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A model-data fusion approach to analyse carbon dynamics in managed grasslands

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Abstract

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Grasslands are an important component of the global carbon (C) cycle, 16 with a strong potential for C sequestration. However, an improved capacity to quantify grassland C stocks and monitor their variation in space and time, particularly in response to management, is needed in order to conserve and enhance grassland C reservoirs. To meet this challenge we outline and test here an approach to combine C cycle modelling with observational data. We implemented an intermediate complexity model, DALEC-Grass, within a probabilistic model-data fusion (MDF) framework, CARDAMOM, at two managed grassland sites (Easter Bush and Crichton) in the UK. We used 3 years (Easter Bush, 2002-2004) of management data and observations of leaf area index (LAI) and Net Ecosystem Exchange (NEE) from eddy covariance to calibrate the distributions of model parameters. Using these refined distri-27 butions, we then assimilated the remaining 7 years (Easter Bush, 2005-2010 and Crichton, 2015) of LAI observations and evaluated the simulated NEE, above and below-ground biomass and other C fluxes against independent data from the two grasslands. Our results show that fusing model predictions with LAI observations allowed the CARDAMOM MDF system to diagnose the effects of grazing and cutting realistically. The overlap of MDF-predicted

- and measured NEE (both sites) and ecosystem respiration (Easter Bush) was
- 92% and 83% respectively while the correlation coefficient (r) was 0.79 for
- 36 both variables. This study lays the foundation for using MDF with satellite
- 37 data on LAI to produce the spatially and temporally-resolved estimates of
- ³⁸ C cycling needed in shaping and monitoring the implementation of relevant
- 39 policies and farm-management decisions.
- 40 Keywords: UK grasslands, primary production, carbon sequestration,
- 41 model-data fusion

Ecosystem carbon accounting abbreviations

Gross Primary Production: GPP
 Autotrophic Respiration: AR
 Heterotrophic Respiration: HR
 Ecosystem Respiration: ER = AR + HR

Net Primary Production: NPP = GPP - AR
Net Ecosystem Exchange: NEE = ER - GPP
Net Ecosystem Production: NEP = GPP - ER

50 1. Introduction

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Grasslands cover a third of the earth's surface and are a major compo-51 nent of the terrestrial biosphere's carbon (C) cycle and a major contributor to global annual fluxes and C stores (Hungate et al., 2017; Friedlingstein et al., 2019; Sollenberger et al., 2019). Temperate grasslands, because of edaphoclimatic conditions and their botanical composition, can transfer and accumulate C in their soils more efficiently than grasslands in warmer and drier regions (Gibson, 2010). Based on this premise, and considering the rise in atmospheric CO₂ concentration, European grasslands have the potential for increased CO₂ sequestration (Chang et al., 2017). Estimates of grassland C balance and its variation in space and time are essential for shaping evidenceinformed climate policies and monitoring progress on Nationally Determined Contributions (NDCs) following the Paris agreement (De Oliveira Silva et al., 2018). Livestock grazing and grass harvesting affect grassland C stocks, typically removing >50\% of vegetation C on an annual basis (Erb et al., 2018). In addition to sustaining livestock farming by providing biomass energy to livestock directly from grazing or as fodder, vegetation also provides inputs to the soil C pool in the form of litter and exudates, as well as indirectly from excrement produced by grazers (Soussana and Lemaire, 2014; Chen et al., 2015; Conant et al., 2017; Abdalla et al., 2018). Because of its dynamic nature, the C balance of vegetation in managed grasslands (i.e. assimilation, allocation, removal and loss) is complex and challenging to monitor and assess.

Detailed, continuous measurements of the C dynamics of grasslands are limited to a few sites globally. Computational approaches are therefore used to extrapolate observed relationships across landscapes. Such computational methods include (1) statistical models that relate climate data and selected

national statistics to grassland productivity and removals (e.g. Smit et al., 2008; Herrero et al., 2013; Qi et al., 2017, 2018); (2) process-based grassland models that simulate C uptake and turnover (e.g. Vuichard et al., 2007; Chang et al., 2013; Snow et al., 2014; Chang et al., 2015; Kipling et al., 2016; Rolinski et al., 2018; Puche et al., 2019; Sándor et al., 2020; van Oijen et al., 81 2020); and (3) processed earth observation (EO) data that map and track key ecological variables, such as leaf area index (e.g. Franke et al., 2012; Dusseux et al., 2014; Asam et al., 2015; Xu and Guo, 2015; Ali et al., 2016; Gómez Giménez et al., 2017; Punalekar et al., 2018). Each method has its strengths and weaknesses. Statistical approaches are strongly grounded on measured data but have low sensitivity to the spatial and temporal variation 87 of system drivers (e.g. climate, management) and have limited explanatory depth (Smit et al., 2008). Process models describe most of the underlying biogeochemical processes, which gives them greater explanatory depth than purely statistical approaches and the capacity to explore the consequences of different management and soil-climate conditions. But process models require observational data for parameter calibration and output error evaluation (Ma et al., 2015; Ehrhardt et al., 2017). Model-based studies tend to present deterministic results, ignoring the role of uncertainties around model inputs, parameters and structure, and observed data (Smith et al., 2012; Kipling et al., 2016). Earth observations increasingly provide snapshots at high temporal and spatial resolution on certain drivers and proxies of C dynamics (e.g. vegetation structure, soil moisture). But these products do not consider the full C budget, particularly below ground, nor diagnose how 100 grassland ecosystems C storage evolves (Ali et al., 2016). 101

Model-data fusion (MDF) is a hybrid approach that combines aspects of the three aforementioned computational approaches (Raupach et al., 2005). MDF uses probabilistic methods to calibrate model parameters and/or to quantify model predictive uncertainty (Gottschalk et al., 2007; Patenaude et al., 2008; Ben Touhami and Bellocchi, 2015; Oenema et al., 2015; van Oijen, 2017). From an ecosystem modelling perspective, MDF can be understood as a framework in which model parameter distributions are calibrated according to a set of observations (observed data assimilation) and model output uncertainty is quantified. The behaviour of simulated fluxes and pools is constrained according to certain rules. For instance, ancillary data from national statistics, land surveys and scientific literature can be integrated in a MDF framework. The strong linkage to observations means that MDF is suitable for quantifying the existing situation and for explaining the

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mechanisms that underlie the functioning of a grassland. Model-data fusion can accommodate models of varying complexity (e.g number of parameters, modules, calculation nodes) but increasing complexity increases the computational cost and can reduce the robustness of the MDF process. The MDF approach has been used in studies focusing on various aspects of terrestrial 119 ecosystem C dynamics (e.g. productivity, biomass, fire emissions) in the past 120 (Wang et al., 2009; Fox et al., 2009; Keenan et al., 2012; Kuppel et al., 2014; Xiao et al., 2014; Kuppel et al., 2014; Bloom and Williams, 2015; Peylin 122 et al., 2016; Smallman et al., 2017; Scholze et al., 2017; Peaucelle et al., 123 2019). Model-data fusion is actively benefiting from the increasing quality 124 and range of EO data and can be used to monitor terrestrial ecosystem C 125 balance at various spatial and temporal scales (Guo et al., 2014; Bloom et al., 126 2016; Ramapriyan and Murphy, 2017; Chen and Wang, 2018).

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Here, for the first time to our knowledge, we apply MDF to analyse ecosystem C cycling in managed grasslands. Previous probabilistic modelbased studies have focused on plant functional type identification and have not considered the role of management on C cycling (Kuppel et al., 2014; Peylin et al., 2016; Peaucelle et al., 2019). In this study, we present a development of the Data Assimilation Linked Ecosystem Carbon model (DALEC) that is tailored for use in MDF for grasslands (DALEC-Grass). DALEC is a C-budget model that is integrated into the Carbon Data Model Framework (CARDAMOM) (Bloom and Williams, 2015; Bloom et al., 2016; Smallman et al., 2017). DALEC and the CARDAMOM MDF framework have been used before in MDF studies on forests and croplands (Revill et al., 2016; Smallman et al., 2017). The aim of the present study is to demonstrate MDF with DALEC-Grass and test its ability to quantify C dynamics in grasslands under variable grazing and cutting regimes. As a first step, we calibrate the distribution of DALEC-Grass parameters using 3 years of measured data on leaf area index (LAI) and net ecosystem exchange (NEE) from a grassland in eastern Scotland (UK). LAI data are routinely estimated from EO systems at fine spatial (<ha) and temporal resolutions (\approx days). Therefore, time series of satellite LAI data have the potential to inform and constrain grassland models effectively at sub-field scales and during critical growth changes and management interventions. As a second step, we tested this assumption by evaluating our model's predictive skill when a limited number of field-measured LAI data are assimilated through the CARDAMOM MDF framework. We assess the model's performance by comparing its outputs to independent eddy flux data on NEE of CO₂, above and below-ground

biomass and soil respiration from chambers. Six additional years of data from the core study site and one year of data from another UK site are used for the validation process. The materials and methods section describes the DALEC-Grass model, the CARDAMOM framework, the characteristics of the grassland sites that are modelled and the methodology that is followed. Finally, we discuss the potential for DALEC-Grass and CARDAMOM to produce landscape analyses of grassland C cycling under varied management systems using earth observation.

¹⁶¹ 2. Materials and methods

2.1. DALEC-Grass

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DALEC-Grass is a development of the DALEC model in which a number of processes related to grass growing, cutting and grazing have been introduced (Smallman et al., 2017). The model is written in fortran and its code is available online at https://github.com/GCEL/DALEC-Grass. DALEC-Grass is a parsimonious terrestrial ecosystem C cycling model of intermediate complexity which tracks the dynamics of three plant C pools and two dead organic matter pools. DALEC-Grass does not resolve explicitly mixtures of grasses and biodiversity, and water and nitrogen (N) cycling are not described. The model is driven by temperature, short-wave radiation, vapour pressure deficit and CO_2 concentration (Table 1). Carbon enters the ecosystem via gross primary production (GPP) which is partitioned into autotrophic respiration or allocated to various plant pools (Fig. 1). The model simulates the turnover of plant C pools to litter and soil organic matter based on both mortality and grazing/cutting. The mineralisation (i.e. heterotrophic respiration) and decomposition of litter and soil organic C pools are temperature dependent first order processes.

DALEC-Grass has 25 parameters impacting photosynthesis, litter production and decomposition, fractional allocation of C, and climate sensitivity of phenology and decomposition. A further 8 parameters relate to initial conditions of C pools and to management impacts of grazing and cutting (see Table 5 in supplementary material). The calculations in DALEC-Grass are performed on a daily basis by default. A component of DALEC-Grass is the Aggregated Canopy Model (ACMv1); a photosynthesis model that emulates a detailed mechanistic model, and that uses daily meteorological data to estimate GPP (Williams et al., 1997). The duration and intensity of the grass growing period is calculated following the growing season index (GSI)

approach (Jolly et al., 2005). The GSI method uses information on vapour pressure deficit (VPD), daylength and daily minimum temperature to adjust the plant's physiological progress (Smallman et al., 2017).

The net primary productivity (NPP) C, which remains after accounting for C losses via autotrophic respiration, is allocated to root, stem and leaf tissues. In DALEC-Grass, the above- to below-ground C allocation balance is dynamic and is calculated on a daily basis using the architecture-dependent strategy presented in Reyes et al. (2017). According to this approach, the C that is transferred to the fine root C pool is linked to above-ground biomass and increases after the plant has grown a sufficient quantity of leaves. This linkage is achieved by the following equation:

$$F_{root_t} = 1 - exp(-1 * P4 * LAI_t) \tag{1}$$

where F_{root_t} is the fraction of NPP C that goes to the root C pool on day t, LAI_t is the LAI of the sward on day t and P4 is a model parameter. The remaining NPP C is allocated to above-ground biomass. Its partitioning to the stem and leaves C pools is based on the idea that increasing stem mass is needed to support increasing leaf mass but the stem to leaf ratio is dynamic and not constant. The C allocation to leaves and stems is calculated using parameter P29 within the following equations:

$$F_{leaf_t} = NPP_t * (1 - (P29 * (LAI_t/LAI_{max})))$$
 (2)

$$F_{stem_t} = NPP_t * (P29 * (LAI_t/LAI_{max}))$$
(3)

where F_{leaf_t} is the fraction of NPP allocated to the leaf C pool on day t, F_{stem_t} is the fraction of NPP allocated to the stem C pool on day t, LAI_t is the LAI of the sward on day t and LAI_{max} (set to 6 m² m⁻²) is a maximum LAI for managed grasslands.

Animal grazing and grass cutting is imposed as a time series forcing. The number of livestock units (LSU) per ha per day determines the animal grazing intensity. The amount of C that one LSU removes from the grassland via grazing is estimated by multiplying the LSU value by a "dry matter demand per weight of 1 LSU" parameter (P31); with the standard weight of one LSU being equal to 650kg. The resulting dry matter (DM) value (in kgDMha⁻¹) is converted to gCm⁻² and removed from the C pool of the foliage. DALEC-Grass has a set of internal mechanisms through which it

can accept/perform or reject/skip a grazing instance. These mechanisms reflect the logical assumption that there is a minimum amount of above-ground biomass that has to remain after grazing for grass to be able to grow the following days; i.e. grazing is not simulated when the simulated above-ground biomass is below a threshold. This minimum biomass threshold is a model parameter (P27) and a similar parameter and concept is applied for cutting (P28); i.e. cutting cannot take place when the simulated above-ground biomass is below a threshold. These mechanisms exist to ensure there are no unrealistic combinations of livestock density and simulated grass biomass.

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DALEC-Grass uses a simple scheme to convert the amount of C in grass into (1) C in animal-respired CO₂; (2) C in methane (CH₄) produced via digestion; and (3) C in animal excrement. Of the total amount of C ($gCm^{-2}d^{-1}$) that is grazed: (1) 54% is lost to the atmosphere as CO_2 ; (2) 4% is lost to the atmosphere as CH₄; (3) 32% returns to the soil as C in excrement; and (4) the remaining 10% stays in the animal's body. In reality the conversion factors of grazed C are dynamic, they depend on animal type, weight and age and vary even between animals that have the same aforementioned characteristics (Vertès et al., 2018; Snow et al., 2014). The grazed C conversion factors used in DALEC-Grass are generic in order to reflect different estimates for beef/dairy cattle and sheep and were extracted from the relevant literature (Bell et al., 2016; Lee et al., 2017; Worrall and Clay, 2012; Parsons et al., 2009). In terms of modelling soil C dynamics, DALEC-Grass uses a simple soil C scheme, in which plant residue and excrement-contained C go into a single litter pool. Litter C undergoes temperature-dependent decomposition with part of the C lost as heterotrophic respiration while the remainder is moved into a single slowly-decomposing soil C pool that represents the soil's organic matter.

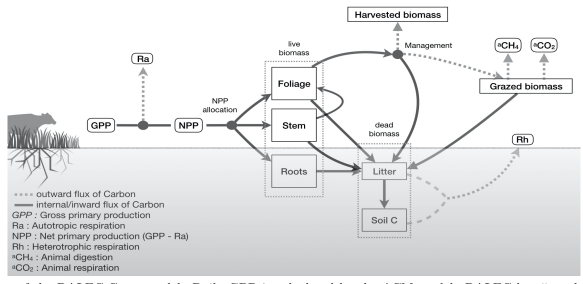


Figure 1: Diagram of the DALEC-Grass model. Daily GPP is calculated by the ACM model. DALEC has 5 pools: leaves, stem, roots, litter and soil organic matter.

Table 1: DALEC meteorological and management inputs

Variable	${ m units}$
Minimum daily temperature	$^{o}\mathrm{C}$
Maximum daily temperature	$^{o}\mathrm{C}$
Short-wave radiation	${ m MJm^{-2}day}$
Atmospheric CO ₂ concentration	ppm
21-day average minimum temperature	\mathbf{C}
21-day average photoperiod	\sec
21-day average vapour pressure deficit	Pa
Animal density	livestock units per ha

2.2. Carbon Data Model Framework

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DALEC-Grass is integrated into the Carbon Data Model Framework (CARDAMOM) (Bloom et al., 2016). CARDAMOM is a MDF framework that uses Bayesian inference to approximate the joint distribution of model parameters. This approximation is done using a function of the likelihood of each sampled parameter vector. The Bayes' theorem forms the basis Bayesian inference:

$$P(\theta|D) = \frac{P(D|\theta)P(\theta)}{P(D)} \tag{4}$$

where θ represents the parameters, D the observed data, $P(\theta|D)$ the posterior distribution of parameters, $P(D|\theta)$ the likelihood of θ given D, $P(\theta)$ the prior distribution of parameters and P(D) the marginal distribution of D 250 (normalisation constant). The effectiveness of Bayesian inference stems from the fact that the posterior distribution of parameters is proportional to the likelihood:

$$P(\theta|D) \propto P(D|\theta)P(\theta)$$
 (5)

Markov Chain Monte Carlo (MCMC) is a group of algorithms that are used to sample from probability distributions (Chib and Greenberg, 1995). For a discussion on MCMC theory we refer to van Ravenzwaaij et al. 2018 and for a description of different MCMC algorithms we refer to Houska et al. 2015. In our implementation of CARDAMOM, the Metropolis-Hastings (MH) MCMC algorithm is used. Metropolis-Hastings creates a Markov chain

by deciding whether each sampled θ is accepted or rejected after comparing its likelihood to that of the last accepted θ . This comparison is done using the acceptance ratio (A):

$$A = \frac{f(\theta')}{f(\theta)} \tag{6}$$

where θ' is the sampled θ under examination and f is a function proportional to $P(\theta|D)$. For this study, MH used a metric that describes the model's predictive skill against observed variables as a surrogate likelihood. This metric is named accuracy and is described in Myrgiotis et al. (2016). Accuracy quantifies the number of simulated data points that fall within the respective measured range (i.e. standard deviation assuming normal distribution for D) while it also considers possible time lags between measured and simulated time series. Accuracy can take any value between 0 (no simulated points within the observed range) and 1 (all simulated points within the observed range). The consideration of time lags when calculating the metric allows CARDAMOM to capture some of the impacts that possible temporal uncertainties in model inputs can have on model outputs (Myrgiotis et al., 2018). Similarly, delayed responses of the grassland's physiology and/or biogeochemistry to driving variables, which could appear due to model formulation uncertainty and/or parametric uncertainty, can also be captured. Moreover, the model's internal mechanisms can lead to instances when e.g. a day's grazing, even though it is specified in the inputs, is not modelled because there is not sufficient simulated grass biomass on that day. This can lead to time lags in LAI fluctuation and affect the level of fit between modelled and measured data.

For each assimilated variable (LAI and NEE) we provide an estimate of uncertainty around the measured data points. For LAI the uncertainty is set equal to $\pm 15\%$ of the mean measured value (Van Wijk and Williams, 2005). Attributing uncertainty levels around measured NEE data is more complicated because most measured datasets depend on a single flux tower and do not provide uncertainty estimates. In this study the uncertainty around the measured NEE data is set equal to $\pm 1~\rm gCm^{-2}$ (Hill et al., 2012; Revill et al., 2016). The overall setup of the implementation of MH in CARDAMOM is the following:

• 10 chains are run in parallel

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• Each chain has 100 million steps

- The initial 10% of all accepted steps is used as burn-in
- At each step of the Markov chain the MH algorithm:
 - 1. obtains a sampled parameter vector
 - 2. calculates the objective function (i.e. accuracy metric)
 - 3. calculates A and :
 - If A > 1 then θ' is accepted
 - If A > a uniform random number [0.3 1] then θ' is accepted
 - If A < a uniform random number [0.3 1] then θ' is rejected

A list of ecological and dynamic constraints (EDCs) is used in CAR-DAMOM to refine the parameter space that the MH sampling explores (Bloom et al., 2016). EDCs are checks of the mathematical, ecological and biogeochemical sanity of the sampled model parameter combinations and model outputs (pools, fluxes). These checks are performed in CARDAMOM before and/or after each run of DALEC-Grass, which is performed to estimate the likelihood of each sampled parameter vector in MH. Altogether, EDCs reflect existing knowledge on grassland ecosystem functioning. Table 2 outlines the EDCs that were used with CARDAMOM in this study. Ecological and dynamic constrains are a key feature of the CARDAMOM MDF framework. Retrieving posterior parameter distributions that are mathematically and theoretically sound depends on the use of appropriate EDCs. In this regard, the present study is a test of CARDAMOM's grassland-specific EDCs.

The assessment of if and when a MCMC algorithm has converged to the stationary distribution of parameters is an essential part of its implementation. The difficulty of convergence assessment increases with the number of parameters and no single convergence diagnostic is generally accepted as being suitable for every application (Brooks and Gelman, 1998). When multiple chains are explored, such as in CARDAMOM, convergence diagnostics based on the comparison of inter and intra-chain variances are appropriate. The Gelman-Rubin (GR, see supplementary material) is one of the most widely used convergence diagnostics of this type and was used to assess chain convergence in this study (Gelman and Rubin, 1992). The equations used for calculating the potential scale reduction factor (PSRF) of the GR method

and the results of chain convergence assessment are presented in the supplementary material.

2.3. Field measured data

Measured data from two managed grassland sites, of contrasting soil and climatic conditions, in eastern (Easter Bush) and southern (Crichton) Scotland are used in this study.

333 2.3.1. Easter Bush

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Easter Bush is located in South East Scotland, 10 km south of Edin-334 burgh (03°02'W, 55°52'N, 190 m above sea level). The mean annual rainfall 335 between 2002 and 2010 was 947 ± 234 mm and the mean annual temperature 336 was 9.0 ± 0.4 °C. The field has been under permanent grassland management 337 for more than 20 years with a species composition of >99\% perennial rye-338 grass (Lolium perenne) and < 0.5% clover (Trifolium repens). The soil type 339 is an imperfectly drained Eutric Cambisol (FAO classification) with a pH of 340 5.1 (in H₂O), a clay fraction of 20-26% (Clayey Loam to Sandy Loam) and 341 a soil organic carbon content of 4% (0-10 cm depth). The grassland was grazed continuously by heifers in calf, ewes and lambs at different stocking 343 densities. The grass was cut for silage in June and August 2002 and in May 2003. Ammonium nitrate fertiliser was applied to the field 3-4 times per 345 year, usually between March and July at an average of 56 kg N ha⁻¹ per application. An additional fifth mineral N application was applied as urea in 347 2008 and organic manure was applied in September 2004 and March 2005 as cattle slurry. Vegetation for above ground biomass and LAI measurements 349 were collected from 4 to 6 quadrats (0.0625 m²) per sampling occasion. The leaf area was analysed using a Li3100 Area meter (LI-COR inc. Lincoln, 351 Nebraska, USA). Fresh weight of biomass samples were recorded before sam-352 ples were dried at 80°C for 24 hours and dry weight was measured. NEE 353 was measured by an eddy covariance system consisting of a fast response 354 3D ultrasonic anemometer (Metek USA-1, Metek GmbH, Elsmhorn, Ger-355 many) and a fast closed path CO₂-H2O analyser (LI-COR 7000 infra-red gas 356 analyzer, LI-COR, Lincoln, NE, USA). Quality control of the eddy covari-357 ance data followed the procedure proposed by Foken and Wichura (1996). 358 Missing NEE data were gap-filled using the online tool developed by Reichstein et al. (2005). Soil respiration rates were measured weekly (297 times 360 between 2003 and 2010) at 4 locations using a closed dynamic chamber (volume 1334 cm³, cover area 78.5 cm², PP-Systems, Hitchin, UK), which was

Table 2: Ecological and Dynamic Constraints

Index Description					
	1	Fluxes cannot be negative			
	2	Pools cannot be negative			
	3	GSI-related minimum parameters cannot be larger than maximum parameters			
	4	Turnover rate of soil organic matter cannot be larger than that of litter			
	5	Initial SOM pool cannot be smaller than the sum of all other pools			
 	6	Annual GPP cannot be more than 2000 g C m ⁻² (Xia et al., 2017; Gilmanov et al., 2007)			
	7	Annual GPP cannot be less than 500 g C m ⁻² (Xia et al., 2017; Gilmanov et al., 2007)			
	8	Daily GPP cannot be more than 20 g C m ⁻² (Xia et al., 2017; Gilmanov et al., 2007)			
	9	Annual ecosystem respiration cannot be more than 2000 g C m ⁻² (Xia et al., 2017; Gilmanov et al., 2007)			
	10	Annual ecosystem respiration cannot be less than 500 g C m ⁻² (Xia et al., 2017; Gilmanov et al., 2007)			
	11	Daily ecosystem respiration cannot be more than 15 g C m ⁻² (Xia et al., 2017; Gilmanov et al., 2007)			
	12	LAI cannot exceed 6 m ² m ⁻²			
	13	Minimum daily estimated root to shoot ratio cannot be less than 1 (Mokany et al., 2006)			
	14	Daily cut grass biomass cannot be more than 300 g C m ⁻² or less than 50 g C m ⁻² (Qi et al., 2017)			

placed onto soil and vegetation. The CO_2 increase within the chamber was monitored over 30-180 s by a portable sensitive infrared gas analyser (EGM 2, PP-Systems). It should be noted that the area covered for the soil respiration measurements also included vegetation and, therefore, measurements are effectively equivalent to ecosystem respiration (the sum of autotrophic and heterotrophic respiration). The data were converted from μ mol CO_2 m⁻¹ s⁻¹ to g CO_2 -C m⁻² d⁻¹ using the daily minimum and maximum temperatures and a Q10 equal to 2 (Meyer et al., 2018; Barba et al., 2018).

2.3.2. Crichton

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The field experiment was located at Crichton Royal Farm, Dumfries (55° 2'3"N, 35° 35'1" W) in South-West Scotland, on a long-term permanent grassland site (6.53 ha) used for intensive dairy production (Bell et al., 2016). The landscape was open grassland dominated (proportion of total harvested biomass > 99%) by perennial ryegrass (Lolium perenne) with white clover (Trifolium repens), creeping buttercup (Ranunculus repens) and chickweed (Stellaria media) being minor sward constituents. The Crichton site is representative of a wet climate zone, with a 30 year (1971-2000) long-term average rainfall of 1140 mm, and mean annual temperature of 9.3 °C. The soil was a Eutric Cambisol (FAO classification) and had a free-draining sandy to sandyloam light texture. The soil organic C concentration (0-10 cm) was 5.25\% (4.3-6.2%), and the pH varied between 5-6.3 at this site. The long term management of the site involved a rotation between cutting (with three cuts per year) and summer grazing. Between March and July 2015, 226 kg N ha⁻¹ were applied as slurry (4 applications) and synthetic fertiliser (2 applications). A sampling grid (20m * 20m) was marked out in the field at the beginning of the measurement campaign in June 2015. Leaf area index (LAI), aboveground biomass and respiration at soil surface (Rs) were made on four occasions, and root biomass on two occasions during June and July 2015. LAI was measured using a LAI-2200C Plant Canopy Analyzer (Licor Biosciences, Lincoln NE) at each point of the sampling grid. Aboveground biomass was measured using a rising plate meter calibrated against destructive biomass sampling. Root biomass was destructively sampled by taking replicate 2 cm diameter soil cores and dividing into 0-10 and 10-20 cm depths from positions adjacent to the collars used for Rs measurements. Soil respiration (Rs) was measured, at midday, on four dates, using a potable PP Systems Infra-red EGM4 Gas Analyser linked to a SRC-1 soil respiration chamber. The chamber (10 cm of diameter and 15 cm height) was equipped

with a fan, and was inserted into bare soil with a basal cutting ring to a depth of 2 cm during measurements. The air from the chamber was send to the analyser at flow rate of 0.2 l min⁻¹. After the chamber equilibrated the CO₂ concentration was measured every 5 seconds and the flux was calculated from the concentration increase over approximately 60 seconds time using a linear regression. Net ecosystem exchange of CO₂ was measured using an eddy covariance tower (EC) sited within the field (11 m height), with a Gill R3 sonic anemometer (Gill Instruments, Lymington UK) and a Licor LI700 CO₂ analyser (Licor Biosciences, Lincoln NE). Velocity measurements were rotated to minimize the mean vertical velocity. A site specific cospectral model was developed, based on sensible heat fluxes. Similarly, sensor specific models of sensor frequency response attenuation were developed and combined with the cospectral models to determine, and then apply, frequency response corrections. The resulting fluxes were screened for plausibility instrument diagnostics and for individual deviations from the group mean by more than two standard deviations.

2.4. Methodology

The volume of field-measured data and the range of measured variables at Easter Bush and Crichton allows us to test DALEC-Grass and CARDAMOM in detail. In designing the methodology of the study we considered two main aspects: (1) the efficient use of the available field-measured data and (2) the ability to relate our computational experiments with the envisioned application of DALEC-Grass in MDF studies. For these reasons, the use of LAI observations has a particular importance. LAI is a physiology-related variable, for which data can be collected rather easily and frequently at the different spatial scales that DALEC-Grass can be applied i.e. farm, land-scape, region. This contrasts with what is the case for the other measured variables examined in this study with the possible exception of aboveground biomass for which satellite data are increasingly available. Because of the lack of accurate satellite-based LAI data for Easter Bush during the simulated period, field-measured LAI data were used in this study.

Initially, all DALEC-Grass parameters have a uniform distribution i.e. only a realistic minimum and maximum value is known for each of them. In order to refine these uniform distributions we drive DALEC-Grass with 3 years of climate and management data for the Easter Bush site while assimilating in-situ NEE and LAI observations (step 1). Through this parameter

calibration step we expect parameter distributions to become more representative of managed (cut and grazed) grasslands. It should be clarified that the term calibration is used to refer to the refinement of the prior distribution of 438 parameters and not the parameters themselves. During calibration, the calculated accuracy metric is the mean of the accuracy for LAI and the accuracy 440 for NEE. In order to test the MDF framework we, then, run DALEC-Grass 441 (step 2) for 6 additional years at Easter Bush, this time assimilating only 442 the available LAI observations. We assess the model's performance at Easter 443 Bush by comparing model outputs with independent in-situ data on NEE (flux tower based) and ecosystem respiration (ER, chamber based). More-445 over, we run DALEC-Grass with one year of climate and management data 446 from Crichton while assimilating the available field-measured LAI data (step 3). The model's performance at Crichton is assessed by comparing model outputs with independent in-situ data on NEE, above and below-ground biomass 440 and soil respiration. The four steps of our computational experiment are:

1. Calibration of DALEC-Grass parameters: Implementation of CAR-DAMOM at Easter Bush by assimilating 3 years (2002-2004) of LAI and NEE observations.

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- 2. MDF at Easter Bush: Using the calibrated distributions (step 1), implementation of CARDAMOM at Easter Bush for 6 years (2005-2010) by assimilating the corresponding LAI observations.
- 3. MDF at Crichton: Using the calibrated distributions (step 1), implementation of CARDAMOM at Crichton for 2015 by assimilating LAI observations.
 - 4. Quantitative assessment of MDF at steps 1,2 and 3 against the assimilated data and against independent data.

In order to assess model accuracy and precision we calculate, for each variable examined, the percentage of observed data points (i.e. mean of observations) that fell within the 95% confidence intervals (CI) produced by the model runs. We refer to this metric as overlap and present it as a percentage that can take a value between 0 and 100. We also calculate (1) the Root Mean Squared Error (RMSE) to quantify the difference between measured and modelled data; (2) the bias in model predictions; and (3) the Pearson correlation coefficient (r) to quantify how well the trends in measured

data are captured by the model (for equations see Myrgiotis et al. (2016)).

It should be noted that —where mentioned— estimates of the uncertainty of
measurements come from using the RMSE equation after replacing base of
the exponent with the sum of 2 standard deviations of each measured data
point.

475 3. Results

3.1. Easter Bush

Three years (2002-2004) of measured LAI and NEE data were assimilated by CARDAMOM to calibrate the distributions of DALEC-Grass parameters. For the calibration period, 25 % of LAI observations and 90 % of NEE observations fell within the 95 % confidence interval of the CARDAMOM analysis while r was 0.25 and 0.56 respectively (Table 3). The calibrated parameter distributions were used to run DALEC-Grass for the subsequent 6 years of measurements (2005-2011); this time assimilating available measured LAI data only.

For the 2005 to 2011 MDF period, 85% of the measured weekly-mean NEE and 82% of the daily LAI data points lied within the 95% CIs. The variation in NEE (r=0.70) and LAI (r=0.74) was well captured. DALEC-Grass tended to overestimate both NEE (bias=0.38 gCm⁻²) and LAI (bias=0.47 m²m⁻²). We used a 15% relative uncertainty around the measured LAI data during the MDF process with CARDAMOM. We found that the RMSE of the measured against modelled LAI data was 10% lower than the mean uncertainty of the measured LAI data. We repeated this process for NEE, and found that the estimated RMSE was equal to the uncertainty attributed to the measured data during the MDF process (i.e. $1~\rm gCm^{-2}$) .

DALEC-Grass was able to capture the patterns and magnitudes in measured ER (Fig. 4). The comparison of modelled and measured ER estimates produced a r of 0.79, which reflects the model's skill in representing ER patterns. In terms of the relative size of ER, 83% of the mean measured ER data were within the modelled 95% CI. The estimated RMSE (1.5 gCm⁻²) was smaller than the average uncertainty of the measured data (1.65 gCm⁻²) (Table 3). The inter-annual patterns in MDF-estimated NEE, ER and GPP mirror the measured data as presented in Jones et al. 2017 (Fig. 5).

The mean simulated harvest $(283 \text{ gCm}^{-2}\text{a}^{-1})$ was just 3% higher than the measured harvest $(270 \text{ gCm}^{-2}\text{a}^{-1})$ in 2002 and 30% lower than the 2003 measured harvest $(170 \text{ gCm}^{-2}\text{a}^{-1})$ (Jones et al., 2017). However, both in

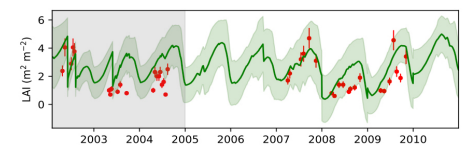


Figure 2: Modelled and measured daily Leaf Area Index (LAI) at the Easter Bush site. Grey-shaded area (2002-2004) shows the parameter calibration period. The unshaded area (2005-2011) represents the LAI data assimilation period. The mean modelled LAI and the CARDAMOM-estimated 95% confidence intervals presented in green. The measured LAI and its 15% relative uncertainty presented in red.

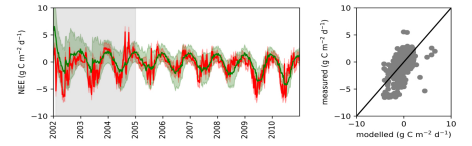


Figure 3: Weekly mean Net Ecosystem Exchange (NEE) at the Easter Bush site. From left to right: (1) Time series of measured (red) and modelled (green) weekly mean NEE. The green-shaded area represents the 95% confidence intervals and the red-shaded area represents the uncertainty around the measured NEE ($1 \text{gCm}^{-2} \text{d}^{-1}$). The parameter calibration period (2002-2004) is shown as grey-shaded area. (2) Scatter plot of measured and corresponding modelled weekly mean NEE.

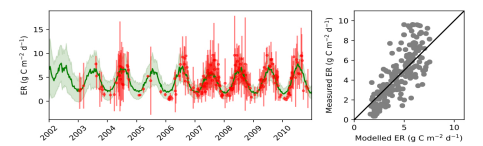


Figure 4: Weekly mean Ecosystem Respiration (ER) at the Easter Bush site. From left to right: (1) Time series of measured (red) and modelled (green) weekly mean ER. The green-shaded area represents the 95% confidence intervals and the red error bars the uncertainty around the measured ER. (2) Scatter plot of measured and corresponding modelled weekly mean ER.

Table 3: Model performance metrics for Easter Bush

Variable	Time period	r	Bias	Overlap	RMSE
LAI	2002-2004 2005-2010	$0.34 \\ 0.74$	$1.33 \\ 0.47$	$25 \% \\ 82 \%$	1.6 1.0
NEE	2002-2004 2005-2010	0.56 0.70	$0.95 \\ 0.38$	90 % 85 %	2.11 1.00
ER	2002-2010	0.79	-0.07	83 %	1.5

Weekly mean data used for Net Ecosystem Exchange (NEE) and ecosystem Respiration (ER). Overlap shows the percentage of observed data that lie within the model-based 95% CIs. Bias and RMSE in gCm $^{-2}$ for ER and NEE and in m 2 m $^{-2}$ for LAI.

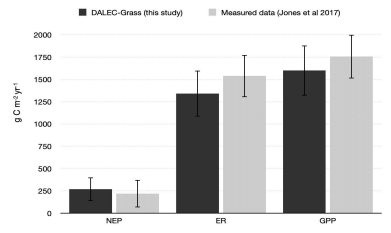


Figure 5: Comparison of MDF-based and measurements-based (after Jones et al. 2017) annual Net Ecosystem Production (NEP), Gross Primary Productivity (GPP) and Ecosystem Respiration (ER). Bars show the mean gCm⁻²yr⁻¹ between 2002 and 2010 and error bars show the inter-annual standard deviation.

2002 and 2003 the measured annual harvest was within the CARDAMOM estimated 95% CIs. The simulated Easter Bush grassland behaved as a typical UK permanent grassland producing an average of 262 gCm⁻² (5.6 tDMha⁻¹) of grass biomass per year. This level of biomass availability and removal is within the expected range (353 \pm 96 gCm⁻²a⁻¹) as estimated in a recent study by Qi et al., 2017. Finally, DALEC-Grass estimated a mean annual input to soils of 710 gCm⁻²a⁻¹ in the form of root and leaf litter.

3.2. Crichton

The calibrated parameter distributions retrieved for Easter Bush (2002-2004) were used as priors for the CARDAMOM analysis at the Crichton site. Four field-measured LAI data points were available for assimilation (Fig. 6). The assimilation of measured LAI data also affected the fit between measured and modelled aboveground biomass and grass harvest. All four measured LAI data points were within the modelled 95% CIs. However, the model did not capture the first of four aboveground biomass measurements. It should be noted that neither the quantity of grass harvested nor the amount of aboveground biomass (at any point in time) were provided to the model during the LAI data assimilation in CARDAMOM. Despite this discrepancy, the two simulated harvests removed 276 gCm⁻² (5.8 tDMha⁻¹) from the

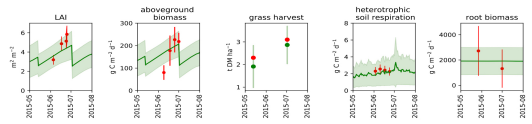


Figure 6: Modelled (green) and measured (red) LAI, above ground biomass, grass harvest, root biomass and heterotrophic respiration at the Crichton site. The red error bars represent the uncertainty around the measured data. The CARDAMOM-estimated 95% confidence intervals are presented in green shading

grassland, and each measured harvest yield was within the corresponding CARDAMOM's 95 % CI (Fig. 6).

DALEC-Grass was successful in reproducing the measured patterns and magnitudes of NEE. The estimated correlation coefficient was 0.88, the mean bias was 0.25 gCm⁻², the RMSE was 0.96 gCm⁻² and all the of the measured data were within the 95% CIs (Fig. 7). Notwithstanding the large variability among the collected samples of root biomass, the mean simulated amount of C contained in grass roots was within the respective measured ranges (Fig. 6). DALEC-Grass currently does not separate autotrophic respiration C into above and belowground fluxes. In order to allow for a comparison between measured and simulated respiration data we assumed that between 40% and 60% of total surface respiration can be attributed to heterotrophic sources (Li et al., 2018). While not directly measured we, henceforth, refer to heterotrophic respiration data as measured data. The comparison between measured and modelled heterotrophic respiration showed that all four measured data points lied within the 95% CI (Fig. 6). Finally, DALEC-Grass estimated that 780 gCm⁻²y⁻¹ were added to Crichton's soil in 2015 in the form of root and leaf litter.

3.3. MDF-retrieved distributions

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Model parameter distributions were calibrated using 3 years of LAI and NEE data from Easter Bush (calibration period). This calibration process led to reductions in the length of the uniform prior distributions that varied according to parameter. The average prior length reduction was 47% with achieved reductions being between 3% and 99%. Details on the application of CARDAMOM for parameter distribution calibration include pos-

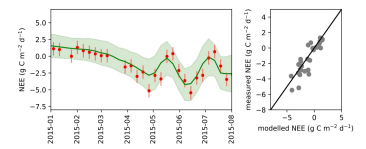


Figure 7: Weekly average Net Ecosystem Exchange (NEE) at Crichton. From left to right: (1) Time series of measured (red) and modelled (green) weekly mean NEE. The green-shaded area represents the 95% confidence intervals and the red error bars represent the uncertainty around the measured NEE ($1 \text{gCm}^{-2} \text{d}^{-1}$). (2) Scatter plot of measured and corresponding modelled weekly mean NEE.

Table 4: Model performance metrics for Crichton

Variable	\mathbf{r}	Bias	Overlap	\mathbf{RMSE}	
NEE (weekly mean)	0.88	0.25	100~%	0.96	

Weekly mean data used for Net Ecosystem Exchange (NEE). Overlap shows the percentage of observed data that lie within the model-based 95% CIs. Bias and RMSE in gCm⁻².

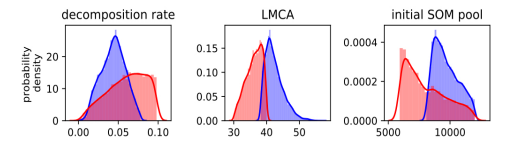


Figure 8: Probability densities of the posterior distributions for three selected DALEC-Grass parameters based on MDF results at Easter Bush (in red) and Crichton (in blue).

terior means, maximum-a-posteriori-probability and prior length reduction, and are presented in Table 5 in supplementary material along with information on MCMC chain convergence assessment (Fig. 9). 552

4. Discussion

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The results of this study show how MDF can quantify C dynamics in 554 grasslands under variable grazing and cutting regimes. A limited number of in-situ LAI observations were assimilated through the CARDAMOM MDF 556 framework in order to test the predictive skill of DALEC-Grass against 7 years of data at two Scottish grasslands. At the grazed Easter Bush grass-558 land 21 in-situ LAI observations were assimilated to produce time series of 559 C fluxes (NEE and ER) that closely matched corresponding measurements 560 (eddy covariance and chamber-based fluxes, respectively). At the harvested Crichton site 4 in-situ LAI observations were assimilated and CARDAMOM produced robust estimates of above and below-ground biomass C pools and NEE C fluxes. The use of CARDAMOM also allowed us to handle observational and model parametric uncertainty and provide a level of predictive uncertainty for the examined variables.

4.1. Model performance

Our MDF approach provides a probabilistic solution to the parameter identification problem. Among the factors that affect the robustness of model analyses are how measurement uncertainty and model parameter identification are assessed. In this study, measured data uncertainty has been considered at the parameter retrieval stage, through the use of the accuracy metric as the cost function, and at the model evaluation stage, through the use of the CIs in quantifying and expressing model prediction skill. As a consequence, the results of simulations are distributions of parameter values and provide a quantitative analysis of the parameter-induced uncertainty around the model's results. The average uncertainty around predicted C fluxes (NEE, ER) was less-than-or-equal to that attributed to NEE (1 gCm⁻²d⁻¹ based on literature) and estimated from ER measurements (1.65 gCm⁻²d⁻¹) (Hill et al., 2012; Revill et al., 2016).

CARDAMOM's key novelty is the use of ecological and dynamical constraints (EDCs) as conditions imposed on the parameter retrieval process. The aim of including EDCs is to ensure that the MDF process respects a set of mathematical, ecological and biogeochemical rules. In practice, the

success of the MDF process can be judged by the level of fit between the outputs of DALEC-Grass and the assimilated measured data. However, in this study, we wanted to have a stricter assessment of DALEC-Grass and CARDAMOM. We used 3 years of measured data on LAI and NEE (Easter Bush) to calibrate the parameter distributions and, thereafter, used the refined prior distributions and LAI data assimilation for the remaining 6 years 590 of data in Easter Bush and the one year of data in Crichton. The evaluation of MDF performance against NEE (Easter Bush and Crichton), ER (Easter Bush), biomass (Crichton) and heterotrophic respiration (Crichton) showed that DALEC-Grass was able to describe the examined C dynamics in managed grasslands with good accuracy. The mean overlap for NEE, ER and LAI 595 during the MDF implementation was equal to 88% and RMSE was less than, or equal to, the respective measurement uncertainty (for NEE and ER). We argue that calculating the overlap along with the RMSE is an effective way 598 to express the level of agreement between modelled and measured data for which uncertainty is quantified. 600

4.2. Retrieved parameter distributions

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The MDF application at Easter Bush and Crichton adjusted the distributions of model parameters to fit the observed LAI at each site. CARDAMOM was able to retrieve parameter distributions that led to model outputs that fit with the available observations while being conceptually and mathematically sound, as proven by the stability of the CIs for all the variables examined in the two grasslands. The parameter distributions obtained for each grassland did not differ for most parameters. However, for three parameters CAR-DAMOM produced informatively different distributions at the two grasslands (Fig. 8). For the "litter decomposition rate" parameter, the addition of animal excrement C to the soil's litter C pool led CARDAMOM to retrieve a higher decomposition rate (i.e. more litter C goes into the SOC pool) in Easter Bush compared to Crichton, where there was no grazing during the measurement period. A higher leaf C per area (LMCA) was retrieved for Crichton compared to Easter Bush. This result suggests that the vegetation of cut Crichton grassland is more C dense than that of the grazed Easter Bush grassland, something that relevant studies confirm (Liu et al., 2017; Zheng et al., 2015; Laliberté et al., 2012). We argue that the ability to infer the relative difference in vegetation C density between grasslands is an important outcome produced by the combination of the model's management related processes and the CARDAMOM EDCs. Moreover, DALEC-Grass depends on a simple soil C scheme and CARDAMOM is given a wide prior range for initial SOC pool size parameter. Despite this, CARDAMOM suggests that, considering productivity levels, allocation patterns and EDCs, Easter Bush had a lower soil C content than Crichton. These results are confirmed by field-measured data, which show that SOC in Easter Bush is around 30% less than SOC in Crichton (COSMOS-UK, 2019).

Based on the DALEC-Grass simulations and the corresponding measured data Easter Bush and Crichton were net sinks of C during the simulated periods. Both grasslands had a simulated annual aboveground biomass productivity that is within the observed limits (i.e. 238-429 gCm⁻²yr⁻¹). The parsimonious mechanisms used in DALEC-Grass to describe grazing and cutting were effective and the retrieved parameter distributions for the relevant parameters are in agreement with relevant observations (Genever and Buckingham, 2016). For Easter Bush, CARDAMOM inferred from the data and modelling that grazing occurs when the total aboveground biomass is $> 38-47 \text{ gCm}^{-2} \text{ (0.8-1.0 tDMha}^{-1})$. The distribution of the minimum precutting above ground biomass for Easter Bush showed that $\approx 120~{\rm gCm^{-2}}$ (2.8 tDMha^{−1}) is the most likely value for this parameter. The minimum precutting and pre-grazing aboveground biomass parameters (P27 and P28) also define how much biomass will be left standing immediately after each cutting and their posterior distributions suggest a minimum harvest of 86 gCm⁻² (1.8 tDMha⁻¹). This is a realistic, albeit low, minimum harvest yield for the UK where grasslands can be cut up to three times per year Qi et al.. 2017. Finally, DALEC-Grass results showed that the annual amount of C added to the soil as root and leaf litter is $\approx 10\%$ more in Crichton compared to that estimated for Easter Bush. This difference is a result of the adaptation of C allocation patterns in response to the presence of grazing animals which also leads to a lower leaf C content in the grazed ecosystem (Easter Bush) (Hao and He, 2019; Chen et al., 2015; Mcsherry and Ritchie, 2013).

4.3. Limitations

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Our analysis highlights areas for potential improvement in both the model and the MDF framework. DALEC-Grass is frugal with its number of parameters and therefore the processes it incorporates. The results of this study showed that there is still a margin for improvement, which can be inferred from the different evaluation metrics. Moreover, the model requires information on grassland management as forcing. This requirement currently limits the applicability of DALEC-Grass to the areas with detailed management

data. The ACMv1 model (i.e. the module used to estimate GPP) used in DALEC-Grass does not consider the impact of plant water availability, and thus drought stress, on GPP. This likely has a limited effect on simulations 661 in Scotland but DALEC-Grass can be updated if needed using a recently 662 developed version of ACM that explicitly considers water cycling on estima-663 tion of GPP (Smallman and Williams, 2019). Furthermore, DALEC-Grass 664 does not, at this stage, have a detailed description of the role of N for grass 665 growth and C allocation. Because Easter Bush and Crichton are amply fer-666 tilised grasslands the results of this study are premised on non N-limited 667 conditions. However, the lack of N cycling representation also means that 668 soil C to N ratio and its role in litter and organic matter decomposition is not 669 considered. The conversion of grazed biomass to C returned to the soil, in the 670 form of excrement, depends on generic conversion factors but the assumption that all the daily-produced animal excrement is deposited on the soil is not 672 realistic. Nevertheless, all livestock-related constants can be converted to parameters with appropriate respective priors, which can, in turn, be refined 674 by CARDAMOM. Finally, in this study, we used field data from two sites in Scotland. These grasslands are representative of grasslands in the UK, and 676 northwest Europe, but testing DALEC-Grass at grasslands across the world 677 is needed for broader application. 678

4.4. Future development

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Notwithstanding the aforementioned limitations, DALEC-Grass has a range of potential applications that vary from gap-filling time-series of C fluxes from micrometeorological towers to farm-level grassland C budgeting and validation of large-scale terrestrial ecosystem models. With appropriate development and testing the model can handle spatially resolved satellite-based data on LAI, which will allow the quantification of grassland productivity and C dynamics at landscape, regional and even national scales. To this end, DALEC-Grass will have to be developed in a way that allows the inference of management at grass-covered pixels of satellite images of the land's surface. Such a spatially-resolved version of DALEC-Grass, when used in CARDAMOM, will allow us to better understand how key factors such as livestock density and cutting intensity affect ecosystem productivity and C sequestration.

5. Conclusions

We demonstrated how a model of C dynamics linked by a model-data 694 fusion framework to observations of LAI generated constrained analyses of 695 grassland ecosystem functioning under management. The results of this 696 study suggest that landscape grassland C cycling can be constrained using 697 LAI data at relevant resolutions and accuracy. An initial parameter calibra-698 tion using eddy flux data constrained key C cycle parameters, leading to a 699 better understanding of grassland productivity and C sequestration capacity. 700 Once this calibration was completed, assimilation of LAI data over time al-701 lowed the model to make robust estimates of the effects of grass grazing and cutting on net CO_2 exchanges. The data assimilation approach meant that 703 the effects of parametric and observation uncertainties could be considered and quantified. We showed that the forecast uncertainty in our predictions 705 was comparable to that of independent observations. We provide evidence 706 that DALEC-Grass is a conceptually sound, structurally robust and compu-707 tationally lightweight model. In the era of EO satellites, and the associated 708 availability of swathes of data, the attributes of the model show its potential 709 to provide in-depth monitoring of managed grasslands across temporal and spatial scales. Our aim is to realise this potential by further appropriate 711 development and testing at landscape scales.

$_{713}$ 6. Acknowledgements

VM and MW devised the study concept. VM developed DALEC-Grass 714 , implemented the MDF and undertook the analysis with support from all authors. Remaining authors provided data from the study sites. VM led the 716 writing, with support from MW and LS. All authors contributed to the text. 717 This study was supported by the Natural Environment Research Council 718 (NERC) of the UK through the Soils Research to deliver Greenhouse Gas 719 REmovals and Abatement Technologies (Soils-R-GGREAT) project, and the 720 NERC GHG Programme GREENHOUSE project. This work was also part 721 funded by the Scottish Government's Strategic Research Programme. We thank Anthony Bloom (JPL-NASA) for his support in development of CAR-DAMOM. We thank Tim Hill (University of Exeter) for access to eddy covariance instrumentation used at Crichton.

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1173 Supplementary material

Table 5: DALEC-Grass parameters. Description, units and calibration results.

Code	Description	Unit	\mathbf{Prior}_{min}	\mathbf{Prior}_{max}	$\operatorname{Posterior}_{mean}$	$\mathbf{Posterior}_{SD}$	MAP	Prior length reduction (%)
P1	Decomposition rate	fraction d^{-1}	1.00e-05	0.3	0.06	0.03	0.08	97
P2	Fraction of GPP that is respired	-	0.4	0.51	0.4427	0.0298	0.4158	3
P3	GSI sensitivity for leaf growth	-	0.75	9	3.968	0.767	4.637	61
P4	NPP belowground allocation	-	0.01	1	0.330	0.068	0.352	60
P5	Maximum GSI for leaf turnover	-	1.00e-07	3	0.214	0.230	0.173	50
P6	Turnover rate of roots	fraction d^{-1}	1.00e-06	0.1	3.45e-03	1.98e-03	5.38e-03	91
P7	Turnover rate of litter	fraction d^{-1}	1.00e-06	0.1	4.39e-03	2.66e-03	3.07e-03	90
P8	Turnover rate of soil organic matter	fraction d^{-1}	1.00e-10	0.01	3.84e-05	2.75e-05	2.44e-06	99
P9	Temperature Q10 factor	-	0.008	0.15	0.0416	0.0162	0.0418	57
P10	Photosynthetic N use efficiency (PNUE)	g C per g N per leaf m ² per day	7	25	15	3	18	31
P11	Maximum GSI for labile/stem turnover	-	0.0001	2	0.634	0.137	0.603	68
P12	Minimum GSI temperature threshold	K	225	330	251	16	264	27
P13	Maximum GSI temperature threshold	K	225	330	304	20	303	17
P14	Minimum GSI photoperiod threshold	seconds	3600	30000	12283	5191	6892	23
P15	Leaf Mass C per Area (LMCA)	g C per m ² of leaf	20	60	46	6	46	56
P16	Initial C in stem/labile pool	${ m g~C~m^{-2}}$	1	300	151	75	52	4
P17	Initial C in foliar pool	${ m g~C~m^{-2}}$	1	300	132	68	43	13
P18	Initial C in roots pool	$ m g~C~m^{-2}$	1	5000	893	800	284	40
P19	Initial C in litter pool	${ m g~C~m^{-2}}$	1	5000	793	561	486	52
P20	Maximum GSI photoperiod threshold	seconds	3600	64800	31917	6262	28880	42
P21	Minimum GSI VPD threshold	Pa	1	5500	1209	929	186	27
P22	Maximum GSI VPD threshold	Pa	1	5500	3376	1156	1457	16
P23	Critical GPP for LAI increase	${ m g~C~m^{-2}~d^{-1}}$	1.00e-05	1	0.30	0.13	0.26	53
P24	GSI sensitivity for leaf senescence	-	0.96	1	0.99	0.00	1.00	52
P25	GSI growing stage indicator	-	0.3	3	1.27	0.13	1.13	82
P26	Initial GSI value	-	0.5	3	1.61	0.23	1.83	61
P27	Minimum vegetation DM for grazing	${ m kg~DM~ha^{-1}}$	500	2000	995	197	1114	30
P28	Minimum vegetation DM for cutting	$kg DM ha^{-1}$	1000	6000	3232	611	2896	52
P29	Leaf to stem allocation parameter	-	0.05	0.9	0.61	0.11	0.66	45
P30	Initial C in SOM pool	${ m g~C~m^{-2}}$	5000	15000	9912	2513	9240	31
P31	DM demand (as $\hat{\%}$ of animal weight)	-	0.01	0.03	0.02	0.01	0.03	2
P32	Post grazing labile/stem loss	-	0.001	0.75	0.16	0.06	0.19	67
P33	Post cutting labile/stem loss	-	0.001	0.75	0.18	0.15	0.08	34

GSI: Growing Season Index, VPD: Vapour Pressure Deficit, SOM: Soil Organic Matter,

OM: Dry Matter, GPP: Gross Primary Productivity, NPP: Net Primary Productivity MAP: Maximum a posteriori probability estimate Prior length reduction: $100*(1-(posterior_{max}-posterior_{min})/(prior_{max}-prior_{min}))$ Estimates for the prior range for parameter P30 come from (Bradley et al., 2006)

1174 Chain convergence assessment

The Gelman-Rubin potential scale reduction factor (PSRF) was calculated using the following equations:

$$B = \frac{N}{M-1} \sum_{m=1}^{M} (\overline{\theta_m} - \overline{\theta})^2$$
 (7)

$$W = \frac{1}{M} \sum_{m=1}^{M} \sigma_m^2 \tag{8}$$

$$\overline{V} = \frac{N-1}{N}W + \frac{M+1}{MN}B\tag{9}$$

$$PSRF = \sqrt{\frac{\overline{V}}{W}} \tag{10}$$

where θ is a model parameter, σ^2 is the variance, M is the number of chains and N is the length of each chain. A PSRF ≈ 1 shows that chain convergence was achieved. The use of EDCs in CARDAMOM means that N was not the same for all chains. For this reason we used the last 10000 values retrieved by the MH algorithm for each chain to calculate the PSRF. The PSRF for each parameter as estimated from results from the calibration period (2002-2004) are presented in Figure 9.

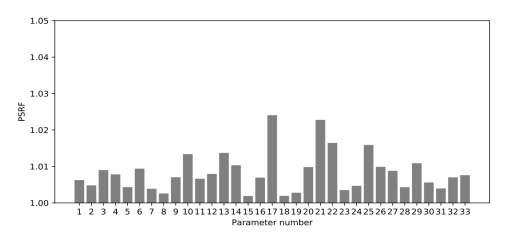


Figure 9: Estimated potential scale reduction factor (PSRF) for each model parameter after parameter calibration. The names (and other information) of each model parameter number are presented in Table 5

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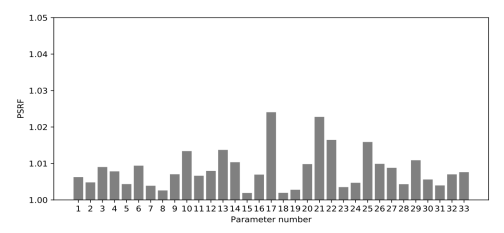


Figure 9: Estimated potential scale reduction factor (PSRF) for each model parameter after parameter calibration. The names (and other information) of each model parameter number are presented in Table 5

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