Multiscale digital Arabidopsis predicts individual organ and whole-organism growth

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Understanding how dynamic molecular networks affect whole-organism physiology, analogous to mapping genotype to phenotype, remains a key challenge in biology. Quantitative models that represent processes at multiple scales and link understanding from several research domains can help to tackle this problem. Such integrated models are more common in crop science and ecophysiology than in the research communities that elucidate molecular networks. Several laboratories have modeled particular aspects of growth in Arabidopsis thaliana, but it was unclear whether these existing models could productively be combined. We test this approach by constructing a multiscale model of Arabidopsis rosette growth. Four existing models were integrated with minimal parameter modification (leaf water content and one flowering parameter used measured data). The resulting framework model links genetic regulation and biochemical dynamics to events at the organ and whole-plant levels, helping to understand the combined effects of endogenous and environmental regulators on Arabidopsis growth. The framework model was validated and tested with metabolic, physiological, and biomass data from two laboratories, for five photoperiods, three accessions, and a transgenic line, highlighting the plasticity of plant growth strategies. The model was extended to include stochastic development. Model simulations gave insight into the developmental control of leaf production and provided a quantitive explanation for the pleiotropic developmental phenotype caused by overexpression of miR156, which was an open question. Modular, multiscale models, assembling knowledge from systems biology to ecophysiology, will help to understand and to engineer plant behavior from the genome to the field.

Our goal is to understand the physiological effects of metabolic and regulatory networks that are now being elucidated at the molecular level. Such networks control the traits, such as drought resistance, that are important both in agriculture and in ecosystem responses to climate change. Molecular genetic approaches, often in model organisms, have uncovered the operating principles and mechanisms for a growing number of physiologically relevant cases. For example, environmental factors such as CO₂ concentration, temperature, and light flux can display coordinated diurnal and seasonal fluctuations (1, 2). For annual plants like the laboratory model species Arabidopsis thaliana, matching the timing of flowering to the favorable season, and thus the associated environment, increases reproductive success (3). This synchronization is achieved by changing gene expression and protein abundance at the molecular level. Arabidopsis FLOWERING LOCUS T (FT) is an example of such an integrator gene that induces flowering in response to environmental signals (4). FT is highly expressed in long (summer) days due to a combination of light and circadian clock regulation (5–8). Such responses collectively enable individual plants to survive in variable conditions. Plants adapt their resource allocation processes to the environmental conditions to optimize growth and biomass accumulation (9). Plants also adjust their architecture to compete for light and nutrient resources (10, 11). Given the multiplicity and interactions of such responses, however, it can be difficult to determine how much a particular molecular change contributes to the effect on the whole plant. To understand physiology and to facilitate predictive biology from the molecular level, there is a well-recognized need for quantitative models that cross biological scales and link understanding from several scientific domains (12–14).

There already exist mathematical models describing various plant processes and their interactions with the environment (13). These models include varying levels of mechanistic detail, starting from simple statistical relationships, and they usually comprise two scales at most (15). Broader, molecular-based models are well advanced in only a few domains of plant science, such as photosynthesis research (16, 17) and root development (18). If the models can be assembled and updated in a modular fashion, then larger, multiscale models might be developed in plant growth model | digital organism | crop modeling | ecology

Significance

Plants respond to environmental change by triggering biochemical and developmental networks across multiple scales. Multiscale models that link genetic input to the whole-plant scale and beyond might therefore improve biological understanding and yield prediction. We report a modular approach to build such models, validated by a framework model of Arabidopsis thaliana comprising four existing mathematical models. Our model brings together gene dynamics, carbon partitioning, organ growth, shoot architecture, and development in response to environmental signals. It predicted the biomass of each leaf in independent data, demonstrated flexible control of photosynthesis across photoperiods, and predicted the pleotropic phenotype of a developmentally misregulated transgenic line. Systems biology, crop science, and ecology might thus be linked productively in a community-based approach to modeling.
a distributed, community approach. Experts in each domain could model a particular aspect of biology in detail and reuse the previously assembled models to represent other aspects, where coarse granularity might be sufficient. This approach follows that developed in other contexts, for example in other areas of biology (19–21) and in the Earth system modeling community. In that case, submodels for the atmosphere, ocean, ice sheets, and the land surface are coupled; their interactions and dynamics are then evaluated against independent observations (22).

Our work was motivated by the multiple challenges of coupling growth models to the molecular level. Computational problems are expected, due to logical or technical incompatibilities among models (23), but historical factors are also common, including the inaccessibility of executable forms or reference data for some published models. These challenges have been overcome to varying extents in modeling frameworks for crop science (24, 25) and for animal physiology (26) but rarely in plant molecular genetics (27) and molecular systems biology (28). In addition, one key biological issue is whether the data that were used in the construction (or calibration) of the constituent models are quantitatively compatible, so the models can be combined with minimal modification. Recalibrating the parameter values of models at many different scales is potentially laborious, requiring coordinated data acquisition by researchers from multiple disciplines. The quantititative compatibility of the constituent models is therefore a key question.

To develop a quantitative model that links multiple, interacting processes from metabolism to development, we implemented the modular modeling approach for *Arabidopsis thaliana*. The *Arabidopsis* research community has abundant molecular and physiological data, but the quantitative variation observed among laboratories is significant (29). We identified four independently developed models that characterize different aspects of plant biology and then combined and extended them to form a framework model (FM) for vegetative growth in *Arabidopsis* (Fig. 1). Key aspects of the models showed very good quantitative agreement. The FM was validated against metabolic, physiological, and biomass data for multiple genetic backgrounds and tested for several growth conditions. Numerical simulations using FM enabled us to understand the physiological relevance of an observed developmental process. Finally, we compared additional data with model predictions to quantitatively explain a developmental phenotype, supporting one of two proposed mechanisms.

**Model Description**

**Components of the FM.** The circadian clock in *Arabidopsis* is one example of a pervasive, molecular regulator where we have substantial understanding of its molecular mechanisms (30). Photosynthetic carbon metabolism, vegetative growth, and flowering time are among the many biological processes controlled by the clock. A multiscale model will be required to understand how the circadian timing (or any other pervasive control) of each of these interacting processes contributes quantitatively to the growth of the whole plant under varying environmental conditions. We therefore integrated four models, each of which originated in a different laboratory:

**i) A carbon dynamic model (CDM) that considers the subcellular processes of photosynthesis and sugar-starch partitioning, as well as carbon (C) allocation between the leaf area and the roots (31). It is assumed that a fixed proportion (12.5%) of C assimilated through photosynthesis is partitioned into starch, with the possibility to accumulate more starch if the remainder of the photosynthate (in the form of sugar) is not used for growth and respiration. At night, starch is degraded at a linear rate, adjusted to the night length, to sustain growth until dawn (32, 33). The rate of starch degradation is set such that 84% of that accumulated in the light period is degraded by dawn. The CDM was a discrete-time model with a 6-s time step, constructed using data of Columbia (Col) WT plants grown under 8-h light (L):16-h dark (D) conditions.**

**ii) A functional-structural plant model (FSPM) that describes individual organ growth and how each organ (leaf) contributes to the above-ground structure for light interception.**

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**Fig. 1.** Overview of the FM. The FM takes environmental data as input (black) to four existing *Arabidopsis* models (blue shadowed boxes), which are (A) a carbon dynamic model (CDM) that describes carbon assimilation and resource partitioning (31); (B) a functional-structural plant model (FSPM) of individual organ growth that determines the rosette structure (green) and the area for light interception (34); (C) a photothermal model (PTM) that predicts flowering time (1); and (D) a photo-periodism model (PPM), which is a gene dynamic model of the circadian clock and the photoperiod response pathway (6). On integration, several original components were discarded (gray), whereas new connections were created (red).
(34). Each of these factors is represented by effective mathematical functions in the model, without mechanistic detail, but in a very concise form that was sufficient to represent *Arabidopsis* shoot growth and structure (34). It was parameterized using data of Col WT plants grown under 12-h L:12-h D conditions. Only a subset of the large original model (34) was applicable to our study. The relevant subset of parameter values and developmental structures was rewritten into a conventional, dynamic form that was compatible with the other submodels, as a discrete-time model with an hourly time step.

iii) A photoperiod model (PTM) that predicts the timing of flowering, based on temperature integrated over time (thermal time) (1). In *Arabidopsis*, flowering time is governed by the photoperiod pathway that enables plants to sense daylength (35), the vernalization pathway that promotes flowering in the spring after a long chilling period over winter (36), and warm ambient temperature (37). Each of these factors is represented by effective mathematical functions in the model, without mechanistic detail, and it was parameterized using field data of various genotypes in the Col and Landsberg erecta (Ler) backgrounds (38). The model was formulated as a discrete-time model with an hourly time step.

iv) A photoperiodism model (PPM), which is a gene dynamic model of the circadian clock (39) and the photoperiod pathway (6). This model was a conventional ordinary differential equation (ODE) model, usually solved with an adaptive time step of minutes or less. The model was parameterized using data from Col and Ler WT plants grown under 16-h L:8-h D and 8-h L:16-h D conditions.

**Model Integration Process.** To link the four models, we first identified the essential variable(s) from each that could act as the connection points. New links and scaling factors were introduced, whereas redundant model components were replaced (Fig. 1 and SI Appendix). Unit conversions were required for compatibility, and two parameter values (viii and ix below) were measured from our experiments. The 124 other parameter values were taken from the original models. A summary of the integration process is as follows:

i) The model’s time step was standardized to 1 h for all except for the PPM, which is solved at shorter, variable time steps. Our model therefore takes hourly meteorological data as input, similar to many crop and ecosystem models (Fig. 2 A–C) and thereby resolves diel behavior.

ii) The simple root-to-shoot carbon allocation ratio in the CDM was directly replaced with the dynamic pattern of demand from individual organs, calculated by the FSPM (Fig. 2 E and F).

iii) To facilitate the replacement step ii above, biomass measures considering only carbon in the CDM were converted to total dry mass using published leaf and root carbon content data (40–42), because not all biomass is carbon.

iv) The simple “big leaf” rosette area for photosynthesis in the CDM was directly replaced by the projected area of the rosette structure from the FSPM.

v) The sugar supply calculated by the CDM, from fine-grained processes such as photosynthesis, respiration, and sugar-starch partitioning, was directly provided to the FSPM as the sugar supply for growth. This variable replaced the empirical light use efficiency (LUE) component, which was previously estimated from experimental data through model inversion (34).

vi) Seedling emergence (43) and flowering time were represented explicitly, in terms of thermal times to emergence and flowering from the PTM. These variables were not previously considered in the CDM and the FSPM.

vii) The simple, piecewise-linear function for photoperiod response in the PTM was replaced by the continuous flowering function driven by the integrated expression of the flowering time gene *FT* in the PPM (6).

viii) The modified photothermal unit (MPTU) threshold in the PPM (threshold, Fig. 2H) was determined using the time of flowering measured in our experiments.

ix) Water content was measured from our experiments to facilitate simulation of fresh biomass, because this is a simpler and more widely available measurement than the dry mass used in both the CDM and the FSPM.

All of the modeling work and analysis were conducted in MATLAB (Mathworks) (SI Appendix). The Plant Systems Modeling (PlaSMo) online model repository (www.plasmo.ed.ac.uk) was developed as a shared portal to disseminate relevant models from systems biology and ecoinformatics. The component models and the FM are publicly accessible from PlaSMo in MATLAB and Simile formats with the following identifiers: CDM, PLM_2; FSPM, PLM_75; PTM, PLM_73; PPM, PLM_74; and FM, PLM_76. Simile provides a visual modeling environment with a graphical user interface, plotting tools, and an animated display of simulated plant growth (Movie S1) (44).

**Results.**

**Model Validation and Testing.** We first examined the performance of the FM in representing the growth of Col, which was the common *Arabidopsis* accession used to create the original models. As the model’s flowering time was calibrated to the data, we focus here on vegetative growth. WT Col plants were grown in 12-h L:12-h D cycles close to 22 °C, because these conditions most closely matched the conditions used for the original models, except for the CDM that was tested using an 8-h photoperiod (31). Highly discriminating data sets were collected for the biomass of the total shoot and individual leaves and for the area of individual leaves at multiple time points after sowing. Using the original parameter values for each submodel, the FM overestimated growth (SI Appendix, Fig. S1). However, the literature shows that *Arabidopsis* grown in an 8-h photoperiod have altered photosynthetic physiology compared with our reference 12-h photoperiod. Specifically, the ratio of maximum electron transport to the maximum rate of carboxylation (*I* <sup>max</sup> *V* <sup>max</sup>) decreases as photoperiod increases (45–47) (SI Appendix, Table S7). The CDM’s original value for *J* <sup>max</sup> *V* <sup>max</sup> has only been tested in an 8-h photoperiod (31). Substituting the value measured in a 12-h photoperiod was sufficient for the FM to fit the Col biomass data (Fig. 2 I–K). The *R* <sup>2</sup> between measured and modeled values of fresh biomass, dry biomass, and area of the rosette were 0.98, 0.99, and 0.98, respectively, with normalized root mean square error (nRMSE) less than 10% (SI Appendix, Table S8). The median values of *R* <sup>2</sup> and nRMSE for all of the data, including individual leaf predictions, were 0.91 and 24.7%, respectively. The dynamic operation of the model in Simile is illustrated in Movie S1.

The FM was next tested by comparison with growth data from other *Arabidopsis* accessions: Ler and Feira-0 (Fei). Accession-specific parameters were measured for the seedling emergence and flowering times, as described above for Col, and for the changing rate of leaf production. Fei was expected to show a higher leaf appearance rate (48), and indeed it showed a larger leaf number compared with Ler at the same time points (SI Appendix, Fig. S24). However, leaf appearance rate in Fei matched the Col rate when plotted against thermal time after seedling emergence (SI Appendix, Fig. S2B). We infer that the principal difference of Fei from Col is actually in the time to emergence, as Fei emerged at half the thermal time for Col (SI Appendix, Table S6). With only these changes, the model’s match to data of Ler and Fei plants was as good as for Col (Fig. 3 A, B, D, and E), with median *R* <sup>2</sup> (and nRMSE) of 0.94 (16%) and 0.95 (17.3%), re-
Fig. 2. The FM’s workflow predicts whole-plant and individual organ growth data. Input data required are hourly light intensity (A), CO₂ level (B), and temperature (C), illustrated for simulated three 12-h light (open):12-h dark (shaded area) cycles. Carbon supply (D) is used as sugar (dashed line) or stored as starch (solid line). Carbon is allocated at each hourly time step according to the organ demand (E and F). The simulated pattern of demand from individual leaves (f, thin blue lines, left axis) is used to calculate the sum of demand (dots) from leaves (thick blue line, right axis) and roots (brown line, left axis). The root-to-shoot allocation ratio (E), calculated dynamically from the FSPM (red line), is similar to the piecewise-linear function prescribed in the CDM (31) (gray dashed line), which it replaces. Times of dawn and dusk (dots in A) affect the level of flowering gene FT mRNA (G) simulated by the PPM, which in turn controls the accumulation of modified photothermal units (MPTU; H). Once the accumulated photothermal units reach the threshold for flowering (dashed lines), rosette growth is terminated in the FSPM (red arrow). Model outputs include biomass of the shoot (I) and individual leaves (J). Simulations for the Col WT (green lines) closely match experimental data for (I) total shoot biomass, (J) leaf biomass, and (K) leaf area at 18 (○), 25 (●), 27 (□), and 38 (■) d after sowing. Leaves are ranked according to the order of appearance. The integrated model uses simulated sizes of individual leaves (K) to calculate the projected rosette area for photosynthesis (red arrow), considering the spiral leaf arrangement (phyllotaxy) and upward (zenithal) angle. Experimental conditions: ∼21.3 °C; 12:12-h light/dark cycle; light intensity, 110 μmol·m⁻²·s⁻¹; mean daytime CO₂ level, 375 ppm. The error bars show the SEs of five plants. The color code links to the model components in Fig. 1.
substituted the literature values for the $J_{\text{max}}V_{\text{cmax}}$ ratio measured in the appropriate photoperiod conditions, assuming upper and lower limits (SI Appendix, Table S7). The simulations also replicated the relevant environmental conditions (SI Appendix, Section 3.11) (51).

Carbon assimilation and respiration rates were slightly underestimated (10.7% and 6% lower in the 12-h photoperiod, for example) on the simulated day corresponding to the day of measurement (Fig. 4A). The resulting net carbon fixation allowed the model to reproduce the full amount of starch accumulation by the end of the 12-h photoperiod (Fig. 4B), but starch levels were underestimated (by 10−26%) in shorter photoperiods (Fig. 4B). The model closely matched the starch levels remaining at the end of the night (Fig. 4B). However, in short photoperiods, the lower amount of starch accumulation in the light meant that the amount of starch mobilized per night was underestimated in the model. Additionally, part of the mobilized starch was used to maintain a higher sucrose level than observed in the data (SI Appendix, Fig. S5A), where sucrose levels decreased progressively as the photoperiod was shortened. These two factors resulted in lower growth per night in the model than in the data (Fig. 4C). The model more closely matched the observed growth increment in the 12-h photoperiod (Fig. 4C), where the simulated starch and sucrose levels matched observations (Fig. 4B and SI Appendix, Fig. S5A). Integrated over the life of the plant, the lower growth at night led the FM to underestimate total rosette biomass for short photoperiods (Fig. 4D). This result indicates that further parameters in addition to $J_{\text{max}}V_{\text{cmax}}$ are important for modeling growth, especially in the extreme 4-h photoperiod. In contrast, the FM accurately predicted the biomass in the 12-h photoperiod protocol, to within the experimental error (Fig. 4D). These results confirm that the FM can closely match the data from independent laboratories in the reference conditions, but the simple CDM did not fully account for the changing starch and sugar dynamics in short photoperiods.

Between a 4- and 12-h photoperiod, biomass increased strongly, and the relative growth rate (RGR, milligrams fresh weight produced per day per unit existing fresh weight) increased almost linearly with light fluence (51). This response would be expected if the conversion efficiency of carbon into biomass was constant. This linear relation between daily light fluence and growth was lost in long photoperiods. Whereas light fluence increased by 50% between the 12- and 18-h photoperiod, RGR increased by only 18% (51). Observed changes in the 18-h photoperiod included higher starch levels at dawn and a reduction in specific leaf area.
(i.e., increased leaf thickness) (51). Both of these are expected to reduce growth rates; incomplete starch mobilization will sequester carbon from growth, whereas increased leaf thickness will mean that less leaf area is generated per unit fixed carbon, which will decrease future light absorption and photosynthesis. Including the slower night-time starch breakdown (to 60% of initial starch rather than 84%) and measured 15–25% increase in leaf thickness in the FM, in addition to substituting $V_{\text{cmax}}$ for $V_{\text{cmax}}$, with the published value for the 14-h photoperiod, reproduced the observed biomass (Fig. 4D). This result was also recapitulated by extrapolating $J_{\text{max}}$: $V_{\text{cmax}}$ below the published value for 14-h photoperiods and reducing starch breakdown, but without considering the increase in leaf thickness (SI Appendix, Fig. S5B). Thus, these three factors are sufficient to account quantitatively for the altered growth rate under long photoperiods, although the balance among them remains to be determined experimentally.

**Model-Guided Understanding: Stochasticity and Tradeoffs in Development.** To explore the model’s potential, we extended the FM to include stochastic development at the organ (leaf) level, adopting a probabilistic organ initiation concept used for describing nonsymmetrical branching in plant architecture (52). Leaves are considered to appear at a regular interval (or growth cycle) with a simple, binomial probability that was estimated at 0.97 from our experimental data on Ler and Fei (SI Appendix, Section 3.12 and Fig. S6). Thus, leaves appear on most growth cycles but not all, reflecting variation in the processes of organ initiation and expansion. This stochastic model explained the variance of leaf number in our samples at every time point ($P > 0.05$; Fig. 3 C and F and SI Appendix, Table S9) while accounting for 11.3% (Ler) and 12.7% (Fei) of the variance of biomass measured at flowering time. The SDs in the timing of leaf appearance (phyllochron) from our simulations [in degree-days: 2.35 (Ler) and 1.86 (Fei)] were, however, lower than the SD in leaf initiation (plastochron) reported in Col (12.72; Discussion) (53). Besides interplant variation, both leaf initiation and leaf appearance rates increase with plant age in Arabidopsis (48, 53). The model reproduces this using a piecewise-linear rate, with a phase transition point at 355 degree-days, around half the vegetative period (Fig. 3 C and F and SI Appendix, Fig. S2). We explored the significance of this developmental timing, by simulating earlier or later transition points (SI Appendix, Fig. S7A). To distinguish the effect of the varying rate, we included controls that generated the same final leaf number at a constant rate. Model simulations with a transition point earlier than the reference, hence a longer interval of rapid leaf production, generated biomass as low as 46.4% of the reference value (Fig. 5A). Most leaves were small: median and third-quartile leaf areas fell to 32.6% and 33.5% of the reference value (Fig. 5C). The high leaf number and smaller size resulted in self-shading that reduced biomass. The varying leaf production rate generally resulted in a larger fraction of functional (photosynthesising) leaves at flowering time than in the controls (Fig. 5B) and, for transition points at 100–400 degree-days, in a greater proportion of large leaves (third quartile area above control; Fig. 5C) that partly escaped shading, resulting in higher biomass than in the controls (Fig. 5A). Simulations with a later transition point, hence a longer interval of slow leaf production, increased biomass (6% increase from transitions at 500–650 degree-days; Fig. 5A). The associated controls increased biomass up to 10.9%. The plant’s observed behavior, represented by the reference transition point, seemed suboptimal. However, the later transition points reduced the percentage of functional leaves at flowering from 88.9% to 81.8% (Fig. 5B). Median leaf area increased by 21.3% with a transition point at 600 degree-days, similar to total biomass, but a few leaves grew very large (third quartile area increased by 73% of the reference, but was only 78.6% of the maximum area; Fig. 5C). Thus, the higher biomass of these simulated plants depended on a smaller number of larger leaves. In contrast, near the reference transition point (300–400 degree-days), the third quartile leaf area was up to 93.8% of the
maximum size, indicating that the proportion of large leaves was high. Taken together, our analysis suggested that increasing the leaf production rate at midvegetative stage incurs a slightly lower total biomass, relative to a later transition point, but reduces the plant’s reliance on a few, large leaves.

Model-Guided Understanding of a Developmental Phenotype. The FM predicted how much rapid leaf production will reduce leaf size (Fig. 5C). This relationship has been described as a dual effect in plants overexpressing microRNA156 (Pro35S:MIR156), which have a short plastochron relative to WT plants (54). To test whether our model could reproduce the behavior of these developmentally altered plants and explain the dual effect, we grew Pro35S:MIR156 plants alongside Col WT for 37 d. Consistent with the previous study, Pro35S:MIR156 plants had a higher leaf production rate and smaller leaves compared with WT (Fig. 5D and SI Appendix, Fig. S7B). The size of the largest leaf in Pro35S:MIR156 was only 57% of that in WT (Fig. 5E). To test whether the leaf production rate alone was sufficient to explain this phenotype, we simulated the growth of the WT and Pro35S:MIR156 for 37 d, fixing the leaf production rate in the model to the measured rates in each genetic background.
notype. With only this change, our model not only replicated the observed size of the largest leaf to within the experimental error (Fig. 5E) but also closely matched the distribution of size ($R^2 = 0.90$; nRMSE = 12.9%) and biomass ($R^2 = 0.92$; nRMSE = 13.3%) for all of the individual leaves in the mutant, including their smaller size relative to the WT (Fig. 5F and SI Appendix, Table S10). As an additional test, we repeated the simulations with the model’s simpler, piecewise linear leaf production rate, using the default values for Col and refitting the piecewise function to the data for Pro35S: MIR156 (SI Appendix, Fig. S7 B and C). The model slightly underestimated leaf number in this experiment, causing an increase in the largest leaf size in both genotypes; nonetheless, the simulated mutant’s largest leaf reached only 65% of the WT value, within the experimental range (Fig. 5E). Our results indicated that the observed, higher leaf production rate in Pro35S: MIR156 plants was sufficient to predict the observed, smaller final size of each leaf, given the normal photosynthetic function and carbon partitioning among organs in the FM.

Discussion

We present methods, examples, and validation for one approach to developing a multiscale, whole-plant model of *A. thaliana*, inspired by crop science, by integrating existing models from different laboratories. The resulting FM closely matched data at multiple levels, acquired by two of our groups in different countries, allowing deeper analysis of experimental results and conceptual growth strategies. Our study suggests that a distributed, community-wide effort could successfully extend and refine the FM by integrating further, focused models into the larger framework.

Our approach stems from the recognition of potential synergies among diverse plant modelers (55), which encouraged us to integrate models from different domains using the modular approach. Ideally the integration process would not have altered the models at all, but this is unrealistic unless the models were originally designed for composition. In practice, unit conversions were required to make the models logically compatible and the FSPM was more substantially rewritten, as our aim was more limited than its original scope. Four redundant components were replaced by new connections. Only two parameter values were calibrated to our experimental data (discussed below). Another measurable parameter, the $\text{C}^{\text{max}}/\text{C}^{\text{e}}$ ratio that describes photosynthetic physiology, was modified using values from the literature for the 12-h photoperiod of our validation experiments. These changes were sufficient for the FM to match our experimental data (Fig. 2), confirming that the models were mutually compatible despite their different origins.

One general concern in mathematical modeling is overfitting, which becomes more significant in models of high complexity. This concern was part of our motivation to maintain the parameter values from the original models, which were already constrained to the most relevant data, instead of reoptimizing them to fit our data. In cases where unit conversions and scaling factors were required or in condition-specific scenarios, e.g., different photoperiods (see above), we adopted values directly from the literature. Although each of the four model components were calibrated and/or optimized with different techniques using independent datasets, the resulting FM matched our experimental data from two different laboratories. This broad predictive performance is generally not displayed in overfitted models.

We conducted a sensitivity analysis to examine the behavior of the FM. This analysis identified 7 of 18 photosynthetic parameters that are highly sensitive, although this number was likely underestimated because parameters related to ribulose-1,5-biphosphate carboxylase/oxygenase were redundant under our light-limiting conditions. In particular, the response of electron transport to temperature appears to have large effects on simulated biomass under our conditions. Indeed, the temperature response of key parameters in the Farquhar model has been the focus of other studies, with model accuracy decreasing when temperatures deviated from the 25 °C condition where the model was originally parameterized (56, 57). These studies proposed different temperature response functions to improve the estimation of photosynthetic parameters, and they can be readily incorporated into the FM in future. We also identified many parameters with large effects on the simulated flowering time and thus biomass at flowering. Our results are consistent with the analysis of many crop models, which revealed high uncertainties in yield predictions at elevated CO$_2$ and increasing temperature, partly due to these models’ simulated phenology and partly caused by the complex interactions between processes such as growth and leaf area (58).

Together, our work and that of others highlight the need for improved systems understanding and mathematical representation to predict plant behavior accurately, for example, in projected, future climates.

The norms of the Arabidopsis research community were obviously beneficial, as each model had independently used the standard, Col accession. Nonetheless, significant variability among laboratories was recently reported even in standardized Arabidopsis studies (29), so compatibility of the models was not assured. The FM accurately predicted CO$_2$ exchange at the population level, as well as biomass and area of both total and individual leaves at various time points during rosette growth, for plants of three accessions grown under 12-h photoperiods (Fig. 3 and SI Appendix, Table S8). Accurate biomass and area predictions depended on simulating the temperature and lighting regimes and the CO$_2$ levels of each experiment and required the joint operation of the CDM and FSPM (Fig. 1; see discussion of miR156). Five or fewer accession-specific parameters were modified based on our data to obtain these results out of a total 126 parameters. These revealed limited variation in water content [88%–92%, in agreement with a previous study (59)], which had only a small effect on the fresh biomass predictions. If calibration is necessary, water content is easily measured. Variation in seedling emergence was discovered (early in Fei), because Fei was selected for its increased leaf number in a previous study (48); its early emergence was sufficient to explain our data without altered leaf appearance rate (phytochron). Phytochron can also easily be determined through observation or automated imaging systems (60, 61) should calibration be required. Using time and CO$_2$ levels in laboratories and accessions is common, indeed the original PTM had four accession-specific parameters (1, 38). Until the sources of variation can be identified, therefore, the flowering threshold (at least) should be calibrated to each laboratory’s data, to test further regulation by the PPM (6) and PTM (1).

The FM also reproduced the measured biomass of plants grown in 12-h photoperiods under slightly different conditions (Fig. 4D), as part of a large, independent dataset testing multiple photoperiods (51). However, in shorter photoperiods, the model underestimated starch accumulation in the light and hence the rate of starch breakdown at night, as well as growth at night and total biomass under these conditions (Fig. 4). These discrepancies highlight how much the plant’s carbon metabolism adapts to different photoperiod conditions. The CDM assumes a fixed relation between photosynthesis and starch accumulation in the day, a fixed proportion of starch mobilization at night, and a fixed minimum sugar level. Although this changes the model’s absolute starch dynamics to some extent under different photoperiods, the responses measured in many plants are even more plastic (9). First, measured starch synthesis is faster in short than in long photoperiods, which contributed to the model’s underestimating the starch level at dusk in short photoperiods (Fig. 4B), consequently underestimating the rate of starch degradation to sugars at night. Sugar dynamics are also flexible; the measured sucrose level was lower at dawn than dusk and was lower at both times under short photoperiods compared with long photo-
periods (SI Appendix, Fig. S5A). Together, these effects caused the model to underestimate growth at night in short photoperiods (Fig. 4C). This error might be compounded, for example, if carbon conversion efficiency was underestimated in the model or maintenance costs were overestimated but these processes were not directly measured. Second, experimental data show that the assumption of almost complete (84%) starch mobilization at night is not always applicable, for example, in long photoperiods when growth is probably sink limited. Indeed, a basic problem of many models is that they assume only source limitation (13, 14). Reducing starch breakdown to the measured level, along with a further change in one (extrapolated $J_{\text{max}}V_{\text{max}}$ ratio) or two ($J_{\text{max}}V_{\text{max}}$ ratio and measured leaf thickness) parameters, matched the data (Fig. 4D). Among many possible extensions, the CDM might in future be supplemented with more detail on the plant's starch dynamics, carbon partitioning, and the relationship of sucrose to growth rate (62–65).

In addition to biochemical regulation, we illustrate the potential of the FM to understand the effect of developmental programs on growth and the final rosette form. First, we introduced stochastic leaf production that reproduced the varying leaf number observed in Arabidopsis rosettes (Fig. 3). However, this developmental variation accounted for rather little (~12%) of the observed variation in rosette biomass. Our simulations of phyllochron (time to leaf appearance) for Ler and Fei had lower SDs compared with the SD of the plastochron (time to leaf initiation) reported in Col (53). However, leaf initiation is a developmental process, whereas leaf appearance also involves growth: variation in growth might thus compensate for variation in development, reducing the observed variance in phyllochron. A field study of sorghum varieties also found a lower deviation in phyllochron compared with plastochron (66), although the two measures were tightly related.

Second, varying the age dependence of the phyllochron (Fig. 5) suggested a tradeoff in the developmentally regulated rate of leaf production and helped us to understand its origins. The measured leaf initiation rate was initially slow and then increased. Constantly rapid leaf production reduced the simulated biomass, because the many, small leaves quickly shaded each other. Conversely, constantly slow leaf production gave a slight advantage in simulated biomass but produced few leaves, many of which were older than in the WTs. Given the risks to leaf function from predation and other damage, this suggested that the plant's strategy maintains almost maximal biomass production, without relying on an aging leaf population. Compared with the biomass-maximizing, slow-production strategy, this developmental program distributes carbon investment (and thus leaf size) more widely, a feature characteristic of bet hedging strategies that could be tested in ecological studies (67, 68).

Third, our model reproduced the smaller leaf size phenotype of the developmentally misregulated Pro35S:MIR156 transgenic line by modifying only the model's leaf production rate (Fig. 5). Two possibilities were proposed to explain this dual effect of miR156 in the original study: (i) the existence of a compensatory mechanism whereby plastochron length and leaf size affect each other reciprocally to reduce changes to the overall plant biomass; or (ii) a common regulator that controls each of the two traits (54). The combined operation of the CDM and FSPM in the FM provides a parsimonious explanation for the dual effect. High leaf production rate requires carbon resources to be shared among more leaves (Fig. 2F), leading to a decrease in individual leaf growth. Using this mechanism alone, the FM matched the mutant leaf size distribution as accurately as it did the WT (Fig. 5F). Partitioning of a given amount of carbon among a larger number of leaves is a sufficient compensatory mechanism (54), although more complex models are of course possible. No common regulator is required to explain the observed relationship between leaf production and organ size. Similar quantitative analysis using the FM might contribute to further research on developmental regulators (such as those targeted by miR156) and sucrose signaling (69–71) to whole-plant phenotypes and might extend to applications that modulate organ size, for example, in pruning (72, 73).

Our results on miR156 again validated the FM, particularly the benefit derived by coupling the CDM and FSPM. The FSPM did not predict growth rate based on the measured experimental conditions but rather used model inversion to learn the light use efficiency from observed plant growth data. This aggregate parameter is not directly measurable, as it combines photosynthesis, sugar-starch partitioning, respiration, and the daily allowable growth rate, which are all separately represented in the CDM. The CDM predicted sugar production and partitioning to starch based on the experimental temperature, light:dark, and CO2 conditions but considered the rosette as one big leaf, whereas the FSPM provided information on the demand and growth of individual organs. We could only predict the biomass and detailed rosette structure in particular experimental conditions by combining these models in the FM. The FM not only explained the relationship of organ number and size in Pro35S:MIR156 plants but also predicts that the measurable parameters of carbon utilization are unaffected in this line.

In conclusion, quantitative dynamic models are valuable both to understand and to engineer organisal growth and physiological processes from the level of molecular and biochemical processes. The FM and the approach used to build it provide a flexible context to expand the detail and scope of component models, for example, to whole-cell models (28), and also to study the dynamic interactions among multiple processes. These resources will be particularly important to understand the pervasive effects of environmental stresses or pleiotropic biological regulators, such as the circadian clock. Finally, multiscale digital plant models might contribute to link systems biologists with ecophysiology and crop science, where significant synergies may be gained.

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