

Integrating membrane transport, signaling, and physiology

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1 2	Integrating membrane transport, signaling and physiology
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18	Membranes constitute a barrier that compartmentalizes cells and organelles
19	from their surrounding environment. Transport across these barriers, mediated in
20	the most part by channels, transporters and pumps, is essential for many aspects of
21	cellular function. In plants, approximately 7% of the encoded proteins host at least
22	four transmembrane spans and are potentially involved in membrane transport.
23	Membrane transport plays key roles in including nutrition, cellular homeostasis,
24	signaling and development, and is a vital component of responses to biotic and
25	abiotic stresses. This Focus Issue brings together updates and research articles that
26	address some of most exciting and recent advances in our understanding of the
27	molecular mechanisms underlying membrane transport, its integration
28	with signaling, and its roles in homeostasis. Here, we highlight the contents of the 14
29	Updates presented in this Focus Issue.
30	
31	Transport integration in signal generation and plant development
32	
33	Plants have evolved many more ATP-binding cassette transporters (ABC)
34	than other organisms. Youngsook Lee's group, that has worked on ABC transporters
35	for more than two decades, comprehensively summarizes the recent developments
36	on ABC transporters, including newly identified functions and regulatory
37	mechanisms (Do et al, 2021). The Update outlines those transport functions that
38	have been deciphered using physiological data and those which have been verified
39	by functional assays in heterologous cell systems. The authors discuss a number of

40 outstanding questions such as how ABC transporters play such diverse roles in

plants and why so many ABC transporter genes have been retained in plant genome.

41

42 43 Metalloids are mineral elements that are often widely present in soils. In their update, Yamaji and Ma (2021) focus on Boron (B), Silicon (Si) and Arsenic (As), 44 45 covering the range of metalloids relevant to plants, whether essential, beneficial, or 46 biotoxic. These three metalloids share overlapping transport pathways for uptake 47 and distribution throughout plants. Central to these pathways is the metalloid 48 influx mediated by highly specialized aquaporins of the NIP subfamily. The NIPs 49 operate with specific efflux transporters displaying distinct polar localizations. 50 Understanding the mechanisms by which substrate specificity of these localizations 51 is determined and their functional divergence among tissues or even plant species 52 (monocots vs. dicots) will provide novel insights into the molecular physiology of 53 metalloids in plants.

54

55 γ -aminobutyric acid (GABA) is a key signaling molecule in animals, but less is 56 known about the roles and actions of GABA in plants. Xu, Sai and Gilliham (2021) 57 update us on the physiological impact of GABA in plants summarizing the current 58 evidence for the roles of aluminum-activated malate transporters (ALMTs) being 59 the targets of GABA signaling in guard cells to regulate water use efficiency, as well 60 as in pollen to regulate fertility. They speculate on other potential targets of GABA in 61 plants, such as H⁺-ATPase, potassium channels, and 14-3-3 proteins. They also 62 discuss the link of GABA to glutamate and hormones, as well as the potential role of 63 GABA for long-distance signaling.

64

65 Cellular phosphate (Pi) homeostasis is regulated by inositol pyrophosphates (PP-InsP), which act as intracellular messengers and are perceived by the SPX 66 67 domains of various proteins involved in Pi transport and signaling. In their Update, 68 Wang et al. (2021) explore the dynamics of Pi and PP-InsP metabolism. They 69 describe the role of PP-InsP as ligands to modulate the interactions between SPX 70 proteins and transcriptional activators in the phosphate starvation response, 71 thereby ultimately coordinating cross-membrane Pi transport and cellular Pi 72 homeostasis.

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Integrating Transport within and between cellular membranes

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Membrane nanodomains are defined as submicron protein and/or lipid
assemblies, whose structural characterization has greatly profited from recent
advances in microscopy techniques. The Update by Martinière and Zelazny (2021)
illustrates the role of membrane nano-organization in regulating of molecule

80 transport in plants. They focus on plasma membrane nanodomain formation, which 81 arises from functional assembly of sterols, sphingolipids, and specific proteins, 82 thereby regulating internalization of transporters in response to stimuli or effecting 83 co-clustering of proteins with their regulators.

84

85 The Update from Mehlhorn, Asseck and Grefen (2021) sets out the 86 background to our understanding of protein maturation and insertion into the lipid 87 bilayers with a particular emphasis on tail-anchored proteins. As they note, the 88 textbook models of protein insertion mechanisms lack credibility for those 89 secretory or membrane proteins which lack an amino-terminal signal sequence or 90 transmembrane domain. They review our current state of knowledge about tail-91 anchored proteins, harboring a single carboxy-terminal transmembrane domain, 92 and they address potential alternative, post-translational insertion routes into the 93 endoplasmic reticulum membrane.

94

95 In recent years, the technologies for exploring the subcellular mechanics of 96 Ca^{2+} signaling have greatly expanded with sensors that can be targeted to specific 97 locations within a plant cell. Two Updates in this Focus Issue evaluate these tools 98 and the information that they yield. The review by He et al (2021) examines 99 transporters likely to be involved in Ca²⁺ signal transmission, as well as those of 100 another divalent (Mn^{2+}), their transport across membranes, and their notable 101 association with glutamate receptor-like proteins (GLRs) and several linked pumps 102 and coupled transporters. Taking an organelle-focused perspective of the 103 interactive mechanics behind Ca²⁺ signaling, Resentini et al. (2021) address 104 questions about how unique characteristics of these signals may have arisen and 105 how the divalent ions are resequestered. They include examples of how latencies in 106 stomatal responsiveness to light and CO_2 have evolved (Jezek et al 2021), as well as 107 temporal properties of Ca²⁺ oscillations.

108 Finally, Tomkins, Hughes and Morris (2021) revisit the fundamentals of ion 109 and solute diffusion, emphasizing on the physics of passive transport systems and 110 the driving forces that give rise to diffusive and bulk flow processes in and out of 111 cells.

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- 113

Transport integration in environmental interactions

114

115 Kv (potassium voltage) channels in plants are important for nutrition by 116 mediating potassium uptake from the soil and distribute it through the whole plant, 117 and also for stomata movements. The Update review by Lefoulon (2021) focuses on 118 how Ky channels work with an emphasis on their mechanistics, our knowledge of 119 plant Ky channel structure, and the structures of closely-related channels in other

organisms. Lefoulon (2021) addresses issues from the selectivity of the Kv channels
to their gating by voltage, and finally how auxiliary proteins can regulate their
function in the face of environmental stress and how the gating properties may be
manipulated to affect plant growth and physiology.

124

125 Plasma membrane H⁺-ATPases and Ca²⁺-ATPases are autoregulated by 126 cytoplasmic concentrations of H⁺ and Ca²⁺, respectively, to generate steep gradients 127 across membranes as a prerequisite for substrate transport and cellular signaling. 128 The Update by Fuglsang and Palmgren (2021) highlights the role of phosphorylated 129 sites in the terminal regulatory domains of these pumps. Although some of these 130 phosphosites appear to be constantly phosphorylated and may function as major 131 switches, others appear to be dynamically phosphorylated to enable gradual 132 reductions in their activity.

133

Kim et al. update us on transport processes, in which sugars and amino acids
are secreted from plant cells by SWEET-type and UmamiT-type exporters,
respectively. Supported by instructive illustrations, key aspects relate to their cell
type-specific expression in phloem parenchyma and companion cells to govern the
distribution of assimilates in plants, as well as to their function in roots and other
organs to feed beneficial or pathogenic microbes.

140

141 Apart from their fascinating and unique biology, carnivorous plants have been highly relevant model systems for exploring long distance electrical signaling 142 143 in plants, since the early work of Charles Darwin. Böhm and Scherzer (2021) review 144 this topic, including their own research on signaling events in Venus flytrap 145 (*Dionaea muscipula* L.) and results from other research groups. The ion channel 146 activities underlying action potentials, their interaction with calcium and links to 147 downstream jasmonate signaling now provide a strong framework for 148 understanding the trap closure mechanism. Many other processes related to trap 149 biomechanics and the respective roles of growth and membrane transport during 150 slow trap reopening remain exciting topics for future research.

151

152 Plants sense the concentration of the major photosynthetic substrate CO₂ 153 within their leaves and, consequently, alter atmospheric connectivity by opening or 154 closing their stomatal pores. Dubeaux et al. (2021) update a current understanding 155 of how plants sense and respond to changing levels of CO₂, focusing on signaling 156 mechanisms that are known or might be involved in stomatal aperture responses to 157 low or high CO_2 concentrations. They also include a brief topical perspective on the 158 effects of rising atmospheric CO₂ concentrations in terms of stomatal densities and 159 their potential impact on crop productivity.

161 Water may constitute up to 98% of the weight of fully turgid plant tissues. Cross-membrane water transport mediated by aquaporins is critical to plant 162 163 expansion/growth, dehydration and germination of pollen and seeds, stomatal 164 closure etc. Accordingly, aquaporins represent central membrane targets of diverse 165 environmental and intrinsic signals. Maurel et al (2021) update us on how 166 aquaporins are regulated by drought, salinity, cold, nutrients, hormones, and other 167 biotic factors to maintain the plant water status. Interestingly, the multiple results 168 on the role of aquaporins in stomatal movement are discussed with explanations 169 proposed: due to their H₂O₂ transport activity, guard cell aquaporins function both 170 as players and targets of ABA and flagellin signalling during stomatal closure.

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160

The combined and considerable efforts of the editorial team over two years have culminated in this Focus Issue. We hope you enjoy reading it as much as we have enjoyed assembling it. Much of the contents of this Focus Issue arise from the open discussions and interactions that took place at the 18th International

176 Workshop in Plant Membrane Biology, in Glasgow in 2019. This Focus Issue serves

as a bridge leading you to the next International Workshop in Plant Membrane

178 Biology, scheduled for October 17-21 of 2022 in Taiwan, where all of the latest

advances in the above areas will be showcased. If nothing else, these Updates and

180 the Workshop show how rapidly the field is now evolving.

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