

## Meetings

# Improving collaborations between empiricists and modelers to advance grassland community dynamics in ecosystem models

Climate change, increasing atmospheric CO<sub>2</sub>, and land use change have altered biogeochemical and hydrologic cycles world-wide, with grassland systems being particularly vulnerable to resulting vegetation shifts (Komatsu *et al.*, 2019). Therefore, incorporating plant community dynamics into ecosystem models is critical for accurate forecasting of ecosystem responses to global change (Levine, 2016). Process-based ecosystem models, which simulate the biogeochemical transfers of mass and energy among biota, the subsurface, and atmosphere, require representation of dynamic composition of organisms within ecosystems. For example, these models simulate leaf and plant-level characteristics, such as electron transport rate and allometry of carbon (C) allocation, to predict how net primary productivity and other ecosystem processes respond to abiotic drivers. These models are particularly useful in scaling from organismal to ecosystem levels but are still underdeveloped in their ability to capture community change, especially in grassland ecosystems. To represent compositional changes, these models must simulate competition, mortality, establishment, and reproduction of plant populations within communities. Yet, current ecosystem modeling approaches to forecast plant community change have derived from studies of forested systems and are either too coarse to capture fine-scale community dynamics (e.g. dynamic global vegetation models (DGVMs)) or too complex to be used at large spatial scales (e.g. forest gap models).

Community ecology often relies on statistical models describing population dynamics or the abundance/frequency of individual species to identify linkages between community dynamics and ecosystem processes. For example, there is a vast literature linking species richness to ecosystem function using statistical models (Hooper *et al.*, 2005; Cardinale *et al.*, 2006). Yet, statistical models are rarely able to scale leaf and plant-level characteristics to ecosystem levels owing to data constraints and the fact that they do not incorporate process knowledge. Process-based ecosystem models are thus needed to predict whole ecosystem function, especially under novel environmental conditions. Yet, as already mentioned, these models often struggle to link plant characteristics to local-scale community dynamics.

To better represent community dynamics in ecosystem models, scientists must identify: (1) how physiological and morphological

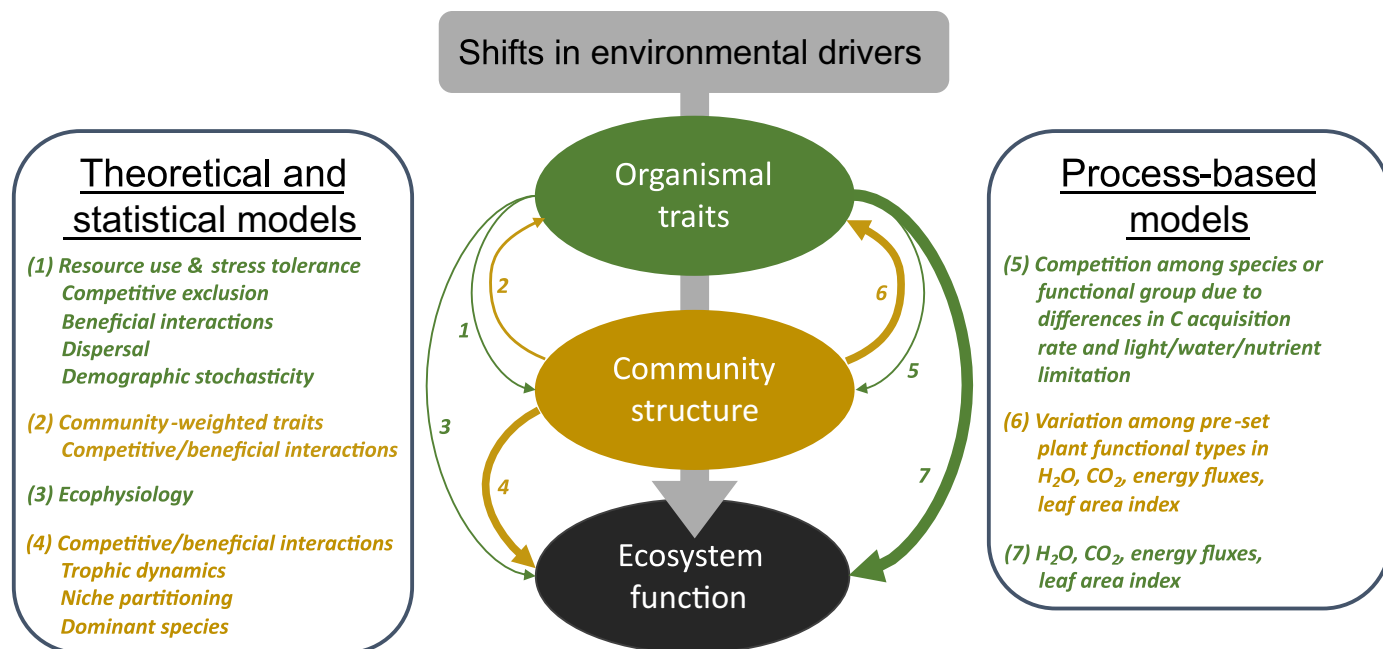
traits of plant species or functional types, and their diversity, can drive changes in community structure (Fig. 1, arrow 1); and (2) how community dynamics alter the distribution of traits across the entire community (Fig. 1, arrow 2); leading to (3) improved ability to simulate shifts in community structure and their concurrent effects on ecosystem functioning (Fig. 1, arrows 6 and 7). Additionally, ecosystem processes, as forced by abiotic drivers, should alter the competitive balance in community dynamics that are sensitive to the coupling between their physiological/morphological traits and the abiotic drivers (Fig. 1, grey arrow). Integration across organismal, community, and ecosystem ecology, as well as between empirical and process-based modeling approaches, is necessary to address this issue. Toward this end, we (K. Komatsu, M. Avolio, K. Wilcox) led a working group funded by the Long-Term Ecological Research Network from 2017 to 2019, where we gathered scientists from diverse fields (C2E Consortium) to identify challenges and formulate directions for better integration of community dynamics in land surface models (LSMs).

## Modelers and empiricists have similar goals

Whether simulating ecosystem processes or conducting field studies, our goal in ecology is often the same: to provide predictions of ecological states and processes under a variety of environmental conditions. Many studies of community dynamics do not collect the data necessary to parameterize process-based models, and models do not represent many of the processes that empiricists consider fundamental. This is often due to limited communication between process-based modelers (hereafter ‘modelers’) and ‘empiricists’ who collect biotic and abiotic data, where both modelers and empiricists lack familiarity with the needs of the other’s approaches. Though there are existing channels of communication, there is room for more discussion between modelers and empiricists focusing on population, community, and ecosystem dynamics. During our meetings, participant modelers and empiricists agreed on three existing knowledge gaps that should be prioritized by the scientific community to better represent critical processes controlling ecosystem dynamics: (1) resource competition; (2) community and population dynamics; and (3) grassland plant processes. To fill these knowledge gaps, empiricists should seek input from modelers when designing experiments, and modelers should consult empiricists when altering or developing models. In other words, let’s all talk more.

## Resource dynamics in dynamic global vegetation models

Current DGVMs are based on the integration of physical (e.g. light penetration through plant canopies, water transport across soil layers), biophysical (coupling of photosynthesis and transpiration),



**Fig. 1** Conceptual figure showing how ecologists bridge hierarchical scales (ovals) using both statistical and process-based models to predict how ecosystem function will respond to shifts in environmental drivers, with arrow thickness representing our interpretation of the general level of understanding and/or frequency of scaling attempts in the field. These scaling attempts focus on various processes and biotic characteristics (examples in italics). Traditionally, statistical models have been used to assess how variation in environmental drivers results in variation in ecosystem function (large gray arrow), as well as how differences in community structure (biodiversity, species composition) result in differences in ecosystem function. Process-based models use information about plant characteristics (i.e. traits) within ecosystems to determine water, energy, and biogeochemical fluxes (arrow 7), sometimes incorporating direct impacts of shifts in abundance of plant functional types or cohorts (arrow 6). More recently, capacity for species-level responses to environmental drivers has been built into process-based models (arrow 5), yet our understanding of how species-level physiological responses translates to changes in community structure (arrow 1), and how these changes alter the physiological/morphological traits of the ecosystem (arrow 2), is still lacking. We propose that future research strives to increase our mechanistic understanding of how organismal responses lead to shifts in community structure so that these mechanisms can be incorporated into process-based models.

and biogeochemical (e.g. allocation of C to growth, uptake of available nutrients by plant roots) processes (Fig. 1, arrows 5 and 6). Whereas leaf biophysics are currently well-modeled in most ecosystem models, scaling up from the leaf to the canopy is challenging. Accounting for demography, plasticity, and other resource dynamics (e.g. nitrogen and water) and their trade-offs is less well developed. For example, stress-tolerant species can outcompete fast-growth species under low resources (*sensu*  $R^*$  theory, Tilman, 1982; Fig. 1, arrow 1). Although this phenomenon would be represented in the demographic component of many models if the process-based physics of C uptake, respiration, and allocation to growth were perfectly described for different species, even the most advanced demographic components of current models struggle to show  $R^*$ -type dynamics due to insufficient data to parameterize the differences between plant types. To enable ecosystem models to better represent plant diversity in resource use and competitive trade-offs, empiricists need to provide modelers with additional data on species characteristics (e.g. demography, physiology, morphology) that define the strategies and trade-offs among plant traits and differential sensitivity to core environmental variables (e.g. soil moisture, temperature) at different temporal scales: sub-diurnally for sensitivity of biota to environmental drivers, daily to seasonally for growth and senescence timing and rates, and interannually for community change. These data will allow for models to better represent differential species responses to

environmental drivers in their acquisition of resources, and how this translates to community structure and ecosystem function.

### Community dynamics in models

Plant community dynamics can have strong impacts on ecosystem function, which has been demonstrated both theoretically (Smith *et al.*, 2009; Langley & Hungate, 2014; Polley *et al.*, 2014) and empirically (Isbell *et al.*, 2013; Reich & Hobbie, 2013; Avolio *et al.*, 2014; Smith *et al.*, 2015; Xu *et al.*, 2015; Fig. 1, arrow 4). Ecosystem function can be altered when community dynamics result in a system having a different collection of plant physiological and morphological traits. Thus, incorporation of community dynamics related to species identities and their traits into process-based models is critical for accurately forecasting ecosystem processes.

The need to incorporate community dynamics into process-based models has been recognized by the scientific community, and increasing efforts are being made to achieve this goal for LSM-coupled DGVMs (Moorcroft, 2006; Evans, 2012; Scheiter *et al.*, 2013; Levine, 2016). Ecosystem models represent community change in a few different ways: DGVMs define suites of traits associated with broad functional groups; gap models assign traits to individual organisms to allow species and cohorts to compete for resources; and trait-based models allow traits to vary plastically with

environmental conditions without explicit representation of species (Fisher *et al.*, 2018). Each of these methods provides unique strengths and challenges. For example, community shifts within functional groups (e.g. shift in dominance from one  $C_4$  grass to another  $C_4$  grass) can have large impacts on ecosystem function (Wilcox *et al.*, 2016), which would not be captured by DGVMs; gap-based models do well to represent cohort competition for light after disturbance, yet they struggle to represent competition for resources such as water and nutrients (Powell *et al.*, 2013); and global or regional-scale trait–environment relationships used within trait-based approaches often make assumptions about space-for-time substitutions, which can be problematic (Burke *et al.*, 1997). Thus, better understanding and representation of mechanisms behind intra-functional group dynamics within ecosystem models is important to improve accuracy of temporal projections under chronically altered environmental scenarios. This requires expanding the number of species or functional groups represented within ecosystem models and a better parameterization of these species/groups.

### Grasslands are not well represented in land surface models

Grasslands, in addition to driving the evolution and rise of hominids (Bobe, 2004; Uno *et al.*, 2016), cover *c.* 40% of the Earth's ice-free terrestrial surface (White *et al.*, 2000) and provide vital ecosystem services, such as forage production, C sequestration, and agriculture (Sala & Paruelo, 1997). Yet, many ecosystem models were initially designed based on forest dynamics and so have difficulty representing important dynamics in grasslands, including  $C_3$  vs  $C_4$  competition, the distribution of mixed grass–tree systems, and intra-functional group competition. Some reasons for these difficulties are variable grassland phenology and grass allometry, and inadequate representation of key drivers of grassland dynamics, such as fire and grazing.

Through our discussions, we identified some important considerations for overcoming current limitations of representing grasslands in ecosystem models. First, grasses and trees have fundamentally different growth forms, and models need to move beyond current approaches that incorporate grasses as either miniature trees or as a green carpet. Instead, ecosystem models should represent the unique attributes of grasses, including strong water limitation, fast tissue turnover rates, multiple aboveground stems from one belowground organ system, and a high prevalence of vegetative reproduction. These are major distinctions important for the C balance and total biomass of grasses. Additionally, moving beyond a 'green carpet' approach may be important for accurate representation of surface roughness in grasslands. Going forward, vegetation demographic models (Moorcroft *et al.*, 2001) are emerging as an approach to solve the potential pitfalls of scaling from individuals to the ecosystem level, as well as from ecosystem patches to the global scale. This approach could be particularly applicable to grasslands, where the number of individuals per square meter makes modeling interactions among individuals computationally unreasonable. Second, the way that fire – a key driver of grassland form and function (Bond & Keeley, 2005) – is

currently modeled in LSMs has also been built with forested ecosystems in mind, leading to poor representation of this driver for grassland ecosystems. For example, although fire can result in mortality in forest systems, thus 'resetting' the ecosystem, fire in grasslands often does not kill individuals, and many dominant grassland species are promoted by frequent fire (Hulbert, 1988).

### Leveraging existing coordinated experiments

One exciting opportunity to realize better understanding and forecasts of community-to-ecosystem linkages is to leverage existing coordinated experiments. Coordinated experiments implement the same experimental design at many sites around the globe, typically involving teams of investigators at each site. Coordination with these existing teams could allow for the measurement of necessary environmental and species physiological/morphological trait data around the globe, which may then be incorporated into models. Examples of such coordinated networks include the Nutrient Network (NutNet; Borer *et al.*, 2014), Drought Network (DroughtNet), and Forest Geological Observatory (ForestGEO; Anderson-Teixeira *et al.*, 2015). For these networks to be leveraged, prioritization of measurements is needed to prevent overloading of network investigators, as these experiments have been designed to limit time investment by any one investigator. Alternatively, the network can be leveraged without relying on many individual investigators if outside funding is available for a small team to make many measurements across many sites. By highlighting the common goals of empiricists and modelers for investigators within coordinated networks, and developing protocols for field measurements of key model parameters, the goal of improved understanding of how community dynamics drive ecosystem processes can be made more achievable.

### Conclusions

Working together, empiricists and modelers can overcome some of the limitations of isolated empirical studies (relatively short timescale, small spatial extent, costs of measuring everything required to mechanistically explain responses) and modeling efforts (unreasonable/simplistic assumptions, limited representation of environmental heterogeneity and ecological complexity). This may be done using data collected from empirical studies for model initialization and parameterization, using updated models to quantitatively test and develop mechanistic ecological hypotheses, and to help explain the processes causing observed community dynamics. We have identified three core areas where both empiricists and modelers could work together to improve understanding of community and ecosystem processes: first, trait diversity in resource use and trade-offs; second, better representation of mechanisms controlling community dynamics; and third, more accurate representation of key grassland processes. Achieving these core goals will require an iterative process where empiricists and modelers work together to develop hypotheses about community dynamics and ecosystem processes, design experiments to generate data to test these hypotheses, and incorporate these data

into ecosystem models for evaluation, which can lead to development of new hypotheses. These types of collaborative research cycles will allow scientists to greatly increase understanding and forecasting capabilities of ecological patterns and processes at broad scales.




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## Author contributions

KRW, KJK and MLA conceived of and organized the meeting. All co-authors attended the meetings. KRW, KJK and MLA wrote the manuscript, with editorial contributions from all other co-authors.

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## References

Anderson-Teixeira KJ, Davies SJ, Bennett AC, Gonzalez-Akre EB, Muller-Landau HC, Jose Wright SJ, Abu Salim K, Almeida Zambrano AM, Alonso A, Baltzer JL *et al.* 2015. CTFS-ForestGEO: a worldwide network monitoring forests in an era of global change. *Global Change Biology* 21: 528–549.

Avolio ML, Koerner SE, La Pierre KJ, Wilcox KR, Wilson GWT, Smith MD, Collins SL. 2014. Changes in plant community composition, not diversity, during a decade of nitrogen and phosphorus additions drive above-ground productivity in a tallgrass prairie. *Journal of Ecology* 102: 1649–1660.

Bohe R. 2004. The expansion of grassland ecosystems in Africa in relation to mammalian evolution and the origin of the genus *Homo*. *Palaeogeography, Palaeoclimatology, Palaeoecology* 207: 399–420.

Bond W, Keeley J. 2005. Fire as a global 'herbivore': the ecology and evolution of flammable ecosystems. *Trends in Ecology & Evolution* 20: 387–394.

Borer ET, Harpole WS, Adler PB, Lind EM, Orrock JL, Seabloom EW, Smith MD. 2014. Finding generality in ecology: a model for globally distributed experiments. *Methods in Ecology and Evolution* 5: 65–73.

Burke IC, Lauenroth WK, Parton WJ. 1997. Regional and temporal variation in net primary production and nitrogen mineralization in grasslands. *Ecology* 78: 1330–1340.

Cardinale BJ, Srivastava DS, Emmett Duffy J, Wright JP, Downing AL, Sankaran M, Jouseau C. 2006. Effects of biodiversity on the functioning of trophic groups and ecosystems. *Nature* 443: 989–992.

Evans MR. 2012. Modelling ecological systems in a changing world. *Philosophical Transactions of the Royal Society B: Biological Sciences* 367: 181–190.

Fisher RA, Koven CD, Anderegg WRL, Christoffersen BO, Dietze MC, Farrior CE, Holm JA, Hurtt GC, Knox RG, Lawrence PJ *et al.* 2018. Vegetation demographics in Earth system models: a review of progress and priorities. *Global Change Biology* 24: 35–54.

Hooper DU, Chapin FS, Ewel JJ, Hector A, Inchausti P, Lavorel S, Lawton JH, Lodge DM, Loreau M, Naeem S *et al.* 2005. Effects of biodiversity on ecosystem functioning: a consensus of current knowledge. *Ecological Monographs* 75: 3–35.

Hulbert LC. 1988. Causes of fire effects in tallgrass prairie. *Ecology* 69: 46–58.

Isbell F, Reich PB, Tilman D, Hobbie SE, Polasky S, Binder S. 2013. Nutrient enrichment, biodiversity loss, and consequent declines in ecosystem productivity. *Proceedings of the National Academy of Sciences, USA* 110: 11911–11916.

Komatsu KJ, Avolio ML, Lemoine NP, Isbell F, Grman E, Houseman GR, Koerner SE, Johnson DS, Wilcox KR, Alatalo JM *et al.* 2019. Global change effects on plant communities are magnified by time and the number of global change factors imposed. *Proceedings of the National Academy of Sciences, USA* 116: 17867–17873.

Langley JA, Hungate BA. 2014. Plant community feedbacks and long-term ecosystem responses to multi-factored global change. *AoB Plants* 6: plu035.

Levine JM. 2016. A trail map for trait-based studies. *Nature* 529: 163–164.

Moorcroft PR. 2006. How close are we to a predictive science of the biosphere? *Trends in Ecology & Evolution* 21: 400–407.

Moorcroft PR, Hurtt GC, Pacala SW. 2001. A method for scaling vegetation dynamics: the ecosystem demography model (ED). *Ecological Monographs* 71: 557–586.

Polley HW, Derner JD, Jackson RB, Wilsey BJ, Fay PA. 2014. Impacts of climate change drivers on C<sub>4</sub> grassland productivity: scaling driver effects through the plant community. *Journal of Experimental Botany* 65: 3415–3424.

Powell TL, Galbraith DR, Christoffersen BO, Harper A, Imbuzeiro HMA, Rowland L, Almeida S, Brando PM, da Costa ACL, Costa MH *et al.* 2013. Confronting model predictions of carbon fluxes with measurements of Amazon forests subjected to experimental drought. *New Phytologist* 200: 350–365.

Reich PB, Hobbie SE. 2013. Decade-long soil nitrogen constraint on the CO<sub>2</sub> fertilization of plant biomass. *Nature Climate Change* 3: 278–282.

Sala OE, Paruelo JM. 1997. Ecosystem services in grasslands. In: Daily GC, ed. *Nature's services: societal dependence on natural ecosystems*. Washington DC, USA: Island Press, 237–251.

Scheiter S, Langan L, Higgins SI. 2013. Next-generation dynamic global vegetation models: learning from community ecology. *New Phytologist* 198: 957–969.

Smith MD, Knapp AK, Collins SL. 2009. A framework for assessing ecosystem dynamics in response to chronic resource alterations induced by global change. *Ecology* 90: 3279–3289.

Smith MD, La Pierre KJ, Collins SL, Knapp AK, Gross KL, Barrett JE, Frey SD, Gough L, Miller RJ, Morris JT *et al.* 2015. Global environmental change and the nature of aboveground net primary productivity responses: insights from long-term experiments. *Oecologia* 177: 935–947.

Tilman DG. 1982. *Resource competition and community structure*. Princeton, NJ, USA: Princeton University Press.

Uno KT, Polissar PJ, Jackson KE, deMenocal PB. 2016. Neogene biomarker record of vegetation change in eastern Africa. *Proceedings of the National Academy of Sciences, USA* 113: 6355–6363.

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- White RP, Murray S, Rohweder M, Prince SD, Thompson KM. 2000. *Grassland ecosystems*. Washington, DC, USA: World Resources Institute.
- Wilcox KR, Blair JM, Smith MD, Knapp AK. 2016. Does ecosystem sensitivity to precipitation at the site-level conform to regional-scale predictions? *Ecology* 97: 561–568.
- Xu Z, Ren H, Li M-H, van Ruijven J, Han X, Wan S, Li H, Yu Q, Jiang Y, Jiang L. 2015. Environmental changes drive the temporal stability of semi-arid natural grasslands through altering species asynchrony. *Journal of Ecology* 103: 1308–1316.A1

## Appendix A1

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