REVIEW



Accelerating the development of a sustainable bioenergy portfolio through stable isotopes

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Abstract

Bioenergy could help limit global warming to 2°C above pre-industrial levels while supplying almost a fourth of the world's renewable energy needs by 2050. However, the deployment of bioenergy raises concerns that adoption at meaningful scales may lead to unintended negative environmental consequences. Meanwhile, the full consolidation of a bioenergy industry is currently challenged by a sufficient, resilient, and resource-efficient biomass supply and an effective conversion process. Here, we provide a comprehensive analysis of how stable isotope approaches have accelerated the development of a robust bioeconomy by advancing knowledge about environmental sustainability, feedstock development, and biological conversion. We show that advances in stable isotope research have generated crucial information to (1) gain mechanistic insight into the potential of bioenergy crops to mitigate climate change as well as their impact on water and nutrient cycling; (2) develop high-yielding, resilient feedstocks that produce

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high-value bioproducts *in planta*; and (3) engineer microbes to enhance feedstock conversion to bioenergy products. Further, we highlight knowledge gaps that could benefit from future research facilitated by stable isotope approaches. We conclude that advances in mechanistic knowledge and innovations within the field of stable isotopes in cross-disciplinary research actions will greatly contribute to breaking down the barriers to establishing a robust bioeconomy.

KEYWORDS

bioeconomy, conversion, environment, feedstock, stable isotopes, sustainability

1 | INTRODUCTION

Bioenergy is central to all socioeconomic pathways compatible with limiting global warming to 2°C above preindustrial levels (Fuss et al., 2014). However, concerns persist over large deployment of biofuels for climate mitigation planning on the grounds of uncertain environmental impacts, the effective delivery of carbon (C) savings and potential to meet energy production targets (Calvin et al., 2021; Field et al., 2020; Reid et al., 2020). Sustainability is broadly defined as development that meets the needs of the present without compromising the ability of future generations to meet their own needs (Brundtland, 1987). Granted the normative nature of the concept, a sustainable bioenergy would secure a sufficient source of renewable energy while minimizing the use of natural resources and impacts on the environment. On this basis, the consolidation of a robust bioeconomy compatible with mitigation ambitions requires: (1) a deep understanding of the interactions between the deployment of dedicated feedstocks and the environment, (2) the engineering of resilient, high-yielding feedstocks that use natural resources—water and nutrients—more efficiently, and (3) the optimization of the energy conversion process through rapid engineering of microbial strains that can efficiently produce diverse, high-value bioenergy molecules and specialty bioproducts.

As with most scientific disciplines, our ability to address critical gaps in knowledge in the field of bioenergy has often been constrained by tools available (Dawson et al., 2002; Ehleringer & Osmond, 2000). Arguably, stable isotopes have recently emerged as one of the most powerful techniques for breaking the barriers to measure otherwise elusive processes in situ. The leverage of stable isotopes lies on a relatively well understood behavior within biogeochemical, physiological, genetic and chemical frameworks, and while limitations exist in their large-scale and cost-effective application (Antoniewicz, 2021; Dawson et al., 2002; Ehleringer & Osmond, 2000), stable isotopes can provide crucial knowledge to accelerate the development of a strong bioeconomy.

Here, we review recent scientific advances that have used stable isotopes to address questions relevant to the effective consolidation of bioenergy within global developmental schemes. Using an integrative cross-disciplinary approach, we explore this through the lens of the inherent challenges associated to sustainable bioenergy systems within three overarching themes: (1) environmental sustainability, (2) feedstock development, and (3) microbial conversion (Figure 1). While relevant to the full assessment of the role of bioenergy in the broader context of sustainable development goals, this manuscript does not attempt to cover the intricacies of a complete life cycle analysis or the potential socioeconomic and political ramifications of bioenergy but to emphasize those aspects in which the use of stable isotopes have significantly contributed to advance science towards a robust bioeconomy. Within this framework, we analyze how stable isotopes can help ascribe causality and elucidate regulating mechanisms and interactions to target research for the improvement of the economic and environmental value of bioenergy feedstocks. Further, we highlight the knowledge gaps that could benefit from future research led by innovative stable isotopes approaches. Finally, we aim to open cross-disciplinary discussions regarding the advantages of an integrative framework across environmental sustainability, feedstock development, and microbial conversion for advances in bioenergy research.

2 | ENVIRONMENTAL SUSTAINABILITY

While designed to lower the pressure on climate, the large-scale deployment of bioenergy crops has been challenged over a potential acceleration of soil C losses and $\operatorname{non-CO}_2$ greenhouse gas (GHG) emissions associated to direct and indirect changes in land use, as well as increased stresses on our already scarce water and nutrient resources. Failing to bring the bioenergy sector to deliver a climate benefit that does not rely on the intensification of

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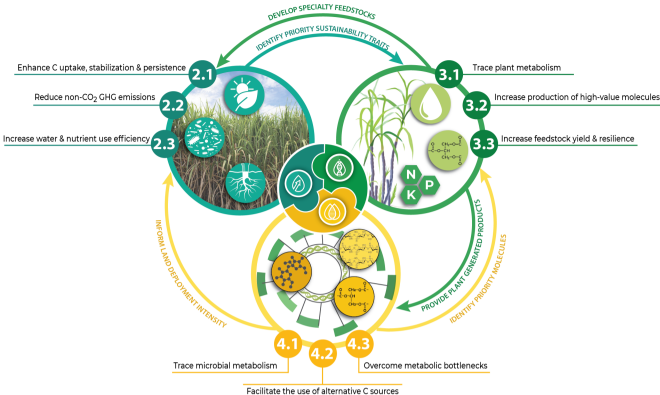


FIGURE 1 Challenges and cross-disciplinary research actions across environmental sustainability, feedstock development and microbial conversion for advances in bioenergy research. Within the context of this manuscript, environmental sustainability is defined as the degree to which bioenergy production may be effectively maintained and deliver sizeable a climate benefit while avoiding the long-term depletion of natural resources; and "land deployment intensity" is defined as how much land needs to be allocated to bioenergy based on feedstock productivity, the efficiency of conversion and target ethanol yields. Numbers link to specific challenges across environmental sustainability, feedstock development and microbial conversion; and arrows indicate cross-disciplinary research actions that integrate the use of stable isotopes.

the use of our natural resources could unintendedly push the Earth system closer to the environmental limits within which humanity can safely operate (Fuss et al., 2014; Hanssen et al., 2020; Heck et al., 2018; Smith et al., 2016).

Key questions remain on the degree to which bioenergy production may be effectively maintained and deliver sizeable climate benefits while avoiding the long-term depletion of natural resources, concept regarded here as environmental sustainability. In this context, beyond considerations of social equity, economic development and political implications, a sustainable bioenergy system, in addition to displacing fossil emissions should: (1) enhance the C uptake, stabilization and persistence of soil C pools (Section 2.1; Figure 1), (2) reduce non-CO₂ GHG emissions (Section 2.2; Figure 1); and (3) increase the water and nutrient use efficiency of the system relative to displaced land uses (Section 2.3; Figure 1).

In the following sections, we take a closer look at how the application of stable isotopes in the plant–soilatmosphere continuum provides key insights into achieving enhanced environmental sustainability of bioenergy cropping systems. We also identify areas in need of a deeper understanding that could benefit from the application of stable isotopes methods.

2.1 | Use of isotopes to investigate C capture, stabilization, and persistence processes

With proven potential for soil organic C (SOC) sequestration, bioenergy systems, particularly perennial lignocellulosic feedstocks, can significantly contribute to negative emissions and deliver significant GHG savings adding to the climate benefit of fossil fuel displacement and potential geologic carbon capture and storage (Harris et al., 2015; Ledo et al., 2020; Whitaker et al., 2018). However, soil type, climate, and previous land use and management can significantly influence SOC sequestration and the net GHG intensity of these systems, suggesting that not all bioenergy systems will likely deliver the savings targeted in some renewable fuel policies (Fuss et al., 2014; Kato

& Yamagata, 2014). Implied in this observation is an urgent need of a deeper understanding of the mechanisms enhancing the C capture, stabilization and persistence of bioenergy crops to confidently predict under which circumstances bioenergy systems will be most effective in decarbonizing the atmosphere.

At the ecosystem scale, a large fraction of C fixed by photosynthesis can be (1) allocated to above- and belowground plant biomass or respired back to the atmosphere by the canopy and roots (i.e., autotrophic respiration); (2) released via rhizodeposition (root sloughing, mucilage and exudates) to the rhizosphere; (3) incorporated into soil organic matter (SOM), including particulate organic matter (POM) and mineral-associated organic matter (MAOM) fractions; and (4) assimilated into microbial biomass or respired by the microbial community (i.e., heterotrophic respiration) (Figure 2). Accurately quantifying CO₂ capture, and C stabilization processes and persistence into biogenic pools, and identifying the mechanisms driving responses to a changing environment (e.g. crop species, soil type, climate and management) is challenging. The C metabolism across the soil-plant-atmosphere continuum

is defined by synergies and thresholds of mechanisms operating simultaneously and at different time-scales in a highly dynamic system. The natural abundance of C isotopes (¹²C and ¹³C) or enriched ¹³C labeling techniques allows tracing C dynamics through the atmosphere-plant-soil continuum in situ and non-destructively, providing a unique opportunity to assess the C budget of bioenergy feedstocks (Box S1; Figure 2).

Photosynthesis and respiration are key determinants of the overall ecosystem C balance of bioenergy cropping systems. Therefore, understanding the drivers of these processes is essential for improving predictions of the C balance of bioenergy systems across the landscape. Combined with micrometeorological techniques and soil gas exchange measurements, ¹³C provides a powerful tool for partitioning ecosystem CO₂ fluxes into source components (Griffis, 2013; Siebers et al., 2021; Voglar et al., 2019) (Table 1). While not yet widely implemented in bioenergy cropping systems, ¹³C-CO₂ exchange measurements can be used to directly partition net ecosystem exchange (NEE) into gross primary productivity (GPP) inputs and ecosystem respiration (Re) outputs (Wehr & Saleska, 2015).

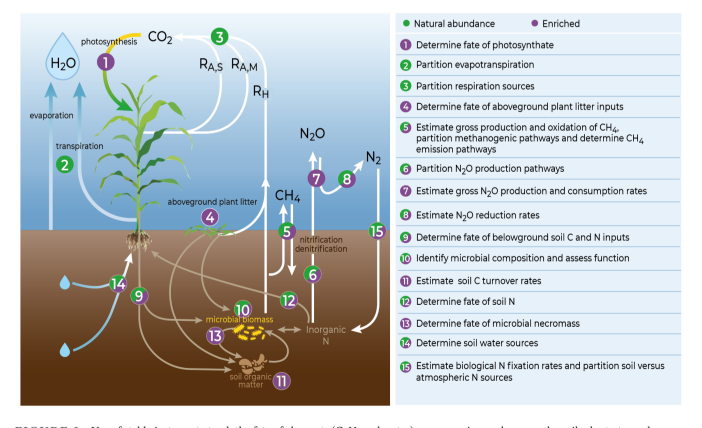


FIGURE 2 Use of stable isotopes to track the fate of elements (C, N, and water) among major pools across the soil–plant-atmosphere continuum, and to determine process rates, flux partition and function. Both natural abundance isotope (green) and enriched isotope (purple) methods used to identify, estimate and determine key processes and properties are shown. Numbers link to specific key processes and properties that determine the environmental sustainability of ecosystems. Arrows reflect water (blue) and C and N fluxes among soil pools (light brown) and with the atmosphere (white). Rh refers to heterotrophic respiration, and Ra,s and Ra,m to autotrophic structural and maintenance respiration, respectively.

TABLE 1 Bioenergy studies where stable C isotopes were used to address how changes in land cover and/or management practices affect C cycling, stabilization and persistence in soils and ecosystems. Soil priming refers to a change in decomposition of soil organic carbon (SOC) in response to fresh C inputs; soil microbial carbon use efficiency (CUE) refers to the amount of organic C allocated to microbial biomass production relative to C consumed and reflects the balance of anabolic and catabolic reactions. The table reflects major findings relevant to advancing science in the field of environmental sustainability through the use of stable isotopes.

Crop	Gained knowledge in bioenergy related with the use of stable C isotopes combined with other methods	References
Miscanthus and switchgrass	Switchgrass contributed a greater proportion of SOC in the root zone than <i>Miscanthus</i> .	Xu et al. (2022)
Miscanthus	Microbial CUE decreased with fertilization and was negatively correlated with root C concentration.	Kane et al. (2022)
Switchgrass	Increased bulk SOC over 10 years was positively associated with switchgrass-derived C across a marginally-productive landscape.	Zegada-Lizarazu et al. (2022)
Sorghum	Sorghum increased total SOC stocks in 2 years under both dry and wet conditions under free-air ${\rm CO_2}$ enrichment (FACE).	Leavitt et al. (2022)
Miscanthus	Miscanthus-derived SOC stocks increased as a function of time for at least 20 years following establishment.	Leifeld et al. (2021)
Miscanthus	Micro-scale SOC dynamics beneath <i>Miscanthus</i> are controlled by slow aggregate turnover, but aggregate turnover is faster on the soil surface compared to deeper soil.	(Vergara Sosa et al., 2021)
Corn, cover crops, fertilized and unfertilized prairie	The conversion of annual grain crops to mixed perennial systems increased the near- surface slow-cycling C pool, whereas the incorporation of cover crops increased the decomposition rates of both fast- and slow-cycling C pools and decreased their pool sizes across all depths.	Ye and Hall (2020)
Switchgrass	Deep-rooted grass cultivars can increase SOC relative to conventional crops in as little as 10 years, while expanding energy biomass production on marginal lands.	Slessarev et al. (2020)
Switchgrass	Extracellular polysaccharides played a prominent role in SOC stabilization.	Sher et al. (2020)
Pennisetum purpureum (Elephant grass)	Biochar increased SOC stocks relative to control in coarse textured soils.	Silveira et al. (2020)
NA	Biochar amendments enhance soil microbial CUE compared to straw-amended soils.	Liu et al. (2020)
Sugarcane	Priming of soil organic carbon induced by sugarcane residues and sugarcane biochar regulated the source of N for plant uptake.	Weng et al. (2020)
Sorghum	The belowground interaction of amended substrate and sorghum genotype influenced sorghum rhizosphere-associated exudates.	Miller et al. (2019)
Miscanthus and willow	Crop type determined the abundance and structure of soil communities as well as which soil resources (root exudates, dead organic matter or microbial derived compounds) were preferentially consumed.	Briones et al. (2019)
Sorghum bicolor	Sorghum root-derived C had higher biochemical recalcitrance which led to more C retention in soil than leaf residues. Root residues resulted in higher particulate organic matter and lower C persistent mineral associated organic matter than leaf residues.	Fulton-Smith and Cotrufo (2019)
Populus	Belowground C inputs contribute the most to soil C accumulation compared to aboveground C inputs. These findings are important in the context of bioenergy crops as aboveground C inputs are often removed for bioenergy production.	Berhongaray et al. (2019)
Miscanthus, willow	Following conversion, the sequestration of soil C under genotypes of willow and Miscanthus is substantial, and fresh C derived from cropping also enhances deep soil C accumulation.	Gregory et al. (2018)
Cover crop in a maize plantation	Cover crop belowground C inputs contributed the most to soil C accumulation compared to aboveground C inputs, and C derived cover crop root was most abundant in the mineral-associated fraction, deemed to be the most persistent fraction in soil.	Austin et al. (2017)
Miscanthus and willow	Crop type determined the dominance of ectomy corrhizal fungi vs bacteria, which in turn affected soil CO_2 fluxes.	Elias et al. (2017)

(Continues)



TABLE 1 (Continued)

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Crop	Gained knowledge in bioenergy related with the use of stable C isotopes combined with other methods	References
Miscanthus	Fresh C inputs do not enhance SOC accumulation after 7 years of <i>Miscanthus</i> growth, and this phenomenon is not explained by soil priming accelerating the decomposition of old SOC.	Robertson et al. (2017)
Switchgrass ecotypes	Variation between switchgrass ecotypes could alter microbial communities (i.e., bacteria vs. fungi) and impact C sequestration and storage as well as potentially other belowground processes.	Roosendaal et al. (2016)
Miscanthus, switchgrass, semi-perennial crops (fescue and alfalfa), annual crop (sorghum and tricale)	New SOC accumulation was higher under semi-perennials than under perennial crops due to variations in C input from crops rather than decreases in mineralization rates. N fertilization applied during the establishment and maturity phases did not impact SOC accumulation.	Ferchaud et al. (2016)
Switchgrass cultivars	Initial (<3 years) soil C losses due to land conversion were quickly offset by the accumulation of newly C from switchgrass. Cultivars with the highest SOC stocks tended to have the highest C concentrations in the persistent mineral soil fraction rather than in the particulate organic matter fraction.	Adkins et al. (2016)
Switchgrass	The ratio of root exudate C inputs to total soil C influences priming effect. The impact of changes in exudate C inputs on the priming of SOC differs in shallow versus deep soil.	de Graaff et al. (2014)
Land conversion to several Miscanthus hybrids and Miscanthus×giganteus	SOC stocks do not change significantly after 6 years of Miscanthus growth, but greater initial SOC decreases were observed in genotypes with higher belowground productivity likely due to enhanced priming effect.	Zatta et al. (2014)
Maize and soybean	Maize and soybean showed different daily patterns of net ecosystem exchange, gross photosynthesis, and respiration using isotope flux partitioning.	Fassbinder et al. (2012)
Miscanthus×giganteus, Cryptomeria japonica	The C input and recalcitrance of <i>Miscanthus</i> × <i>giganteus</i> (a fast-growing grass) litter was higher relative to <i>Cryptomeria japonica</i> (a fast-growing conifer) suggesting longer persistence of C under the studied grass system compared to the forest plantation.	Toma et al. (2012)
$Miscanthus \times giganteus$	A study of the priming effect; decomposition of old SOC was induced only when fresh C exceeded C content in soil microbes	Blagodatskaya et al. (2011)
Switchgrass	After 5 years of cropping, SOC stocks in low C soils increased substantially due to newly C derived from switchgrass	Collins et al. (2010)
Maize	Plant activity regulated root rhizosphere respiration.	Rochette et al. (1999)

For example, under controlled conditions, a recent study demonstrated that peak daily GPP was about 50% greater in maize than soybean, but peak daily Re was similar between species, resulting in greater NEE in maize than soybean (Fassbinder et al., 2012). Compared with isotopic partitioning of NEE, partitioning of soil respiration into autotrophic (root and rhizosphere) and heterotrophic (microbe and fauna) components via ¹³C has been more widely used (Table 1; Box S1). A study (Rochette et al., 1999) used natural abundance ¹³C in a maize (C4) system planted on a C3-derived soil to show that autotrophic respiration was driven primarily by changes in plant photosynthetic activity whereas heterotrophic respiration varied with soil temperature. Although technical limitations still exist (e.g., cross-sensitivity between target gases, concentration dependent non-linearities and ¹³C-CO₂ values, or the temperature sensitivity of the analyzer stability), recent

improvements in situ ¹³C-CO₂ instrumentation such as the development of cavity ring-down spectroscopy (CRDS) and integrated cavity output spectroscopy (ICOS) technologies for continuous, high precision measures, coupled with decreasing costs (Griffis, 2013; Voglar et al., 2019) will help facilitate future studies of ecosystem CO₂ fluxes in bioenergy systems.

Agricultural activities have historically caused the loss of up to 133 Pg C from soils globally, the rate of loss increasing dramatically in the past 200 years (Sanderman et al., 2017). Thus, building soil organic matter (SOM) provides a key step toward climate stabilization (Smith et al., 2016). While ecosystem scale measurements can paint a broad picture of C inputs, outputs and balance, understanding the detailed dynamics of SOM storage requires more targeted approaches. Stable isotopes can help to identify predominant SOM inputs, assess the relative

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persistence of SOM, and trace the interactions between plant and microbial components (Table 1). For instance, Berhongaray et al. (2019) used natural abundance ¹³C to show that belowground rather than aboveground inputs dominated the SOM pool of a poplar plantation. In another study, Austin et al. (2017) used ¹³C-CO₂ pulse labeling to demonstrate that rhizodeposition from a rye cover crop accounted for a third of total belowground inputs in a maize system. Because SOM is known to comprise different pools of varying recalcitrance and vulnerability to decomposition (e.g., particulate organic matter vs. mineral-associated organic matter), it is critical to elucidate links between in-plant C allocation to contributions to SOM pools and persistence in soils. In this light, Fulton-Smith and Cotrufo (2019) found that root litter was less effective at forming more stable mineral-associated SOM compared to aboveground litter in a sorghum system, highlighting the need to consider residue management when targeting SOC storage.

Also relevant to the vulnerability of recently fixed and old organic C pools is the residue-induced priming of SOC, defined as the change in the mineralization rates of SOC in response to fresh SOM inputs. A major challenge in investigating the mechanisms of priming is the partitioning of CO₂ emissions from multiple sources in a highly dynamic system (i.e., root respiration and the mineralization of labile and stable SOC). The combination of natural ¹³C abundance and ¹³C-labeled approaches have been proven instrumental in identifying the mechanisms of priming in response to fresh organic inputs and the environment. For example, Hall et al. (2019), capitalized on the distinctive natural abundance of ¹³C in C3 and C4 plants to reveal that soybean residue stimulated microbial growth and favored priming over the following corn phase in cornsoybean rotations. In another study, Weng et al. (2020) used a dual-isotope approach with two levels of ¹³C enrichment to show that in-soil incorporation caused higher positive SOC priming (acceleration of mineralization rates) than surface applied residue in a sugarcane system. Further, Weng et al. (2020) showed that in-soil biochar incorporation led to negative SOC priming (deceleration of mineralization rates).

Growing evidence reveals a prominent role of the microbiome on the regulation of the SOC budget. The contribution of microbial necromass to the total SOC pool is greater than previously anticipated, and microbial ecology (i.e., identification and function) has been proven critical to the stabilization and persistence of C in soils (Li et al., 2021). By enabling in situ C tracing, stable isotopes offer a unique opportunity to ascribe functionality and unveil the role of distinctive microbial taxa on C sequestration and stabilization in soils. For instance, using a combination of differences in the natural abundances of ¹³C

and ¹³C-pulse chase approaches, recent studies showed that Miscanthus led to greater ¹³C enrichment in bacterial phospholipid fatty acids (PLFA) whereas willow allocated greater C to ectomycorrhizal fungi and more shallow roots (Briones et al., 2019; Elias et al., 2017). These observations accompanied less ¹³C-CO₂ released in *Miscanthus*, casting some light into the role of fungal and bacterial communities in the microbial C assimilation and SOC stabilization pathways of bioenergy systems (Table 1). In another study using stable isotope probing of PLFAs (Box S2), Roosendaal et al. (2016) showed that switchgrass root architecture can influence fungal versus bacterial activity, which in turn affects SOC sequestration. Sher et al. (2020) used a ¹³C-CO₂ tracer to show that switchgrass stimulated microbes producing more extracellular polymeric substances, which have been linked to more soil aggregation and thereby increases in SOC. In another recent study, Ridgeway et al. (2023) using ¹³C labeled litter coupled measures of nitrogen (N) mineralization, showed that Miscanthus roots primed microbes to extract N and stabilize SOM. Therefore, ¹³C-CO₂ labeling experiments can help elucidate the interdependencies between crops species, microbial C turnover and soil C stabilization, paving the path to identifying desired traits for the development of a sustainable bioenergy portfolio (Table 1).

Results from some stable C isotopes studies have already been incorporated into modern bioenergy models, such as FUN-BioCROP or DAYCENT-CABBI (Juice et al., 2022), and the significance of findings from ¹³C tracing research is increasingly recognized for benchmarking SOC dynamics in global models (Camino-Serrano et al., 2019). Nevertheless, additional empirical work in this field is direly needed to further constrain SOM dynamics across a wide array of scenarios.

2.2 | Use of isotopes to explore the regulation of non-CO₂ GHG (i.e. CH_4 and N_2O) emissions

Amplified global warming potential, high sensitivity to both biotic and abiotic factors, and a strong dependence on local parameters make unintended increases of non- CO_2 GHG from bioenergy cropping systems determinant of the capacity of these systems to mitigate climate change. Curbing methane (CH₄) emissions is critical to short-term actions for climate stabilization (Ocko et al., 2021). Bioenergy crops in temperate, semi-arid and arid climates are usually net sinks of CH_4 (Drewer et al., 2012; Gauder et al., 2012; Oertel et al., 2016; Toma et al., 2011). However, increasing trends in the frequency of flash flood events and the expansion of bioenergy crops into marginal land and tropical and subtropical regions could decrease or

reverse this sink potential (Denmead et al., 2010; Gomez-Casanovas et al., 2018; Weier, 1999) (Box S3).

The net emission of CH₄ to the atmosphere results from a dynamic equilibrium among CH₄ production (methanogenesis), CH₄ oxidation (methanotrophy), plant-mediated CH₄ transport and physical processes (i.e., ebullition and diffusion) (Bridgham et al., 2013). Further, methanogens produce CH₄ by two major pathways, acetate fermentation and CO₂ reduction. All these processes operate simultaneously, and their tight interdependencies and distinct kinetic efficiencies and sensitivities to both biotic (e.g. predominant vegetation, substrate quality and availability, microbial abundance and distribution) and abiotic (e.g. soil volumetric water content and air-filled pore space, and soil redox, pH and temperature) factors make the identification and quantification of individual contributors to net CH₄ fluxes crucial, particularly in changing environmental conditions (Zhang et al., 2021). For example, fertilizers and high soil pH can either increase or decrease CH₄ emissions in response to changes in water management, fertilizer type, and the baseline N status of ecosystems (Blanc-Betes et al., 2016; Gomez-Casanovas et al., 2016, 2020). The in situ assessment of CH₄ production, oxidation, and transport rates with minimal or no disturbance is therefore critical to determine the net CH₄ source or sink strength (Banger et al., 2012) (Figure 2).

Studies using natural abundance of ¹³C-CH₄ in bioenergy crops to partition net CH₄ emissions into gross fluxes (oxidation vs. methanogenesis) are rare, particularly in marginal lands and in tropical and subtropical regions (Table S1). The stable isotope trace gas pool dilution technique, an iterative model that solves for gross production rates based on the isotopic dilution of the isotopically enriched chamber headspace pool of CH4 by natural abundance CH₄ emitted by the soil, is an underutilized approach to the partitioning gross CH₄ fluxes. Using this technique, Yang and Silver (2016) revealed substantial gross CH₄ production and oxidation occurring within the soils of corn fields with negligible CH₄ emissions. Evaluations of the predominance of the CH₄ metabolic pathways (i.e. acetate fermentation and CO2 reduction) in bioenergy systems, typically determined using natural abundance C isotopic composition of CH₄ and CO₂ in surface emissions and soil water pores, are also scarce (Table S1). Similarly, predominant CH₄ transport mechanisms and contributions to emitted CH₄ from bioenergy systems may be elucidated based on the discriminations against the heavy isotope (13C) of the different transport pathways (Table S1). A recent study using natural abundance of 13C-CH₄, found substantial CH₄ transport through *Populus spp.* stems, whereas transport through leaves was negligible, improving the accuracy of predictions of CH₄ emissions from this crop (Kutschera et al., 2016). Studies focusing on gross CH₄ production,

oxidation, and transport pathways could be extremely valuable to inform climate smart decisions on the deployment and management of bioenergy systems to reduce the CH₄ footprint of bioenergy systems in a climate change context.

Agriculture accounts for over half of anthropogenic N_2O emissions to the atmosphere (Tian et al., 2020). Understanding of mechanisms regulating soil-atmosphere N₂O fluxes is therefore, critical to policy making and management decisions aimed to minimize a collateral forcing on climate that may significantly reduce the efficacy of biofuels to abate global warming (Reay et al., 2012). With lower fertilizer requirements, prioritizing biomass- over grain-based feedstocks could significantly reduce the N₂O feedback from the bioenergy sector (Oates et al., 2016; Ruan et al., 2016), but further emission reductions could potentially be achieved (Table 2). Designing high-yielding bioenergy landscapes with minimal soil N₂O emissions requires resolving the partitioning of N₂O production and reduction metabolic pathways and regulatory mechanisms (Shan et al., 2021; Yu et al., 2020; Zhu-Barker et al., 2015). This can be achieved using both labeling and natural abundance stable isotope approaches (Figure 2).

Unlike more qualitative source partitioning approaches, stable isotope techniques can avoid the uncertainty inherent in direct measurements to accurately identify the dominant processes driving N₂O production. For example, N₂O producing metabolism may be partitioned into nitrification and denitrification, by tracking labeled ¹⁵N recovery in ¹⁵N₂O from ¹⁵N-labeled ammonium, a dominant substrate for nitrification, relative to ¹⁵N-labeled nitrate, the main substrate for denitrification (Krichels et al., 2019; Morse & Bernhardt, 2013). While labeling experiments are useful for identifying predominant pathways and potential activities, trace substrate additions may alter microbial function and limit experimentation in uncontrolled environments (Yang et al., 2014). Natural abundance stable isotope approaches avoid disturbance allowing for the direct determination of in situ activities under field conditions. Most promising is the use of N₂O isotopomers, the difference between $\delta^{15}N$ of the alpha and beta N atoms in N₂O (i.e., the central or terminal position on the asymmetrical N₂O molecule), to differentiate between bacterial denitrification versus nitrification and fungal denitrification (Ostrom & Ostrom, 2017). This approach has revealed that denitrification prevails over nitrification as the N₂O source in switchgrass soils (Ostrom et al., 2021), directing mitigation efforts to manipulating conditions for the anaerobic, heterotrophic process of denitrification rather than the aerobic, chemolithoautotrophic process of nitrification. Molecular approaches, such as functional gene transcript analysis, used in conjunction with stable isotope approaches can potentially reveal the importance of other microbial N₂O source processes, such

TABLE 2 Strategies to reduce soil N_2O emissions. Both tracer and natural abundance stable isotopes are used to partition the many microbial and abiotic processes involved in N_2O production and consumption in soil, which is necessary due to the differing controls on these processes.

	Strategies to reduce soil N_2O emissions	References
Management practices	Using alternative fertilizers, designed to release N gradually during periods of plant growth	Shoji et al. (2001)
	Enhancing the synchrony between N applications and crop N demands	Venterea et al. (2012)
	Optimizing the physical placement of fertilizer	Nkebiwe et al. (2016)
	Inhibiting soil N transformations with chemical additions or with precision application of fertilizer	Paustian et al. (2016)
	Using soil amendments including biochar and basalt	Blanc-Betes et al. (2020); Gomez- Casanovas et al. (2021)
	Using species with root exudates containing biological nitrification inhibition compounds (BNIs)	Luo et al. (2018)
Bioengineering and microbial inoculants	Manipulating plant-microbe interactions including BNI enabled species	Calvo et al. (2016); Subbarao et al. (2021); Usyskin-Tonne et al. (2019)

as dissimilatory nitrate reduction to ammonium, that cannot be differentiated from denitrification and nitrification based on stable isotopes alone.

Soil N₂O reduction rates are notoriously challenging to quantify (Groffman et al., 2006) but recent advances in stable isotope approaches are beginning to provide muchneeded field estimates of these rates that are lacking across all upland terrestrial ecosystems (Almaraz et al., 2020). The stable isotope trace gas pool dilution technique has been used for in situ surface flux measurements of gross N₂O production and consumption (Yang et al., 2011), showing that on average, roughly one-third of N₂O produced in a well-drained agricultural field can be reduced to N₂ before emitted to the atmosphere (Yang & Silver, 2016). Natural abundance stable isotope approaches for measuring in situ N₂O reduction rates have more recently been developed, for instance using clumped isotopes of ¹⁵N₂ along soil profiles (Yeung et al., 2019). Increased access to the instrumentation and training to use these advanced approaches will broaden their use across ecosystems, including in bioenergy cropping systems (Almaraz et al., 2020).

2.3 | Use of isotopes to examine the use of water and nutrient resources at the ecosystem scale

2.3.1 | Use of isotopes to investigate the use of water resources

Rising water demand and reduced supply is increasing global water stress, which can be further aggravated by climate change. At present, agriculture consumption represents

87% of global freshwater use (D'Odorico et al., 2020; Wu et al., 2022). Even limiting deployment to unirrigated bioenergy would increase evapotranspiration (ET; i.e. water loss at the ecosystem scale) by ~5% over the average short vegetation, leading to an additional water use of ~3% of global agricultural water appropriation for a targeted minimum deployment of biomass production compatible with <2°C scenarios (Smith et al., 2016; Wu et al., 2022). Moreover, high biomass targets would require the expansion into irrigation zones more than doubling agricultural water withdrawals (Heck et al., 2018; Stenzel et al., 2021).

ET is defined by two fluxes, plant transpiration (T) and evaporation (E) of water from plant surfaces and the soil (Bernacchi & VanLoocke, 2015). A key metric of the sustainable use of water resources is the water use efficiency (WUE; i.e. amount of C assimilated per unit of water lost) (Table 3). Among the diverse WUE metrics, a crucial parameter is the T to ET ratio (Table 3), that can be enhanced by genetic improvement or by adopting practices that reduce E and divert more water into T (Table 4). Given the strong link between T and plant productivity and the role of ET in determining water loss and availability, achieving a sustainable bioenergy portfolio requires understanding how a variety of bioenergy crops and management strategies affect both the T and E components of ET, and how these components respond to changes in climate (Kool et al., 2014; Xiao et al., 2018).

Water isotopes provide invaluable information for a more efficient use of freshwater resources. They can be used to directly partition ET in situ without disturbing the normal functioning of an ecosystem given the difference in isotopic fractionation between E and T (Rothfuss et al., 2021; Xiao et al., 2018) (Figure 2; Annex S1). While many studies have

TABLE 3 Summary of water use efficiency (efficiency in the use of water; WUE) metrics. The definition, methodology and description of each metric are also shown. Methodological approaches to measure water cycle related processes include biometric (i.e., productivity and yield), gas exchange and eddy covariance (EC) methods, and water stable isotopes (Annex 1).

Metric	Definition	Methodology	Description
Productivity or Yield, WUE_T	Aboveground productivity or yield: T	Biometric; stable isotopes	Emphasizes agronomic efficiency
			Reflects the consumption of water through plants
Canopy or Ecosystem, WUE_T	C exchange: T	Gas exchange; EC; stable isotopes	Emphasizes environmental efficiency
			Reflects the consumption of water through plants
Ecosystem WUE_{ET}	Gross Primary Productivity or Net Ecosystem Production: ET	EC	Emphasizes environmental efficiency
			Reflects the consumption of water through both plants and soil evaporation
Productivity or Yield, WUE_{ET}	Aboveground productivity or yield: ET	Biometric; EC	Emphasizes agronomic efficiency
			Reflects the consumption of water through both plants and soil evaporation
Biome WUE _{ET} (BWUE)	Net Ecosystem Production— Harvest: ET	Biometric; EC	Emphasizes environmental efficiency
			Reflects the consumption of water through both plants and soil evaporation

TABLE 4 Strategies to enhance the transpiration (T) to evaporation (ET) ratio by genetic crop improvement and by adopting practices that reduce E and divert more water into T. These strategies are not mutually exclusive, and their implementation to enhance water use efficiency (WUE) could be combined. T fluxes reflect the consumption of water through plants, and it is associated with productivity, whereas E does not contribute to it. Agrivoltaics refers to the novel strategy of collocating solar panels in agricultural land for the simultaneous production of energy and food.

	Strategy	Potential desired impact	References
Management practices	Crop residue management, mulching, row spacing	Reduce the amount of E	Farooq et al. (2019); Hatfield and Dold (2019)
	Soil fertility	Enhance the resilience of ecosystems to drought	Farooq et al. (2019); Kantola et al. (2022)
	Transformative strategies such as Agrivoltaics	Enhance WUE and decrease ET	Gomez-Casanovas et al. (2021)
		Tighter regulation of water if rainfall collected from the panels is routed to irrigated areas under or adjacent to the panels	
Genetic crop improvement of plant traits	Canopy size, leaf orientation morphology	Reduce the amount of E	Condon (2020)
	Root architecture and deeper root length	With deeper roots, plants can extract subsurface water available.	Condon (2020); York et al. (2022)
		With certain root architectures, the resistance to water movement decreases from soil to root (e.g. longer root hairs) and within the roots (e.g. larger xylem diameter).	

isolated the T flux in ET using water isotopes in cropping systems and grasslands, only a few focused on bioenergy crops including maize, *Populus* and sorghum (Rothfuss et al., 2021), suggesting that more studies are needed to improve the sustainable use of water by bioenergy crops. Using this isotopic approach, Lu et al. (2017) partitioned ET in an irrigated *Sorghum bicolor* plantation for bioenergy production. They found that of all the irrigated water, only 28%–39% was used by T, and that E accounted for as much

as 54% of ecosystem ET. These results suggest that management and bioengineering strategies to improve WUE in sorghum will play a major role in the sustainable use of water by this lignocellulosic biofuel system.

In addition to partitioning ET, stable isotopes can be used to better understand the source of the water consumed by plants as recently reviewed (von Freyberg et al., 2020), guiding both genetic engineering efforts and management decisions to optimize bioenergy use of water resources.

Under the assumption that the isotopic composition of xylem water reflects the mixture of water sources used by roots (Oerter et al., 2019; Penna et al., 2018), studies on this topic could resolve interactions between root architecture and drought tolerance, leading efforts towards the development of a sustainable bioenergy portfolio that builds resilience in the face of climate change (Joo et al., 2016).

2.3.2 | Use of isotopes to investigate the use of nutrient resources

Bioenergy crop production is plagued by the same tradeoff as food crop production. Routine biomass removal results in the depletion of soil nutrients and has a toll on soil health and future productivity. Fertilizer inputs to maximize yields leads to N losses that contribute to both air and water pollution (Tilman et al., 2011). Perennial bioenergy crops require less N inputs than annual crops due to nutrient retranslocation to the root system at senescence and reduced nutrient losses (Lawrence et al., 2021; Oates et al., 2016). However, the N mass balance of perennial bioenergy crops suggest greater N removals with harvest than inputs (Smith et al., 2013), indicating the need of additional fertilizer N inputs to sustain yields in the long-term (Sharma et al., 2022). Therefore, sustainable production of bioenergy crops requires strategies that minimize anthropogenic nutrient inputs.

Increased reliance on biological N fixation (BNF; i.e., atmospheric N2 assimilation into biologically useable ammonia by endophytic or free-living diazotrophic bacteria and archaea), internal nutrient cycling such as mineralization and translocation, and minimizing N losses due to leaching or gaseous fluxes can improve the sustainability of plant nutrient provisioning. There is a tradeoff between N mineralization and BNF as potential N sources for bioenergy crops. With 55-60 million tons of atmospheric N₂ fixed annually by agricultural crops, BNF provides a significant N supply to agroecosystems, replacing N lost through biomass removal or biogeochemical processes (Davies-Barnard & Friedlingstein, 2020). Contrary to N mineralization, which may deplete soil N over time, BNF can be considered a "renewable" source of N, that may be inhibited by other "nonrenewable" sources of readily available N (e.g., fertilizer or mineralized N). Increasing plant nutrient provisioning from BNF should be prioritized for long-term sustainability. However, the relative contribution of fertilizer N, N fixation, and N mineralization as sources of N to support the productivity of bioenergy crops is poorly characterized. Stable isotopes of N can be used for quantifying process rates to address these major gaps in understanding of N mass balances that underlie the N cycling of bioenergy cropping systems (Wewalwela et al., 2020).

Abundant indirect evidence suggests that BNF supports the productivity of perennial bioenergy grasses (Davis et al., 2010; Dohleman & Long, 2009; Heaton et al., 2004), but quantifying BNF at relevant time and spatial scales has proved challenging. The isotope dilution approach has been used in annual cropping systems to quantify BNF rates and monitor plant N assimilation dynamics over entire growing seasons following an initial isotope (15N) enrichment (Boddey et al., 1990; Iniguez et al., 2004; Rennie, 1982) (Figure 2). Plants that benefit from N₂ fixation will incorporate unlabeled atmospheric ¹⁴N₂, resulting in a dilution of the ¹⁵N-¹⁴N ratio of the enriched soil N pool, compared to plants without N₂ fixation. However, homogenous incorporation of ¹⁵N-enriched fertilizer to the volume of soil occupied by perennial grass systems is infeasible due to their extensive and heterogenous roots and rhizomes, leading to uncertainties in BNF estimates (Chalk, 2016; Keymer & Kent, 2014; Miranda et al., 1990; Miranda & Boddey, 1987). Approaches that take advantage of ¹⁵N natural abundance in soil (Fuertes-Mendizábal et al., 2018; Houngnandan et al., 2008; Urquiaga et al., 2012; Wewalwela et al., 2020) avoid the pitfalls of isotope tracer incorporation, but these methods have other limitations (Table 5). The identification of reference plants that do not benefit from N2 fixation is particularly challenging in non-legume systems, as many grasses benefit from associative or free-living diazotrophs. In this case, differences in yields could be wrongly attributed to endophytic N₂ fixation. Yield-independent isotope dilution models (Chalk, 2016; Witty, 1983) have been used to address this limitation in Miscanthus (Keymer & Kent, 2014), but this is not a panacea. Alternatively, BNF can be quantified by measuring incorporation of ¹⁵N into plant tissues or soil incubated with ¹⁵N-enriched dinitrogen gas (Chalk, 2016). However, the high cost and logistical constraints of this approach (Table 5) limits its use for fieldscale measurements and for capturing intra- and interannual variability in BNF, as observed in switchgrass (Roley et al., 2018, 2019; Wewalwela et al., 2020). Despite the challenges of using these stable isotope approaches for quantifying BNF (Chalk & Craswell, 2018), these remain the best approaches available and should be used more broadly to address the uncertainty in the current role of BNF in bioenergy cropping systems and to advance strategies to enhance BNF for sustainability benefits (Chalk, 2016).

3 | FEEDSTOCK DEVELOPMENT

The great challenge that lies ahead of feedstock development is the need to design crops to support an environmentally and economically robust bioeconomy. This entails selecting for and/or adding traits that improve yields while modifying the cell wall to be more easily converted to bioproducts

TABLE 5 Stable isotope methods for assessing biological N fixation (BNF) in bioenergy crops, rationale, benefits and limitations.

Method	Benefits	Limitations in bioenergy crops
¹⁵ N isotope dilution	Allows soil N sources to be distinguished from atmospheric N sources	Difficult to evenly incorporate label
Alters the isotopic signature of soil inorganic N (enriching ¹⁵ N) to allow source partitioning.		Label must be incorporated into all soil sampled by roots
N derived from the atmosphere (as ¹⁴ N ₂) will "dilute" out the ¹⁵ N signal from the soil		Isotope incorporation is especially challenging for established deep rooting perennial crops
derived N.		Challenging to identify reference plants that do not benefit from N_2 fixation
		Variance is large
¹⁵ N natural abundance	Allows soil N sources to be distinguished from atmospheric N sources	Challenging to identify reference plants that do not benefit from N_2 fixation.
Uses the different isotopic signatures of soil N sources (naturally ¹⁵ N enriched) and atmospheric N sources to allow source partitioning. N derived from the atmosphere (as ¹⁴ N ₂) will "dilute" out the ¹⁵ N signal from the soil derived N.	Does not require incorporation of ¹⁵ N label. Allows long-term studies	Variance is large, making it challenging to identify and measure N fixation
15N-enriched dinitrogen gas	Direct measurement of N inputs from atmospheric N allow ready assessment of BNF inputs	Limited to short term incubations
Alters the isotopic signature of atmospheric N ₂ (enriching ¹⁵ N) to allow source partitioning. Plants or soil benefiting from N fixation will be enriched in ¹⁵ N.		Challenging to deploy in the field Expense of ¹⁵ N ₂ -enriched gas Limitations to plant size Requires use of a gas-tight chamber

and introducing *in planta* production of high-value, energy-dense molecules including lipids (Amthor et al., 2019; Baligar et al., 2001; Gomez-Casanovas et al., 2007; Ort et al., 2015; Zhu et al., 2010). The development of optimal crops and specialty bioproducts requires developing state-of-the-art tools to trace photosynthetically fixed C into end-products (Allen & Young, 2020). In the next section, we focus on how stable isotopes can help us explore the plant primary metabolism (Section 3.1; Figure 1), a crucial first step to develop robust high-yielding bioenergy crops, and how they can be used to better inform the production of important bioproducts (Section 3.2; Figure 1) as well as to enhance feedstock productivity and resilience by improving water and nutrient use at the plant level (Section 3.3; Figure 1).

3.1 | Use of isotopes to explore plant primary metabolism

Mapping the flow of C through primary metabolism, including the Calvin-Benson-Bassham (CBB) cycle is

essential to determine allocation to bioenergy products. A suitable experimental approach, together with a powerful computational algorithm, have been developed in what is known as Isotopically Nonstationary ¹³C-Metabolic Flux Analysis (INST-MFA) (Jazmin et al., 2014; Young et al., 2011). INST-MFA is typically performed on photosynthetically active leaves under light. The experiment starts by changing the C source of a leaf from unlabeled to ¹³C-labeled CO₂. The ensuing transient in label signatures of CBB cycle intermediates is traced through a series of time points at which tissues are sampled and isotopomer signatures are measured in central metabolism intermediates. Since different pathways lead to different atomic rearrangements in the C chains of CBB cycle intermediates, flux-dependent transients in isotopomer signatures can be computationally resolved into a flux map representing the primary metabolism (Jazmin et al., 2014). As an example, using INST-MFA, Ma et al. (2014) revealed a tight regulation of photorespiration through acclimation to light levels on 4-week-old Arabidopsis thaliana labeled plants. Another study using INST-MFA on

Camelina sativa leaves attributed additional respiratory C losses to operation of the 6-phosphogluconate shunt (Xu et al., 2021)—a recently described metabolic cycle where photo-assimilated CO₂ is released by the oxidative pentose phosphate pathway (Sharkey & Weise, 2016). While the INST-MFA approach can produce a fairly detailed flux map of the central metabolism, it may also be the case that co-occurring fluxes can only be resolved with large statistical uncertainties in their values. Therefore, kinetic flux profiling (KFP) was developed as an alternative approach that is more explorative and makes use of the isotope tracer data to derive metabolic rates for individual reactions (Szecowka et al., 2013; Yuan et al., 2008).

Both INST-MFA and KFP approaches are well suited to monitor the primary metabolism under various physiological conditions on the premise that the cell population of the test sample is representative of one uniform metabolic state. For plants, which are complex multicellular organisms, this assumption may often not hold true. The tissue under study may present marked cellular and subcellular metabolic heterogeneity. If labelled transients from a labeling experiment result from the operation of distinct metabolic processes in different cell types, the analysis of tissue-extracted metabolites represents an averaged analytical readout and the flux modeling process might give a distorted or unreliable picture of cellular metabolism. Thus, the effects of metabolic heterogeneity must be diagnosed and considered by the metabolic modeling process in order to avoid misinterpretations (Chu et al., 2022; Szecowka et al., 2013). Cellular heterogeneity in metabolism can be resolved if the different cell types are separated prior to extraction and analysis of metabolites. For example, quite tedious non-aqueous fractionation techniques have been applied to separate fractions enriched in mesophyll and bundle sheath cells of C₄ plants prior to extraction to capture the pronounced metabolic differences between the two cell types (Arrivault et al., 2017). Alternatively, spatial heterogeneity of metabolism can be explored by mass spectrometry imaging (MSI) techniques (Boughton et al., 2016). If combined with isotope tracer techniques, MSI might allow developing a dynamic approach to analyze metabolic flux in situ with spatial resolution. Towards that goal, Romsdahl et al. (2021) presented a workflow combining isotope labeling of developing embryos of oilseed crops with MSI analysis of the major membrane lipid component phosphatidylcholine. Their analysis suggests different flux patterns of fatty acid elongation and desaturation in cotyledon vs. embryo axis tissue. In the future, the integration of ¹³CO₂ labeling protocols of photoautotrophic tissue and MSI techniques should improve our ability to spatially resolve metabolism in leaves and other photosynthetic plant tissues. Efforts to improve

bioenergy crops should benefit from the resulting refined understanding of multicellular plant systems.

3.2 | Use of stable isotopes to assess the production of high-value biofuel products

Much of the C fixed by photo-assimilation is programmed for storage in stems as low-value lignocellulose. There is a significant interest to re-route some of this C into high-value end-products. This is the case of energydense lipid-based biofuels that promote triacylglycerol (TAG) accumulation in plant biomass by the upregulation of lipid synthesis in vegetative tissues (Vanhercke et al., 2019; Xu & Shanklin, 2016). It has been estimated that if plants could be engineered to accumulate TAG in all the above-ground plant biomass at a 10% w/dw level, oil yields per acre could be substantially higher than achievable for any conventional oilseed crop (Ohlrogge & Chapman, 2011). However, vegetative plant tissues like leaves tend to have low intrinsic capacity to produce and store TAG at high levels (Chapman et al., 2013; Xu & Shanklin, 2016). Engineering strategies in the leaf must therefore aim at complex metabolic reprogramming. In fully grown mature leaves, CO2 is assimilated mainly into sucrose and starch (Figure 3). While sucrose is exported from the leaf to other tissues, most of the starch accumulated during the day is remobilized during the night to sustain energy demands in the dark (Figure 3). This transient diurnal accumulation pattern is often pronounced. Therefore, the engineering strategy could be to divert the C flow away from sucrose export towards oil synthesis (Figure 3). A recent theoretical modeling study on Sorghum bicolor found that, based on daily rates of net CO2 photo-assimilation observed for mature leaves under field conditions, TAG could accumulate to a 20% (w/dw) level in less than a month (Clark & Schwender, 2022). However, this goal could only be accomplished if 5% of the photo-assimilated C was diverted towards oil accumulation, and futile cycles of lipid biosynthesis and degradation operated on a massive scale, which would significantly impede the overall photo-assimilation (Clark & Schwender, 2022). In many cases, reported TAG levels are substantially below a 20% w/dw level, although some plants such as Nicotiana tabacum have been engineered to produce TAG levels above 30% w/dw level (Box S4). It can be expected that genetic manipulation of metabolic enzymes involved in TAG biosynthesis in leaves regularly results in unforeseen effects like lipid futile cycles. Nevertheless, we envision that isotopic tracer experiments will play a major role in diagnosing such roadblocks, allowing for further refinements of metabolic engineering strategies.

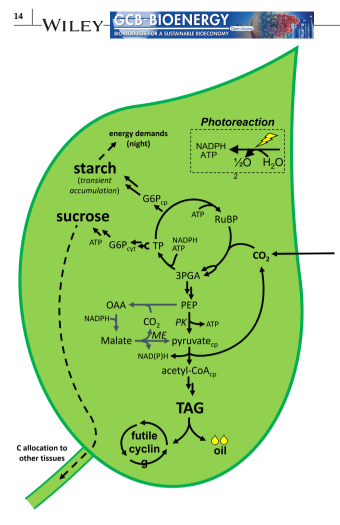


FIGURE 3 Assimilation of CO_2 via the Calvin-Benson-Bassham (CBB) cycle into sucrose, starch or TAG. Core reactions for these processes are outlined and the metabolic shunt for bypassing pyruvate kinase is shown in blue. 3PGA, 3-phosphoglycerate; G6P, D-glucose-6-phosphate; ME, malic enzyme; OAA, oxaloacetate; PEP, phosphoenolpyruvate; PK, pyruvate kinase; RuBP, ribulose 1,5-bisphosphate; TP, triose phosphates.

3.3 | Use of stable isotopes to investigate water and nutrient use efficiency at the plant level

As described above, water stress is increasing worldwide as a result of the large water footprint of agriculture—and other sectors—and shrinking water resources due to climate change. Increasing WUE at the plant level is therefore critical for the development of a strong bioeconomy. From a physiological standpoint, high WUE means that high rates of photosynthetic C fixation can be achieved with minimal water loss. Quantifying WUE requires direct measurements of transpiration rates with gas exchange methods. This approach, however, is not easily applicable on a high-throughput scale, which is crucial for fast identification of best-performing engineered crop

species (Ellsworth & Cousins, 2016; Vadez et al., 2014). Isotopes can help us with rapid quantification of plant WUE as shifts in the $^{13}\mathrm{C}/^{12}\mathrm{C}$ isotope ratio in leaf biomass relative to atmospheric CO2 reflect the plant water state serving as a proxy for plant WUE estimates (Ellsworth & Cousins, 2016; Seibt et al., 2008). While rapid WUE quantification method is yet to be applied in bioenergy research, it has been successfully used in conjunction with quantitative trait loci analysis to identify genes that promote enhanced WUE in cereals (Chen et al., 2011). The application of these emerging methods along with standardized approaches that facilitate the process of estimating WUE (Mathias & Hudiburg, 2022) could substantially expedite the identification of transgenic bioenergy crops for high WUE traits.

A major component of plant biomass and a key yield limiting factor is N. The engineering of dedicated crops with a more efficient uptake and utilization of N would reduce fertilization requirements and limit environmental pollution from nitrous oxides and nitrate leaching (Vitousek et al., 2009). Improving N use efficiency requires detailed understanding of the complexity of N metabolism at both cellular and plant scales (Beatty et al., 2016). To that end, ¹⁵N tracers can be used to trace N uptake and its movement among plant organs (Malagoli et al., 2005), and when coupled with ¹³C analyses they can provide a tool for metabolic profiling of distinct engineered phenotypes. For instance, Dersch et al. (2016) combined ¹³C and ¹⁵N tracers with subsequent labeling analysis of entire plants, plant tissues and metabolites to quantify assimilate movements between leaf and root. In this study, the design of the labeling chambers allowed tracing both elements in situ under controlled environmental conditions, providing a technology that allows for fast screening and metabolic profiling. Although this technology has not been applied to bioenergy crops yet, this approach could provide a tool for high-throughput plant metabolic phenotyping.

4 BIOLOGICAL CONVERSION

A major challenge in the field of conversion is to develop robust microorganisms for the efficient bioconversion of plant biomass into biofuels and bioproducts at large scale with improved energy yields and titer of products (Jullesson et al., 2015). In this light, enhancing the efficiency of the conversion process requires exploitation of novel microorganisms with less-understood metabolism as well as rewiring their native metabolism towards producing non-natural compounds. This often proceeds in iteration, where diagnosing metabolic bottlenecks in engineered cells informs next-round design. Further,

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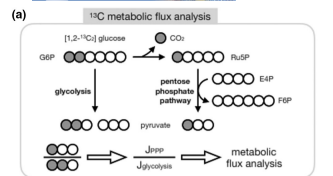
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bioconversion will also benefit from the efficient utilization of alternative C sources for the high-performance generation of value-added compounds (An et al., 2021). Stable isotopes are powerful tools that can contribute to achieving these two goals.

By rewiring the metabolism of yeast and other microorganisms, metabolic engineering facilitates the production of a greater diversity of high-value products. Engineering designs often build on a thorough knowledge of the microorganism's native metabolism, which makes model organisms successful microbial factories (e.g., Saccharomyces cerevisiae and Escherichia coli). However, wide diversity of less studied organisms may be better platforms for the production of certain chemicals. These microorganisms are increasingly accessible for metabolic engineering as genomic sequence and genetic editing toolboxes become available (Fatma et al., 2020). Understanding metabolic activity or flux (i.e. rate of substrate conversion to product per unit time) is the necessary first step in pathway design (Keasling et al., 2021). Metabolic activity, however, is not a physical entity which can be captured in a test tube and measured. Rather, it must be inferred, and isotope tracers combined with a suitable detection method for the labeled metabolites (e.g., Nuclear Magnetic Resonance or mass spectrometry) provide the most powerful tool for such inference (Figure 4a). We dedicate the following sections to explain how stable isotopes can help us understand metabolism of microorganisms (Section 4.1; Figure 1), measure activity in C co-utilization pathways (Section 4.2; Figure 1), and diagnose metabolic bottlenecks in engineered cells (Section 4.3; Figure 1).

4.1 Use of stable isotopes to infer metabolic microbial flux

Isotope tracing allows metabolic flux to be resolved accurately. The most common tracer to assess metabolic flux is ¹³C, in the form of ¹³C Metabolic Flux Analysis (¹³C MFA) (Box S5) (McAtee et al., 2015; Wiechert, 2001). A typical experiment involves culturing the microbes in ¹³C labeled C source(s) to a steady state, and then measuring isotope labeling in metabolites or proteinogenic amino acids by NMR or mass spectrometry. With proper tracer choice, valuable information on pathway usage can be obtained (Figure 4a) and used to constrain flux (Antoniewicz et al., 2007; Blank et al., 2005) either for a subset of metabolism of particular interest—typically central metabolic reactions—or at the genome scale (Gopalakrishnan & Maranas, 2015; Martín et al., 2015; Suthers et al., 2007).



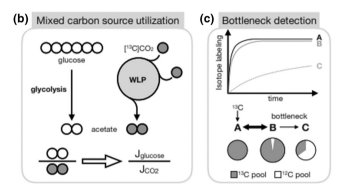


FIGURE 4 Isotope tracing in microbial metabolic engineering. (a) Isotope tracing reveals internal fluxes. A classic example is that [1,2-13C₂] glucose produces different isotope labeling in pyruvate, depending on whether it is metabolized through glycolysis or pentose phosphate pathway. (b) Isotope tracing reveals alternative carbon source utilization. For example, in mixotrophs, ¹³CO₂ tracing can reveal relatively how much CO₂ is fixed into a product like acetate. (c) Dynamic isotope tracing reveals metabolic bottlenecks. G6P, glucose-6-phosphate; Ru5P, ribulose-5-phosphate; E4P, erythrose-4-phosphate; F6P, fructose-6-phosphate; JPPP, pentose phosphate pathway metabolic flux; Jglycolisis, glycolysis metabolic flux; J glucose, glucose etabolic flux; JCO₂, CO₂ metabolic flux; WLP, Wood-Ljungdahl pathway.

The flux distribution in wild type cells provides the initial point for optimization algorithms for engineering design. These algorithms test deletion or activation of reactions (or combinations thereof) in silico and identify the most promising strategy for optimal production. Strategies for optimal production often eliminate competitive or redundant reactions but may occasionally involve nonintuitive mechanisms or suggest coupling target production to biomass generation thereby enabling growth-based lab selection and adaptation (Burgard et al., 2003; Patil et al., 2005; Ranganathan et al., 2010; Rocha et al., 2010). A notable example is severing gluconeogenesis by deleting phosphoglycerate mutase and thereby forcing CO₂ fixation as essential reaction for gluconeogenesis in E. coli (Antonovsky et al., 2016). The effectiveness of all these approaches is greatest when starting from a rigorous

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knowledge of the endogenous metabolism, enabled by isotope tracing.

Usually after testing the initial metabolic design, metabolic flux analysis can be a useful part of "learning" in the classical design-build-test-learn (DBTL) cycle of metabolic engineering to overcome the obstacles associated with driving biological systems to produce non-natural compounds. The DBTL cycle represents a framework that helps systematize metabolic engineering and increase its efficacy and generalizability. Potential next-round engineering targets can be identified by looking for reactions which flux is positively correlated with the improved production performance (McAtee et al., 2015). Following this principle, the oxidative pentose phosphate pathway was identified as a target for improving lipogenesis in an oleaginous yeast; pyruvate carboxylase for an engineered lysine producer; pyruvate kinase in cyanobacterial isobutyraldehyde production; and TCA cycle flux for protein production in Schizosaccharomyces pombe (Jazmin et al., 2014; Klein et al., 2014; Koffas & Stephanopoulos, 2005; Wasylenko et al., 2015).

4.2 | Use of stable isotopes to enable the utilization of alternative C sources

An important engineering objective, crucial to a sustainable bioenergy production, is enabling the utilization of C sources, other than glucose, that are less preferred, to enhance the microbial metabolic conversion process. With this context, stable isotope tracing can help us validate the assimilation of alternative C sources independent of benefits in growth. These less preferred substrates include abundant components in plant tissues such as lignocellulose and xylose.13C isotope tracing provides a powerful tool to monitor the in vivo activity of the alternative substrate utilization relative to glucose. Lignocellulosic hydrolysate is produced by enzymatic or thermochemical pretreatment of plant mass and contains a complex mixture of hexose (C6 sugars), pentose (C5 sugars), and acetate. In particular, xylose is the second most abundant sugar in lignocellulosic hydrolysates besides glucose, thus efficient xylose co-utilization is highly demanded (Stephanopoulos, 2007). The ¹³C labeled xylose and acetate have been used with glucose to confirm co-assimilation by the microorganisms, and further elucidate the pathway activity (Table 6).

Another important engineering objective is the development of conversion processes that assimilate ${\rm CO_2}$, motivated by the goal of developing C-neutral microbes. Tracing $^{13}{\rm CO_2}$ can be used to assess how much fixed ${\rm CO_2}$ is used for bioproduction compared to glucose (Figure 4b). Non-photosynthetic organisms cannot live on ${\rm CO_2}$ as the

6 Use of stable isotopes to establish native or engineered pathways for alternative C source utilization. TABLE

Reference	Aristilde et al. (2015); Liu et al. (2012); Xiong et al. (2015)	Wei et al. (2013)	Jones et al. (2016)	Antonovsky et al. (2016); Gleizer et al. (2019); Meyer et al. (2018); Yishai et al. (2017)
Relevance in advancing science in the area of conversion	Facilitated understanding of the native sugar catabolic pathway	Confirmed flux through (engineered) pathway during substrate co-feeding	Confirmed flux through (engineered) pathway during substrate co-feeding	Confirmed flux through (engineered) pathway during substrate co-feeding
Discovery	Phosphoketolase pathway has major contribution to pentose catabolism in addition to pentose phosphate pathway	Engineered acetate reduction pathway is active	${\rm CO}_2$ assimilation occurs in mixotrophy	Engineered CO ₂ , methanol or formate assimilation pathway operates in a non-C-fixation host
Method	Tracing $[1^{-13}C]$ xylose or $[1,2^{-13}C_2]$ xylose	Tracing ¹³ C-labeled acetate into ethanol	Tracing ¹³ CO ₂ into acetate	Tracing ¹³ CO ₂ , ¹³ C-methanol or ¹³ C-formate into biomass or central metabolic pathway intermediates
Microbe	Clostridium acetobutylicum and cyanobacteria	Engineered Saccharomyces cerevisiae	Clostridium ljungdahlii	Escherichia coli

Yishai et al., 2017).

only C source (autotrophy), hence CO₂ is often co-utilized with simple sugars (mixotrophy). In Clostridium ljungdahlii, a non-photosynthetic mixotrophy, CO₂ can be assimilated through the native Wood-Ljungdahl pathway (Jones et al., 2016). This occurs in parallel with glucose catabolism, and both processes make acetate as the final product. In this case, ¹³CO₂ tracing was used to confirm active CO2 assimilation, and reveal that CO2 has major contribution to acetate. It was further shown that coutilization of CO₂ improves the mass yield compared to glucose alone. Besides monitoring a native pathway in mixotrophs, CO2 assimilation has also been achieved through a non-native Calvin-Benson cycle (CBC) in *E.coli*, as revealed by ¹³C tracing into biomass components (Czajka et al., 2020). (Antonovsky et al., 2016; Gleizer et al., 2019). Similar to CO₂, 1C chemicals such as methanol and formate are also

Use of stable isotopes to overcome metabolic bottlenecks

promising renewable C sources as they can be effectively obtained by electrochemical reduction of CO₂. Therefore,

monitoring 1C assimilation with isotope tracing can guide pathway design and optimization (Meyer et al., 2018;

Use of C stable isotopes to overcome metabolic bottlenecks

Stable isotopes can also be used to reveal pathway mechanisms and overcome metabolic bottlenecks (i.e., key reactions that limit conversion efficiency) (Figure 4c). Production of the desired engineered chemical often involves many metabolic or transport reactions, and pathway flux is often limited by suboptimal operation of one or a few enzymatic steps (Figure 4c). Pre-steady-state isotope tracing, such as kinetic profiling or nonstationary metabolic flux analysis, is often applied to resolve metabolic flux when steady-state labeling is not informative, such as CO2 tracing in autotrophic organisms (McAtee et al., 2015; Young, 2014). The core concept is simple: all else being equal, fast labeling means high flux. By following labeling kinetics through a pathway, one could potentially identify where flux is blocked. An important caveat is that labeling into large pools is slow, and therefore it is important to figure out if apparent bottlenecks are due to actual flux impairment or just large metabolite pool size. Despite this complication, the approach has proven valuable for metabolic engineering. For example, kinetic ¹³CO₂ labeling with nonstationary ¹³C flux analysis in wild type cyanobacteria reveals a bifurcated tricarboxylic acid (TCA) cycle (Young et al., 2011), with low flux mainly serving the purpose of biosynthesis. To draw C from the



TCA pathways, Xiong et al., 2015 introduced an ethylene forming enzyme that also converts oxoglutarate to succinate, effectively producing ethylene and activating the TCA cycle as confirmed by kinetic ¹³CO₂ labeling. Kinetic ¹³C labeling has also been applied to reveal potential metabolic channeling, where metabolites are transferred between enzymes without mixing with the bulk pool, and pulse ¹³C labeling can also be used to reveal shift in pathway activity in the course of fermentation (Abernathy et al., 2019). For example, in an engineered strain that produces α-Ionone, the production is driven by glucose in the early phase of fermentation and shifts to re-assimilation of an overflow metabolite, mevalonate, in the later phase

Use of non-C stable isotopes to overcome metabolic bottlenecks

We have focused our conversion efficiency section on C stable isotopes. However, isotopes other than C have also been used to enhance the conversion efficiency of microbes. For example, deuterium isotopes are useful to reveal the metabolic sources of the key reductive cofactors, NADH and NADPH (Chen et al., 2019; Fan et al., 2014). Deuterium tracers are also well suited to quantify reaction reversibility, and thereby thermodynamics. For metabolic pathways to run efficiently, it is desirable for most steps to be moderately (e.g. 2-10 kJ/mol) thermodynamically forward driven. Smaller driving force may translate into high enzyme requirements and hence the cost of sustaining the desired forward flux. Canonical glycolysis contains several steps operating near thermodynamic equilibrium, but this is tolerated because those steps involve intrinsically very fast enzymes, while the Entner-Doudoroff pathway is a more thermodynamically favored glucose catabolism branch (Jacobson et al., 2019; Park et al., 2019). Larger driving force disadvantageously wastes energy, and isotope tracers can be used to diagnose engineered pathway energetics and thereby optimize them.

CROSS-DISCIPLINARY RESEARCH ACROSS **ENVIRONMENTAL** SUSTAINABILITY, FEEDSTOCK DEVELOPMENT, AND BIOLOGICAL **CONVERSION**

The grand challenge of achieving a robust bioenergy economy will only be possible with a shift towards an integrative framework that encompasses simultaneous advances to enhance the environmental sustainability of bioenergy

crops while developing highly productive, resilient and resource-efficient feedstock crops and optimizing biological conversion processes (Figure 1). We have shown that stable isotopes contribute feedstock development in the provision of plant-generated products, whereas new systems biology and synthetic biology approaches draw from isotopic tracing to inform developments in conversion efficiency. Further, stable isotope methods in biological conversion guide feedstock engineering targeting high-value molecules and inform land requirements for bioenergy deployment based on the assessments of bioconversion efficiencies of plant biomass into biofuels and bioproducts, and subsequent environmental evaluation (Figure 1). Finally, stable isotopes provide invaluable insight into the bioenergy interactions with the environment, identify critical tradeoffs and guide new avenues of research in feedstock development and conversion designs (Figure 1).

6 | OUTLOOK AND FUTURE DIRECTIONS

Stable isotopes have generated crucial discoveries urgently needed for the effective incorporation of bioenergy into global developmental schemes. They have been used to generate mechanistic understanding to evaluate the role of bioenergy crops as climate mitigation strategies, to assess the potential impacts of a large-scale deployment in the use of our already scarce natural resources (i.e., nutrients, water) and to identify research priorities that guide the development of a sustainable bioenergy and bioproducts portfolio. These powerful tools have also facilitated the development of optimized crops and specialty bio-products, for instance by engineering C allocation patterns to produce energy-dense lipids or increasing resource use efficiency at the plant level. Stable isotope tracing has also found broad applications in numerous non-model organisms and has inspired pathway design to enhance the microbial metabolic conversion process.

Challenges remain in the path toward a robust bioeconomy. There is an urgent need for in situ and undisturbed determination of the regulatory mechanisms behind the impacts of changes in crop choice, management practices, and climate on process rates (e.g., gross production, BNF), flux partition (e.g., evaporation vs. transpiration, nitrification vs. denitrification) and the fate of key elements in sustainability research (C, N, and water). With continuous high-frequency technology and improved instrumentation becoming more available, stable isotopes are deemed to continue playing a prominent role in the development of a sustainable bioenergy portfolio. Our current interpretation of the primary metabolism of plants is still constrained by technical difficulties and future advances

using a combination of stable isotopes and other techniques will expedite the development of higher-yielding bioenergy crops. Future research using stable isotopes will allow rapidly refining engineering strategies to re-route C in bioenergy crops to produce higher-value bioenergy products (e.g., TAG) and higher-nutrient and water use efficient crops. As more high-throughput engineering tools are made available, we envision a rapid development for isotope tracing techniques adapted for high-throughput analysis, thus accelerating the design-build-test-learn cycle. Finally, the advances that can be made for a robust bioeconomy by integrating the use of stable isotopes within a cross-disciplinary environmental sustainability, feedstock development, and microbial conversion framework are auspicious and deserve further attention.

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CONFLICT OF INTEREST STATEMENT

The authors declare no competing interests.

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