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Citation for published version:

Millar, AJ, Urquiza Garcia, J, Freeman, P, Hume, A, Plotkin, GD, Sorokina, O, Zardilis, A & Zielinski, T 2019, 'Practical steps to digital organism models, from laboratory model species to 'Crops *in silico*", *Journal* of Experimental Botany, vol. 70, no. 9, JEXBOT/2018/235879, pp. 2403–2418. https://doi.org/10.1093/jxb/ery435

Digital Object Identifier (DOI):

10.1093/jxb/ery435

Link:

Link to publication record in Edinburgh Research Explorer

Document Version:

Peer reviewed version

Published In:

Journal of Experimental Botany

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1 Practical steps to digital organism models, from laboratory

2 model species to 'Crops in silico'.

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- 6 Running title: Realising digital plant models
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27 Submitted 25 November 2018; Main text: 7700 words; 1 table; 5 figures.

- 29 **Keywords:** Arabidopsis thaliana; biochemical kinetics; community standards; computational
- 30 modelling; data science; whole cell modelling.

Highlight [<30 words]

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- 32 Combining models of biology across scales, for fundamental understanding and crop
- improvement, presents multiple challenges. We review practical experiences and promising
- 34 approaches in the pursuit of digital organism models.

Abstract [198 words]

- 36 A recent initiative named "Crops in silico" proposes that multi-scale models "have the potential
- 37 to fill in missing mechanistic details and generate new hypotheses to prioritize directed
- engineering efforts" in plant science, particularly directed to crop species. To that end, the group
- 39 called for "a paradigm shift in plant modelling, from largely isolated efforts to a connected
- 40 community" (Marshall-Colon et al., 2017). 'Wet' (experimental) research has been especially
- 41 productive in plant science, since the adoption of *Arabidopsis thaliana* as a laboratory model
- species allowed the emergence of an Arabidopsis research community. Parts of this community
- invested in 'dry' (theoretical) research, under the rubric of Systems Biology. Our past research
- combined concepts from systems biology and crop modelling (Chew et al., 2017; Chew et al.,
- 45 2014b). Here we outline the approaches that seem most relevant to connected, 'digital organism'
- initiatives. We illustrate the scale of experimental research required, by collecting the kinetic
- 47 parameter values that are required for a quantitative, dynamic model of a gene regulatory
- 48 network. By comparison to the SBML community, we note computational resources and
- 49 community structures that will help to realise the potential for plant systems biology to connect
- with a broader crop science community.

Introduction

- What distinguishes crop modellers from systems biologists, one of us was told ten years ago, is
- some responsibility to feed the world population. Systems Biology aims to understand the
- 54 interactions among the component parts of a living system and the emergent properties that arise
- from such interactions (Alberghina and Westerhoff, 2005; Kitano, 2002). Its aspiration was to
- include components across multiple scales from the molecular to at least the organism. In
- 57 practice the research started from intracellular pathways and only gradually intersected with
- 58 physiological, organism-level approaches; most often, the organism in mind was a human
- 59 (Kitano, 2015). Readers seeking to pin down systems biology, to a claim for novelty or
- otherwise, should consult earlier commentaries (Bothwell, 2006; Hammer et al., 2004; Marcum,
- 61 2008). The holistic, systems approach led to a meeting with mission-orientated research in crop
- science, though the whole-plant scale to which Systems Biology aspired was then at the lower

bound for crop models. The approach also distinguished Systems Biology from much research focusing on the properties of individual, biological components.

Along with the move from reductionism towards holism came a need for the 'dry' methods of formal modelling, because the unaided human brain is quite inept in reasoning quantitatively about dynamical systems as complex as those in biology. Several areas of plant science (cell physiology and ecology, to name but two) and crop science, have been 'amphibious' for decades, mixing 'wet' (experimental) and 'dry' (theoretical) approaches. The benefits of interfacing plant systems biology with crop modelling were recognised over a decade ago (GARNet Advisory Committee, 2006; Thomas, 2007), not only for modelling expertise but also for the real-world impacts. Crop models are regularly used by growers, breeders and Earth scientists, amongst others. Ten years later, an initiative named "Crops in silico" proposed that multi-scale models "have the potential to fill in missing mechanistic details and generate new hypotheses to prioritize directed engineering efforts" in plant science, particularly directed to crop species. To that end, the group (including A.J.M.) called for "a paradigm shift in plant modelling, from largely isolated efforts to a connected community" (Marshall-Colon et al., 2017; Zhu et al., 2016). However, formal models have been largely absent from the training of plant biologists, so this seemingly-natural interface has emerged only slowly. The diversity of models may also be less obvious for plant researchers, though it is arguably as great as the diversity of experimental methods. Crops in silico aims to link several, current approaches, such as functional-structural plant models that have organ-scale spatial resolution and process-based crop models with lower spatial resolution.

Dealing with diverse models is inevitable in the holistic agenda of Systems Biology. This article outlines some types of model that seem valuable for a community initiative such as "Crops *in silico*". Our experiences, tools and approaches to combine and use them arose particularly from joint work on the Framework Model for Arabidopsis growth (Chew *et al.*, 2017; Chew *et al.*, 2014b), which in part followed practices from crop modelling. *Arabidopsis thaliana* emerged as the laboratory model species for plant science, with an open research community (Ankeny and Leonelli, 2011; Leonelli, 2007), about fifteen years before Systems Biology emerged as a research field (Vermeulen, 2017). We illustrate results, resources and social organisation of Arabidopsis research that are benefitting plant Systems Biology, and could further contribute to and benefit from the interaction with crop science. The challenge is to ensure that actual researchers with particular skill sets are motivated and able to complete research in realistic time,

97 and to make the results comprehensible, useful and reproducible for others. We point to current,

omputational tools and resources that will help to realise this potential.

Standpoint

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100 The authors represent a spectrum of systems biology research, spanning plant science, molecular

biology, computer science, research management, software engineering and advanced

computation. We are linked by research in or associated with SynthSys, the centre for Synthetic

and Systems Biology at the University of Edinburgh, which has a long association with Systems

Biology (Bard, 2008) and with Science, Technology and Innovation Studies in social science

(Henry, 2008). A.J.M. previously coordinated GARNet, the UK community organisation for

Arabidopsis researchers (see Box 1) and contributed to the "Crops in silico" proposals.

The diversity of "models"

108 A biologist's "model" often describes the contemporary understanding of a biological process,

expressed in text, or as a diagram or cartoon (Figure 1A). Such descriptions are informal and

very useful as a distillation of biological knowledge, but they are fatally flexible, ultimately

ambiguous and difficult to reuse in a formal context. In contrast, mathematical models are formal

and unambiguous, inflexibly imposing a rigour of description that often exposes serious gaps in

biological knowledge. Identifying such gaps can be extremely valuable to direct ongoing work

but the gaps must be bridged with assumptions in order to complete a model.

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We summarise below some modelling approaches used in Systems Biology, based broadly upon

their explanatory ability. An explanatory model can illuminate the mechanisms of a biological

system and its principles of operation, whereas a descriptive model simply aims to predict the

behaviour of the system based upon its past behaviour, irrespective of the biological

mechanisms. Models in crop science and in systems biology each span this range. Models of

"Crops in silico" will usually combine several approaches, so more detailed classification is

difficult (Coveney and Fowler, 2005). Rather, we highlight opportunities for each model type in

building complex models in plant and crop science. Detailed spatial models of plant

development have been reviewed elsewhere (Ndour et al., 2017; Prusinkiewicz and Runions,

125 2012; Truskina and Vernoux, 2018). Despite omitting this area for brevity, we note that models

of cellular processes at the shoot apical meristem (Jonsson et al., 2006; Kierzkowski et al., 2012)

or in lateral root formation (Dyson et al., 2014; Xuan et al., 2016) have often combined multiple

model types.

Graphical models

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130 A useful, formal description of a biological process can start without equations or computer 131 programming, because a diagram can be formal (as can a text description). A defined vocabulary 132 of graphical symbols (glyphs) can represent the various types of biological components as nodes 133 in the diagram, with a defined set of connecting arcs to represent the processes by which the 134 components interact. Drawing such a diagram can reveal gaps in understanding and record the 135 assumptions made to bridge the gaps, as noted above. Maps of the metabolic network are a 136 familiar example but complex models need to represent much more than metabolism. The 137 Systems Biology Graphical Notation (SBGN) is a community standard for drawing intracellular 138 pathways (Le Novere et al., 2009), representing various types of molecules, their modifications, 139 complexes, compartments and so on. SBGN is supported by free software tools, such as 140 VANTED (Rohn et al., 2012) and Cytoscape (Goncalves et al., 2013). These can be extended to 141 support other notations, for example for plant structures. Several online repositories provide 142 SBGN diagrams of pathway information or models for download (Buchel et al., 2013; Naithani 143 et al., 2017). A diagram of this type can comprehensively represent the state of knowledge, as a 144 valuable addition to a review publication. A hand-curated diagram of mTOR response pathways 145 included 964 molecular components, for example (Caron et al., 2010) but such a large diagram is 146 difficult to read in practice. Moving from a diagram to a quantitative model requires additional 147 stoichiometry and parameter values, which can be added in graphical modelling software such as 148 Cell Designer (Funahashi et al., 2008) and Simile (Muetzelfeldt and Massheder, 2003). 149 150 For a diverse and growing community like "Crops in silico", investing in graphical models offers 151 three advantages. A non-modeller should be able to find, download and start to modify an 152 existing diagram to represent their process of interest within 30 minutes, without prior 153 preparation. This is the fastest route to modifying a model, similar in approach to the graphical 154 languages used to teach computer programming (Marji, 2014). An expert modeller could use 155 such a diagram as a starting point for detailed modelling of an unfamiliar process, similar to the 156 pseudo-code that is used to sketch software functions prior to full coding. For experts and non-157 experts alike, the diagrams also offer a human-readable format to orient themselves quickly 158 within a model.

Data-driven modelling

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High-throughput technologies such as automated phenotyping platforms capture information on many components of a system simultaneously. Analysis of high-throughput data involves modelling with statistical techniques such as clustering, principal component analysis (PCA) and

regression (Jagaman and Danuser, 2006). Similar methods can apply to the meta-analysis of data 163 164 curated from the literature (Poorter et al., 2012), with very broad scope (Diaz et al., 2016). These 165 data-driven methods can use little or no prior knowledge about the system and overlap with the 166 expanding range of machine learning approaches, such as neural networks (reviewed in Ma et 167 al., 2014). Data-driven methods are usually descriptive and can inform simple, mathematical 168 relationships that are used in many models where more detail is unavailable or undesirable. They 169 represent a relevant process concisely, in sufficient detail to lead to the formation of specific 170 hypotheses, for example about the mechanisms that underlie the differences between clusters 171 (Janes and Yaffe, 2006) or the connections among variables (Dalchau et al., 2011; Onoda et al., 172 2017). Thus advanced analysis by data-driven methods grades into conceptual modelling 173 (Valladares et al., 2014). In a spatial context, Mundermann et al. (Mundermann et al., 2005) 174 modelled the development of the Arabidopsis shoot in the L-studio software, using 175 measurements of architectural parameters to support detailed simulation and realistic 176 visualisation of plant growth (Figure 3). 177 178 The articles by Dalchau et al. and Mundermann et al. used data generated by the same labs that 179 conducted the modelling, which is common in small or emerging fields that use laborious assays. 180 In contrast, the work of Poorter and colleagues allows meta-analysis of many data sets from 181 well-established, eco-physiological assays (Poorter et al., 2010). The more data is required for a 182 modelling project, the more data availability can limit its progress and the career prospects of the 183 modellers. The Open Research movement, with its FAIR and Open data principles, deserves 184 their wholehearted support (see final section). 185 186 Baker et al. (2018) argue that data-driven methods' rapid focus on results may be more attractive 187 for research that is close to professional practice (clinical medicine in their case), whereas other 188 disciplines emphasise explanatory power. Several benefits can clearly follow from integrating 189 these approaches. Our work on the circadian clock encountered some practical difficulties in this 190 process. Data-driven approaches to learn the gene circuit structure were hampered by the very 191 non-linearity, time-dependency and density of interactions that had originally motivated us to 192 initiate modelling studies, remaining difficult even with a series of new methods (for example, 193 Aderhold A., 2013; Grzegorczyk et al., 2008; Higham and Husmeier, 2013). In contrast, data-194 driven connections of the clock to metabolism were published (Grzegorczyk et al., 2015) and 195 personnel had moved on, years before the follow-up experimental studies were complete (Flis et 196 al., 2015; Flis et al., 2018).

Qualitative modelling

Whereas data-driven models can represent detailed data with little explanatory power, qualitative models offer explanatory power with limited detail. Boolean models are the most common type, where components and connections are represented as present or absent, and this coarse state of the system may change over time. These models test hypotheses about the logical and causal relationship between events, stimuli and system responses (De Jong, 2002). An early example in plant science represented the network of transcription factors that specify organ identity during Arabidopsis flower development. The model's logical rules tested (and supported) the conceptual "ABC model" of gene interactions (Espinosa-Soto et al., 2004). Complex waveforms can be represented by allowing a time delay between the activation of one component and the next, yet the models remain attractively concise. A time-delay model (Figure 2) allowed us to test all possible connections among the genes of the Arabidopsis circadian clock (Akman et al., 2012), for example, highlighting a new circuit that explained the experimental data better than the circuit proposed at the time. This qualitative model's circuit was independently confirmed by new data and in a more detailed, quantitative model from our lab (Pokhilko et al., 2013). Note that we could not have tested all possible circuits in the quantitative model in a reasonable computation time.

For Crops *in silico*, Boolean models (and other qualitative models) might be the easiest way to incorporate large gene-regulatory networks. They do, however, risk discarding information for the best-studied components, which may have sufficient data for more detailed treatment. Hybrid models are then natural, where some components are represented in qualitative and others in quantitative form. For example, a binary representation of (unmeasured) transcriptional activation of a reporter gene allowed us to test several possible gene circuits in an algal clock, combined with a continuous, quantitative model for the levels of a luminescent reporter protein that reproduced experimental data (Ocone *et al.*, 2013). The software to support logic models is growing, exemplified by development of the Systems Biology Markup Language (SBML) "qual" standard for model exchange (see below)(Buchel *et al.*, 2013). Software tools can also help in converting qualitative models to quantitative forms (Wittmann *et al.*, 2009), which is not yet a common path (Ortiz-Gutierrez *et al.*, 2015) but might become a natural progression for Crops *in silico* as more data becomes available (Le Novere, 2015).

Constraint-based modelling

Even dynamic, biological systems can be treated as being in steady state, when their homeostatic

mechanisms buffer changes, at least substantially. The numbers of some molecule being

231 generated and degraded are equal, for example, so its level is almost constant in time. 232 Additionally, the time scale for metabolic events (seconds) is typically much faster than for 233 genetic regulation (hours): from the perspective of genetic regulation, the metabolic system is 234 always in steady state. The characteristics of this constant state depend on the structure of the 235 system (the related biochemical reactions and their stoichiometry), general thermodynamics laws 236 and external parameters, such as the cellular energy supply. Where a metabolic network is well 237 understood, for example, constraint-based analysis is able to identify a set of fluxes through the 238 network that are compatible with the observed steady state, to predict missing reactions and 239 alternative pathways, and to find steady states that become accessible under different conditions. 240 More prior knowledge is required than for qualitative models, and the models have greater 241 explanatory power. In the areas relevant to Crops in silico, De Reuille et al. used constraint-242 based modelling to create the geometry of the shoot apical meristem, subsequently using this 243 geometry as a constraint for auxin transport to evaluate the distribution of auxin fluxes (Reuille 244 et al., 2006). The approach can be extended to represent data that change over time, such as day 245 and night states of central carbon metabolism (Cheung et al., 2014) or the hourly dynamics of 246 the starch pathway (Sorokina et al., 2011). These extensions for dynamic systems are limited and 247 development is ongoing. They are attractive in principle for Crops in silico, because constraint-248 based models are computationally tractable and do not require the detailed kinetic parameters of 249 full, quantitative models.

Quantitative modelling

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251 Quantitative modelling techniques represent the most detailed explanation of the underlying 252 mechanisms and allow the most extensive numerical comparison of simulation results with 253 experimental data. Correspondingly, they require the most prior information on the system 254 (illustrated below). Where changes over time (dynamics) are of interest in the biology, for 255 example in the cell cycle or the circadian clock, these methods have given impressive results (Bujdoso and Davis, 2013; Novak and Tyson, 2008; Tyson and Novak, 2015). Systems of 256 257 ordinary differential equations (ODE) are a popular approach where time is continuous, as are 258 the equivalent, difference equations with discrete time steps. Each equation describes the change 259 in one variable (organ mass, protein concentration etc.) as a sum of reactions (synthesis, 260 destruction, transport etc.) that are represented with empirical, kinetic terms (law of mass action, 261 Michaelis-Menten approximation, piecewise-linear functions etc.). Variables can justifiably be 262 continuous, implying an infinite number of intermediate concentrations, if molecular numbers 263 are in fact large, reactions are frequent and the system behaves reproducibly. This style of 264 modelling is common in plant Systems Biology and has been reviewed elsewhere (Chew et al.,

265 2014a; Middleton et al., 2012). However, data at the single-cell level increasingly reveals 266 components that are present in small numbers (Libault et al., 2017), where the continuous, 267 deterministic approach is inaccurate and instead discrete, stochastic models describe the 268 probabilities of each reaction event (Shahrezaei and Swain, 2008). Stochastic models of the plant 269 clock circuit suggested that circadian timing would be variable at the single-cell level, for 270 example (Guerriero et al., 2012), as recently confirmed experimentally (Gould et al., 2018). 271 Multi-model frameworks like Crops in silico must therefore anticipate stochasticity at this micro-272 scale, in addition to the formation of discrete organs in a plant model, or germination of 273 individual weeds in a field model. 274 275 Multiple types of model are as natural in a digital organism as the many biological processes that 276 contribute to a physical organism (or the many research perspectives to understand it). 277 Integrating these diverse model types is by no means only a technical topic. In the example of 278 data-driven and quantitative modelling approaches to the circadian clock (above), flexible 279 management was required (Balmer et al., 2016) to reconcile the timelines of each modelling 280 approach and their different concepts of the "publishable unit" of research. New approaches to 281 research dissemination could be adopted in a Crops in silico community, as preprints, data 282 publications, model archive files, and institutional innovations such as "inside-out" libraries 283 (Bergmann et al., 2014; Dempsey, 2013; Leitner et al., 2016; Schloss, 2017) offer more 284 flexibility in what constitutes a "unit" for dissemination. We return to these social factors in the 285 context of community standards, below, and in the final section.

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Modelling frameworks and languages

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289 The technical challenge to link heterogeneous models is long-standing and well recognised 290 (Adam et al., 2012; Ghosh et al., 2011; Goldberg et al., 2018; Macklin et al., 2014; Marshall-291 Colon et al., 2017; Pradal et al., 2008). The approaches can be simplified to two extremes, either 292 to rewrite all the models in a common modelling language or to devise an integration system that 293 links the models in their diverse, native forms, as loosely-coupled "black boxes" (Figure 3). 294 Tightly woven into this problem is the distinction between declarative and procedural models. 295 Declarative models are a formal specification of the model, such as its mathematical definition. 296 Separate software is then required to simulate the model, leading to advantages described 297 elsewhere (Muetzelfeldt, 2007). If in addition a declarative model uses a standardised format,

298 then the model becomes easy to exchange between software tools (discussed in the following 299 section), and therefore easier to understand and modify. 300 301 In contrast, implementing the model in a programming language is procedural (or 'imperative'): 302 the model specification is also the computer code for simulation, whether it is in a scripting 303 language such as python or R, a high-level language such as Matlab, or a general-purpose 304 language such as C++. Good programming conventions can separate the declarative part of the 305 model but there is no guarantee of this. The code may then be executable but obscure, making 306 the model a black box. Modelling procedures are clearly important as well as the models. Open-307 source, well-documented code makes these more accessible than a closed-source or 308 undocumented modelling framework. The importance of open-source software for reproducible 309 research is discussed elsewhere (Mendes, 2018). 310 311 To illustrate these general considerations with a detailed example, we consider the development 312 of the Arabidopsis Framework Model from four previously-separate models (Chew et al., 313 2014b). Rewriting each of the constituent sub-models into a common language in the Simile 314 modelling environment, then re-validating them in numerical simulation, was a major effort 315 (Muetzelfeldt and Massheder, 2003). A preliminary project, PlaSMo, first collected likely 316 component models from idiosyncratic computing code (Davey et al., 2009). The refactoring 317 process depended on access to the model files. Files for one model had been deleted online and 318 were only available from the Google cache. The commercial, Simile environment was selected 319 for refactoring because it offered a rich, graphical interface and supported a declarative, XML 320 model format, SimileXMLv3 (see Box 1). Like SBML, this was based on the widely-used 321 MathML standard (Hucka et al., 2003). In practice, refactoring the various model codes required 322 unusually broad skills. As benefits of this investment, the component models in a web portal (see 323 Box 1) became more readily and uniformly accessible for future work, and the process of model 324 curation and re-validation provided stringent quality control. Among the challenges were IF ... 325 ELSE ... conditions: standard programming tools, which might distinguish parts of a model that 326 are used at different stages of plant development. These effectively, and very concisely, embed 327 multiple, alternative models within the same procedural code. Rewriting such models could 328 involve untangling a web of conditional statements, improving clarity but expanding the model 329 description. The Agricultural Model Exchange Initiative (Martre et al., 2018) are currently 330 embarking on a similar approach, with contemporary software tools (see Box 1). 331

332 The "black box" approach is initially faster, at least for a small number of models. The L-studio 333 framework, for example, can call external model codes (Figure 3), and the emerging Crops in 334 silico interface links models in four programming languages (see Box 1). More ambitious model 335 integration systems have been applied in projects (Marshall-Colon et al., 2017; Zhu et al., 2016) 336 such as the European agricultural assessment project SEAMLESS (van Ittersum et al., 2008). 337 The promise of this loose coupling is that modellers continue to develop their diverse, 338 component models independently, and yet can still interact with the ensemble. The practical risk 339 is that their unencumbered innovation flies beyond the reach of the integration system, so the 340 ensemble can no longer be simulated. More dangerously for the long term, a growing set of 341 'black box' models is harder for any individual to understand, frustrating the need for modellers 342 to refine and revise the component models. This seems to be an opportunity for biology to 343 inspire new computer science, for example using domain-specific languages that naturally 344 express the relevant biology (Honorato-Zimmer et al., 2017; Kniemeyer et al., 2007; Zardilis et 345 al., 2019) and meta-languages that integrate these models and control their simulation 346 (Mjolsness, 2018).

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Standards-based modelling for Crops in silico

349 If a growing number of plant modellers are to understand and use a wider range of model types, 350 investing in a standards-based approach can speed up the process. Systems Biology uses several 351 modelling standards, notably Systems Biology Markup Language (7) and Cell Markup Language 352 (CellML). SBML is a standard for constraint-based and quantitative models (Hucka *et al.*, 2018). 353 CellML adds support for various cellular interactions (Lloyd et al., 2004). These machine-354 readable, model exchange formats (Figure 1C) that have spurred investment in a mutually-355 reinforcing economy of online repositories and software tools that use the standard format as 356 input and/or output. For example, storing a private SBML model file in the self-service 357 FAIRDOM data repository (Wolstencroft et al., 2015) automatically allows simulation of the 358 model at the JWS-online resource (Snoep and Olivier, 2002). Complementary standards are 359 growing the economy. The Simulation Experiment Description Markup Language (SED-ML), 360 for example, describes how a particular SBML model simulation was run (Waltemath et al., 361 2011). Uploading a SED-ML file to an online resource can exactly reproduce a published 362 simulation figure. The file specifies how the resource should retrieve a model file from an online 363 repository, send it to an online simulator and plot the relevant part of the simulation results. This 364 level of transparency and replicability is a highly attractive product of the global SBML

economy (Mendes, 2018). Given these potential advantages, we considered how SBML would represent a plant growth model that might arise from Crops *in silico*.

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368 The plant growth use case highlighted three main issues for SBML: input weather data, 369 expressing some key concepts, and simulators for multi-models. First, systems biology models 370 usually reflect controlled, laboratory conditions. The Input Signal Step Function in SBML 371 represents step and cyclic experimental manipulations (Adams et al., 2012), for example, 372 motivated by the light-dark cycle in a plant growth chamber. Most crop models, in contrast, read 373 in timeseries of fluctuating weather data during the simulation. SBML does support custom-374 defined functions, including splines and piecewise-linear functions. These can represent input 375 timeseries data as new variables in the SBML model file, interpolating between timepoints to 376 make environmental data available at any point in the simulation. Simple SBML Data Tools 377 were therefore created to support such modification of SBML files, for crop and other models 378 (see Box 1). Secondly, core SBML cannot represent the creation of compartments during a 379 simulation, as required to model the formation of new plant organs. SBML development was 380 revised in 2010 to extend the core (Hucka et al., 2018) with specialised, modular packages, 381 which are proposed by the community ("qual" was noted above). Three packages were 382 particularly relevant for the Arabidopsis Framework Model, which would be representative for 383 many plant-level models: arrays, dynamic processes (the package known as "dyn") and 384 hierarchical model composition ("comp"), among a larger set that was discussed earlier 385 (Muetzelfeldt, 2010). Productive interaction with any such community effort needs some 386 understanding of the community norms. The packages are at varying stages of development 387 (SMBL community, 2017). SBML community rules focus their resources on the exchange of 388 models between software tools, where there is demand for the exchange and support for its 389 standardisation (Hucka et al., 2015; Schreiber et al., 2015). To be formally adopted, new SBML 390 packages must be implemented in two, independent software products. A potential drawback of 391 the modular approach is that, even if each of the three packages mentioned is fully developed in 392 SBML, there is no guarantee that any simulation software will support all three together. 393 Engaging with SBML models offers a bridge to Systems Biology but the sensible norm that 394 demand and software tools together lead the development of SBML standards, as noted above, 395 has a significant repercussion. Both demand and tools will initially be limited, when an initiative 396 such as Crops in silico aims to lead a field. Engagement with community standards might 397 therefore be a later step. Lastly, controlling disparate simulation timesteps and reconciling the 398 availability of shared resources among competing sub-models were considered at a workshop in 399 2015, which tested the representation of a landmark "whole cell" model (Karr et al., 2012) in a

400 standardised form (Waltemath et al., 2016). One option considered for modular, multipart models was a model-control system, using a standard akin to SED-ML. This approach might be 402 equally relevant to integrating diverse models for Crops in silico. However, the workshop report 403 coyly notes that "Significant effort will also be needed to develop an efficient, parallelized, 404 multi-algorithm simulator." (Waltemath et al., 2016).

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After a suitable modelling approach has been selected, the modellers must represent the biological processes of interest with enough detail to address the relevant issues. The question of "what's in the model" (specifying the model's variables) usually has many reasonable answers, which provoke debate rather than consternation. If the biological issues require a quantitative model, however, specifying the rates that are associated with each process (the values of the model's parameters) can be an overwhelming and contentious task. We next provide a specific example that illustrates this challenge.

Parameter values for a quantitative model

414 The 24-hour, circadian clock in Arabidopsis thaliana has been a paradigmatic system for studies of dynamic gene regulation over 20 years (Millar, 2016). Because timing was the critical, 415 416 biological issue, quantitative, dynamic models were a natural approach (Bujdoso and Davis, 417 2013). They operated with time in real hours and their success was judged on whether the 418 simulated waveforms of rhythmic gene expression helped to understand (explain and predict) the 419 experimental timeseries data, in various conditions. The RNAs and proteins of the dozen or so 420 clock genes were represented with arbitrary concentration units, in contrast to the real hours. 421 These models were built to understand results from molecular genetic assays, which often uses 422 relative or arbitrary units, rather than biochemical kinetics, where absolute units are more 423 common. Models in absolute units are advantageous, however (as outlined below). We therefore 424 summarise the parameter values that would be required to convert a model of a plant gene 425 regulatory network, such as P2011 (Pokhilko et al., 2012), to absolute concentration units. The 426 values described are listed in Table 1, extending similar resources of parameter estimates for 427 other organisms (Milo et al., 2010).

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Macromolecular synthesis and degradation

- 430 Most of the models deal with the birth and death of the clock gene RNAs and proteins. However,
- 431 absolute RNA transcription rates have not been measured in plants. Sidaway-Lee et al.
- 432 (Sidaway-Lee et al., 2014) measured the distribution of nucleotide incorporation rates in

433 Arabidopsis and their temperature-dependence. The results were reported in microarray 434 fluorescence units per hour. We are therefore limited to estimating a maximum transcription rate 435 for eukaryotes in general, from a maximum RNA polymerase II elongation rate of 5 kbp/minute 436 in human cell lines (Danko et al., 2013) and 4.5 kpb/min in zebrafish (Hanisch et al., 2013), and 437 occupancy of typically one RNA polymerase complex per gene (Zenklusen et al., 2008). Maximal transcription rate is then 2min⁻¹ for a 2.5kb RNA, for example, ignoring short-term 438 439 transcriptional bursting (Harper et al., 2011). RNA degradation rates have been measured in 440 large-scale studies (Narsai et al., 2007; Sidaway-Lee et al., 2014), either after transcriptional 441 inhibition or by inference from the nucleotide incorporation data. Mean RNA half-life was 5.9h 442 in plant cell cultures at 22°C (Narsai et al., 2007), or 1.9h (at 27°C) to 5.0h in plants (17°C, 443 Sidaway-Lee et al., 2014). The microarray readout signals were less reliable for rare and 444 unstable RNAs, however, and RNAs with daily rhythms must be unstable. Specific analyses of 445 clock-relevant RNAs are therefore important, again using inhibitors (Lidder et al., 2005) or by 446 inference from statistical timeseries models without inhibition (Finkenstadt et al., 2008). Note 447 that the inhibitors could give paradoxical results (Finkenstadt et al., 2008): if the degradation of a 448 target RNA is regulated by an RNA mediator that is itself unstable, then rapid depletion of the 449 mediator during a transcriptional block may stabilize the target RNA. 450 451 Protein translation rates were measured by Piques et al. (Piques et al., 2009) for a set of 452 metabolic-related genes in Arabidopsis, using calibrated qRT-PCR assays to measure the 453 absolute number of transcripts in free RNA or bound to ribosomes. The fraction of transcripts 454 engaged in translation can be calculated, yielding a range of 0.56-0.9, mean 0.77. A ribosome 455 translation velocity of 3 amino acids/second and density of 6.6 ribosomes/kb of coding sequence 456 (CDS), based on data from bacteria (Brandt et al., 2009) were then used to estimate protein 457 synthesis rates (mol protein g⁻¹FWh⁻¹) and their increase in the light compared to the dark period 458 (Ishihara et al., 2015; Piques et al., 2009). Protein degradation rates have been measured in large 459 studies following metabolic labelling (Li et al., 2017), though the mass spectrometry methods 460 involved are biased towards abundant and therefore often stable proteins and the dynamics of 461 amino acid pools introduce further limitations (Ishihara et al., 2015). The median half-life of 6 462 days (Li et al., 2017) clearly does not represent the clock regulators with high-amplitude, daily 463 rhythms. However, constraints on the possible protein degradation rates can be estimated from 464 the available timeseries data, where the clock protein has been detected as a tagged fusion 465 protein or with antibodies to the native protein (for example, Knowles et al., 2008; Nakamichi et 466 al., 2010).

467 **Volume and transport** 468 Given these synthesis and degradation rates, various models can estimate molecular copy 469 number per cell. The next critical values are the volumes of the relevant cellular compartments, 470 to convert copy number estimates to concentrations. Koffler et al. (Koffler et al., 2013) 471 quantified the volumes of A. thaliana mesophyll cells in young and old leaves, reporting each 472 compartment as a fraction of total cellular volume. For example, the mean volume occupied by 473 the nucleus was 0.16% of the cell volume in an older leaf. Wuyts et al. (Wuyts et al., 2010) 474 report the distribution of volumes for palisade mesophyll cells, with a mean cell volume of 475 73,000µm³. Combining these gives a nuclear volume of 117µm³. This is reassuringly close to an estimate of $113\mu m^3$ that we calculate from the nuclear diameter of $5.99 \pm 0.72\mu m$ measured by 476 3D-FISH (Tirichine et al., 2009), assuming a spherical nucleus. 477 478 479 Finally, model components must be transported among cellular compartments; in our case the 480 nucleus is particularly relevant. No data is present for the size, number or distribution of A. 481 thaliana nuclear pore complexes (NPCs), the route for such transport. Data on tobacco BY-2 cell cultures showed around 50 NPCs per µm² of nuclear envelope (Fiserova et al., 2009). 482 483 Furthermore, in human cultured HeLa cells the transport rates of NTF2 and Transportin are 170 484 and 140 molecules/s/NPC respectively (Kubitscheck et al., 2005). If we assume that similar 485 transport rates are achievable in A. thaliana, using the nuclear diameter above suggests possible 486 transport rates up to 960,000 molecules/s into the nucleus. These are unlikely to affect dynamics 487 on a circadian timescale of multiple hours, unless nuclear transport is specifically regulated. 488 Binding affinity 489 Clock proteins function in the model by interacting either with each other or with the DNA in a 490 clock gene's promoter. The affinity (K_d) of each interaction affects the model's behavior but 491 almost none of the specific values have been measured. General (Kastritis et al., 2011; Kumar 492 and Gromiha, 2006) or more specific (Stiffler et al., 2007) databases describe protein-protein 493 interactions in other species. Wide variation in even the median K_d (233nM, 12nM and 14 μ M, 494 respectively) in part reflects the inclusion of protein classes such as high-affinity antibodies, 495 emphasizing the importance of more targeted resources. A sample of 42 published DNA-protein 496 affinities for plant DNA-binding proteins gives median K_d of 20nM (Figure 4A) (Aggarwal et al., 497 2010; Hao et al., 1998; Hofr et al., 2009; Izawa et al., 1993; Liang et al., 2008; Moyroud et al., 498 2009; O'Neill et al., 2011; Prouse and Campbell, 2013; Reymond et al., 2012). A similar

collection of plant protein-protein interactions (n=45) suggested a median K_d of 86nM (Figure

4B) (Ballut et al., 2005; Bauer et al., 2013; Bernal-Bayard et al., 2014; Bisson and Groth, 2010;

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501 Dong et al., 2010; Fuglsang et al., 2003; Hao et al., 2011; Levskaya et al., 2009; Li et al., 1999;

Liu et al., 2007; Luoni et al., 2006; Mantovani et al., 2014; Ogawa et al., 2008).

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Means and ends of detailed models with absolute parameterisation

504 One advantage of a model species such as Arabidopsis is the concentration of research effort, 505 resulting in measured values for parameter such as the nuclear volume (above). Nonetheless, 506 building a quantitative model of a plant gene regulatory network such as the P2011 clock model 507 seems to demand more parameter values than have been measured. Parameter fitting is one 508 means to overcome the incomplete parameter measurement, and was used extensively to 509 construct past clock models (Bujdoso and Davis, 2013). Rather than being constrained by input 510 parameters alone, the model outputs were constrained to match functional data, in this case the 511 detailed waveforms of rhythmic timeseries. The data in Fig. 1D would help to constrain the clock 512 model, for example. Timeseries data have been published by many research groups for tens of 513 light-dark conditions and clock-mutant plants. Each timeseries typically has 10-100 data points. 514 Public, reference data sets are available (Flis et al., 2015), only for Arabidopsis, to ease the 515 burden of data collation (Fogelmark and Troein, 2014). Mathematical analysis suggests that the 516 clock might be particularly tractable to parameter fitting, because the interlocked, negative-517 feedback loops of gene regulation constrain the system's dynamic behaviour (Rand et al., 2006). 518 Regulatory networks of this form have much less flexible behaviour than a modeler might expect 519 to gain from the many parameters, so correspondingly fewer sets of parameter values can 520 produce model outputs that match the timeseries data. Indeed, detailed measurements in 521 Arabidopsis have subsequently validated some of the fitted parameter estimates of clock models 522 (Pudasaini et al., 2017), suggesting that more such measurements could further validate the 523 approach. 524 525 Model development still required searching a high-dimensional space (several 10's of 526 parameters) to discover sets of parameter values that were consistent with the data, which is 527 computationally demanding. We have shown that open data, free software (Alves et al., 2006) 528 and public computational resources can make this process accessible (Flis et al., 2015) but 529 experts in advanced computation will remain important contributors to Crops in silico. Absolute 530 parameter estimates (above) are valuable here too, in limiting the range of values that the search 531 algorithms must explore, speeding the parameter search. Moreover, qRT-PCR assays calibrated 532 to absolute RNA copy numbers are now providing the first gene expression timeseries data that 533 naturally match the simulation outputs from models with absolute parameter values (Baudry et 534 al., 2010; Flis et al., 2015; Piques et al., 2009).

Modelling with absolute biochemical units should benefit our understanding of the clock, judging by earlier examples in biology. We should discover whether the models' arbitrary units concealed some processes that required unusual or impossible parameter values, suggesting that the plant uses a different biochemical mechanism to achieve that aspect of its circadian timing. Unrelated studies (including high-throughput surveys) will more easily test parts of the model, by measuring a relevant biochemical parameter value or the level of a model component, compared to the model's predicted value (as noted above, Pudasaini et al., 2017). The most important benefit may come not in fundamental understanding but in engineering. The models in absolute units should better represent particular manipulations, such as altering the K_d for a particular clock protein binding to a particular promoter. This is the level of understanding that the Crops in silico initiative and others propose for some key processes in crop growth, in order to apply molecular genetic tools most powerfully to crop improvement (Zhu et al., 2016). Detailed models will be required to design interventions in those processes, such as the comprehensive, OnGuard stomatal physiology model (Hills et al., 2012) or the ePhotosynthesis model (Zhu et al., 2013). The biochemical and biophysical parameter values in ePhotosynthesis derive from many species but none is from Arabidopsis. In part, this reflects the technical challenges that a very small plant presents for photosynthesis research (Stitt et al., 2010). However, the (excellent) researcher who most directly measured parameter values for our clock models rated that as their most boring work ever, hinting at the social factors that also shape research. Process and Pizzazz for a digital plant community Crops in silico aims to link discovery science that is far from agricultural production, with crop

Crops *in silico* aims to link discovery science that is far from agricultural production, with crop models that are closely linked to practice (Figure 5). Such different research areas bring distinct types of social organisation, as Vermeulen pointed out in another context: "In (post-)genomics research understanding is geared towards innovation, which requires higher levels of integration [among research groups], while ecology research is primarily oriented towards understanding nature and environmental change, allowing more decoupled forms of organisation. This different orientation of molecular biology and ecology also causes a difference in financial resources for collaboration, as the goal of improving human health attracts more research funding than increased understanding of basic environmental processes." (Vermeulen *et al.*, 2013). The Crops *in silico* initiative foresees a substantial effort in social organisation, drawing from examples including SBML, the Physiome and "virtual organism" initiatives such as the Virtual Rat or

569 Virtual Physiological Human (Marshall-Colon et al., 2017). These networked, interdisciplinary 570 research organisations are an active domain for social science research, which is generating 571 results and concepts that seem relevant for practitioners (Freeman and Millar, 2017). The 572 "Community of Practice", for example, links members who share a common goal across the 573 boundaries of previously-separate fields: Crops in silico seeks to establish such a community. 574 One challenge is to attract members. The relative youth of the Arabidopsis field might offer 575 some advantage here, in providing new members to an emerging plant modelling community 576 (see final section). 577 The promise and challenge of shared resources 578 "Boundary Organisations" can also support the emerging community, particularly if they 579 manage "Boundary Objects" (Star and Griesemer, 1989). These can be physical: the high-580 throughput plant phenotyping facilities and the EMPHASIS network that coordinates them in the 581 EU form one example (Roy et al., 2017). The Biomodels repository of models (Glont et al., 582 2018) is such an Object from the Systems Biology community, and its original focus was on 583 models in SBML format. Biomodels addresses a practical need specific to that community, 584 attracts investments from different constituencies (models from biologists and software tools 585 from computer science) and thereby creates a form of shared, social capital. Plant science is not, 586 however, a major component: 38 models include Arabidopsis components or literature, of a total 587 1649 published models (in mid-2018); 2 models include maize references; 0 for wheat or barley. 588 Biomodels policy is now to accept models in any format, increasing its relevance for crop 589 models. It seems relevant that Biomodels is hosted by the European Bioinformatics Institute 590 (EBI), itself part of the inter-governmental, treaty organisation EMBL (established 1974). One or 591 more anchor institutions with stable mission and funding will be extremely beneficial for the 592 risky, long-term development of complex plant models and their associated communities. 593 594 Crops in silico must link very diverse data with the diverse models, so resources to manage the 595 data might form another, helpful, Boundary Object. Alongside the experimental phenotyping 596 facilities mentioned above, data resources have been developed to manage and share large-scale 597 plant phenotyping data (Neveu et al., 2018). The Agricultural Models Intercomparison and 598 Improvement Project (AgMIP) has worked to assemble benchmark data as well as crop models, 599 for example (Asseng et al., 2013; Rosenzweig et al., 2013). Systems biology models, in contrast, 600 are too rarely benchmarked: open, community-based benchmarking would help to give credit for 601 model improvements. However, many of the data that we need are acquired at the single-project 602 scale (as in Table 1), where data sharing is still not routine.

603 604 The Open Research movement (The Royal Society, 2012) promotes sharing of data (Open Data), 605 as well as publications (Open Access), software (Open Source) and in some cases, even lab 606 notebooks (Open Notebook Science). "Data" is very broadly conceived, including protocols, 607 analysis or visualisation scripts, and models, as well as experimental data. The principles of 608 FAIR data are more recent but equally important for Crops in silico, as they promote data that 609 are Findable, Accessible, Interoperable and Re-usable (Wilkinson et al., 2016). FAIR data need 610 not be Open, but if access is granted then they should be easier to use. In contrast, Open data that 611 is not FAIR might be unusable. FAIR is therefore being proposed as a guiding principle for 612 international initiatives such as the European Open Science Cloud and the US NIH Data 613 Commons (see Box 1). 614 615 To get FAIR data beyond the principles and into common research practice, we need easy-to-use 616 software tools and resources. Resources to manage the "long-tail" data (Ferguson et al., 2014) 617 that is required for detailed modelling can in theory be "explicitly created to meet the 618 researchers' needs, support extensive curation, and embody a heightened awareness of what it 619 takes to make data re-useable by others" (Leonelli et al., 2013). Although this is clearly 620 desirable, few biology groups have such data management resources, or the software skills to 621 customise them for their needs, or much appetite to add data curation to their overloaded 622 schedules. The data curated in Table 1, for example, were assembled only because they were 623 required for a specific research project. The software that might underpin such resources is 624 fragmented (Kwok, 2018), except where research funders have coordinated internationally as in 625 the AgMIP and FAIRDOM projects (see Box 1) (Rosenzweig et al., 2013; Wolstencroft et al., 626 2017). Coordination among funders, including direct funding for data curation, will be essential 627 to get beyond pilot, example models and create a broadly-based digital organism framework that 628 is regularly updated and refined with new information, in turn supporting the careers of a new 629 generation of modellers. **Conclusion** 630 631 No one should be surprised that such major research problems are relatively neglected, if 632 funders, researchers and their institutions recognise and reward individual lab heads catching 633 transient, project awards, like superhero characters in a video game. We have argued that 634 projects should be valued, rather than individuals (Freeman and Millar, 2017). This requires the 635 intellectual platform, capability and leadership to manage such projects, which is itself an area

for rich debate (Mazzucato, 2014; Rip, 2000; Weber et al., 2016). Large projects in this area

require international, community-wide effort but this does not imply that they should be 637 638 monolithic. Rather they need particular infrastructure, with funding mechanisms suited to 639 infrastructure, to integrate the results from distributed projects that might be independently 640 funded. 641 642 This article focussed on the need for digital organism initiatives to create and integrate a network 643 of diverse models, and practical steps towards integration (summarised in Figure 5). Model 644 diversity will always be with us, due to the variety of biological, chemical and physical processes 645 involved, the uneven states of knowledge, mathematical and computational tools, and the 646 differing aims of model users. Digital organism initiatives recognise both the model integration 647 tasks and the parallel challenge of managing diverse data. We touched on the technical 648 infrastructure that is required but community structures and community dynamics also contribute 649 to the operation and governance of such research networks (Freeman and Millar, 2017). Social 650 infrastructure therefore has a key role and might require parallel, infrastructural funding, which 651 will change over time. Community organisation might initially focus on understanding and 652 testing pilot model integrations, for example, whereas standardisation might be a later stage, as 653 we noted in the case of SBML. 654 655 In a landscape of this complexity, engaging multiple research and stakeholder communities, 656 projects like Crops in silico will be demanding of their leadership. The social sciences may 657 contribute useful strategies (Balmer et al., 2016) but these do little to mitigate the risks for junior 658 faculty, until concerns over lower funding and recognition for interdisciplinary research are 659 resolved (Bromham et al., 2016; Rafols et al., 2012; Yegros-Yegros et al., 2015). We might 660 rather harness the motivation of our youngest researchers. The success of the student-led 661 International Genetically Engineered Machines competition (iGEM) brought a definite buzz to 662 Synthetic Biology (Matheson, 2017), by giving them tools, keeping an open competition, and 663 making it fun. 664 **Acknowledgements** 665 666 We gratefully acknowledge discussion with Prof. Vincent Danos (Informatics, University of 667 Edinburgh) and Dr. Robert Muetzelfeldt (Simulistics Ltd., Edinburgh), Dr. Paul. E Brown 668 (University of Warwick) for the simulations in Figure 3, along with funding from the UK

Biotechnology and Biological Sciences Research Council (awards C006658, D019621, F010583,

- 670 F005237, M017605, M018040) and the European Commission (FP7 TiMet, award 245143), and
- research studentships from BBSRC (iCASE K011294) and CONACYT (Mexico).

Text Box 1

Box 1: Online Resources and Software

- Agricultural Models Exchange Initiative (AMEI), repository of models and resources for model exchange in CropML, by Pierre Martre, Christophe Pradal *et al*. https://github.com/AgriculturalModelExchangeInitiative.
- Agricultural Models Intercomparison and Improvement Project (AgMIP), international programme of data format interconversion and model comparison for crop models, http://www.agmip.org.
- cis_interface, software tools to link "black box" models, by Meagan Lang (National Centre for Supercomputing Applications, Illinois, USA) https://github.com/cropsinsilico/cis_interface.
- European Open Science Cloud, high-level initiative in Open Research that includes FAIR data principles, https://ec.europa.eu/research/openscience/.
- FAIRDOM, international project developing software for "long-tail" research data management and advocating Open and FAIR data, https://fair-dom.org.
- FAIRDOMHub, instance of FAIRDOM software providing a self-service commons for public or private data, models and protocols, https://fairdomhub.org.
- GARNet (previously the Genomics Arabidopsis Research Network), organization representing the UK Arabidopsis research community; several relevant reports online: http://www.garnetcommunity.org.uk.
- NIH Data Commons, pilot project (2017-2020) including FAIR data principles, https://commonfund.nih.gov/commons.
- Plant Systems Modelling (PlaSMo), repository of plant growth models in several formats, https://www.plasmo.ed.ac.uk; now migrated to the FAIRDOMHub commons.
- SBMLDataTools, software tools to add external timeseries data as a function in an SBML model, by Alastair Hume (EPCC, Edinburgh, UK). https://github.com/allyhume/SBMLDataTools.
- SimileXMLv3, XML schema for Simile models, with a model conversion tool. http://www.simulistics.com/book/similexml/simile-markup-languages/similexmlv3 [the PlaSMo project presented a dozen models, refactored into this standard; Simile software support had lapsed at the time of writing].

Table 1. Parameter values for detailed modelling were collated from the literature.

¹ PMID, PubMed identifier of the publication.

				_	Publication reference				
Component	Process	Sample	Value	units	PMID ¹	First Author	Year	Data display	Comments .
Cytosol	Volume	A. thaliana leaf	4.1	% of cell volume	23265941	Koffler BE	2013	Table 1	
Mitochondria	Volume	A. thaliana leaf	0.47	% of cell volume	23265941	Koffler BE	2013	Table 1	
Chloroplasts	Volume	A. thaliana leaf	15.63	% of cell volume	23265941	Koffler BE	2013	Table 1	
Nucleus	Volume	A. thaliana leaf	0.16	% of cell volume	23265941	Koffler BE	2013	Table 1	
Peroxisomes	Volume	A. thaliana leaf	0.14	% of cell volume	23265941	Koffler BE	2013	Table 1	
Vacuole	Volume	A. thaliana leaf	79.19	% of cell volume	23265941	Koffler BE	2013	Table 1	
Nucleus	Diameter	A. thaliana leaf	5.99	μm	19650905	Tirichine L	2009		
Cell	Volume	A. thaliana leaf	73000	μm^3	20598116	Wuyts N	2010	Fig. 8, left bottom	Mean value for palisade mesophyll cells. Reported range is 2-30
Gene transcription	transcription rate	Yeast	2 - 30	mRNA/hour	21103382	Pelechano V	2010	Abstract	mRNA/hour.
RNA Polymerase II	density on DNA	Yeast	0.078	Pol II molecules/kb	21103382	Pelechano V	2010		
RNA Polymerase II	density on DNA	Yeast	2	pol II/gene	19011635	Zenklusen D	2008		
RNA Polymerase II	elongation rate	Yeast	0.56	kb/min	24103494	Miguel A	2013	Fig. 1A	21°C
RNA Polymerase II	elongation rate	Mammalian cells	4	kb/min	21264352	Brody	2011		
RNA Polymerase II	elongation rate	Zebrafish	4.8	kb/min	23250218	Hanisch A	2013	Abstract	Measured at 28.5 °C.
Ribosome density Nuclear Pore Complex	Translation density on nuclear	E. coli	11 ± 2	ribosomes/RNA	19167328	Brandt F	2009	Fig. 2G	In polysomes translating firefly Luciferase
(NPC)	envelope density on nuclear	lymphocytes	2 - 4	NPCs/µm²	19392704	Fiserova	2009		
NPC	envelope	Mature Xenopus oocytes	60	NPCs/µm²	19392704	Fiserova	2009		40 E0 for 2 day old calls
NPC	density on nuclear envelope Nuclear	Tobacco cell cultures	50	NPC/µm²	19392704	Fiserova	2009		40-50 for 3-day-old cells; 50 for 10-day-old cells.
Transportin protein	translocation rate Nuclear	Mammalian (HeLa) cells	140	molecules/s/NPC	15657394	Kubitscheck U.	2005		
NTF2 protein Nucleoplasmin core	translocation rate Nuclear	Mammalian (HeLa) cells	170	molecules/s/NPC	15657394	Kubitscheck U.	2006		
domain fusion protein	translocation rate	Mammalian (HeLa) cells	17	MDal/s/NPC	11250898	Ribbeck K.	2001		

Figure legends

Fig. 1. A model can usefully be represented in several forms.

3 4 (A 5 in 6 (S 7 h 8 p

(A) A simple model of the circadian clock gene circuit (Locke et~al., 2005) is shown as an informal diagram, linking four genes (helices) via their proteins (ovals), with inputs from light (sun). (B) The differential equation for changes in cytosolic LHY protein (cL_c) in the model is human-readable (and declarative). This equation involves LHY mRNA (cL_m), a translation rate parameter (p_1), RNA degradation rate parameters (m_2 , k_2), and translocation of nuclear LHY protein (cL_n) with rates r_1 , r_2 . (C) A fragment of SBML represents the equation with the same names but is now machine-readable. The first line provides a stable reference to interpret its MathML format. (D) Timeseries simulation of the SBML model in suitable software provided a model output for the RNA level of gene Y (Y fit; red, open symbols; timepoints selected to match data), for comparison to RNA data acquired for a candidate gene in Arabidopsis (GI data, filled symbols). After a dark night (-12h to 0h), dawn light transiently induces both the hypothetical Y and candidate gene GI; the simulation continues in constant light. The comparison of model to data leads to future model refinement (dashed arrow) in the iterative

Fig. 2. The simple, qualitative form of a model can retain key behaviours.

cycle of systems biology. Adapted from (Locke et al., 2005).

(A) Simulation outputs show RNA levels changing continuously, from the simple clock model (Locke *et al.*, 2005) in quantitative form (differential equations, as in Figure 1B). (B) RNAs are either expressed (1) or not (0) in the qualitative form of the same model (Akman *et al.*, 2012). The binary, time-delay model still shows bimodal peaks of RNA expression from gene *Y* (green), with light induction after dawn (as in Figs. 1D, 2A). Levels are slightly offset for clarity in (B). Time 0h is midnight. Open box, light interval; filled box, dark interval.

Fig. 3. New capabilities arise from a "black-box" combination of models.

The circadian clock model shown in Figure 1 (Locke *et al.*, 2005) can communicate to the Arabidopsis architectural model (Mundermann *et al.*, 2005) running in L-studio software. A version of the clock model in Matlab software was automatically compiled into the C programming language (creating a 'black box'), in order to interact as a black box with the lpfg programme of L-studio. TOC1 protein level from the clock model controlled a leaf angle parameter in the architectural model, creating a simple simulation of rhythmic leaf movement in Arabidopsis over day/night cycle. The clock model's light:dark setting also darkened plant colour at night (16h, 20h). Image generated by Paul E. Brown and A.J. Millar.

Fig. 4. Published parameter values can inform detailed modelling.

(A) Distribution of published K_d values for plant protein-protein interaction affinities. (B) Distribution of published K_d values for plant protein-protein interaction affinities. In the (many) cases where an interaction of interest has not been measured directly, data such as these help to constrain the range of parameter values that computational, parameter-fitting procedures should explore. Publication references are listed in the main text.

Fig. 5. Linking Systems Biology with Crop Science models.

The solid line links the concepts of biology, first from genome sequence *via* genotype, biochemical parameters and molecular regulation to whole-organism phenotype in a particular environment (yellow area); then from phenotypes to field traits and adaptation or to yield under

particular management (green area); finally, given genetic variation, through natural selection or artificial selection in crop breeding, to the evolution of genome sequences (adapted from Millar, 2016). Initiatives like Crops *in silico* will deal with the whole cycle, by linking several models (coloured arcs) into a seamless, causal chain. The top line of graphics locate the topics considered in the main text with reference to this cycle. The arcs suggest current types of model, in systems biology (indigo), crop science (cyan) and evolution (dark blue). The dimensions that are often considered in such models are capitalized (G, P, E, M). Underpinning infrastructures (grey) help to bridge these disciplines. 'Anchor' institutions are shown (buildings), which might provide major experimental facilities, digital infrastructure or a focus for social infrastructure, such as training or standardisation workshops.

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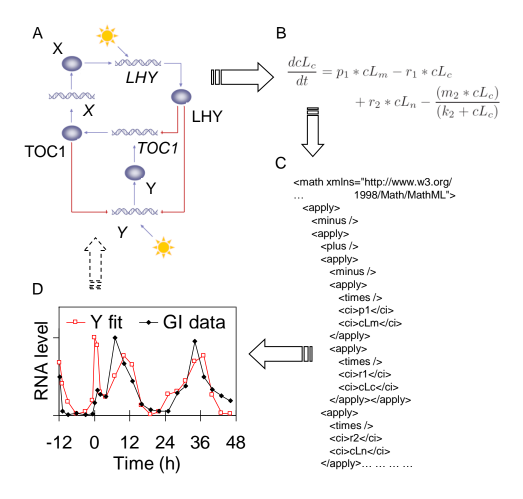


Fig. 1. A model can usefully be represented in several forms.

(A) A simple model of the circadian clock gene circuit {Locke, 2005} is shown as an informal diagram, linking four genes (helices) via their proteins (ovals), with inputs from light (sun). (B) The differential equation for changes in cytosolic LHY protein (cL_c) in the model is human-readable (and declarative). This equation also involves LHY mRNA (cL_m), a translation rate parameter (p_1), RNA degradation rate parameters (m_2, k_2) , and translocation of nuclear LHY protein (cL_n) with rates r_1 , r_2 . (C) A fragment of SBML represents the equation with the same names but is now machine-readable. The first line provides a stable reference to interpret its MathML format. (D) Timeseries simulation of the SBML model in suitable software provided a model output for the RNA level of gene Y (Y fit; red, open symbols; timepoints selected to match data), for comparison to RNA data acquired for a candidate gene in Arabidopsis (GI data, filled symbols). After a dark night (-12h to 0h), dawn light transiently induces both the hypothetical Y and candidate gene GI; the simulation continues in constant light. The comparison of model to data leads to future model refinement (dashed arrow) in the iterative cycle of systems biology. Adapted from {Locke, 2005}.

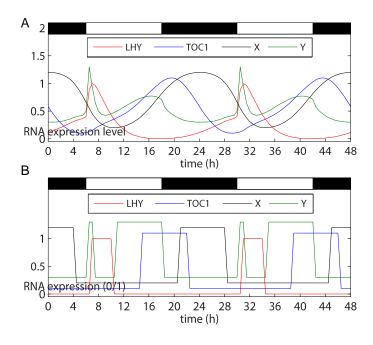


Fig. 2. The simple, qualitative form of a model can retain key behaviours.

(A) Simulation outputs show RNA levels changing continuously, from the simple clock model {Locke, 2005} in quantitative form (differential equations, as in Figure 1B). (B) RNAs are either expressed (1) or not (0) in the qualitative form of the same model {akman, 2012}. The binary, time-delay model still shows bimodal peaks of RNA expression from gene Y (green), with light induction after dawn (as in Figs. 1D, 2A). Levels are slightly offset for clarity in (B). Time 0h is midnight. Open box, light interval; filled box, dark interval.



Fig. 3. New capabilities arise from a "black-box" combination of models.

The circadian clock model shown in Figure 1 {Locke, 2005} can communicate to the Arabidopsis architectural model {Mundermann, 2005} running in L-studio software. A version of the clock model in Matlab software was compiled into the C programming language, in order to interact with the lpfg programme of L-studio. A clock protein level from the clock model controlled leaf angle in the architectural model, creating a simple simulation of rhythmic leaf movement in Arabidopsis over day/night cycle. The clock model's light:dark setting also darkened plant colour at night (16h, 20h). Simulation by Paul E. Brown.

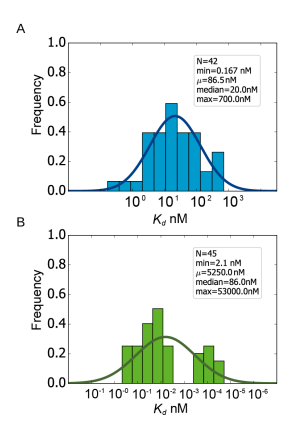


Fig. 4. Published parameter values can inform detailed modelling.

(A) Distribution of published K_d values for plant DNA-interaction affinities. (B) Distribution of published K_d values for plant protein-protein interaction affinities. Data such as these help to constrain the range of parameter values that parameter fitting procedures should explore. Please see main text for publication references.

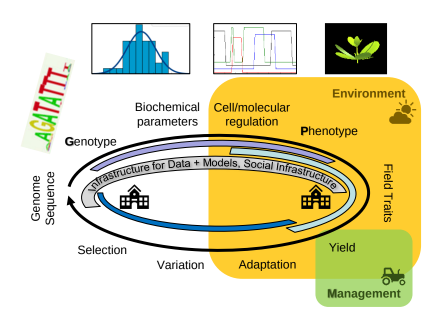


Fig. 5. Linking Systems Biology with Crop Science models.

The solid line links the concepts of biology, first from genome sequence via genotype, biochemical parameters and molecular regulation to whole-organism phenotype in a particular environment (yellow area); then from phenotypes to field traits and adaptation or to yield under particular management (green area); finally, given genetic variation, through natural selection or artificial selection in crop breeding, to the evolution of genome sequences {adapted from \Millar, 2016). Initiatives like Crops in silico will deal with the whole cycle, by linking several models (coloured arcs) into a seamless, causal chain. The top line of graphics locate the topics considered in the main text with reference to this cycle. The arcs suggest current types of model, in systems biology (indigo), crop science (cyan) and evolution (dark blue). The dimensions that are often considered in such models are capitalized (G, P, E, M). Underpinning infrastructures (grey) help to bridge these disciplines. 'Anchor' institutions are shown (buildings), which might provide major experimental facilities, digital infrastructure or a focus for social infrastructure, such as training or standardisation workshops.