 878 | NSME | Nash-Sutcliffe model efficiency |
| 879 | PDF | Probability density function |
| 880 | RMS | Relative model score |
| 881 | SEP | Skew exponential power distribution |
| 882 | SEP-AC | Skew exponential power distribution with autocorrelation |
| 883 | SLS | Standard least square |
| 884 | SLS-AC | Standard least square with autocorrelation |
| 885 | SOC | Soil organic carbon |
| 886 | WLS | Weighted least squared |
| 887 | WLS-AC | Weight least square with autocorrelation |
| 888 | WSEP | Weighted skew exponential power distribution |
| 889 | WSEP-AC | Weighted skew exponential power distribution with autocorrelation |
| | | |

891 Code and data availability

The data and codes and models used to produce this paper are available on contact of the corresponding author at mye@fsu.edu. We cannot publicly share the workflow because MT-DREAM_(ZS) code (Laloy and Vrugt, 2012), which is a main component in the workflow, is in the process of becoming a commercial code.

896 Author contributions

897 ASE developed and implemented the code for the eight data models for soil respiration modeling,

- and prepared the manuscript with contribution of all co-authors. MY developed the research idea
- and outline, and supervised the research implementation when ASE was a post-doc at Florida State
- 900 <u>University</u>. GN developed the soil respiration models. GAB collected and processed the eddy-
- 901 covariance data used for model calibration.
- 902 Competing interests
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- Figure 1. Diagram of model 6C representing the processes of (1) degradation of soil organic carbon 1267 (SOC) to dissolved organic carbon (DOC) through catalysis of enzymes (ENZ) produced by 1268 microbes (MIC), (2) MIC uptake of DOC, and (3) microbial (MIC) respiration to produce CO₂ 1269 1270 (CUE is the carbon use efficiency). SOC degradation and microbial uptake rates are controlled by water saturation (θ / θ_{a}) . The DOC and ENZ pools are split into two subpools, one for the wet zone 1271 and the other for the dry zone of the soil pore space. Microbial uptake of DOC occurs only in the 1272 wet zone, and the uptake rate is linearly related to θ/θ_s . Catalysis through ENZ in the wet zone is 1273 proportional to θ/θ_s , while that in the dry zone is proportional to $1 - \theta/\theta_s$. V_{max} (s⁻¹) is the maximum 1274 rate, and K_m is the half-saturation concentration. 1275
- 1276



Figure 2. Time series of soil moisture and efflux observations. The dashed line marks the divideof the dataset into calibration and validation periods.





Figure 3. Residual analysis of the best realization (among multiple MCMC realizations) for model 6C using data models (a-c) SLS and (d-f) WSEP-AC.



Figure 4. Residual quantile-quantile (Q-Q) plots of the best realization (among multiple MCMC
realizations) for the three soil respiration models and eight data models.



1291 Figure 5. Marginal posterior parameter density of carbon use efficiency (CUE) for the three soil

1292 respiration models and eight data models.



Figure 6. Observation data (blue dots) and mean prediction (green line) and 95% credible intervals (red line) of prediction ensembles for (a)-(f) the calibration period and (g)-(l) the validation period. The plots are for the three soil respiration models using data models SLS and WSEP-AC. *The prediction ensembles are generated to consider parametric uncertainty of the soil respiration models only*.





Figure 7. (a-b) Nash-Sutcliffe model efficiency (NSME), (c)-(d) sharpness, (e)-(f) predictive coverage, and (g)-(h) relative model score for measuring predictive performance of the three soil respiration models and the eight data models during the calibration and cross-validation periods. *The statistics are evaluated from the prediction ensembles generated to consider parametric uncertainty of the soil respiration models only.*



1310 Figure 8. Observation data (blue dots) and mean prediction (green line) and 95% credible intervals

1311 (red line) of prediction ensembles for (a)-(f) the calibration period and (g)-(l) the validation period.

1312 The plots are for the three soil respiration models using data models SLS and WSEP-AC. *The*

1313 prediction ensembles are generated to consider parametric uncertainty of not only the soil

1314 *respiration models but also the data models.*





Figure 9. (a-b) Nash-Sutcliffe model efficiency (NSME), (c)-(d) sharpness, (e)-(f) predictive coverage, and (g)-(h) relative model score for measuring predictive performance of the three soil respiration models and the eight data models during the calibration and cross-validation periods. *The statistics are evaluated from the prediction ensembles generated to consider parametric uncertainty of not only the soil respiration models but also the data models*.



Figure 10. Observation data (blue dots) and mean prediction (green line) and 95% credible intervals (red line) for 6C for the eight likelihood functions during the calibration period (a)-(h) and the validation period (i)-(p). *The prediction ensembles are generated to consider parametric uncertainty of not only the soil respiration models but also the data models*.



| Residual | Likelihood function | Data model | Residuals | Variance | Likelihood functionData |
|------------------------------------------------------------|-----------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------|--------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------|
| Assumptions | | | | | model parameters |
| | <u>Generic data model</u> | Generic data model | \mathcal{E}_t | σ_{t} | |
| | | $a_{t} = \frac{\varepsilon_{t}}{\sigma_{t}} \qquad a_{t} \sim X$ $a_{t} = \frac{\varepsilon_{t}}{\sigma_{t}} \qquad a_{t} \sim N(0, 1)$ | | | |
| Independent, normally distributed, and homoscedastic | Standard least square (SLS) | $a_t = \frac{\varepsilon_t}{\sigma_0}$ $a_t \sim N(0,1)$ | $\varepsilon_t = d_t - Y_t$ | $\sigma_t = \sigma_0$ | Constant $\sigma_{_0}$ |
| Independent, and homoscedastic | Skew exponential power (SEP) | $a_t = \frac{\varepsilon_t}{\sigma_0}$ $a_t \sim SEP(0, 1, \xi, \beta)$ | $\varepsilon_t = d_t - Y_t$ | $\sigma_t = \sigma_0$ | Constant $\sigma_{_0}$ Skewness ξ , Kurtosis eta |
| Independent and normally distributed | Weighted least square (WLS) | $a_t = \frac{\varepsilon_t}{\sigma + \sigma Y}$ $a_t \sim N(0, 1)$ | | $\sigma_t = \sigma_0 + \sigma_1 Y_t$ | Heteroscedasticity model parameters $\sigma_{\scriptscriptstyle 0}$, σ |
| Independent | Weighted skew exponential power (WSEP) | $a_t = \frac{\varepsilon_t}{\sigma_0 + \sigma_1 Y_t} \qquad a_t \sim SEP(0, 1, \xi, \beta)$ | $\varepsilon_t = d_t - Y_t$ | $\sigma_t = \sigma_0 + \sigma_1 Y_t$ | Heteroscedasticity model parameters $\sigma_{_0}$, σ Skewness ξ , Kurtosis eta |
| Normally distributed, and homoscedastic | Standard least square with auto- correlation (SLS-AC) | $a_{t} = \frac{\varepsilon_{t} - \sum_{i=1}^{p} \phi_{i} \varepsilon_{t-i}}{\sigma_{0}} \qquad a_{t} \sim N(0, 1)$ | $\varepsilon_t - \sum_{i=1}^p \phi_i \varepsilon_{t-i}$ | $\sigma_t = \sigma_0$ | Constant $\sigma_{\scriptscriptstyle 0}$, Autoregressive model parameters $\phi_{\scriptscriptstyle i}$ |
| Homoscedastic | Skew exponential power with auto- correlation (SEP-AC) | $a_{t} = \frac{\varepsilon_{t} - \sum_{i=1}^{p} \phi_{i} \varepsilon_{t-1}}{\sigma_{0}} \qquad a_{t} \sim SEP(0, 1, \xi, \beta)$ | $\varepsilon_t - \sum_{i=1}^p \phi_i \varepsilon_{t-i}$ | $\sigma_t = \sigma_0$ | Constant σ_0 , Autoregressive model parameters ϕ_i Skewness ξ ,Kurtosis eta |
| Normally distributed | Weighted least square with auto- correlation (WLS- AC) | $a_{t} = \frac{\varepsilon_{t} - \sum_{i=1}^{p} \phi_{i} \varepsilon_{t-1}}{\sigma_{0} + \sigma_{1} Y_{t}} \qquad a_{t} \sim N(0, 1)$ | 1=1 | $\sigma_t = \sigma_0 + \sigma_1 Y_t$ | Heteroscedasticity model parameters σ_0 , σ_0 Autoregressive model parameters ϕ_i |
| | Generalized likelihood function (WSEP-AC) | $a_{t} = \frac{\varepsilon_{t} - \sum_{i=1}^{p} \phi_{i} \varepsilon_{t-1}}{\sigma_{0} + \sigma_{1} Y_{t}} \qquad a_{t} \sim SEP(0, 1, \xi, \beta)$ | $\varepsilon_t - \sum_{i=1}^p \phi_i \varepsilon_{t-i}$ | $\sigma_t = \sigma_0 + \sigma_1 Y_t$ | Heteroscedasticity model parameters σ_0 , c Autoregressive model parameters ϕ_i , Skewness ξ , Kurtosis β |

Supplementary Figure 1. Workflow scheme

