



# Divergent carbon use efficiency-growth rate tradeoff in popular biological growth models

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Received: 4 November 2025 – Discussion started: 17 December 2025

Revised: 18 May 2026 – Accepted: 29 May 2026 – Published: 19 June 2026

**Abstract.** Carbon use efficiency (CUE) is an important trait emerging from processes regulating biological growth. CUE can be computed either based on the growth of structural biomass or total biomass divided by substrate uptake rate. Nonequilibrium thermodynamics and observations suggest that, for an exponentially growing population of cells, structural biomass CUE should first increase, then peak, and finally decrease with specific growth rate; meanwhile, total biomass CUE increases asymptotically with specific growth rate. We compared predictions from six popular models that are often used for plant and microbial growth in existing ecosystem models. We found that, for an exponentially growing population of biological cells, (1) the source-driven Pirt and Compromise models predict that structural biomass CUE increase asymptotically with growth rate; (2) the apparent sink-driven modified Droop model predicts that structural biomass CUE decreases with growth rate; and (3) the sink-driven variable internal storage model and two dynamic energy budget models predict that structural biomass CUE first increases, then peaks, and finally decreases with growth rate. Moreover, the modified Droop model predicts that total biomass CUE is constant with growth rate, while all other five models predict that total biomass CUE increases with growth rate asymptotically. For non-exponential biological growth, we show that there is no static relationship between total biomass CUE or structural biomass CUE with respect to either growth rate or temperature. Therefore, we contend that biological growth models should explicitly represent interactions between substrate acquisition, substrate transformation, and maintenance respiration to better capture observed CUE dynamics, and the sink-driven model should be preferred for general ecosystem biogeochemistry modeling.

## 1 Introduction

For a growing biological organism, be it a microbe, a plant, or even an animal, carbon use efficiency (CUE) is usually defined as the fraction of a unit mass of newly captured carbon that is retained as newly synthesized biomass (for simplicity, our discussion here regards all excreted organic polymers, such as exoenzymes, as part of the total biomass), while the remaining carbon is respired to support this growth (e.g., Manzoni et al., 2018). Because the carbon used to support new biomass synthesis also fulfills the metabolic needs of maintaining existing biomass, and, due to environmental fluctuations, the cost of such maintenance is generally dynamic, CUE is inferred to be dynamic even for an organism that is of fixed chemical elemental stoichiometry. Nevertheless, biogeochemical (BGC) models differ widely in their representation of microbial CUE, for example, as a constant parameter (Wieder et al., 2015), a static function of various regulating factors (e.g., temperature, macronutrient concentration; Allison et al., 2010), or an emergent variable that changes according to the chosen way of analysis (Tang and Riley, 2015; Manzoni et al., 2018; Allison, 2025). Moreover, based on observations that biological activity, such as the hatching of an egg or the germination of a plant seed, continues for a significant period of time in the absence of carbon uptake from the environment, variable internal storage models differentiated between structural and reserve biomass (e.g., Tang and Riley, 2015; Kooijman, 2009; Nev and van den Berg, 2017). Specifically, the structural biomass refers to DNA, cell wall and membrane material, and any biomass that requires maintenance to support an organism's normal function, while reserve biomass includes lipids, glycogen, cir-

culating metabolites, short-lived RNA and any biomass that stores energy and acts as a precursor of structural biomass. Consequently, the variable internal storage models reasoned that, since only the structural biomass requires maintenance, CUE can be either defined with respect to the growth of structural biomass or total biomass. Further, because plants drive growth using photosynthates (that is already stored in cells), the variable internal storage models fit plant growth naturally. However, many models do not differentiate between structural biomass and reserve biomass (e.g., (1) all microbe-implicit soil BGC models; Koven et al., 2013; Parton et al., 1988; Jenkinson, 1990), (2) the Pirt model (Pirt, 1982), and (3) the Compromise model (Beefink et al., 1990; Wang and Post, 2012)), and implicitly assume an equal CUE for structural and total biomass growth.

When the variable internal storage models are applied to microbes, structural biomass represents the effect of microbial population size on substrate uptake. Thus, it is important to recognize how the method of CUE computation affects the interpretation of CUE dynamics. Specifically, is it the structural biomass CUE that designates population growth, or is it the total biomass CUE that designates biomass growth? Among the existing empirical methods measuring microbial CUE, for the respiration induced by a given amount of substrate addition, the chloroform fumigation-extraction method and the  $^{13}\text{C}$  or  $^{14}\text{C}$ -labeling method focus on total biomass increase after substrate addition, thus they derive CUE conditioned on total biomass (Geyer et al., 2019; Hagerty et al., 2022). In contrast, the PLFA (phospholipid fatty acids) approach quantifies structural biomass growth by measuring how much new membrane is synthesized upon the substrate addition (Ibekwe and Kennedy, 1998; Liu et al., 2020). The  $^{18}\text{O}$ - $\text{H}_2\text{O}$  labeling method and  $^{13}\text{C}$ -DNA stable isotope probing (SIP) method quantify the amount of microbial growth based on new DNA synthesis (Xu et al., 2024; Sayre et al., 2026). However, DNA synthesis is not equivalent to increase in cell population size. Therefore, only the PLFA method measures the structural biomass CUE, while the  $^{18}\text{O}$ - $\text{H}_2\text{O}$  method and  $^{13}\text{C}$ -DNA SIP method measure what is more accurately described as “apparent” structural biomass CUE. Despite the difference in those approaches, the measured microbial CUE values are often treated as equivalent in the literature. Meanwhile, the plant CUE is often defined as the ratio between net primary productivity and gross primary productivity, signifying the total biomass CUE (Sinsabaugh et al., 2017).

In models of soil organic matter decomposition dynamics, CUE defined for the representative microbe in a population is regarded as an essential parameter to determine how much carbon can be retained in soils after respiration. Recently, Tao et al. (2023), adopting the model structure from Allison et al. (2010), highlighted microbial CUE as the most important parameter to be better constrained to improve the estimation of global soil carbon storage. Overall, microbe-explicit soil BGC models are demonstrating very diverse

variations of predicted microbial CUE (some of their prototypes are analyzed later in this manuscript). In general, these microbe-explicit models can be organized into four classes: (1) the Herbert model (Dawes and Ribbons, 1964), (2) the Pirt-model (Pirt, 1982, 1965), (3) the Compromise model (Beefink et al., 1990), and (4) variable internal storage models, including the plant growth model by Thornley (1972), dynamic energy budget models (Tang and Riley, 2015, 2023; Kooijman, 2009), and the Droop model (Thingstad, 1987). Due to the missing of reserve biomass, the first three models equate total biomass CUE to structural biomass CUE, while the fourth group of models can differentiate total biomass CUE from structural biomass CUE.

As another group of biological organisms essential for ecosystem biogeochemical models, plants are structurally more complex than microbes, and, naturally, the CUE dynamics during plant growth is also more complicated. However, since plants are constrained by the same physical rules as nature enforces onto microbes, plant and microbial CUEs should be treated similarly both in modeling and measurements. As it is argued in Tang et al. (2024), a consistent treatment of similar processes in different contexts will make the resultant ecosystem models more robust by improving the coupling coherency between the models' different components. Theoretically, the whole plant CUE should be defined as the fraction of allocated carbon to various organs that becomes newly synthesized biomass (Thornley, 1972). Because the relative carbon allocation to each plant organ varies with environmental conditions and different organs usually have different CUEs, the whole plant CUE is generally dynamic (Manzoni et al., 2018). Even when all plant organs are assumed to have the same CUE, the dynamics of relative carbon allocation for maintenance and growth will still result in dynamic whole plant CUE. In the literature, the mean whole plant CUE has been mapped for some regions using remote sensing imagery (Bloom et al., 2016; Zhang et al., 2009) and field data (Liu et al., 2022).

Conceptually, the growth of any biological organism is thermodynamically controlled and therefore naturally resembles how a thermal or mechanical engine works with its supplied energy (Roach et al., 2018). For a Carnot thermal engine, classical thermodynamics has shown that its energy use efficiency has an upper bound of  $1 - T_L/T_H$ , with  $T_L$  and  $T_H$  being the low and high temperatures of two heat reservoirs, respectively (Feynman et al., 2011). However, this classical thermodynamic result assumes that all processes are reversible. That is, the engine is working so slowly that its power is de facto zero. This assumption disagrees with the empirical observation (and desire) that a working engine should be able to deliver a finite amount of work during any given time. Thus, nonequilibrium thermodynamics was developed to analyze the efficiency of various engines (Bejan, 1996; Andresen et al., 1977, 1984; Onsager, 1931a, b). One important inference from nonequilibrium thermodynamics is that there exists a tradeoff between power and energy effi-

ciency of an engine, so that one sub-optimal efficiency may correspond to both a low and high input power (Roach et al., 2018; Yuan et al., 2022).

The power-yield tradeoff mentioned above can be understood using the real-life experience of driving a vehicle. In this example, the yield is fuel use efficiency (FUE), i.e., distance traveled per volume fuel consumed by an internal combustion vehicle (e.g., km per liter gas). When a vehicle's engine is off, its output power is zero, and the vehicle has zero FUE. When the engine is on but the car is not moving, FUE is still zero (and the fuel use just keeps the engine idling). When the vehicle gradually accelerates on a flat road, FUE gradually increases, and then stabilizes at a value that is more or less as advertised by the vehicle manufacturer. When the driver is overtaking another vehicle or climbing a slope, greater power is demanded. This is done by pressing the pedal (and probably shifting the gear into sport mode at the same time), so that the vehicle may pick up further speed (with the greater power) at the cost of lower FUE. However, in the real-world, because conditions where a car operates usually fluctuate widely, one is more likely to find the car's power-yield tradeoff varies continuously (and there is not a static relationship for this tradeoff). We note that internal combustion engines generally cannot store the energy produced from burning the fuel, thus this energy is either wasted or used for moving the vehicle. A hybrid vehicle may store some of the kinetic energy generated from fuel burning for later use. As we will see later, biological cells work more like a hybrid vehicle. Consistent with this analogy, for mean values based on 398 observations of foreign and US domestic automobiles from the model years 1970 to 1982, Boland and Schreyer (2024) found a clear decrease of FUE with increasing horsepower.

Although biological organisms and vehicles' mechanical engines use energy in different forms, they must adhere to the same thermodynamic principles. Therefore, it is reasonable to infer that the CUE of biological growth should also be a dynamic variable, and follow a similar power-yield tradeoff. This pattern has been hypothesized for microbial CUE by Lipson (2015) based on a synthesis of empirical studies from the literature, and was predicted by the dynamic energy budget models (Tang and Riley, 2023, 2025). Nonetheless, we do expect the CUE dynamics of biological growth to be more complex than the energy use efficiency dynamics of thermal engines. Thermal engines, like internal combustion engines, generally do not include energy storage. In contrast, biological growth is often compounded with processes like dormancy, adaptation and acclimation to environmental changes, giving rise to hysteretic CUE dynamics. Particularly for trees in winter, their roots can grow vigorously while the shoots are dormant (Marchand et al., 2025). Still, these complex CUE dynamics can analogously be conceptualized as the coupling of multiple thermal engines, such as the coupling of an electric motor and an internal combustion engine in hybrid vehicles, which are more resilient in terms

of energy expenditure than either electric vehicles or internal combustion engine vehicles (under mixed disruption conditions of energy supply; Orecchini et al., 2018).

Considering the significance of CUE in predicting ecosystem biogeochemistry, it is thus important to answer the following questions: are the formulations commonly used by existing ecosystem biogeochemical models representing (microbial and plant) CUE dynamics consistently with thermodynamics? If not, what important features could have been missed by those formulations? In the following sections we analyze the tradeoff between growth rate and CUE in six mathematical formulations, some of which underlie most existing ecosystem biogeochemical models. We then highlight the importance of resolving the growth rate-CUE tradeoff dynamically and discuss how it can be properly formulated in future models.

## 2 Theory

We first introduce a modified version of the model used by Odum and Pinkerton (1955) for their analysis of diverse systems that involve the coupling between energy supply and energy use. We describe that model to illustrate the generic features of coupled energy supply and use as implied by nonequilibrium thermodynamics. Then we introduce the biological growth models that are used in many ecosystem models, and analyze the relationship between growth rate and CUE implied by those growth models. All mathematical symbols are described in Table A1 in the Appendix.

### 2.1 The modified Odum-Pinkerton model

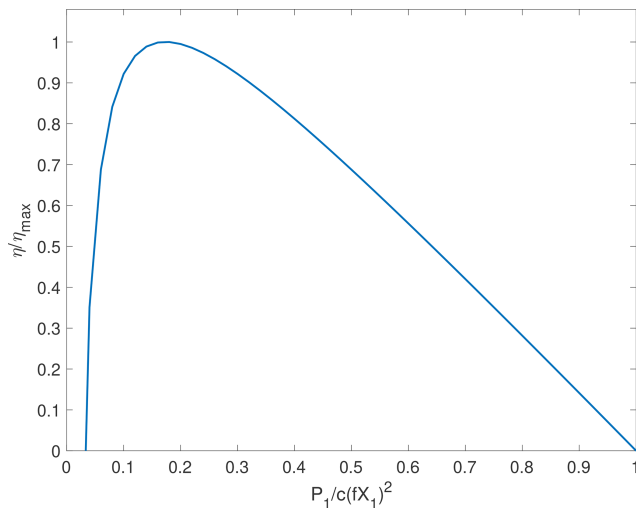
The model is formulated using Onsager's reciprocal relations (Onsager, 1931b):

$$J_1 = (l + cf^2) X_1 - cf X_2, \quad (1)$$

$$J_2 = -cf X_1 + (c + \epsilon) X_2, \quad (2)$$

$$T \frac{dS_e}{dt} = J_1 X_1 + J_2 X_2. \quad (3)$$

In the above model,  $l$ ,  $f$ ,  $c$  and  $\epsilon$  are constant parameters (whose definitions depend on the application),  $X_1$  and  $X_2$  are forces,  $J_1$  is the supply flux into the system under the influence of force  $X_1$ , and  $J_2$  is the output flux with force  $X_2$ . The sum of products of fluxes and forces contributes to the heat dissipation of the system as measured by the entropy production rate  $dS_e/dt$  multiplied with temperature  $T$  in Eq. (3). Note that Eq. (3) follows directly from the first law of thermodynamics (aka energy conservation), so that given the supply power  $P_1 = J_1 X_1$ , and the heat dissipation rate  $T \frac{dS_e}{dt}$ , the useful output power is  $P_2 = -J_2 X_2$ . Consequently, the



**Figure 1.** The power-yield tradeoff predicted by the modified Odum-Pinkerton model. Here, we set  $l = 0$  and  $\epsilon = 0.03$ , so that the model is describing the 7th example in Odum and Pinkerton (1955), which represents an organism that captures (via  $J_1$ ) and uses food for its maintenance (signified by  $\epsilon$ ) and growth (via  $J_2$ ), except that we here assume that the maintenance cost is paid by  $X_2$  (analogous to the reserve biomass in the variable internal storage models, e.g., the DEB models in this study). While we do not plot negative values of  $\eta$ , it may occur resembling to a starving organism that keeps respiring at the expense of biomass under insufficient food supply.

system's energy use efficiency ( $\eta$ ) is

$$\eta = \frac{P_2}{P_1} = \left(1 + \frac{l}{cf^2} - p\right) \left[1 - \frac{l}{pcf^2} - \frac{\epsilon}{c} \left(\frac{l+cf^2}{pcf^2} - 1\right)\right], \quad (4)$$

where  $p = P_1/c(fX_1)^2$ . Further, we note that

$$p = \frac{P_1}{c(fX_1)^2} = \left(1 + \frac{l}{cf^2}\right) - \frac{X_2}{fX_1} < 1 + \frac{l}{cf^2}, \quad (5)$$

so that  $\eta$  may vary from being negative to a positive maximum value, depending on the value of terms in  $[\ ]$ . Particularly, when  $\eta$  is plotted against  $p$ , we obtain the non-monotonic tradeoff curve (Fig. 1).

Odum and Pinkerton (1955) (hereafter OP1955) used the above model to analyze (1) Atwood's machine (e.g., Monteiro et al., 2015), (2) a water wheel turning a grindstone, (3) one battery charging another battery, (4) a thermocouple running an electric motor, (5) a thermal diffusion engine, (6) the metabolism of a pseudo-organism (with no self-repair), (7) food capture by an organism for its maintenance and growth, (8) a model of photosynthesis, (9) primary production in a self-sustaining climax community, and (10) growth and maintenance of a civilization. Among them, the 7th example is closest to the biological growth we discuss here. Moreover, the fact that these equations can describe so many

types of systems suggests what can be inferred from these equations for the energy use efficiency is a universal feature that is guaranteed by nonequilibrium thermodynamics. Consequently, the 7th example and the subject of our analysis – biological growth – should have the same characteristics in terms of energy use. Following the interpretation by OP1955, in this example,  $J_1$  describes the rate of food supply,  $X_1$  corresponds to the energy drop in metabolism of captured food units (thus  $X_1$  is equivalent to Gibbs free energy, e.g. that of glucose, representing the energy quality of the substrate taken from the environment),  $J_2$  represents the rate of effective food use for biomass synthesis,  $X_2$  is the energy drop inherent in the metabolism of a unit of food (thus  $X_2$  is analogous to the Gibbs free energy of reserve biomass in the DEB models),  $l$  designates the basal metabolism spent in self-repair (as paid by  $X_1$  in the original OP1955 formulation),  $c$  is related to the effectiveness of food concentrating method, and  $f$  equals to  $X_2/X_1$ , measuring the metabolic efficiency of the food capture process. However, considering that biological organisms may keep respiring in the absence of food uptake (aka when  $J_1 = 0$ ; e.g., Dawes and Ribbons, 1964), it is more reasonable to pay the basal metabolism with  $X_2$ , so that  $l = 0$  while  $\epsilon > 0$ . Nonetheless, this change does not alter the non-monotonic tradeoff between efficiency  $\eta$  and the input power  $P_1$  (Fig. 1). Moreover, in the OP1955 model, all input power  $P_1$  goes to output power  $P_2$  and waste, thus there is no change in storage energy density (aka  $X_2$ ). Therefore, the implied non-monotonic power-yield tradeoff corresponds to exponential biological growth, as discussed below.

## 2.2 Biological growth models

We next analyze six biological growth models: (1) Pirt model (Pirt, 1965), (2) Compromise model (Beetink et al., 1990), (3) Modified Droop (mDroop) model (Thingstad, 1987), (4) Variable Internal Storage (VIS) model modified from Thornley's plant model (Thornley, 1972), (5) the standard dynamic energy budget (sDEB) model (Kooijman, 2009), and (6) the modified dynamic energy budget (mDEB) model (Tang and Riley, 2025). Both the Pirt and Compromise models consider a single biomass pool, and treat growth and maintenance as independent processes, having the same priority in using the substrate. As biological growth in the Pirt model and Compromise model is directly driven by substrate uptake, they are in the category of source-driven growth models (Fatichi et al., 2014). The other four models assume biomass consists of structural and reserve biomass. Among them, the VIS, sDEB and mDEB models drive structural biomass growth using reserve biomass, thus they are in the category of sink-driven models (Fatichi et al., 2014). Following Thingstad (1987), the mDroop model has the most parameters, and uses the total biomass to compute the growth rate (through the total-biomass-based substrate quota, even though it does have reserve biomass), thus it is a sink-driven model only apparently. For the Pirt and Compromise models under general

conditions, their equations stay the same as in Table 1. The general forms of the other four models differ from those (for exponential growth) in Table 1 and are detailed (in Sects. S1 to S4) in the Supplement.

We compared the six models' predictions of CUE dynamics for (1) an exponentially growing cell population under constant carbon inputs using equations in Table 1, (2) 50-year simulations using equations from Sect. S5 in the Supplement that mimic microbe-driven soil carbon dynamics under monthly varying but annually constant carbon inputs (with a rate of  $350 \text{ gC m}^{-2} \text{ yr}^{-1}$ ; red line in Fig. S1a), and (3) 50-year simulations that further consider the influence of smoothed daily variable temperatures (red line in Fig. S1c) on related temperature-dependent kinetic parameters. For models used in the 50-year simulations of soil carbon dynamics, we detailed their mathematical formulations in Sect. S5 and parameterizations in Sect. S6 in the Supplement, and gave here only a high-level description. Specifically, each of the six models describes the dynamics of one microbial population that feeds on one fast C pool and one slow C pool. Note that the use of fast and slow pools here simply means the former carbon substrate is kinetically more favorable than the latter to the microbes, and they do not have the same meaning as in microbe-implicit models, which are directly associated with turnover times (Parton et al., 1988). We formulated the substrate uptake based on the argument in (Tang et al., 2024), i.e. a form that resembles the competitive Michaelis-Menten kinetics as simplified from the equilibrium chemistry approximation (ECA) kinetics (Tang and Riley, 2013). In these models, the fast and slow C pools increase due to external input, and all microbial biomass from mortality is added to the slow pool C. Additionally, since the Pirt model, Thornley's model and the sDEB have also been adopted for modeling plant growth (Thornley, 1972; Russo et al., 2022; Thornley, 2011), the assertions we draw below can also be more generally extended to plants.

### 3 Results of model comparisons

#### 3.1 Predicted CUE-growth rate tradeoffs for exponentially growing populations

For an exponentially growing population of biological cells, predictions by these six models can be sorted into three groups: (1) the Pirt and Compromise models predict that (both structural and total biomass) CUE increases asymptotically with increasing specific growth rate; (2) the mDroop model predicts that the total biomass CUE is insensitive to specific growth rate, while the structural biomass CUE decreases with increasing specific growth rate; and (3) the VIS, sDEB, and mDEB models predict that structural biomass CUE first increases then decreases with specific growth rate, while the total biomass CUE increases asymptotically with specific growth rate (Fig. 2). Particularly, the increasing to-

tal biomass CUE as a function of specific growth rate aligns well with empirical measurements (e.g., Zheng et al., 2019; Hu et al., 2025; Collado et al., 2014), and the Pirt and Compromise models were specifically created to capture this feature observed in batch experiments (Pirt, 1965; Beftink et al., 1990). However, the Compromise model is argued to be more realistic than the Pirt model by allowing negative growth resulting from biomass degradation under low to no substrate conditions (Beftink et al., 1990; Wang and Post, 2012), while the Pirt model forces the growth rate to be zero under the no-supply substrate condition.

#### 3.2 Dynamic CUE in transient soil carbon dynamics under constant temperature

Among the six models, predicted relationships between total biomass CUE and specific growth rate can also be organized in three groups (Fig. 3): (a) the Pirt and Compromise models predict essentially identical, almost linearly increasing, relationships (Fig. 3a); (b) the mDroop model predicts a curve that gradually closes on itself (Fig. 3b); and (c) the VIS, sDEB, and mDEB models predict relationships look like ellipses that are slanting slightly up and to the right, and those by the two DEB models are nearly overlapping each other (Fig. 3c). (The predicted soil carbon dynamics are presented in Fig. S2. Even though the temporal variability of fast pool C, slow pool C, and total biomass are different but arguably quite consistent among all six models by our choice to share as many parameters as possible among the models.)

Since the Pirt and Compromise models each have only one biomass pool, their predicted microbes have only one CUE for each specific growth rate (Fig. 3a). In contrast, the representation of reserve biomass makes the other four models predict multiple total biomass CUE values for each specific growth rate (Fig. 3b, c), which gradually converge to a closed curve due to repeating growth pattern (Fig. S2d) driven by the temporal response of reserve biomass to the fixed seasonal pattern of carbon input (Figs. S1b and S2a).

When the structural biomass CUE is analyzed with respect to the specific growth rate (Fig. 4), the mDroop model predicts this relationship to have a left-leaning asymmetric oval shape (Fig. 4a). Meanwhile, the VIS, sDEB, and mDEB models all predict the relationship to be of more vertically aligned asymmetric oval shapes (Fig. 4b, c, d). In all, just like that for the total biomass CUE, these four models predict that there is not a one-to-one relationship between structural biomass CUE and specific growth rate. Rather, the relationship traces in curves that gradually close on themselves as enabled by the temporal response of reserve biomass to the constant temperature and monthly varying but annually constant carbon input (Figs. S1b and S2a).

**Table 1.** Models for exponential biological growth  $\mu(S)$  as a function of substrate uptake  $q(S)$ . Listed in parentheses by each model's name are key model parameters, whose definitions are detailed in the nomenclature table in Appendix A. Notably,  $Y_V$  and  $Y_B$  are CUEs for structural biomass and total biomass, respectively, both of which are normalized with substrate assimilation efficiency  $Y_S$ . More detailed model descriptions are in the Supplement.

Pirt model: $(\mu_{\max}, m_V, Y_S), h(S)$	Pirt (1965, 1982)
$\mu(S) = \mu_{\max} h(S)$ $q(S) = \frac{\mu}{Y_S} + \frac{m_V}{Y_S} = \frac{\mu}{Y_S} \left(1 + \frac{m_V}{\mu}\right)$ $\frac{Y_V}{Y_S} = \frac{Y_B}{Y_S} = \frac{\mu(S)}{q(S)Y_S} = \frac{\mu}{\mu + m_V}$ <p>Description: This source-driven model assumes that specific growth rate <math>\mu(S)</math> is down-regulated linearly from the maximum growth rate <math>\mu_{\max}</math> by the substrate response function <math>h(S)</math>. Substrate uptake <math>q(S)</math> is partitioned between that for biomass synthesis <math>\mu/Y_S</math> and maintenance <math>m_V/Y_S</math>. As a single biomass compartment model, its structural biomass yield <math>Y_V</math> and total biomass yield <math>Y_B</math> are equal.</p>	
Compromise (Comp) model: $(\mu_{\max}, m_V, Y_S), h(S)$	Wang and Post (2012)
$\mu(S) = \mu_{\max} h(S) - m_V [1 - h(S)]$ $q(S) = \frac{\mu}{Y_S} + \frac{m_V}{Y_S}$ $\frac{Y_V}{Y_S} = \frac{Y_B}{Y_S} = \frac{\mu(S)}{q(S)Y_S} = \frac{\mu}{\mu + m_V}$ <p>Description: This source-driven model assumes that specific growth rate <math>\mu(S)</math> consists of two parts, where the first part linearly scales the maximum growth rate <math>\mu_{\max}</math> using the substrate response function <math>h(S)</math>, and the second part signifies negative growth due to maintenance <math>m_V [1 - h(S)]</math> that decreases linearly with the substrate response function <math>h(S)</math>. Substrate uptake <math>q(S)</math> is partitioned between that for biomass synthesis <math>\mu/Y_S</math> and maintenance <math>m_V/Y_S</math>. As a single-biomass-compartment model, its structural biomass yield <math>Y_V</math> and total biomass <math>Y_B</math> are equal.</p>	
Modified Droop (mDroop) model: $(\mu_{\max}, c_1, c_2, Q_{\min}, Q_{\max}, Y_S) j_A(S)$	Thingstad (1987)
$\mu(S) = \frac{\mu_{\max} j_A (Q_{\max} - Q_{\min})}{j_A Q_{\max} + [c_2 + (1 + c_1) \mu_{\max}] Q_{\min} (Q_{\max} - Q_{\min})}$ $q(S) = \frac{(1 + c_1) \mu_{\max} + c_2}{\mu_{\max} - \mu} \frac{Q_{\min}}{Y_S} \mu$ $\frac{Y_V}{Y_S} = \frac{\mu_{\max} - \mu}{(1 + c_1) \mu_{\max} + c_2} \frac{1}{Q_{\min}}$ $\frac{Y_B}{Y_S} = \frac{\mu_{\max}}{(1 + c_1) \mu_{\max} + c_2}$ <p>Description: This apparent sink-driven model assumes that specific growth rate <math>\mu(S)</math> is linearly down-regulated from the maximum growth rate <math>\mu_{\max}</math> by the carbon quota function <math>1 - Q_{\min}/Q</math>, where the carbon quota <math>Q</math> is defined as the ratio between total biomass <math>B_T</math> and structural biomass <math>B_V</math> (see Eq. S3 in Sect. S1 in the Supplement). The total biomass <math>B_T</math> increases due to substrate uptake <math>j_A \frac{Q_{\max} - Q}{Q_{\max} - Q_{\min}} B_V</math>, and decreases due to respiration <math>R B_V</math> (see Eq. S2 in the Supplement), where the specific respiration <math>R</math> is a linear function of specific growth rate <math>\mu(S)</math>, whose coefficients depend linearly on carbon quota (Eq. S5 in the Supplement). As a two-biomass-compartment model, its total biomass yield <math>Y_B</math> and structural biomass yield <math>Y_V</math> differ, and they can be analytically represented as a function of specific growth rate <math>\mu</math> only for an exponentially growing population.</p>	
Variable-internal-stores (VIS) model: $(v_E, m_V, Y_S, Y_{RV}) j_A(S)$	Williams (1967), Thornley (1972)
$\mu(S) = \frac{v_E}{2} \left( -1 + \sqrt{1 + \frac{4(j_A Y_S - m_V) Y_{RV}}{v_E}} \right)$ $q(S) = \frac{\mu}{Y_S Y_{RV}} \left( 1 + \frac{\mu}{v_E} \right) + \frac{m_V}{Y_S}$ $\frac{Y_V}{Y_S} = \frac{v_E \mu}{(v_E + \mu) \mu + v_E m_V Y_{RV}} Y_{RV}$ $\frac{Y_B}{Y_S} = \frac{(v_E Y_{RV} + \mu) \mu}{(v_E + \mu) \mu + v_E m_V Y_{RV}} Y_S$ <p>Description: This sink-driven model assumes that reserve biomass <math>B_X</math> is mobilized to parallelly meet the demand from maintenance and growth of structural biomass <math>B_V</math> (see Eq. S14 in Sect. S2 in the Supplement). As a two-biomass-compartment model, its total biomass yield <math>Y_B</math> and structural biomass yield <math>Y_V</math> differ, and they can be analytically represented as a function of specific growth rate <math>\mu</math> only for an exponentially growing population.</p>	

Table 1. Continued.

Standard dynamic energy budget (sDEB) model: $(v_E, m_V, Y_S, Y_{RV}) j_A(S)$	Tang and Riley (2023), Kooijman (2009)
$\mu(S) = \frac{Y_{RV} Y_S j_A - m_V}{1 + Y_{RV} Y_S j_A / v_E}$ $q(S) = \frac{v_E (m_V + \mu)}{Y_S Y_{RV} (v_E - \mu)}$ $\frac{Y_V}{Y_S} = \frac{\mu}{\mu + m_V} \left( 1 - \frac{\mu}{v_E} \right) Y_{RV}$ $\frac{Y_B}{Y_S} = \left( \frac{\mu}{\mu + m_V} \right) \left( \frac{Y_{RV} v_E + m_V + (1 - Y_{RV}) \mu}{v_E} \right)$	
<p>Description: This sink-driven model assumes that, when the reserve biomass <math>B_X</math> is mobilized to meet the demand from maintenance first and then the growth of structural biomass <math>B_V</math>, the reserve density (i.e. <math>B_X/B_V</math>) follows the first order kinetics (Eq. S26 in Sect. S3 in the Supplement). As a two-biomass-compartment model, its total biomass yield <math>Y_B</math> and structural biomass yield <math>Y_V</math> differ, and they can be analytically represented as a function of specific growth rate <math>\mu</math> only for an exponentially growing population.</p>	
Modified dynamic energy budget (mDEB) model: $(v_E, m_V, Y_S, Y_{RV}) j_A(S)$	Tang and Riley (2025)
$\mu(S) = \frac{m_V + v_E}{2} \left( -1 + \sqrt{1 + \frac{4v_E(j_A Y_S Y_{RV} - m_V)}{(m_V + v_E)^2}} \right)$ $q(S) = \left( 1 + \frac{\mu}{v_E} \right) \left( \frac{\mu + m_V}{Y_{RV} Y_S} \right)$ $\frac{Y_V}{Y_S} = \left( \frac{\mu}{\mu + m_V} \right) \left( \frac{v_E}{v_E + \mu} \right) Y_{RV}$ $\frac{Y_B}{Y_S} = \left( \frac{\mu}{\mu + m_V} \right) \left( \frac{Y_{RV} v_E + \mu + m_V}{v_E + \mu} \right)$	
<p>Description: This sink-driven model assumes that reserve biomass <math>B_X</math> is mobilized kinetically to meet the demand from maintenance first and then the growth of structural biomass <math>B_V</math>. Unlike the sDEB model, it does not restrict the reserve density (i.e. <math>B_X/B_V</math>) to follow the first order kinetics. As a two-biomass-compartment model, its total biomass yield <math>Y_B</math> and structural biomass yield <math>Y_V</math> differ, and they can be analytically represented as a function of specific growth rate <math>\mu</math> only for an exponentially growing population.</p>	

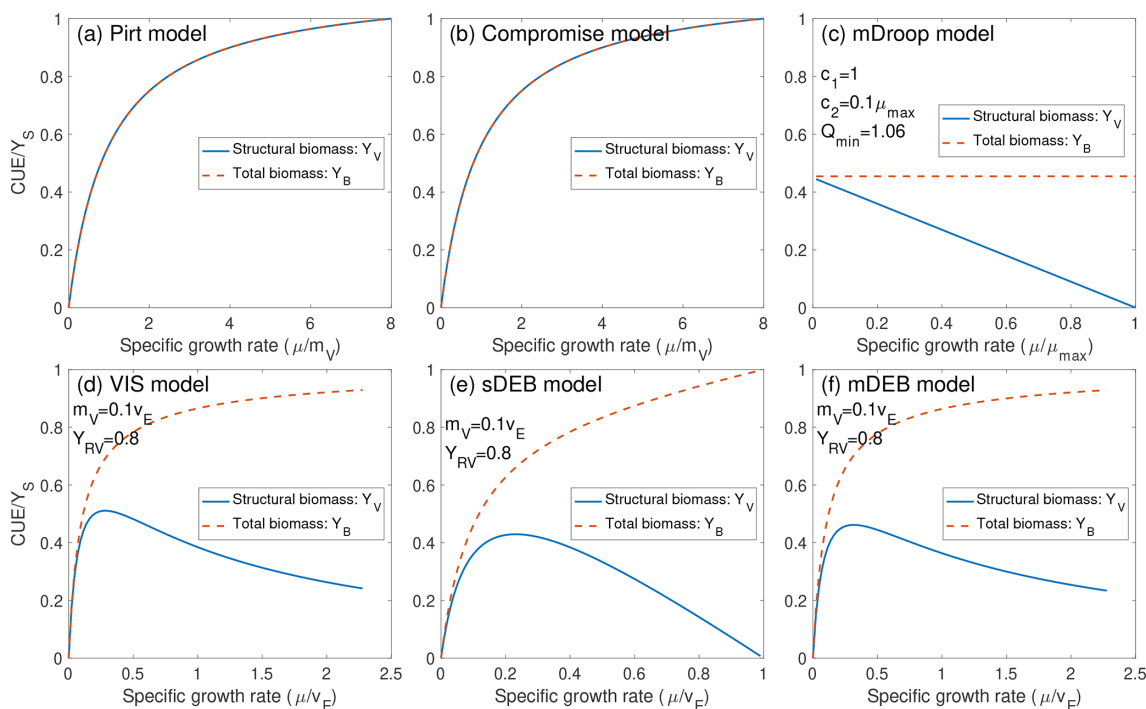
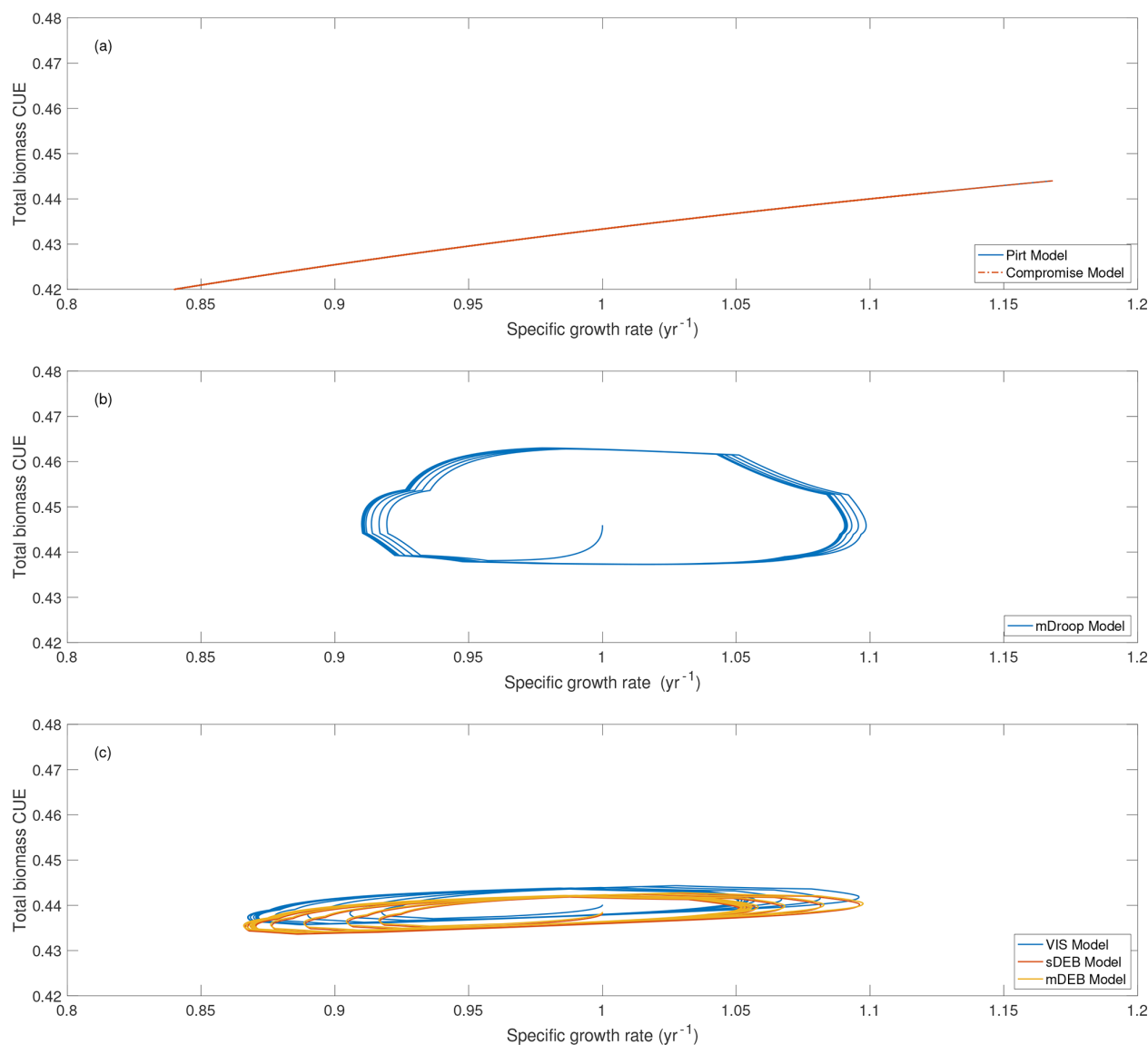


Figure 2. Comparison of relationships between CUE and specific growth rate predicted by the six models. Shown in panels (c–f) are parameter values to construct the plots. Structural biomass CUE  $Y_V$  is defined as the fraction of captured carbon substrate that is retained as structural biomass. Total biomass CUE  $Y_B$  is defined as the fraction of captured carbon substrate that is retained in structural and reserve biomass. Note that the specific growth rate for each model is normalized by parameters chosen by convenience, while CUE is normalized by the substrate assimilation efficiency  $Y_S$ .

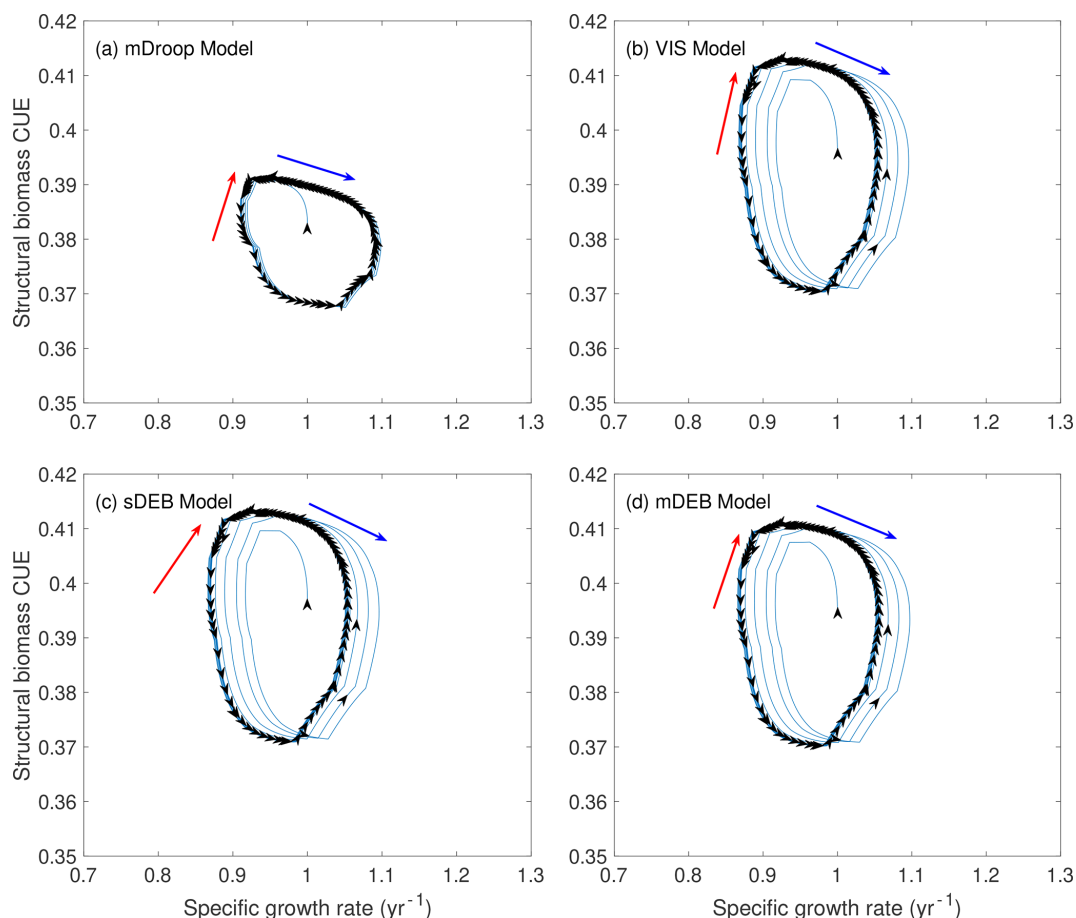


**Figure 3.** Relationship between total biomass CUE and specific growth rate. The evolution of different microbial carbon pools and total microbial biomass is presented in Fig. S2 in the Supplement. Note that results for the Pirt model and the Compromise model overlap each other in panel (a).

### 3.3 Dynamic CUE in transient state soil carbon dynamics under monthly varying but annually constant carbon input and day-to-day varying temperatures

When the temperature effect on kinetic parameters is further considered (on top of the 50-year monthly variable but annually constant carbon input), we find much richer variation of the relationship between total biomass CUE and specific growth rate. Still, the predicted relationships can be organized into the same three groups: (a) those by the Pirt and Compromise models (Fig. 5a), which look like a fish-hook, and no longer overlap, (b) that by the mDroop model

(Fig. 5b), which looks like a thread with an open knot, and (c) those by the VIS, sDEB, and mDEB models (Fig. 5c), which look like entangled threads. Among these three groups, the relationship predicted by the mDroop model has the largest spread. In particular, the mDroop model predicts significant occurrence of negative CUE, where excessive maintenance cost leads to the loss of biomass in the presence of active carbon substrate uptake. Meanwhile, predictions by the VIS, sDEB, and mDEB models show a much tighter spread, and, due to the chosen parameter combinations, do not involve negative CUE (Fig. 5c) as also found for the Pirt and Compromise models (Fig. 5a). Nonetheless, by considering temporally varying carbon inputs and temperatures, one specific



**Figure 4.** Relationship between structural biomass carbon use efficiency (CUE) and specific growth rate. Across all panels, blue arrows indicate the period when structural biomass CUE decreases with specific growth rate, while red arrows indicate the period when structural biomass CUE increases with specific growth rate. Black arrows in the figure panels indicate the temporal progression in the simulations.

growth rate can correspond to many values of CUE. However, the curves still gradually close on themselves due to the temporal response of reserve biomass to the fixed seasonal pattern of carbon input and temperatures.

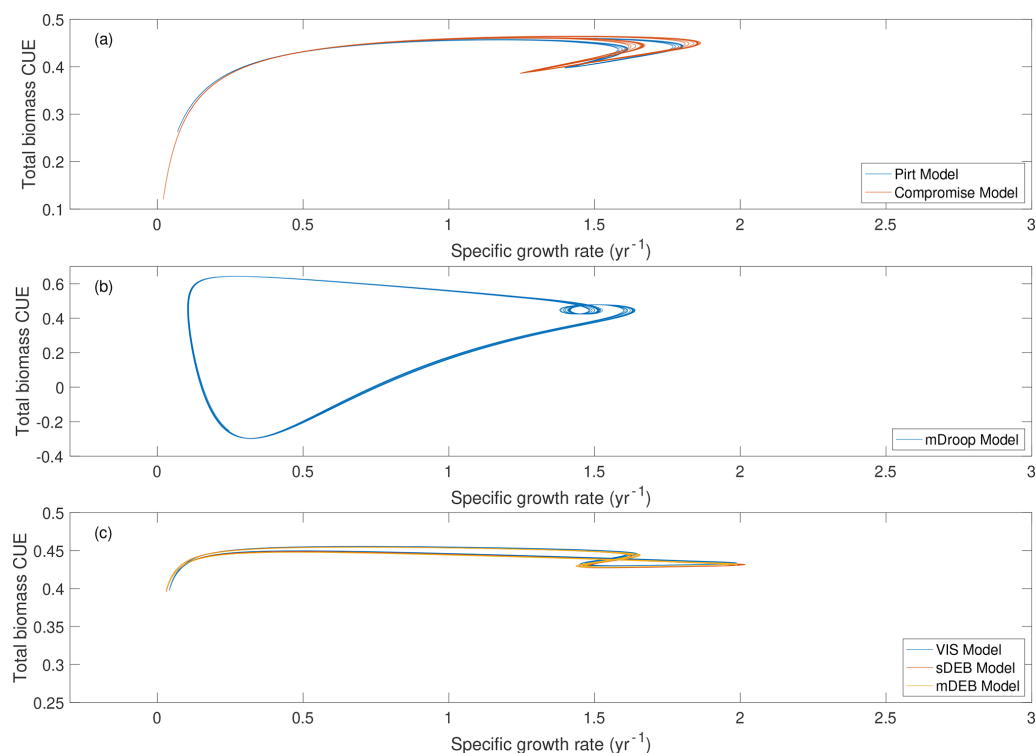
When the relationship between total biomass CUE and temperature is analyzed, the patterns obtained are even more complex (Fig. 6). (Note that temperature was varied daily, and repeated over the 50-year simulation period.) The Pirt and Compromise models both show a pattern of larger spread at low and high temperatures, and a general pattern that first increases and then decreases with temperature (Fig. 6a). The relationships predicted by the VIS, sDEB, and mDEB models have a much tighter pattern and smaller variation (Fig. 6c), a phenomenon that is consistent with observations that biological organisms evolutionarily develop reserve dynamics to stabilize performance (thereby less variable CUE) in the presence of fluctuating environmental conditions (Kooijman, 2009). Predictions by the mDroop model show the largest variation: although the CUE-temperature curve is well-defined, its spread are largest under low temperature, followed by intermediate and high temperatures with a cross

over at around 22.5 °C (Fig. 6b). Overall, all models predict that there are multiple CUE values almost at any given temperature.

## 4 Discussion

### 4.1 No static one-to-one relationship of CUE with respect to specific growth rate, or other environmental factors

It has long been recognized that CUE has a strong control on carbon and nutrient cycling through biological organisms and their host systems. However, many existing models treat microbial CUE as either a constant or a function depending statically on its controlling factors (Allison et al., 2010; Wang et al., 2021; He et al., 2015; Wieder et al., 2015). Our analysis here suggests that due to the indirect connection between maintenance respiration and substrate uptake, under time-varying conditions (like the varying carbon input and temperature considered here), there is not likely a static rela-

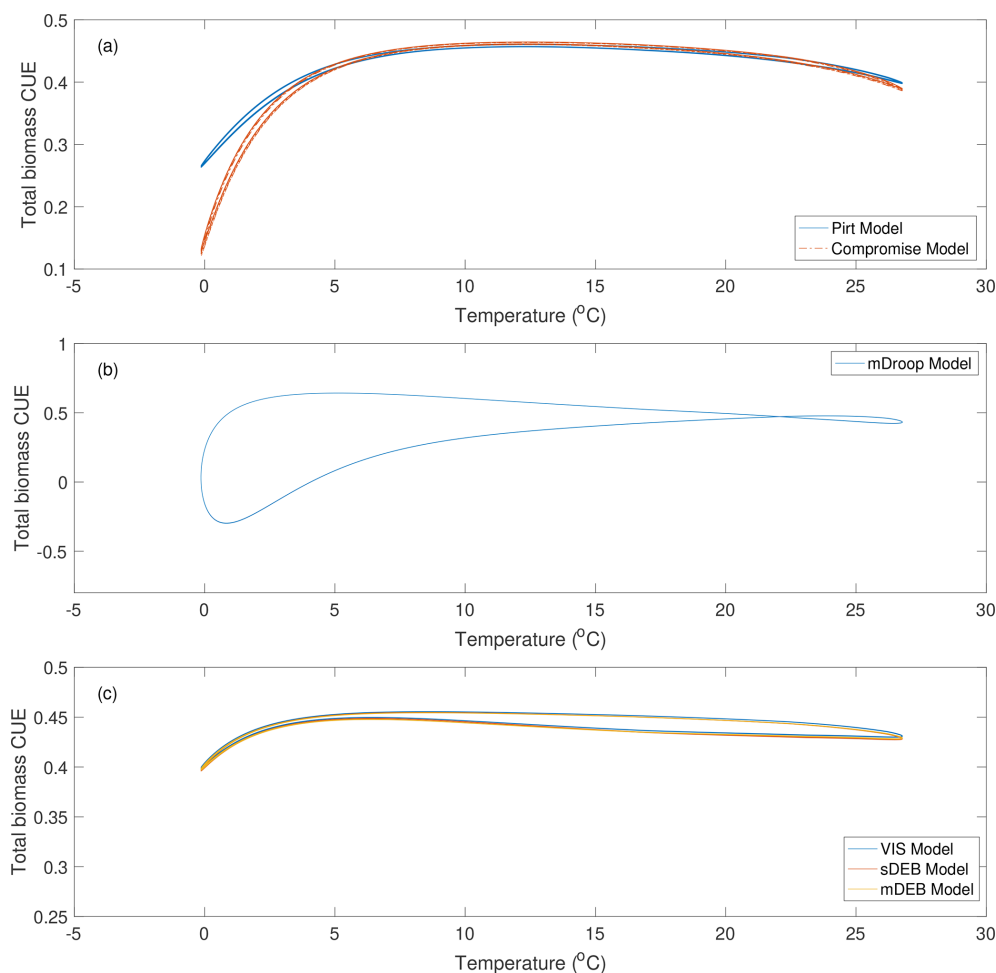


**Figure 5.** Variation of total biomass CUE with respect to specific growth rate. Data are extracted from the last 40 years of the 50-year simulations driven by monthly varying but annually constant carbon input and day-to-day varying temperatures. The corresponding soil carbon dynamics are shown in Fig. S3 in the Supplement.

relationship between CUE and its controlling factors or specific growth rate. Note, in systems that couple multiple groups of microorganisms (Graham et al., 2007), we expect that CUE dynamics could even be chaotic. Therefore, depending on the sampling time and location (e.g. the line segments indicated by blue or red arrows in Fig. 4), as well as the temporal averaging method involved in analysis (Tang and Riley, 2015), we may find CUE to have various trends with respect to the controlling variable being investigated (He et al., 2024). While it is not investigated here how much the simulated soil organic matter dynamics will change if the CUE is otherwise parameterized with a static function, based on results from Tang and Riley (2015), we expect the change will be significant. For test, when we drive the models with monthly and annually variable carbon input and non-smoothed daily variable temperatures, the spread of CUE curves becomes much wider (Fig. S5). In all, we contend that CUE should be resolved dynamically by explicitly representing the acquired carbon use for processes other than assimilation. This representation has been conceptually adopted in plant growth models, and soil biogeochemical models should also adopt it to ensure the mechanistic coherency of the overall ecosystem models. However, as we argue below, current plant growth models are generally formulated inappropriately.

#### 4.2 The source-driven vs. sink-driven models of biological growth

In the literature, due to their simple mathematical formulation and relatively good performance, the source-driven Pirt and Compromise models have been used much more frequently than the other four models in modeling microbes and plants. For instance, the land component (ELM) of the Energy Exascale Earth Model (Zhu et al., 2019) represents plant growth in a form that resembles the Compromise model, where new structural growth is driven by the residual gross primary productivity flux after subtracting the carbon requirement for maintenance and storage-replenishment. This source-driven approach is also adopted for plant growth by the BiomE model (Weng et al., 2019) and the FATES model (Knox et al., 2024). Sierra et al. (2022) criticized that this source-driven approach may lead to overly fast plant carbon turnover as indicated by a modeled too young radiocarbon signal in plant respiration compared to observations. Indeed, these source-driven models assume that carbon storage is negligible, so that biological growth is dictated by carbon supply. For plants, this approach contrasts with the existence of nonstructural carbon, which leads to observed phenomenon like delayed growth under nutrient limitation (Li et al., 2021; Boussadia et al., 2010), or coarse root growth during the dormant winter (Marchand et al., 2025). In the



**Figure 6.** Variation of total biomass CUE with respect to temperature for different models. Results were plotted using the last 40 years of the 50-year simulations (shown in Fig. S3).

relative demand configuration of ELM, in order to match observed nighttime root growth (when carbon supply from photosynthesis is zero), its source-driven approach forces the model to introduce a carbon storage pool that may often (unrealistically) become negative at night and be replenished by new photosynthates in the following daytime (Burrows et al., 2020). When these source-driven models are extended to include nitrogen and phosphorus regulation of plant growth, they may be forced to adopt the law of the minimum (Yang et al., 2014), making predictions that contradict the often-observed multiple nutrients co-limited plant growth (Fay et al., 2015). Additionally, the source-driven models may also be forced to adopt the carbon overflow mechanism under nutrient limitation and forbid luxury nutrient uptake under carbon limitation (Jarrell and Beverly, 1981), both artificially accelerating the carbon cycling. When microbes are also modeled with the source-driven approach, such as in the traditional CENTURY-like models (Koven et al., 2013; Parton et al., 1988), or the more recent MIMICS and Millennium models (Wieder et al., 2014; Abramoff et al., 2018), by in-

duction, they will also predict too-fast soil carbon cycling unless being compensated by physically unrealistic model parameter values.

It has long been advocated that plant growth is better modeled using the sink-driven approach, where photosynthesis product is first stored as nonstructural carbon, which is then mobilized to drive the growth of plant organs (Fatichi et al., 2014; Fourcaud et al., 2008). Our analysis here suggests that the Pirt and Compromise models, although treating growth as driven directly by substrate uptake, are able to produce very rich variation in microbial CUE when analyzed with respect to either specific growth rates or temperatures (Figs. 5a and 6a). Interestingly but not surprisingly, their predicted CUE patterns vary more widely than those predicted by the VIS, sDEB, and mDEB models (Figs. 5c and 6c), where structural biomass growth is sink-driven as fueled by reserve biomass, which is evolutionarily developed by biological organisms to stabilize their metabolic performance in the face of environmental fluctuations (Kooijman, 2009). From the greater CUE variation by the Pirt and Com-

promise models, we deduce that, due to the missing of buffering effect from the reserve biomass, the source-driven models may very likely overestimate the plant and microbial response to environmental change, such as elevated CO<sub>2</sub>, nutrient limitation or warming. (Some support is shown in Figs. S4 and S5, which show that by increasing the variability of carbon input and temperature, the source-driven models respond stronger.) Among the four sink-driven models, the VIS model considers maintenance respiration and growth as two-parallel processes of equal priority, while the sDEB and mDEB models regard maintenance respiration to have priority over growth. Thus, at the cellular level, the VIS model seems to be mechanistically less reasonable by triggering earlier death through maintenance and growth competition. The mDroop model represents biological growth as sink-driven only apparently (because it computes the carbon quota based on total biomass, rather than reserve biomass, to drive the growth), and predicted transient CUE dynamics that are quite different from the three truly sink-driven models (Figs. 5 and 6, and also Figs. S4 and S5). We believe that predictions by the mDroop model are likely to be more easily falsified empirically.

#### 4.3 The mDEB model seems to be superior for representing general biological growth

Our analysis indicates that only the sink-driven models (i.e. the VIS model and two DEB models) are able to capture the feature that structural biomass CUE first increases and decreases with specific growth rate as inferred from the synthesis of empirical observations (Lipson, 2015). Nonetheless, when exponential growth is considered, the VIS, sDEB, and mDEB models are found to be more consistent with the Odum-Pinkerton model (Odum and Pinkerton, 1955) that is formulated using non-equilibrium thermodynamics. Our numerical analysis here shows that the VIS, sDEB and mDEB models are equally good (as evidenced by their very similar performance for all cases analyzed here), because maintenance respiration is usually a small fraction of total respiration. As the VIS model was initially proposed to model plant growth (Thornley, 1972), we hypothesize that the sDEB and mDEB models would be comparably good for modeling plant growth. (Some prototype applications of the sDEB model to plant growth are in Russo et al. 2022; Kooijman, 2009.) However, if we consider previous comparisons between the sDEB and mDEB models for microbes (Tang and Riley, 2025), then the mDEB model has better mechanistic coherency with the first law of thermodynamics, and is numerically much easier to incorporate multiple nutrients co-limited growth. Therefore, mDEB model seems to be a better formulation for representing biological growth under general conditions. In particular, the mDEB model will enable the growth of plant shoots and roots to be modeled separately, while the two are modulated with shoot-plant exchange of water, carbon, and nutrients. Although the resultant model

may appear to be more sophisticated, counter-intuitively (as we observed in EcoSIM), it will be better constrained by using more directly measurable parameters (e.g. nutrient kinetic parameters, root morphological size and hydraulic resistance) and state and flux variables (e.g. phloem transport rate of carbon and nitrogen, root biomass and morphology profile). For instance, the coarse root dynamics can then be better represented, while the source-driven models forbid coarse root growth in the absence of photosynthesis, a structural deficiency unlikely to be compensated by parameter tuning.

## 5 Conclusions

By starting from non-equilibrium thermodynamics, we infer that under no change of internal energy storage density (as usually is the case for internal combustion engines), the power use efficiency of a thermal engine as defined by the ratio of useful power output and power input will first increase, then plateau and finally decrease with increasing input power. Following this, we deduced that (which was also inferred by Lipson (2015) based on a synthesis of empirical observations), for an exponentially growing population of cells, the relationship between structural biomass carbon use efficiency and specific growth rate should follow the same pattern. In our analysis of six biological growth models adopted in currently popular ecosystem biogeochemical models, we found that only the VIS, sDEB, and mDEB models correctly capture this relationship. Considering transient carbon inputs and temperatures, all six models suggest that there is not a static relationship of carbon use efficiency with respect to specific growth rate, substrate uptake rate, carbon inputs, or temperature. Therefore, considering that our previous analysis based on the sDEB model (Tang and Riley, 2015) showed that treating the temperature effect as a multiplier led to significant overestimation of warming induced soil carbon loss (see Fig. 3 there), we contend that, in order to robustly simulate the emergent carbon use efficiency dynamics and its consequent influence on ecosystem biogeochemistry, models should represent biological growth as sink-driven, and the effect of controlling factors like temperature and moisture should not be applied as multiplier functions, rather they should be applied to directly modify the kinetic parameters of different metabolic processes. Our analysis shows that the mDEB model is a good candidate framework to meet all these needs, and will potentially help ecosystem biogeochemistry models to improve their simulated variability of ecosystem dynamics in response to changing driving conditions.

## Appendix A: Nomenclature

**Table A1.** Definition of symbols used in the modified Odum-Pinkerton model and models in Table 1.

Symbol	Unit	Description
$c$	Application dependent	Parameter for the modified Odum-Pinkerton model
$c_1$	None	Scaling parameter for growth respiration in mDroop model
$c_2$	year <sup>-1</sup>	Scaling parameter for maintenance respiration in mDroop model
$f$	Application dependent	Parameter for the modified Odum-Pinkerton model
$h(S)$	None	Substrate response function in Pirt model and Compromise model
$j_A(S)$	year <sup>-1</sup>	Specific substrate uptake in the mDroop model, VIS model and DEB models.
$l$	Application dependent	Parameter for the modified Odum-Pinkerton model
$m_V$	year <sup>-1</sup>	Specific maintenance respiration rate for Pirt model, Compromise model, VIS model, and DEB models
$p$	None	Normalized input power in the modified Odum-Pinkerton model
$q(S)$	year <sup>-1</sup>	Specific substrate uptake rate in all growth models
$v_E$	year <sup>-1</sup>	Specific reserve mobilization rate in the VIS model and DEB models
$B_T$	gC m <sup>-3</sup>	Total biomass C for mDroop model
$B_X$	gC m <sup>-3</sup>	Reserve biomass C for VIS model, sDEB model and mDEB model
$B_V$	gC m <sup>-3</sup>	Structural biomass C
$J_1$	Application dependent	Input flux in the modified Odum-Pinkerton model
$J_2$	Application dependent	Input flux in the modified Odum-Pinkerton model
$P_1$	Application dependent	Input power in the modified Odum-Pinkerton model
$P_2$	Application dependent	Useful output power in the modified Odum-Pinkerton model
$Q_{\min}$	None	Minimum carbon quota for growth in the mDroop model
$Q_{\max}$	None	Maximum carbon quota for growth in the mDroop model
$R$	year <sup>-1</sup>	Specific respiration in the mDroop model
$S$	gC m <sup>-3</sup>	Substrate concentration in the growth models
$S_e$	J K <sup>-1</sup>	Entropy
$T$	K	Temperature
$X_1, X_2$	Application dependent	Forces in the modified Odum-Pinkerton model
$Y_{RV}$	None	Conversion efficiency from reserve biomass to structural biomass in VIS model and DEB models
$Y_S$	None	Substrate assimilation efficiency in all models
$Y_V$	None	Structural biomass carbon use efficiency in all models
$Y_B$	None	Total biomass carbon use efficiency in all models
$\epsilon$	Application dependent	Parameter for the modified Odum-Pinkerton model
$\eta$	None	Power efficiency in the modified Odum-Pinkerton model
$\mu_{\max}$	year <sup>-1</sup>	Maximum specific growth rate in the Pirt model, Compromise model and mDroop model
$\mu(S)$	year <sup>-1</sup>	Specific growth rate in all growth models

**Code and data availability.** The code and data used in this study are available at: [https://github.com/jinyun1tang/cue\\_paper](https://github.com/jinyun1tang/cue_paper) (last access: 10 June 2026; <https://doi.org/10.5281/zenodo.20629498>, Tang, 2026).

**Supplement.** The supplement related to this article is available online at <https://doi.org/10.5194/bg-23-3995-2026-supplement>.

**Author contributions.** JYT formulated the idea, conduct the analysis and wrote the paper. WJR, GM, ELB discussed the results and edited the manuscript.

**Competing interests.** The contact author has declared that none of the authors has any competing interests.

**Disclaimer.** Financial support does not constitute an endorsement by the Department of Energy and National Science Foundation of the views expressed in this study.

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**Acknowledgements.** The authors appreciate the careful review of the paper by the two reviewers and the handling editor, their suggestions greatly improved its clarity.

**Financial support.** This research has been supported by the director of the Office of Science, Office of Biological and Environmental Research, of the US Department of Energy (contract no. DEAC02-05CH11231) and the National Science Foundation (grant no. 2125069).

**Review statement.** This paper was edited by Bertrand Guenet and reviewed by Xianjin He and Stefano Manzoni.

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