Opinion

Embracing 3D Complexity in Leaf Carbon–Water Exchange

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Leaves are a nexus for the exchange of water, carbon, and energy between terrestrial plants and the atmosphere. Research in recent decades has highlighted the critical importance of the underlying biophysical and anatomical determinants of CO2 and H2O transport, but a quantitative understanding of how detailed 3D leaf anatomy mediates within-leaf transport has been hindered by the lack of a consensus framework for analyzing or simulating transport and its spatial and temporal dynamics realistically, and by the difficulty of measuring within-leaf transport at the appropriate scales. We discuss how recent technological advancements now make a spatially explicit 3D leaf analysis possible, through new imaging and modeling tools that will allow us to address long-standing questions related to plant carbon–water exchange.

Why Does 3D Matter For Leaf Function?
The leaves of green plants are a striking example of the nexus of structure and function, and of physiology and environment. Here carbon, water, and energy dance through multiple phases, tissues, and scales in a complex 3D landscape that has evolved and diversified under selection for effective exchange in contrasting environments. Yet plant biologists have often resorted to simple leaf-scale models to study these processes – evading the difficult, confounding, and beautiful 3D reality.

It is now well established that CO2 supply to the sites of photosynthesis (i.e., inside chloroplasts) is limited not only by stomata, but also by a 3D network of resistances within the leaf, collectively termed mesophyll resistance; the pathways for CO2 diffusion are further complicated by (photo) respiratory CO2 release in the mitochondria. Yet the precise locations and dynamics of resistances within the mesophyll, their sensitivities to the internal and external environment of the leaf, and their influence on the regulation of net photosynthetic rate remain poorly understood. Similarly, recent research on the 3D architecture of leaf water transport has established the existence of large water potential gradients outside the leaf xylem minor veins. These gradients develop as a result of transport resistances within the leaf, are influenced by temperature and radiation absorption, and may contribute to stomatal dynamics and leaf vulnerability to dehydration.

Imaging and simulation technology can now reproduce the inner reality of the leaf in a 3D image (see Glossary) with a clarity and resolution inconceivable a generation ago (Figure 1). Continued progress in understanding how leaf structure affects function hinges on embracing the structural complexity of real leaves using technologies now widely available (Table 1). The

Highlights

Plant biologists have long resorted to highly simplified 1D or 2D imaging methods and modeling to study fundamentally 3D leaf processes of CO2 and H2O transport.

Recent advances in imaging and computational technology are enabling a data-rich scientific pipeline that integrates leaf 3D measurement, anatomical modeling, and biophysical simulation.

Adopting a 3D approach is not only critical for testing when dimensionality reduction is reliable and accurate, but also promises to deliver insights about: (i) fundamental processes of leaf CO2 and H2O transport and exchange, (ii) the translation of leaf anatomical diversity to functional diversity, and (iii) fine-scale CO2 and H2O exchange processes in broader-scale models.

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potential payoffs of a focused and coordinated effort on these problems are great, both for understanding how anatomical diversity translates into functional diversity and for harnessing that knowledge to improve the photosynthetic performance of crops.

**Limitations of a Bulk-Leaf Paradigm for Leaf CO₂ and H₂O Transport**

Photosynthesis, respiration, and transpiration are governed by biochemical and transport processes at several spatial scales within and among multiple tissue types throughout the leaf. Most commonly, however, the leaf has been implicitly treated as a single point with no spatial structure, representing a CO₂ sink during photosynthesis (or source during darkness) and an H₂O source during transpiration (or sink during dewfall). These point sources and sinks are connected to the atmosphere through resistances arranged in series (e.g., through stomata and the leaf boundary layer), and Fick’s first law of diffusion is used to calculate rates of gas exchange with the atmosphere. A biochemical model is used to simulate the photosynthetic CO₂ sink [1], and the H₂O source is modeled as a wet surface in equilibrium with intercellular air at the measured leaf temperature.

This approach of aggregating transport and biochemistry at the leaf scale has driven tremendous advances by providing a simple bridge between processes at the cell and tissue scales and observations at the leaf scale. However, a bulk-leaf approach precludes understanding how the geometry and biochemistry of different tissue types within the leaf influence CO₂ and H₂O exchange. The photosynthetic rate of a single chloroplast depends on light intensity, [CO₂], temperature, and photosynthetic capacity, all of which vary throughout the leaf because of the gradients generated by the interplay of transport with leaf structure. Thus, any single chloroplast is influenced by 3D leaf structure in ways that simple models cannot easily resolve [2,3]. Similarly, complex spatial gradients in metabolites, the allocation of resources such as nitrogen, and temperature and water potential can influence water transport, stomatal function, and metabolism [4–8].

**Spatial Aggregation Conflates Structural Features with Transport Processes**

Because leaf CO₂ transport and metabolism are not neatly organized into macroscopic compartments, leaf-scale models and measurements inevitably mask a great deal of structural and process complexity. For example, mesophyll conductance (gₘ), which describes 3D diffusional pathways between the intercellular airspaces and the chloroplast, is typically computed from bulk-leaf estimates of intercellular CO₂ concentration (cᵢ) chloroplast CO₂ concentration (cₘ), and leaf net CO₂ assimilation rate (A) [9] and traditionally thought to be strongly related to averaged mesophyll anatomy traits including cell wall thickness and chloroplast surface area [10]. **Apparent mesophyll conductance** [11; gₘ,app] is a better term because in real leaves, both cᵢ and cₘ vary considerably within a leaf and are dependent on the 3D distribution and biochemical properties of CO₂ sinks and sources [11–16] – features that can vary in ways not consistent with the conceptualization of gₘ as describing fixed pathways. As a result, some effects of environmental conditions on bulk-leaf gₘ,app may not reflect shifts in intrinsic transport properties, but may instead result from averaging fluxes and concentrations across the 3D leaf structure or from fine-scale positioning of CO₂ sources relative to sinks.

For example, an effect of irradiance on gₘ,app can emerge from changes in the contributions of different mesophyll layers to CO₂ uptake, despite constant transport properties in each layer [17], and chloroplast movement in response to light can alter gₘ,app by changing the spatial relationship between supply pathways and reactive demand for CO₂ [18]. Similarly, CO₂ released from mitochondria can be rebathed by chloroplasts, which can enhance gₘ,app and photosynthesis by providing a CO₂ source close to the photosynthetic sink [19]. Yet, the
likelihood of refixation depends on the relative positions of mitochondria and chloroplasts, which can shift over time [20,21]. Clearly these patterns and dynamics need to be accounted for in future models seeking to predict \( g_{m,\text{app}} \) at the leaf scale.

Similar issues arise in relation to functional traits other than \( g_m \). For example, a mismatch between light absorption and photosynthetic capacity among mesophyll layers can affect the response of CO\(_2\) uptake to irradiance and light quality [22–24] and hinder the interpretation of chlorophyll fluorescence [25]. Although transdermally explicit models of photosynthesis have been used to address these questions [26,27], the role of leaf anatomical diversity remains largely unexplored. In addition, the distribution of enzymes and metabolites influences metabolic fluxes measured at the leaf level [28]. A full understanding of photosynthetic function in intact leaves thus requires high resolution and/or spatially explicit treatments of CO\(_2\) transport to understand estimates of \( g_m \) using both the stable isotope and the fluorescence methods [2].

### Aggregating to the Tissue Scale Limits Understanding

Some important parameters that affect water transport in leaves remain poorly known, because they depend on fine-scale features that cannot be accurately measured with traditional imaging or experimental methods, and this hinders attempts to model water transport [6,7,29–31]. One example is tangential water flow through cell walls outside the xylem, which depends on anatomical features that are difficult to discern in 2D light micrographs, such as cell wall thickness and the location and extent of hydrophobic barriers to water flow such as lignin or suberin, and forms a major component of leaf hydraulic conductance. The lateral connectivity or contact area between cells also affects transport but is difficult to estimate accurately from 2D light micrographs, particularly for cells with highly variable shapes within the vascular parenchyma, bundle sheath, and mesophyll. Our ability to model water movement within the leaf xylem may also be improved by 3D approaches that capture the arrangement and connections between xylem conduits. The complexity of water movement through the xylem is increased as some conduits in the leaf vein network become gas filled (embolized) due to water stress [32].

Another process which depends on 3D tissue arrangement is water vapor transport, which is typically modeled by assuming the air is in chemical and thermal equilibrium with the nearest liquid water surface, generally a cell wall. This assumption implies that vapor concentration adjacent to the cell surface equals equilibrium vapor pressure of that surface, which can be calculated from its water potential and temperature [6,30,31]. However, because this ‘local equilibrium’ approximation neglects the vapor gradients that must exist across air-filled pore spaces between cells, it will tend to underestimate the total resistance to vapor transport within the leaf, especially as intercellular airspaces become large relative to cell size. Precise quantification of multidirectional vapor transport at these scales requires equally precise resolution of the actual tissue geometries. Indeed, interpretation of recent experiments using 0D models of gas exchange and isotopic enrichment has recently challenged the assumption [33] that the airspaces on the interior side of a stomatal pore are saturated with water vapor at the leaf surface temperature, a dogma that is generally used to define stomatal conductance in gas exchange systems. Fully 3D models of both processes, together with spatially resolved knowledge of cell wall material properties, could play a decisive role in resolving this controversy.

### Reducing Dimensionality Can Generate Bias

It is common to simplify descriptions of leaf anatomy to facilitate modeling and analysis. This simplification discards potentially important data, limiting insights and generating bias.
Intracellular organization

Cell wall, plasmalemma, cytosol, chloroplast envelope and stroma ultrastructure, organelle placement and biochemistry

Intercellular organization

Organelle and cell quantity, position, biochemistry, and dynamics

Tissue-level organization

Intercellular airspace (IAS) geometry; mesophyll surface area exposed to the IAS; stomatal–IAS linkage; epidermal anatomy vein geometry

Heat transfer
Light propagation
H₂O transport
CO₂ reaction–diffusion

Path length (μm)

0 150 15 μm

(See figure legend on the bottom of the next page.)
Table 1. Advanced 3D Technologies for Imaging Leaves

<table>
<thead>
<tr>
<th>Example Ref</th>
<th>Confocal/MP/LS microscopy*</th>
<th>X-ray CT*</th>
<th>SBF-SEM*</th>
<th>FIB-SEM*</th>
<th>TEM tomography</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample preparation</td>
<td>Live or fixed, fluorochrome labeling</td>
<td>Live or fixed, heavy-metal stain, whole mount</td>
<td>Fixed, heavy-metal stain, resin embed</td>
<td>Fixed, heavy-metal stain, resin embed</td>
<td>Fixed, heavy-metal stain, resin embed</td>
</tr>
<tr>
<td>Sample preparation time</td>
<td>0–2 days</td>
<td>0–2 days</td>
<td>7–14 days</td>
<td>3–5 days</td>
<td>3–5 days</td>
</tr>
<tr>
<td>Lateral resolution</td>
<td>200–300 nm</td>
<td>1–40 μm (μCT)</td>
<td>5–10 mm</td>
<td>2 nm</td>
<td>1 nm</td>
</tr>
<tr>
<td>Axial resolution</td>
<td>500 nm</td>
<td>1–40 μm (μCT)</td>
<td>30–50 nm</td>
<td>3–5 nm</td>
<td>2–3 nm</td>
</tr>
<tr>
<td>Field of view</td>
<td>160 μm to 4 mm</td>
<td>1–40 mm</td>
<td>50–300 μm</td>
<td>10–100 μm</td>
<td>0.5–10 μm</td>
</tr>
<tr>
<td>Sample size</td>
<td>1 × 1 mm</td>
<td>up to 50 × 50 mm</td>
<td>1 × 1 mm</td>
<td>10 × 10 mm</td>
<td>up to 200 × 200 nm</td>
</tr>
<tr>
<td>Sample thickness</td>
<td>~300 μm</td>
<td>1–50 mm (μCT)</td>
<td>600 μm</td>
<td>10 mm</td>
<td>&lt;200 nm</td>
</tr>
<tr>
<td>Imaging time</td>
<td>1–8 h</td>
<td>Minutes to days</td>
<td>1–5 days</td>
<td>1–10 h</td>
<td>1–4 h</td>
</tr>
<tr>
<td>Data set size</td>
<td>1–100 MB</td>
<td>5–50 GB</td>
<td>50–500 GB</td>
<td>~100 MB</td>
<td>~1 GB</td>
</tr>
<tr>
<td>Visible leaf structure</td>
<td>Mitochondria, chloroplasts, airspace network, cell, cell network, leaf</td>
<td>Cell, cell network, airspace network, leaf</td>
<td>Plasmodesmata, cell membranes, cell wall, mitochondria, chloroplasts</td>
<td>Plasmodesmata, cell membranes, cell wall, mitochondria, chloroplasts</td>
<td>Plasmodesmata, cell membranes, cell wall, mitochondria, chloroplasts</td>
</tr>
</tbody>
</table>

*Abbreviations: confocal/MP/LS microscopy, confocal, multiphoton, and/or light sheet microscopy; FIB-SEM, focused ion beam scanning electron microscopy; SBF-SEM, serial block face scanning electron microscopy; TEM, transmission electron microscopy; X-ray CT, X-ray computed tomography.

Mesophyll conductance is again an illustrative example. \( g_m \) includes resistances across intercellular airspaces [2], cell walls [34,35], cell and chloroplast membranes [36–38], and chloroplast stroma [34], and experimental measurements of \( g_m \) integrate across all of these elements. Although isotopic methods can help distinguish the components of \( g_m \) [39,40], there are no experimental methods to directly quantify their spatial variation, which must instead be inferred by applying anatomical measurements to spatially explicit models [25]. Such measurements are commonly simplified to bulk-leaf descriptors – for example, the 3D geometry of leaf intercellular airspaces is often summarized in a single scalar, the porosity (volumetric air fraction), which cannot describe how the multitude of 3D cell shapes and tissue geometries [41,42] impact path length, lateral diffusivity, tortuosity, and airspace connectivity [2,43], nor how the spatial distribution of stomata and mesophyll surfaces influences CO\(_2\) diffusion into chloroplasts [20]. The surface areas of mesophyll cells and chloroplasts exposed to airspaces also influence \( g_m \) [44] and vary considerably [45], but are commonly estimated from 2D sections
based on simple models for cell shape. Complex cell geometries can confound such estimates and complicate allometric scaling among cell dimensions [17,46].

**Stable Isotope Discrimination May Depend on 3D Leaf Anatomy**
Understanding how bulk-leaf stable isotope discrimination arises from diffusion and exchange at smaller scales would also benefit from a high-resolution, spatially explicit approach. Stable isotopes are important tools for plant physiologists as tracers of atoms through systems that record processes such as carboxylation, mesophyll conductance, and transpiration [47]. Theoretical models describing the underlying biophysics and biochemistry exist, but the most widely applied are spatially aggregated at the leaf level [48]. Examples of attempts to improve on this include adding a conceptual dimension by analyzing CO₂ fluxes in and out of the leaf in parallel [19], adding a spatial dimension by exploring radial isotope effects in leaf water [49–52], or adding a temporal dimension by probing nonsteady state leaf water isotope enrichment [53]. However, there remains no spatially resolved and anatomically accurate 3D model of stable isotope fractionation [54], so we have no means to quantify the influence of 3D anatomy on isotope processes. 3D transport of 13C CO₂ within leaves is of particular interest because one of the primary techniques to estimate gₑ requires a thorough understanding of carbon isotope discrimination within photosynthesizing leaves, and isotopic discrimination (during carboxylation, respiration, photorespiration, retrodiffusion, and refixation) would be influenced by the 3D structure of leaf airspaces, which governs the relationship between diffusion paths and exchanging surfaces, and hence the relative rates of gross and net exchange to and from those surfaces. Similar issues are inherent in interpretation of photosynthetic C¹⁸O₁⁶O discrimination [40] and leaf water isotopes [54].

**Toward a High-Resolution, Spatially Explicit Approach**
Applied models of plant-atmosphere CO₂ and H₂O exchange do not typically represent the leaf interior in a spatially explicit way, because it is impractical with current technology to parameterize and apply microscale spatially explicit models to address questions that integrate over large scales. This practical constraint should inform basic research that takes place within a spatially explicit paradigm (Box 1). In basic research, however, a mechanistically accurate understanding of leaf carbon and water exchange requires that we move beyond the spatially aggregated ‘bulk-leaf’ paradigm in imaging and modeling. Such a high-resolution 3D approach will improve the reliability of leaf-scale models, thus informing their application at larger scales.

An obvious first step is to adopt spatially explicit models in research. Many insights have been generated using such models, with simplified cell, organelle, and tissue geometries, often

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**Box 1. Scaling to the Leaf and Canopy Level**
Reliable prediction of global plant-atmosphere interactions [65] and crop yield [66] inevitably involves interaction of modeling efforts at different scales [67]. Complex, mechanistic models must ultimately be simplified for application at larger scales, and likewise, understanding gained from a 3D anatomically explicit paradigm for modeling leaf processes must eventually be shaped into a form that can be applied practically in the context of lower-dimensional or scaled models. Similar considerations have helped the application of leaf-level photosynthesis models to the crop canopy level as a useful tool for estimating consequences of photosynthetic perturbations at the biochemical level [22,68]. Analogously, upscaling of physiological processes that have been improved by 3D analyses and subsequently simplified for the desired purpose will benefit our understanding of their effects on the canopy scale. This perspective should inform research design; for example, in studies using fine-scale 3D-driven imaging and modeling, hypotheses should be framed in terms of anatomical parameters that can be linked to commonly measured leaf traits (e.g., leaf mass per unit area, leaf airspace fraction, or apparent mesophyll conductance). This ensures an immediate conduit for extension of new knowledge from the 3D paradigm to the vast body of existing leaf trait data [69]. Of particular benefit are analyses focused explicitly on anatomical scaling [70,71] and on strategies for mapping 3D transport processes to widely useful 1D models [72] or inferring 3D properties from 1D or 2D imaging data (e.g., tortuosity and porosity).
reconstructed from 2D cross-sectional images, to simulate light propagation [55], CO₂ reaction–diffusion [3,13,56–58], and leaf water and energy transport [6,30,31]. However, these approaches have typically depended on simplifications of anatomy that may affect model predictions. Thus, a critical and more challenging step is to directly generate 3D tissue models from imaging data, thereby circumventing the abstractions and loss of spatial resolution inherent in simplified models.

Recent advances in microscopy have streamlined the acquisition of 3D volumes at high resolution (e.g., serial block face scanning electron microscopy and focused ion beam scanning electron microscopy; Table 1), and modern computational tools are available for spatially explicit modeling of matter and energy transport (Table 2). Advanced imaging and computation can now quantify tissue geometry directly, alleviating the tedious of extracting measurements by hand from 2D micrographs and the uncertainty created by using assumptions about 3D geometry to infer tissue properties from 2D images. Such studies have indicated that 2D estimates of mesophyll surface area exposed to airspace can be 15–30% lower than 3D measures [59], and inclusion of 3D measurements of airspace tortuosity and lateral path lengthening reduced estimates of diffusional conductance within the intercellular airspace of bromeliad leaves by 37%, on average [60]. Visually segmenting and measuring distinct tissue types and cellular properties require measuring these traits at multiple scales with different microscopy methods (Table 1). Each method presents unique challenges, especially regarding specimen viability, but their spatial and temporal resolution have already delivered novel insight into the effects of 3D tissue and organelle arrangement on photosynthesis [61], viral transmission [62], and salinity stress [63].

Table 2. 3D Biophysical Leaf CO₂ and H₂O Transport

<table>
<thead>
<tr>
<th>Biophysical phenomena</th>
<th>Biophysical and computational model</th>
<th>Biophysical traits</th>
<th>Traits measured/inferred at the bulk-leaf level</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heat transfer</td>
<td>3D temperature distribution; depends on radiative, conductive, convective, and latent energy transfer and external and internal energy sources; interacts with light propagation; solved via heat transfer partial differential equations on geometric mesh or finite-element analysis</td>
<td>3D cellular and intercellular airspace network geometry, photosystem light absorption and heat dissipation, H₂O liquid–vapor transport patterns and interactions</td>
<td>Leaf temperature, leaf hydraulic conductance, apparent mesophyll conductance, respiration, photosynthesis</td>
</tr>
<tr>
<td>Light propagation</td>
<td>3D distribution of light intensity and spectral/directional quality; depends on internal absorption and scattering, and external light sources; solved via Monte Carlo ray tracing photon simulation on geometric mesh</td>
<td>Spatiotemporal distribution and geometry of organelles, cells, and tissues/airspace; optical properties of organelle, cell, and tissue material types with respect to light intensity and spectral/directional quality</td>
<td>Apparent mesophyll conductance; respiration; photosynthesis; reflectance, absorptance, and transmittance; chlorophyll fluorescence</td>
</tr>
<tr>
<td>H₂O transport</td>
<td>3D distribution of water potential; depends on bulk flow, diffusion, and evaporation; interacts with heat transfer; solved via mass transport partial differential equations on geometric mesh or infinite-element analysis</td>
<td>3D cellular and intercellular airspace network geometry, aquaporin activity, temperature distribution, leaf liquid water transport conductance inside and outside xylem, leaf water vapor transport, stomatal function</td>
<td>Leaf water potential, transpiration rate, leaf hydraulic conductance, stomatal conductance</td>
</tr>
<tr>
<td>CO₂ and metabolite reaction–diffusion</td>
<td>3D distribution of [CO₂]; depends on reactive photosynthetic demand, (photo)respiratory supply, and diffusion; interacts strongly with heat transfer and light propagation and weakly with H₂O vapor diffusion</td>
<td>Spatiotemporal distribution and geometry of organelles, cells, and tissues/airspace; 3D cellular and intercellular airspace network geometry, liquid phase diffusivities; metabolic network structure and flux</td>
<td>Apparent mesophyll conductance, respiration, photosynthesis, intercellular and chloroplast [CO₂], metabolite concentrations</td>
</tr>
</tbody>
</table>
A recent study provides a landmark example of integrated, anatomically explicit 3D modeling of leaf transport [3]. That study applied 3D leaf microstructure data [based on micro-computed tomography (microCT) imaging; Table 1] to modules that calculated light and CO2 and HCO3− exchange in intercellular airspaces and within cells, with different scenarios for 3D distribution of photosynthetic capacity. This fusion of 3D imaging and modeling technologies enabled several critical insights about fixation of CO2, the role of carbonic anhydrase, and the economy of photosynthetic capacity distribution, especially when predictions from the 3D-explicit approach differed systemically from an earlier 2D model [64]. For example, the more realistic representation of the 3D interconnectivity within the intercellular airspaces resulted in 61.7% of (photo)respired CO2 estimated to be re-fixed by RuBisCO. An analogous 3D explicit model could provide a more exact description of vapor transport through the intercellular airspaces. Notably, 3D imaging at current resolution will not be sufficient, as some features of transport, such as liquid phase water transport through cell walls and membranes, depend on structural details and material properties that must be determined empirically [6] or built up from nanoscale models of wall, membrane, and cell structure [29].

It is likely that a 3D paradigm will substantially alter our understanding of many of the nuanced questions associated with leaf anatomy and function. Currently, it is unclear a priori whether 3D analysis for any given process would lead to qualitatively different understanding. However, the technological advances outlined here are rapidly obviating any justification for adhering to the status quo (i.e., the spatially aggregated approach to studying leaf transport) without first, or concurrently, evaluating the results of analyses informed by high-resolution, spatially explicit approaches. A reasonable goal that could be achieved within the next decade with a coordinated, collaborative approach is to characterize internal leaf anatomy in detail for a few model species representing a range of plant functional types, and to apply the resulting data to a 3D model of leaf transport. Comparing predictions to leaf-scale gas exchange measurements would help determine how diversity of leaf structure affects photosynthesis and transpiration, and how to account for these effects in traditional leaf-scale models. Thus, the leaf biology communities should embrace a 3D approach with all its complexity, and all its promise (see also Outstanding Questions).

Acknowledgments

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References


Outstanding Questions

Light propagation and absorption: How does the 3D anatomical, biochemical, and biophysical arrangement of organelles, cells, airspace, and tissue types influence light propagation, absorption, and ultimately CO2 and H2O transport?

Interorganellar CO2 exchange: How does the spatial arrangement of organelles, such as chloroplasts, mitochondria, and vacuoles, influence CO2 transport through the liquid phase of the cell? What is the resulting effect on CO2 fixation, release, and refixation associated with photosynthesis and respiration? To which degree do plants modify these anatomical properties, and do these modifications convey a carbon benefit in given environments?

H2O transport pathways: What is the influence of the 3D anatomical arrangement of membrane transporters, organelles, cells, airspace, and tissue types on water potential gradients within the leaf? What is the resulting effect on stomatal dynamics and transpiration?

Spatial coordination of intraleaf CO2 and H2O sources and sinks: How does the geometric arrangement of stomata, organelles, and intercellular airspaces influence light absorption and affect leaf internal CO2 concentration gradients within leaves? What are the resulting effects on photosynthetic rate at the whole leaf level? To what extent does internal leaf geometry modulate water use efficiency of leaves? What are the evolutionary and adaptive solutions of plants to deal with environmental heterogeneity?

Isotope discrimination: How does leaf internal 3D anatomical arrangement affect stable isotope fractionation and what are the implications of these effects for estimates of mesophyll conductance?
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