Carbon fluxes and environmental interactions during legume development, with a specific focus on *Pisum sativum*

Amélie Morin | Laurence Marousset | Cécile Vriet | Rémi Lemoine | Joan Doidy | Nathalie Pourtau

Université de Poitiers, UMR CNRS 7267, EBI “Ecologie et Biologie des Interactions”, Poitiers, France

Correspondence
Nathalie Pourtau, Université de Poitiers, UMR CNRS 7267, EBI “Ecologie et Biologie des Interactions,” Poitiers, France.
Email: nathalie.pourtau@univ-poitiers.fr

Funding Information
Centre National de la Recherche Scientifique; Grand Poitiers; Université de Poitiers

Edited by: R. Le Hir

Abstract
Grain legumes are major food crops cultivated worldwide for their seeds with high nutritional content. To answer the growing concern about food safety and protein autonomy, legume cultivation must increase in the coming years. In parallel, current agricultural practices are facing environmental challenges, including global temperature increase and more frequent and severe episodes of drought stress. Crop yield directly relies on carbon allocation and is particularly affected by these global changes. We review the current knowledge on source-sink relationships and carbon resource allocation at all developmental stages, from germination to vegetative growth and seed production in grain legumes, focusing on pea (*Pisum sativum*). We also discuss how these source-sink relationships and carbon fluxes are influenced by biotic and abiotic factors. Major agronomic traits, including seed yield and quality, are particularly impacted by drought, temperatures, salinity, waterlogging, or pathogens and can be improved through the promotion of beneficial soil microorganisms or through optimized plant carbon resource allocation. Altogether, our review highlights the need for a better understanding of the cellular and molecular mechanisms regulating carbon fluxes from source leaves to sink organs, roots, and seeds. These advancements will further improve our understanding of yield stability and stress tolerance and contribute to the selection of climate-resilient crops.

1 | INTRODUCTION

Legumes have been consumed for millennia and remain a staple food cultivated worldwide for their high nutritional value. Rich in proteins and starch, legumes make an essential food for human diet and animal forage. In this review, we mainly focus on the pea crop (*Pisum sativum*), which is particularly rich in various nutrients, notably proteins. Its seeds contain starch (50%), proteins (25%), soluble sugars (5%), fibers (5%), and also vitamins and minerals (Bastianelli et al., 1998). Plant-based proteins are a promising alternative to meat consumption and could reduce the global impact of food production on the environment (Sandberg, 2011). As a result, the international market for pea proteins is booming and opportunities are rising for the market of starch valorization (Yu et al., 2021). This societal challenge towards more sustainable practices in the context of global environmental changes implies a reconsideration of the current agricultural system.

The yield of many crops rarely reaches its maximum potential of production. This is particularly the case for grain legumes which have benefited from fewer agronomic improvement programs than cereals (Foyer et al., 2016). For instance, yields of wheat in France increased on average by 92 kg ha$^{-1}$ year$^{-1}$ between 1961 and 2018, while those for pea only increased by 37 kg ha$^{-1}$ year$^{-1}$ over the same period (FAOSTAT, 2021). This large difference is in part attributable to unfavorable environmental conditions. Indeed, grain legume production is particularly affected by environmental factors, both biotic and abiotic, causing an unstable production with large variability in yield (Bénézit et al., 2017; Smith et al., 2018). Several stresses negatively impact plant production but drought, temperatures, salinity, and
waterlogging are the major abiotic stresses implicated in crop yield destabilization (Rane et al., 2021; Zander et al., 2016). Drought stress is the main constraint that affects crop yield and quality. A low water availability associated with high temperatures can also result in higher soil salt concentration (Rane et al., 2021; Saberi Riseh et al., 2021). Moreover, because of climate change, dry periods alternate with wet episodes leading to waterlogging and submergence of the aerial part (Rane et al., 2021). Global climate change, associated with a degrading environment, amplifies these major stresses, impacting crop development, and productivity. Developing new crop varieties more resilient to climate change and ensuring yield stability to feed the world’s growing population, while complying with environmental sustainability, represent a huge challenge (Dhankher & Foyer, 2018; Martignago et al., 2020; Smith et al., 2018). To this end, breeders and researchers must focus on understanding how the environment influences yields in crops (Bénézit et al., 2017; Smith et al., 2018; Strydhorst et al., 2015).

Crop yield directly relies on carbon fluxes, which is particularly affected by global changes. To counteract the effects of environmental stresses, a significant part of the energy captured by photosynthesis is redirected, which generates yield losses. Indeed, in optimal conditions, the carbon fixed by photosynthesis will be primarily allocated towards sink organs (roots, young leaves, flowers, pods, and seeds), also mediating a flow of nitrogenous substances to these organs. Altogether, this contributes to the optimal nutritional filling of carbon (starch) and nitrogen (proteins) into seeds. Conversely, if the plant is subjected to biotic or abiotic stresses, a significant part of the carbon will be diverted, and thereby lost for sink organs (Chen et al., 2006; Durand et al., 2016; Fougeroux et al., 1997; Guillioni et al., 1997; Henriet et al., 2019). A better understanding of the modulation of these carbon fluxes during both biotic and abiotic stresses and uncovering the genes involved in these biological processes will contribute to improving legume productivity.

Carbon fluxes are mediated by sugar transport proteins of the SUCROSE TRANSPORTER (SUT), MONOSACCHARIDE TRANSPORTER (MST), and SWEET (SUGARS WILL EVENTUALLY BE EXPORTED TRANSPORTER) families. Notably, SUTs, MSTs, and SWEETs control the loading of photosynthates from source leaves and coordinate carbon fluxes towards sink organs (e.g., root and seed), thereby determining the yield and nutritional quality of seed crops (Sosso et al., 2015). Their crucial roles in response to environmental factors have also been highlighted in model crops (Lemoine et al., 2013), but their contribution to stress tolerance has been overlooked in legumes. In addition, sugar transport and carbon metabolism genes have potential applications as biomarkers in plant breeding programs (Schroeder et al., 2013; Sosso et al., 2015; Wang et al., 2008). Today, pea enters the genomic era with a reference genome recently sequenced (Kreplak et al., 2019). This represents a major advance and should help identify genetic markers for yield and stress tolerance in pea. We recently identified major genes mediating sugar transport (PsSUT, PsMST, and PsSWEET) and carbon metabolism genes (PsINV) in P. sativum, focusing on plant productivity in response to drought (Doidy et al., 2019; Morin et al., 2022).

Here, we review major research advances on carbon partitioning in grain legumes, with a specific focus on pea crop and in response to environmental challenges. We cover the properties and processes governing carbon allocation and sugar transport during legume development, from germination to seed production. We also highlight current knowledge on the biological pathways at the molecular level and propose hypotheses to improve the resilience of legume crops towards major environmental changes.

2 | CARBON FLUXES DURING SEED GERMINATION AND SEEDLING ESTABLISHMENT

Germination is a key developmental process including imbibition and metabolism recovery of the seed resulting in the emergence of the root and the shoot (Figure 1A). Dry pea storage compounds mainly consist of starch, proteins, soluble sugars, and fibers. The most abundant soluble sugars in dry pea seeds are sucrose and RFOs, mainly stachyose and verbascose (Kuo et al., 1988; Vidal-Valverde et al., 2002). To sustain growth and seedling establishment, these reserves must be remobilized. The role of RFOs catabolism during early pea germination has been highlighted by a 25% decrease in the germination rates when α-galactosidase was chemically inhibited (Blöchl et al., 2007). Degradation of RFOs leads to an increase in sucrose, glucose, and fructose (Vidal-Valverde et al., 2002), and when the radicle emerges, 50% of the RFOs are already metabolized. Starch remobilization occurs later on as amylase activity is enhanced between 4 and 10 days after imbibition (Yomo & Varner, 1973). Pea storage compounds are degraded and transported from the cotyledons to the embryo axis to be directed to the root and the shoot (Figure 3). In germinating pea seeds, several SUTs, SWEETs, and CWINVs (cell wall invertases) genes are strongly expressed in the first days following imbibition (Jameson et al., 2016). For instance, the authors showed a strong induction of PsSUT3 (renamed PsSUT2), PsSWEET12 and PsCWINV6 (renamed PsCWINV2.1) after 4 h of imbibition. This suggests a transport of hexoses and sucrose from the cotyledons to the growing sink organs during the germination process (Figures 1A and 3). Indeed, cotyledons are the major source organs at the beginning of seedling development. By removing one cotyledon (source organ) or by manipulating the sink strength by adding sucrose, Schulz (1994) showed that the translocation of sugars is sink-regulated in pea seedlings, mainly through the apical root region (Schulz, 1994). The main gene that seems involved in sugar translocation is PsSUT1 (renamed PsSUT1.1) (Figure 3), which is strongly expressed in shoot and root (Jameson et al., 2016). This demonstrates that source-sink relationships are rapidly established after imbibition (Figure 1A) and subsequent degradation of the cotyledon storage compounds is linked to a transition from germination and seed reserve dependency to autotrophy and sugar production through photosynthesis.

During their development, crops must face various abiotic stresses (Figure 1A). Major stresses include drought stress,
temperature, salinity, and waterlogging. In pea, germination and seedling establishment are sensitive to thermic stress, which affects seed germination rate. Even though pea seeds are rather highly tolerant to cold and can germinate on ice (Macherel et al., 2007), germination rates tend to increase with temperatures, reaching an optimum between 21°C and 27°C (Raveneau et al., 2011). These optimal temperatures probably allow a better activity of the hydrolytic enzymes and sugar transporters into the cotyledons. Water deficit also affects germination (Figure 1A) (Raveneau et al., 2011) (Table 1). By adding PEG to germinating seeds, germination percentages in three pea varieties were dramatically reduced from 100% to 5% at an osmotic potential of −0.8 MPa (Okçu et al., 2005) (Table 1).
Moreover, seedlings grown under different soil water holding capacities contained more proteins and carbohydrates, with a proportional increase related to the stress intensity (Al-Quraan et al., 2021) (Table 1). Sucrose accumulation is a common response of seedlings to dehydration, probably thanks to the increased activity of sucrose synthase. During water stress, sucrose content increases earlier and is higher in pea roots than in epicotyls (but not in cotyledons) and these changes are associated with a lower accumulation of galactinol and sugar alcohol in the phloem sap (Devi et al., 2007). However, sucrose synthesis was not affected in source cotyledons, but the sucrose content was modified in the sink radicle due to an inhibition of sucrose transport (Table 1). This could be linked to seedling growth reduction as well as variations in carbohydrate composition (Devi et al., 2007).

In addition to abiotic constraints, pathogenic microorganisms can also be detrimental to pea germination and early seedling growth (Figure 1A). Indeed, pea is sensitive to several pathogens, and this from the start of its lifecycle. For instance, inoculation of a pathogenic strain of Rhodococcus fascians on germinated pea plants resulted in reduced root growth and multiple shoot formation with intact cotyledons (Dhandapani et al., 2017). In cotyledons, the expression of PsSWEET, PsSUT, and PsINV genes was higher than in the noninfected control, suggesting changes in the source-sink relationship induced by the addition of cadmium resulted in decreased α- and β-amylase activities leading to a reduction of starch degradation in cotyledons (Devi et al., 2007). However, sucrose synthesis was not affected in source cotyledons, but the sucrose content was modified in the sink radicle and shoot. The negative effect of cadmium could therefore mainly be due to an inhibition of sucrose transport (Table 1). This could be linked to seedling growth reduction as well as variations in carbohydrate composition (Devi et al., 2007).

### Table 1: Overview of the main results on pea responses to challenging environments

| Stress | Developmental phase | Experimental results | Reference |
|--------|---------------------|----------------------|-----------|
| Abiotic stress | Seed germination | Probable inhibition of sucrose transport | Devi et al. (2007) |
| Cadmium | Vegetative stage | Lower chlorophyll content with subsequent reduced photosynthesis | Sandalo et al. (2001) |
| Cold | Seed germination | Suboptimal activity of hydrolytic enzymes reducing germination rates | Raveneau et al. (2011) |
| | Vegetative stage | Increased leaf and root soluble sugars levels/ temporary starch storage to compensate for loss of photosynthetic activity in a pea winter variety | Bourion et al. (2003) |
| Drought | Seedling stage | Accumulation of carbohydrates and proteins | Al-Quran et al. (2021) |
| | Vegetative stage | Osmotic adjustment through soluble sugar accumulation in leaves | Kumar et al. (2021), Lahuta et al. (2022), Sanchez et al. (1998, 2004) |
| | Vegetative stage | Increased osmotic potential in roots | Kumar et al. (2021), Lahuta et al. (2022) |
| | Vegetative stage | Increased content of sucrose, glucose, and sugar alcohol in the phloem sap | Blichartz et al. (2021) |
| Flowering stage | Decrease of plant biomass | Prudent et al. (2016) |
| Flowering stage | Decrease of flower and pod number | Nadeem et al. (2019) |
| Seed development | Reduction in individual seed weight and seed number | Ney et al. (1993), Guilloni et al. (2003), Nadeem et al. (2019) |
| Heat stress | Flowering stage | Cessation of reproductive organ formation and subsequent flower abortion | Guilloni et al. (1997) |
| Osmotic stress | Seed germination | Decrease of germination percentages | Okçu et al. (2005) |
| Waterlogging | Vegetative stage | ROS production, reduced plant growth, and seed yield | Pampana et al. (2016a), Rane et al. (2021) |
| Pathogens | Rhodococcus fascians | Seed germination | Changes in source-sink relationship between cotyledons and shoot/root | Dhandapani et al. (2017) |
| | Erysiphe pisi | Vegetative stage | Increased invertase enzyme activity | Storr and Hall. (1992) |
| | Erysiphe pisi | Vegetative stage | Promotion of resistance through overexpression of MtSTP13.1 | Gupta et al. (2021a, 2021b) |
| | Ascochyta blight (not specified) | Seed development | Yield loss | Bretag et al. (1995) |
| Mycosphaerella pinodes | Seed development | Decreased seed starch content | Garry et al. (1996) |
pathogen, with cotyledons maintained as sinks for its benefit rather than evolving to a source tissue sustaining plant growth (Table 1).

Altogether, data in the literature point to a strong dependency between seed sugar remobilization for seedling development and their adaptation to environmental constraints. When the seedling’s environment enables its establishment, vegetative development proceeds and source-sink relationships evolve, with mature leaves becoming the main source organs.

3 | Carbon fluxes in shoot during vegetative development

3.1 | Sugar fluxes and signals for axillary bud growth

Pea plants have an indeterminate growth habit with the successive addition of phytomers defining the main stem. At each node, an axillary bud has the capacity to develop in a lateral branch, according to the same scheme as the main axis. Lateral branches will support the growth of more flowers and pods. Therefore, the number of lateral branches will have a positive impact on yield (Patrick & Colyvas, 2014). In pea, lateral branches are usually located on the lowest vegetative nodes (Doré, 1994) but growing conditions and genotype can affect the number and position of branches. However, the number of axillary branches depends heavily on photosynthate (mainly sucrose) availability (Patrick & Colyvas, 2014). Pea has been a model for studying branching, allowing the discovery of the role of auxin, cytokinins, and more recently strigolactones in the release of apical dominance (for review see Rameau et al., 2015). The role of sucrose as an antagonist of auxin on axillary bud outgrowth has been described in the past decade. In pea, Mason et al. (2014) demonstrated that sucrose is “both necessary and sufficient” for the initial release of axillary buds from apical dominance before the auxin concentration in the bud is affected. This initial effect could of course be linked to the trophic role of sucrose; however, it was also possible to induce a similar rapid release of bud growth with nonmetabolized analogs of sucrose. Thus, Barbier et al. (2015) suggested that sucrose is also involved in a signaling pathway. However, hexoses derived from sucrose may also be involved considering the recently discovered role of HEXOKINASE 1 (HXK1) in regulating shoot branching (including in pea; Barbier et al., 2021). Bertheloot et al. (2020) proposed a quantitative model for the interactions of sucrose with the hormonal pathways regulating bud outgrowth. This model, in rose and in pea, suggests that plant sugar status modulates the auxin-controlled apical dominance. This latter can be maintained only if the sugar status of the plant is low. On the opposite, if the plant’s sugar status is high, apical dominance is low and axillary buds develop with, in the case of pea, a positive effect on yield as mentioned above. The link between the sugar status of the plant and the development of axillary buds may represent an adaptative response to the environment (Bertheloot et al., 2020).

Trehalose-6-phosphate (Tre6P), a sugar signaling molecule, has recently been implicated in shoot branching in Arabidopsis (Fichtner & Lunn, 2021). Increasing the level of Tre6P in the vasculature had a positive effect on branching. Interestingly, the expression of the sucrose transporter genes AtSWEET11 and AtSWEET12 was also increased. In Rosa hybrida, RhSUC2 expression was increased, together with an influx of sucrose, in the bud during bud burst (Henry et al., 2011). In Chrysanthemum morifolium, CmSWEET17 expression was increased during bud outgrowth, and its overexpression promoted bud growth (Liu et al., 2020). Taken together, these results indicate that increased expression of sucrose transporter genes is associated with an increased flux of sucrose towards the bud, thus promoting bud outgrowth.

3.2 | Carbon fluxes in source leaves

Pea leaves are pinnate and are composed of a basal pair of stipules, 2–3 pairs of basal leaflets and a terminal tendrill. In pea, a wide variety of genotypes are available, some carrying mutations leading to different arrangements of the leaf parts. The most described is the afila mutation, where all leaflets are replaced by tendrils (Figure 1B). The leaf is therefore reduced to a pair of basal stipules and distal pairs of tendrils that are branched, forming together the tendril complex (Côté et al., 1992a). In conventional (leafy) cultivars, it was assumed that the contribution of tendrils was neglectable. However, it has been demonstrated that mature tendrils act as source organs and export photo-assimilates through the phloem (Côté et al., 1992a). Further studies by Coté et al. (1992a) specified that the sink/source transition occurs when tendrils begin to coil. The authors calculated that 65%–70% of the carbon gained per day can be attributed to the tendrils in an afila genotype (Côté et al., 1992b).

Little is known about sugar transport at the molecular level in different pea genotypes. In Arabidopsis, it is commonly accepted that sucrose is transported from the mesophyll cells, where it is produced, to the phloem parenchyma cells through symplastic transport (Lalonde et al., 2003) (Figure 3). Sucrose is then unloaded into the apoplasm near the conductor complex (companion cell/sieve element) by two facilitators: AtSWEET11 and AtSWEET12 (Chen et al., 2012) (Figure 3). As sucrose concentration of the conducting complex is much higher than in the apoplast, the influx of sucrose requires active transport by the sucrose transporter AtSUC2 (Ayre, 2011; Dinant & Lemoine, 2010; Lalonde et al., 2004). P. sativum also seems to possess a similar phloem loading configuration (Van Bel & Gamalei, 1992). Therefore, we could postulate that PsSWEET12 (AtSWEET12 homolog) mediates the efflux of sucrose into the apoplasm, while PsSUT1.1 (AtSUC2 ortholog) is responsible for loading sucrose into the phloem (Doidy et al., 2019) (Figure 3). Ninan et al. (2019) studied the expression of PsSUT and PsCWINV genes during leaf development and senescence in two cultivars: Bolero (leafy) and Bohatyr (afila). PsSUT1 (renamed PsSUT1.1) and PsSUT2 (renamed PsSUT1.4) were identified as the most expressed SUT genes, both in mature and senescent leaves (Figure 3), but with a lower expression in leaves of the Bolero cultivar. However, the authors did not investigate the different parts (stipules, leaflets, tendrils) independently. In
Bolero, PsSWEET12 was highly expressed in mature and senescent leaves (Figure 3). In Bohatyr, clade III SWEET genes were expressed in sink and mature leaves but at a very low level in senescent leaves. In both cultivars, high expression of PsCWINV genes was measured in sink leaves as expected for organs with a high mitotic activity.

3.3 | Carbon fluxes in shoots in response to environmental constraints

Drought is the most important constraint limiting legumes growth and subsequent yield (Figure 1B). Water deficit leads to deep modifications of the plant primary and secondary metabolisms (Kumar...
et al., 2021; Lahuta et al., 2022) (Table 1), as it decreases net photosynthesis and nutrient uptake. Hence, when facing drought, a major issue for the plant is to produce enough carbohydrates by photosynthesis to sustain sink growth (Kumar et al., 2021; Nadeem et al., 2019). A study on the tolerance to water deficit of 49 pea cultivars indicated that soluble sugars could take part in leaf osmotic...
adjustment and turgor maintenance to a bigger extent than proline (Sánchez et al., 1998). This study highlighted that sugars have a major role in osmotic adjustment since a positive correlation among the cultivars studied was found between the accumulation of soluble carbohydrates, especially myo-inositol and raffinose, their osmotic capacity and level of tolerance to stress (Lahuta et al., 2022; Sánchez et al., 1998; Sánchez et al., 2004) (Table 1).

In soybean, a study applying drought stress during the vegetative stage led to differential expression of genes involved in sugar metabolism and transport (Chen et al., 2016). Among them, glycolytic enzymes (hexokinase and fructokinase), as well as SWEET genes, were induced by drought. However, this induction of glycolytic enzymes was not observed in chickpea (Khanna et al., 2014), which suggests variability of response among legumes. In Arabidopsis, an enhanced export of sugar, linked to higher expression of AtSUC2, AtSWEET11, and 12—three genes associated with phloem loading (Figure 3)—was observed in response to water deficit (Durand et al., 2016). The same kind of investigations are still lacking in pea, and more studies on this topic are needed to uncover the mechanisms involved in sugar homeostasis and transport in response to drought. Nevertheless, some studies indicate that afila cultivars, with reduced leaf surface area, are less sensitive to drought than leafy cultivars thanks to a better ability to remobilize carbon to sinks (Baigorri et al., 1999; Couchoud et al., 2020). For instance, it was shown that in response to water stress, the semi-leafless cultivar (Solara) displayed an arrest in vegetative growth and an induction of leaf senescence, and its pod and seed dry weights did not decrease in comparison to the control condition (Baigorri et al., 1999). Furthermore, in pea senescing leaves, which are exporting carbohydrates, an increased expression of several clade II and III SWEETs as well as PsSUT1 and PsSUT2 genes (renamed PsSUT1.1 and PsSUT1.4) was observed (Ninan et al., 2019).

Drought stress, and other abiotic stresses, such as high temperature and salinity (Figure 1B), lead to cellular dehydration in plants (Rane et al., 2021). To counteract this effect, plants synthesize osmolytes, such as sugars, which play different roles in carbon allocation, energy storage, signaling, or maintaining membrane integrity (Kumar et al., 2021; Rane et al., 2021) (Table 1). Moreover, to face abiotic stress, plants produce reactive oxygen species (ROS), which induce oxidative stress and increase the accumulation of sugars, such as trehalose and sucrose to protect plant cells (Figueroa & Lunn, 2016; Kumar et al., 2021; Lin et al., 2019). A study on trehalose-6-phosphate phosphatase (TPP) overexpressing lines, the last enzyme implied in trehalose synthesis pathway, indicated that trehalose-6-phosphate (Tre6P) plays an important role in tolerance to several abiotic stresses, such as anaerobic conditions, salt, chilling, and cold stress (Figure 1B), in different plant species (Figueroa & Lunn, 2016). More precisely, Tre6P, a phosphorylated intermediate of trehalose biosynthesis, present in low concentration, seems to induce changes in sucrose concentrations, thus, promoting drought stress resistance (Figueroa & Lunn, 2016; Lin et al., 2019).

Even though cold stress can have severe effects on plant growth and subsequent yields at maturity (Figure 1B), pea, like most plants, has developed mechanisms of cold acclimation (Baldwin et al., 2014). Cold acclimation responses to chilling (0°C–15°C) and freezing temperatures (<0°C) have notably been studied by comparing behaviors of winter versus spring varieties. First, their response depends on light intensity. Under regular lighting, winter peas presented a higher freezing tolerance than spring peas (Bourion et al., 2003). Moreover, a close relationship was observed between the soluble sugar concentrations in leaves and temporary starch storage (Bourion et al., 2003) (Table 1). However, such acclimation response was not observed in spring peas. Leaf architecture also seems to play a role in cold tolerance, specifically in new leaves developing during the cold period, according to a study comparing the photosynthetic activity of a leafy winter pea, two afila spring pea and an afila winter pea cultivar subjected to cold stress. The leafy winter pea (the most resistant cultivar) displayed the most rapid decrease in photosynthesis activity, which may be linked to its survival strategy (Husičková et al., 2019).

Waterlogging (or submergence) is detrimental to growth and yield and induces ROS accumulation because of oxygen deprivation stress (Pampana et al., 2016a, 2016b; Rane et al., 2021) (Figure 1B and

---

**Figure 3**  Schematic representation of sugar fluxes from source to sink organs mediated by sugar transporters during pea development. During germination, cotyledons are the main source organ and reserve degradation enables sucrose allocation towards sink radicle and shoots. When the first leaves are formed, photosynthesis is initiated and sucrose is released in the apoplast by SWEETs and retrieved in the companion cell, most likely by PsSUT1.1. Phloem loading is necessary to sustain the growth of sink organs. During the vegetative stage, roots are the main sinks. Sucrose is unloaded in roots by SUT and retrieved in root cells by SWEETs. Noteworthy, legumes can develop tripartite symbiosis. In AM symbiosis, sugars are exported by SWEETs towards the symbiotic interface across the periarbuscular membrane. Sucrose and monosaccharides can be retrieved back into the plant cell by MST (STP in *Medicago truncatula*) or SUT. Sugar transporters are also likely to be involved in plant-Rhizobia symbiosis, mainly by SWEETs (e.g., MtSWEET11 in *M. truncatula*). Vacular transporters, such as SUT4-type, are also induced, especially in AM roots (Hennion et al., 2019), suggesting that remobilization of intracellular reserves occurs in colonized roots. During reproductive development, new sinks (flowers and seeds) are formed. Sugar unloading in the seed coat involves a symplastic pathway. Sucrose is released in the apoplast by SWEETs and SUT transporters. During embryogenesis, CWINV hydrolyses sucrose in glucose and fructose, which might be either loaded in the embryo and the endoderm either by MSTs or SWEETs hexose transporters. The expression of PsSUT1.1 in the endoderm suggests that sucrose can be directly loaded in this tissue and escapes invertase hydrolysis. Sucrose can be exported from the endoderm by SWEETs (GmSWEET15 in soybean). Cotyledonary cell expansion results in the progressive disappearance of the endoderm, which characterizes the transition from embryogenesis to seed-filling. During early seed-filling, invertase expression and activity decrease, and sucrose is directly imported to the cotyledons by PsSUT1.1, as starch and protein storage begin.
Table 1). In relation to climatic fluctuations observed these last years, waterlogging is becoming a critical problem. By restricting oxygen diffusion in submerged tissues, aerobic respiration is impaired, and a shift occurs to anaerobic ethanolic fermentation (Phukan et al., 2018; Rane et al., 2021). Waterlogging impacts plant development by reducing photosynthesis, plant growth, nodule activity, and grain yield. Waterlogging damages increase as plants become older and advance towards the reproductive stage (Pampana et al., 2016a, 2016b; Rane et al., 2021). Photosynthesis is decreased by stomatal closure and the weak ATP production by anaerobic fermentation slows down plant growth. Sugar availability, metabolism, and transport are important components of the response to waterlogging, as for many other abiotic stresses (Phukan et al., 2018; Rane et al., 2021). In rice and pigeon pea, tolerant varieties accumulate a larger amount of soluble sugars than susceptible ones (Phukan et al., 2018). During waterlogging, Arabidopsis transgenic lines over-expressing a waterlogging-responsive ethylene response factor (MaRAP2-4) regulating AtSWEET10 showed a higher stress tolerance in comparison to the wild type (Phukan et al., 2018). Indeed, to face waterlogging, plants could accumulate sugars, but this strategy might not be sufficient to survive under prolonged waterlogging stress. Plants may also use sugar resources to get their leaves out of the water through the rapid growth of the shoot system. These strategies are under the tight control of the source/sink relationship regulating carbon allocation (Phukan et al., 2018; Rane et al., 2021).

Waterlogging also induces anoxia in the soil, which can lead to an increase in toxic compound concentrations (Rane et al., 2021). The impact of cadmium on plant growth is mainly due to a lower chlorophyll content leading to a reduction of photosynthetic rate and an accumulation of oxygen free radicals, leading to possible leaf senescence (Sandallo et al., 2001) (Table 1). Soluble sugars could play a role in Cd stress response since a study on sensitive maize demonstrated that the more plants produced soluble sugars the higher the Cd concentrations were, indicating that this variety uses more carbohydrates to defend itself against Cd rather than to sustain its growth (Li et al., 2020).

Legumes are also affected by different fungal, bacterial, and viral diseases, as well as insect and nematode infections (Van Emden et al., 1988). The major pathogen infecting pea during early vegetative development is the Oomycete Aphanomyces euteiches, the causing agent of root rot disease (Figure 1B). Numerous pathogens, such as Pythium sp., Phytophthora sp., Rhizoctonia solani, or Botrytis cinerea are also responsible for “damping-off;” killing or weakening pea seedlings before or after their germination. Notably, these heterotrophic pests infect plants and feed on sugars. As such, pathogens have evolved an arsenal of sugar-splitting enzymes and sugar transporters to gain access to plant photosynthates (Doidy et al., 2012). For instance, the biotrophic rust fungus Uromyces fabae possesses a hexose transporter, UHHT1, specially targeted to the plant-fungal interface (haustoria), thereby feeding on Vicia faba hexose (Voegele et al., 2001). The activity of invertase enzymes in P. sativum also increased in response to Erysiphe pisi infection (powdery mildew), and the leaf starch content was also impacted (Storr & Hall, 1992) (Table 1). In response, plant transporters are regulated to retrieve photosynthates back to the plant cell. For instance, the monosaccharide transporter STP13 has long been shown to be regulated by multiple stresses, both biotic and abiotic. Notably, its expression is induced by several pathogens and contributes to plant resistance in different species (Yamada et al., 2016).

In legumes, MtSTP13.1 is involved in sugar competition when infected with the pea powdery mildew Erysiphe pisi and over-expression of MtSTP13.1 promoted pathogen resistance in pea (Gupta et al., 2021a) (Table 1). Finally, SWEET transporters also play a key role in plant disease resistance. More precisely, a growing number of studies have shown that pathogens are able to modulate the expression of these sugar exporters to redirect sugar fluxes towards them (Gupta et al., 2021b; Lemoine et al., 2013). SWEET transporters represent a promising target for the development of resistant crop cultivars (Gupta et al., 2021b, see Section 6) and we identified 22 PsSWEET in P. sativum (Doidy et al., 2019). In conclusion, sugar transporters play crucial roles in plant-microbe interactions and legume species offer an excellent model to study them (Chandran, 2015; also see Section 3.3).

4 | CARBON FLUXES IN ROOTS DURING VEGETATIVE DEVELOPMENT

4.1 | Sugar unloading in root

Roots pattern establishment is divided into three steps. First, an initial lag phase during germination, followed by an early rapid growth during shoot development and a linear phase after flowering (Mitchell & Russell, 1971; Thorup-Kristensen, 1998; Vocanson et al., 2006). Root legume development starts with the rapid taproot elongation followed by lateral root emergence (Mitchell & Russell, 1971; Tricot et al., 1997). A decrease in root elongation rate was observed when C allocation starts from source leaves to roots, probably in link with the exhaustion of seed reserves (Tricot et al., 1997). The transition from heterotrophy to autotrophy seems to affect firstly lateral root development and later the taproot. Indeed, C supply sustains the growth of lateral root primordia, and depletion in C allocation reduces lateral root elongation and root branching before the taproot growth rate is affected (Tricot et al., 1997). This result indicates that the taproot displays a higher priority for C allocation compared to lateral roots (Tricot et al., 1997) (Figure 1B).

During the vegetative stage, roots and young leaves are major sinks (Wardlaw, 1990) and the hierarchy between sinks is determined by the strength of the sink (Ho, 1988). Sink strength corresponds to the ability to reduce photoassimilates in cells of the sieve tube at the sink level, thereby creating a hydrostatic pressure gradient favorable to long-distance transport of sucrose and unloading towards this sink (Wardlaw, 1990). The first step of sucrose unloading in sink organs involves a symplasmic pathway (Patrick, 1997) (Figure 3), as observed in Arabidopsis root tips (Oparka et al., 1994). Beyond the plasmol unloading zone, the type of transport varies according to the type of sink and the stage of development, but transport of sugars via the apoplastic pathway is possible and involves active SUT transporters.
and SWEET facilitators (Chen, 2014; Durand et al., 2018; Lalonde et al., 2004) (Figure 3). Moreover, sugar transporters’ expression is modulated by adverse conditions, highlighting that source-sink regulation for sucrose is affected by environmental factors (Durand et al., 2016; Lemoine et al., 2013).

Although carbon allocation to the roots is important during vegetative development (Voisin, 2003), some authors suggested that root growth stops during reproductive development due to reduced sink strength (Jeuffroy & Warembo, 1991). Other studies showed that during seed-filling, a slow rate of root growth also takes place, which is associated with a decrease in lateral root emergence and root biomass (Armstrong et al., 1994; Salter & Drew, 1965).

### 4.2 Carbon fluxes in challenging soil conditions

Root growth and development are strongly influenced by environmental factors, such as soil structure and climatic conditions (Ali-Khan & Snoad, 1977) (Figure 1B). When occurring at the start of the vegetative stage, drought negatively impacts seed germination and seedling growth with a reduction of shoot and root dry weights in different species, including in pea (Fahad et al., 2017). To face such water stress, the first plant responses include a decrease in photosynthesis and transpiration activities followed by an accumulation of osmoprotectants (non-reducing sugars, polyols, amino acids, etc.) in both roots and leaves (Kumar et al., 2021; Lahuta et al., 2022). This higher osmotic potential maintains root water uptake from the soil (Kumar et al., 2021; Lahuta et al., 2022) (Table 1). In seven-day-old pea seedlings, an earlier accumulation of sucrose was observed in roots than in epicotyl after 24 h of dehydration, followed by an accumulation of galactinol and raffinose (RFOs family) in both organs (Lahuta et al., 2014). Therefore, drought stress increases the content of sucrose, glucose, and sugar alcohol in the phloem sap of pea to sustain osmotic potential, water content and transport, and maintain cell metabolism (Blicharz et al., 2021) (Table 1). Thus, water deficit response induces metabolic modifications, some of which are linked to source/sink relationships (Lemoine et al., 2013). In Arabidopsis, a preferential C allocation towards roots was observed during water deficit to sustain root growth. This phenomenon was associated with an induction of AtSWEET11, AtSWEET12, and AtSUC2 genes, which are responsible for phloem loading in leaves and might also be involved in phloem unloading in roots (Durand et al., 2016) (Figure 3).

In field conditions, a drought episode may be followed by strong rainfall leading to flooding. Autumnal grain legumes present a higher risk of being exposed to such submergence (Pampana et al., 2016a). During waterlogging, inhibition of aerobic respiration and TCA dysfunction are observed in submerged tissues associated with ROS increase, leading to a decrease in shoot and root growth (Table 1) (Pampana et al., 2016b; Rane et al., 2021). Limiting oxygen availability observed during waterlogging enhances alcoholic fermentation, an adaptive mechanism to provide energy to the plants facing this stress. Moreover, waterlogging induces root architecture arrangement with the formation of aerenchyma in the cortex for storing O$_2$, stem hypertrophy, and formation of adventitious roots. This stress also impacts photosynthesis activity, plant development, seed productivity, and nodule integrity (Pampana et al., 2016a). Both synthesis of fermentable sugars and release of hydrolyzed sugars contribute to waterlogging tolerance in crop plants (Goyal et al., 2020).

### 4.3 Carbon fluxes in beneficial microorganism interactions

Legumes are notably cultivated for their agroecological value through their beneficial associations with microorganisms. Symbiosis of legumes with Rhizobia leads to an improved nutritional status, mainly in nitrogen (Figure 1B), which in turn leads to increased photosynthetic rates and beneficial effects on plant growth and yield. Not only is pea crop able to establish symbioses with N-fixing bacteria, but it simultaneously interacts with arbuscular mycorrhizal fungi (AMF) supplying phosphate (Figure 1B). In agroecology, this symbiotic interaction is called “tripartite” because it implies three partners: the plant, the bacteria, and the fungi. Legumes exploit this synergistic biofertilization from both the bacteria and fungi, thereby improving yield and grain quality.

In return, legumes provide a significant amount of its fixed photosynthetic carbon to the symbiont (Kaschuk et al., 2009). Noteworthy, the increased photosynthetic rates not only result from the increased nutrient uptake provided by these symbionts, but also from sink stimulation (Kaschuk et al., 2009). To our knowledge, no study has estimated plant carbon fluxes in tripartite symbiosis, but authors rather focused on the improved seed quality and yield gained through synergistic symbioses (Geneva et al., 2011; Shinde & Thakur, 2016; Xavier & Germida, 2003). In both mycorrhizal and rhizobial interactions, sugar transport systems are also involved, and we previously speculated that a common transport pathway might be shared between both symbioses (Hennion et al., 2019). Indeed, the expression of several sugar transporters is regulated in mycorrhizal and Rhizobia interactions (Banasia et al., 2021; Doidy et al., 2012), suggesting that they may participate in sucrose distribution towards the symbionts. Just to name a few, this concerns several SUTs and SWEETs in legumes: MtSUTs (Doidy et al., 2012), GmSUT1 (Deng et al., 2021); MtSWEET1b (An et al., 2019) (Figure 3), MtSWEET11 (Kryvoruchko et al., 2016), and LjSWEET3 (Sugiyama et al., 2017). Orthologous genes in pea also seem regulated in plant–microbe interactions (Alves-Carvalho et al., 2015; Doidy et al., 2019; Zhou et al., 2007). Very few studies have so far focused on the molecular players involved in the synergistic co-inoculation of bacterial and mycorrhizal fungi. Up to now, the only information comes from an elegant split-root system, showing that legumes allocated more carbon to Rhizobia under nitrogen demand, but more carbon to the fungal partner when nitrogen is available in the fungal compartment (Kafel, 2018). This report also highlighted several SWEETs (MtSWEET1b, MtSWEET6, MtSWEET11, MtSWEET12, MtSWEET15c, and MtSWEET15d) regulated in Medicago truncatula roots depending on the colonization with different root symbionts subjected to different nutrient supply conditions.
Plants, including legume species, also interact with a diverse set of non-pathogenic rhizobacteria species (Figure 1B), commonly named Plant Growth Promoting Rhizobacteria (PGPR). PGPR can confer to plants an improved growth and/or tolerance to various biotic and abiotic stresses and present a high potential for agricultural applications. Currently, this prospect is limited by the lack of information on the molecular mechanisms involved in these biological processes (Vacheron et al., 2013). Interactions of plants with these beneficial soil microorganisms involve the liberation of plant organic compounds via rhizodeposition, which is itself dependent on the plant carbon budget (Lepinay et al., 2012). As mentioned above, a growing body of evidence demonstrates the importance of sugar transport in plant-pathogen resistance and in plant-microorganism mutualistic symbioses. In contrast, the role and regulation of sugar transporter activities in plant-PGPR interactions remain largely unknown (Hennion et al., 2019). Using the model plant Arabidopsis and a collection of PGPR strains, it was recently shown that PGPR can induce major transcriptional changes in plant sugar transport (Desrut et al., 2020; Desrut et al., 2021). Moreover, it was shown that AtSWEET11 and AtSWEET12, two sugar transporter genes whose expression is repressed by the well-characterized PGPR strain Pseudomonas simiae WCS417r, are functionally involved in its plant growth-promoting effects, possibly by controlling the allocation of carbon resources (Desrut et al., 2020).

In addition to their positive effects on plant productivity under optimal growth conditions, evidence in the literature shows that beneficial microorganisms, including Rhizobia, AMF, and PGPR, are also efficient in alleviating the harmful effects of drought and other abiotic stress in legumes (reviewed in Nadeem et al., 2019). Such research area should help improve our understanding of plant resilience to its environment. Although many studies have focused on sugar transporters’ expression in N-fixing and AM symbioses, much more attention should be paid to their role in stress acclimatation.

5 | CARBON FLUXES DURING FLOWERING

The transition from vegetative to reproductive development implies the emergence of new sink organs (flowers and fruits) (Figure 2) and occurs in response to environmental factors and internal stimuli (for a review see Weller & Ortega, 2015). Flowering time is influenced by photoperiod (Lejeune-Hénaut et al., 1999) and temperature (Alcalde et al., 2000; Murfet & Reid, 1973). In legumes, the relationship between sugars and induction of flowering remains to be uncovered, although resource availability is tightly linked to flower abortion (Guilioni et al., 1997). In eudicots, floral meristem initiation, floral bud development, and transition to mature flowers depend on C provided by photosynthesis (Patrick & Colynas, 2014). Indeed, in several species, the concentration of sucrose in the phloem sap increases to supply shoot apical meristem with sucrose and induces floral transition (Cho et al., 2018; Yoon et al., 2021). In soybean, abortion occurs more frequently during anthesis, which corresponds to a decrease in photo-assimilates imported into floral structures (Patrick & Colynas, 2014). A study on the late-flowering phenotype of the FT mutant in Arabidopsis further demonstrates that sucrose functions downstream of CONSTANS (CO) and upstream of FLOWERING LOCUS T (FT) (Cho et al., 2018; Yoon et al., 2021). Moreover, in Arabidopsis, Tre6P, which is synthesized in the phloem companion-cell-sieve element complex, acts as a signal for sucrose availability. Indeed, Tre6P favors sucrose demand of growing sink organs and its involvement in the flowering process has been recently identified (for review see Fichtner & Lunn, 2021).

Sucrose demand by sink organs is controlled by SUT and SWEET transporter families (Chen et al., 2010; Schneider et al., 2012; Sivitz et al., 2007). Several of them are specific to the reproductive organs and their high level of expression during flowering and fertilization indicates the importance of sugar exchanges in these processes (Andrés et al., 2020; Durand et al., 2018; Gu et al., 2020; Iftikhar et al., 2020). In pea, multiple sugar transporter and invertase genes were specifically induced in flowers (Doidy et al., 2019; Morin et al., 2022), but their expression pattern has been much more detailed in other model species. For instance, a study in petunia (Petunia axillaris) at five flowering stages indicates an increased expression of two SUC genes (PaSUT1 and PaSUT3) and five SWEET genes (PaSWEETS13c, PaSWEET9a, PaSWEET1d, PaSWEET5a, and PaSWEETS14a) during floral development. Other SUT and SWEET genes are expressed during specific stages of flowering development (Iftikhar et al., 2020). As in Arabidopsis (AtSWEET9, Lin et al., 2014), PaSWEET9c is specifically expressed in nectaries of petunia (Iftikhar et al., 2020). Moreover, PaSWEET10a and PaSWEET10b display a higher expression in the early flowering stage in petunia (Iftikhar et al., 2020) like AtSWEET10, their ortholog in Arabidopsis, whose expression has been found induced by FLOWERING LOCUS T (FT) (Andrés et al., 2020).

Drought stress mostly occurs during flowering time in pea (Figure 2). Indeed, since winter and spring varieties flower, respectively, at the beginning of April and early June, pea often faces drought events during its reproductive development. Legumes are highly vulnerable to drought during the flowering phase, which leads to a shorter flowering period, fewer flower, and pod numbers, and a significant reduction in seed yield (Nadeem et al., 2019) (Table 1). Water stress applied at the flower onset also led to a marked decrease in biomass of non-reproductive organs in pea (Prudent et al., 2016) (Table 1). In pigeon pea (Cajanus cajan), water deficit applied during flowering caused over 50% reduction in seed yield, probably due to reduced photosynthesis activity leading to a disturbance of assimilate partitioning (Fahad et al., 2017). Floral organs are very strong sinks, especially during anther development when sugar import is ensured by cell wall invertases (CWINV), sucrose synthases (SUS), and sugar transporters (STP) (Sinha et al., 2021; Yu et al., 2019). Water deficit notably reduces invertase activity, preventing pollen from metabolizing sucrose to hexose, and thus, leading to sterility (Yu et al., 2019).

As high temperatures are often associated with water deficit, photosynthesis activity, and grain yield are negatively impacted by these stresses (Guilioni et al., 1997; Rane et al., 2021) (Table 1). Both drought and heat stress impact source/sink relationship by decreasing...
photosynthesis rate and sucrose export to sink organs (Fahad et al., 2017; Hageman & Van Volkenburgh, 2021). This induces impairment in sugar availability (import and utilization) during reproductive growth by reducing the enzyme activity involved in sugar metabolism and transport (Li et al., 2015; Sinha et al., 2021; Yu et al., 2019). Interestingly, drought or heat stress-tolerant genotypes are able to maintain sugar concentration and partitioning (Dong & Beckles, 2019; Li et al., 2015; Sinha et al., 2021).

6 | CARBON FLUXES DURING REPRODUCTIVE DEVELOPMENT

6.1 | Filling carbon nutrients and sugar signals during seed development

Once flowers are pollinated, seed development occurs in a series of temporally and spatially defined steps. In Fabaceae, seed development can be divided into three phases: embryogenesis, a phase of cell division, followed by seed-filling, a phase of cell expansion, and finally a phase of desiccation/maturation (Ruan et al., 2012; Wang et al., 2019; Weber et al., 2005). Seed development highly depends on nutrient remobilization through source-sink long-distance transport driven by seed sink strength (Hageman & Van Volkenburgh, 2021; Lu et al., 2020; Pampana et al., 2016a, 2016b; Weber et al., 1997b). This latter is promoted by embryonic cell division, which occurs quicker in seeds from distal nodes, where mitotic activity is higher than in proximal ones (Munier-Jolain & Ney, 1998; Munier-Jolain & Salon, 2003). This is potentially linked to the higher sink strength of distal nodes (Munier-Jolain & Salon, 2003). As seed abortion is directly linked to C availability, seeds from distal nodes tend to abort more due to their higher demand (Guillioni et al., 2003; Ney et al., 1993), as shown by the decrease of seed number in these nodes (Jeuffroy & Devienne, 1995) (Figure 2).

Seed development is highly dependent on carbon remobilization and allocation mediated by sugar transporters. One day after fertilization in pea, an increase in the expression of PsSUT2 (renamed PsSUT1.4) and PsSUT5 (renamed PsSUT1.2) was observed in flowers, suggesting that these transporters may potentially be involved in sucrose transport to sustain mitotic activity and maintain osmotic pressure for embryonic growth (Smitha Ninan et al., 2017). First, in pea (Tegeder et al., 1999) and in faba bean (Vicia faba) (Offler et al., 1989), nutrients are unloaded into the seed coat from the chalazal vein through a symplastic pathway (Patrick, 1997; Van Dongen et al., 2003) (Figure 3). Then, in developing seeds, maternal tissues are physically separated from filial tissues, which implies the contribution of sugars transport systems (Lu et al., 2020; Tegeder et al., 1999; Weber et al., 1997a; Zhou et al., 2007). Here, export of sugars from the seed coat to the apoplast may involve both a passive (de Jong et al., 1997; Zhou et al., 2007) and an active component (Figure 3). At the molecular level, sugar provision towards the embryo is highly regulated by sugar transporters and invertases (Doidy et al., 2019; Morin et al., 2022) (Figure 3). During the early embryonic development, a cell-wall bound invertase activity has been observed in the seed coat of legumes. In pea and in faba bean, this activity is associated with a high level of glucose and fructose in the apoplast surrounding the embryo (Weber et al., 1996) (Figure 3). Thus, invertase controls the ratio of sucrose to hexose and is likely a signal for the high mitotic divisions required during embryogenesis (Morin et al., 2022; Weber et al., 1995). During the later stages of embryogenesis, it is commonly accepted that sucrose is released from the seed coat. In pea, several studies have shown that sucrose facilitators PsSUT1 (renamed PsSUT1.2) and PsSUF4 (renamed PsSUT4.2) are expressed in transfer cells of the seed coat parenchyma to allow sucrose unloading to the seed apoplast (Zhang et al., 2007; Zhou et al., 2007) (Figure 3).

After its translocation in the apoplast, the outcome of sucrose depends on the phase of seed development. During the endosperm cellularization (heart and torpedo stages), it has been shown in faba bean (Weber et al., 1996) that acid invertases cleave sucrose into hexoses (Figure 3), creating a favorable sucrose gradient between the seed coat and the apoplast, and thus drive sucrose unloading (Weber et al., 1995). Expression of PsSUT1.1 during this step also suggests that part of sucrose escapes invertase hydrolysis and is directly imported into transient endosperm and embryo cells (Melkus et al., 2009; Rosche et al., 2002) (Figure 3). During the cotyledon development stage, which starts at the torpedo stage, sucrose is taken up from the apoplast into the cotyledons thanks to PsSUT1.1, a proton-coupled sucrose transporter localized in the cotyledon epidermal cells (Tegeder et al., 1999; Weber et al., 1997a; Zhou et al., 2007) (Figure 3). Interestingly, PsSUT1.1 overexpression resulted in increased sucrose content in developing seeds, and plants eventually produced more protein and starch, as well as higher seed yield (Lu et al., 2020; Zhou et al., 2007).

In soybean and faba bean, GmSWEET15a/b mediates sucrose transport from the endosperm to the embryo (Figure 3), while the high hexose status driving the embryonic growth through cell division is maintained by VfCWINV1 (Wang et al., 2019; Weber et al., 1996). In Arabidopsis, SWEET genes of the clade III, AtSWEET11 and AtSWEET15, are also implicated in this sucrose transfer from the endosperm to the embryo (Chen et al., 2015). In pea, we identified the gene orthologs, PsSWEET15 (Doidy et al., 2019) and PsCWIN1.2 (Morin et al., 2022), as highly expressed during the embryonic stage and regulated by water stress (Figure 3). When embryogenesis ends, the Final Stage in Seed Abortion (FSSA) is passed, thus the seed can no longer abort (Duthion & Pigéaire, 1991). Altogether this emphasizes the important role of SWEET and INV during early seed development, thus controlling abortion rates and final seed size (Morin et al., 2022; Wang et al., 2019; Weber et al., 1996).

During the seed-filling stage, growth continues through cell expansion, and simultaneously sucrose assimilation increases, and soluble protein and starch synthesis begin (Patrick & Offler, 2001). A decrease in hexose to sucrose ratio is marked by a sharp decrease in PsINV gene expression and enzymatic activities (Morin et al., 2022; Weber et al., 2005). Notably, this temporal downregulation of PsINV activity results from the progressive mechanical crushing of seed coat cells expressing the enzyme (Weber et al., 1995). Sucrose is released.
from the seed coat and taken up directly by the cotyledons through PsSUT1.1 and stored as starch (Hageman & Van Volkenburgh, 2021) (Figure 3). In pea, sucrose fluxes from the seed coat to the embryo are coordinated through turgor homeostasis, detecting variation of the apoplastic sucrose pool through turgor modifications, and regulating PsSUT1.1 expression (Zhou et al., 2009). This enables a balanced control between sucrose supply and demand of the embryo. While invertase activity declines, sucrose synthase and sucrose synthase phosphatase are strongly expressed in the cotyledons, thereby promoting the onset of starch synthesis and accumulation (Lu et al., 2020; Weber et al., 1997a; Yang et al., 2019; Yu et al., 2020).

In pea, starch synthesis during seed-filling is promoted by Tre6P and there is evidence that the plant hormone auxin plays an important role in controlling sucrose utilization (McAdam et al., 2017). Indeed, it has been observed that Tre6P stimulates auxin synthesis through the induction of TRYPTOPHANE AMINOTRANSFERASE RELATED2 (TAR2). This study concludes that a certain auxin level is needed for the action of Tre6P for promoting seed-filling and auxin might act downstream of Tre6P to accelerate storage processes (Meitzel et al., 2021). The last stage of seed development, dormancy, is dependent on FLOWERING LOCUS T in pea, and FT is also regulated by Tre6P signaling pathway in Arabidopsis silique (Fichtner & Lunn, 2021; Meitzel et al., 2021). Moreover, in Arabidopsis, ABA is implicated in seed dormancy driven through the ABA INSENSITIVE4 gene expression associated with AtTPS1 (TREHALOSE-6-PHOSPHATE SYNTHASE 1) gene expression. These results indicate that Tre6p might have a role in seed maturation since trehalose pathway mutants had lost the ABA sensitivity and could not enter the dormancy stage (for review see Fichtner & Lunn, 2021).

### 6.2 Carbon fluxes towards seeds are impacted by environmental changes

As discussed above, drought negatively impacts seed yield (Sánchez et al., 1998; Sousa-Majer et al., 2004; Strydhorst et al., 2015) (Figure 2) because of an increase in flower and pod abortions reducing seed number (Guillioni et al., 1997) (Table 1). Water deficit also directly impacts seed development, and this more strongly during early seed development (prior FSSA) than at later stage of seed-filling (Mahieu et al., 2009). When drought stress occurs during the filling stage, individual seed weight and phytotherm number decrease (Guillioni et al., 2003; Nadeem et al., 2019; Ney et al., 1993) (Table 1) while leaf senescence is accelerated (Rane et al., 2021), therefore affecting source-sink relationships (Chen et al., 2006). Drought (and heat stress) changes the distribution of pea seeds along the stem (Guillioni et al., 2003). Indeed, pods located on basal phytomers had more seeds in response to stress conditions than upper nodes. This trend is likely linked to the higher demand of upper nodes (Munier-Jolain & Salon, 2003) (Figure 2), which may not be fulfilled because of a decrease in net photosynthesis during stress conditions (Guillioni et al., 2003).

Interestingly in common bean, sucrose content in seeds increased more in the drought-tolerant variety than in the sensitive one when water deficit was applied during early seed-filling (Gebeyehu et al., 2011). However, both varieties displayed lower starch accumulation in seeds, potentially due to limitations in assimilate availability by the source. However, in soybean, drought applied during reproductive development increased soluble sugars in leaves and reduced photosynthetic rates (Du et al., 2020). This increase of sugars in leaves may increase the amount of sucrose loaded into the phloem for the nutrition of sink organs. In parallel, activities of sucrose phosphate synthase, sucrose synthase and acid invertases, as well as the expression of their respective encoding genes, increased in leaves (Du et al., 2020). The authors also showed higher expression of genes related to sugar metabolism and transport in developing seeds, suggesting maintenance of the push and pull mechanism during drought stress. Indeed, in common bean, tolerance to water deficit may be linked to both sink strength maintenance and the capacity of the source organs to produce and export carbohydrates towards sink organs (Hageman & Van Volkenburgh, 2021). However, Hageman & Van Volkenburgh (2021) describe that source strength is not a good indicator for yield (unlike the pod harvest index), both under normal growing conditions and during water stress.

During drought stress, stomatal closure in leaves leads to a reduction in photosynthesis in green peas, which limits C availability and causes yield losses (Nemeskéri et al., 2015). In addition to its effect on stomatal closure during water deficit, ABA also acts on the loading of resources (Hageman & Van Volkenburgh, 2021). For instance, ABA is known to increase the expression of cell wall invertase inhibitor genes resulting in decreased activity of CWIs involved in maintaining sucrose gradient and sink strength, which at term impact seed size (Ruan, 2014; Ruan et al., 2010; Westgate et al., 1996).

Pathogens also influence the proper course of reproductive development and source-sink relationships. For instance, Ascochyta blight (Figure 2) is caused by a fungus (Didymella piniodes) that penetrates through stomatal apertures in the spring and its severity increases after flowering, causing severe yield loss, especially in early-maturing varieties (Bretag et al., 1995) (Table 1). Up to now, breeding for resistant cultivars with suitable agronomic traits has had limited success (Khan et al., 2013). The C/N ratio in seeds is also affected by an increased protein concentration and a decreased starch content in response to Ascochyta blight pathogen (Garry et al., 1996) (Table 1). Results from a comprehensive study comparing the metabolic and proteomic profiles of the susceptible pea cultivar Messire with the more tolerant cultivar Protecta revealed that enhanced tolerance towards D. piniodes is notably associated with higher sugar levels, including a significant increase in maltose and fructose as well as in the TCA cycle compounds, pyruvate and citrate (Turetschek et al., 2017). Altogether, these results suggest that, during the defense response against necrotrophs, plant resistance is enhanced if cell death is not triggered, which limits the release of sugars to the pathogen. Plant sugar levels then remain high because they cannot be metabolized by the fungi (Turetschek et al., 2017). In a study focusing on unraveling resistance genes to this pathogen in the Cicer genus, the authors identified a probable sugar transporter (Newman et al., 2021). This transporter may be involved in retrieving sugars.
from the apoplastic to limit extracellular sugar pools available for the pathogen. To our knowledge, very little data are available to date on pea sugar homeostasis and transporter in relation to these different environmental conditions. Thus, research strategies are currently being developed to unravel the mechanisms that should contribute to the creation or selection of new pea varieties more resilient to the present and future challenging environments.

Several studies have shown that beneficial microorganisms, previously described in this work, often result in improved yield and seed quality at maturity (Geneva et al., 2011; Shinde & Thakur, 2016; Xavier & Germida, 2003), and this occurs through their effect during the different stages of plant development (Figures 1 and 2) and in response to various abiotic stresses (Nadeem et al., 2019). Therefore, exploiting such associations with beneficial microbes represents a promising strategy to increase legume crop resilience towards environmental changes, especially during their crucial developmental phase of seed production.

7 FUTURE MILESTONES: ENGINEERING SUGAR TRANSPORTERS TO IMPROVE SEED YIELD, QUALITY, AND STRESS RESILIENCE IN LEGUMES

We reviewed here the importance of C fluxes and resource allocation during key developmental processes of P. sativum and its multifactorial interactions with the environment. Selection of new pea genotypes for higher yield stability should now consider the genotype-environment interactions and their impact on shoot (source), root and seed (sink) developments (Vocanson et al., 2006).

A first breeding strategy should focus on the phenotypic diversity of pea. Indeed, pea plant architecture is an important factor in determining yield and recent works highlight possible targets to increase plant productivity and stress resistance (Guo et al., 2020). In pea, the impact of water deficit seems to be tightly linked to leaf architecture with a better resistance for afila than leafy varieties (Baigorri et al., 1999). In fact, the absence of leaflets in afila genotypes does not negatively impact photosynthates export towards pod (Harvey, 1974). The author hypothesized that a better illumination, and so a higher photosynthesis of the pods in the afila genotypes, could explain this lack of difference. Therefore, the afila genotypes (widely cultivated nowadays) seem comparable to classical leafy phenotypes, in terms of C assimilation and export to the pod (Baigorri et al., 1999). The root system architecture is also a key feature to take into account in the next breeding programs to improve stress resilience and nutrient remobilization, since yield stability in field conditions was correlated to the rooting depth (Bertholdsson, 1989). Moreover, genotypes displaying longer roots during early development showed increased root growth time and water use efficiency, leading to a deeper rooting at later developmental stages (Thorup-Kristensen, 1998). This observation could be linked to a better use of starch by the roots. As starch is the main form of C storage, starch metabolism could be a possible target to explore adaptive changes in the source-sink allocation of plants to face abiotic stress (Dong & Beckles, 2019; Li et al., 2015).

Secondly, exploiting Pisum genetic variability should also enable breeders to create novel pea varieties more productive and stress-resilient. Future research strategy needs to focus on the regulation of assimilate partitioning and carbon metabolism in sink organs to improve grain filling under adverse conditions (Dong & Beckles, 2019; Fahad et al., 2017; Hageman & Van Volkenburgh, 2021; Li et al., 2015). Thus, characterization of the molecular actors involved in carbon partitioning between source and sink organs is key to achieving this goal. Multiple mutants in carbon utilization have been described with defects in the starch metabolic pathway in pea (Yu et al., 2020). For instance, molecular biologists have re-discovered in the 90s that the wrinkled trait (rr), initially observed by Gregor Mendel, was due to a disruption in SBE1, a gene responsible for branching starch molecules (Bhattacharyya et al., 1990). However, to the best of our knowledge, only a single mutant is available for sugar transporters in pea. Lines overexpressing PsSUT1.1 showed increased sucrose phloem loading and nutrient fluxes towards sink organs (Figure 3) and produced higher seed yield with more protein and starch (Lu et al., 2020). Sugar transporters like PsSUT1.1 (Lu et al., 2020) that play a central role in phloem loading in source organs and control sink carbon storage and seed yield (Figure 3), represent therefore key targets for breeders and farmers. The potential of harnessing carbon fluxes is just starting with the recent identification of sugar transport systems in legume genomes (Doidy et al., 2019).

In addition, uncovering the influence of environmental factors along the plant lifecycle is an important area of research. In pea, in response to drought stress, vegetative growth was stopped and leaf senescence was induced so that similar partitioning of dry weight to pod and seeds was maintained as in control conditions (Baigorri et al., 1999). During this last stage of development, an increased expression of several clade II and III SWEETs associated with a high level of SUT1 and 2 was noticed in pea leaves (Ninan et al., 2019). Functional characterization of these genes might help improve our understanding of carbon flux regulation in response to water stress. Moreover, Hageman and Van Volkenburgh (2021) propose that genetic lines with decreased sensitivity to ABA, which could maintain a strong sink strength and a higher yield, might represent a promising way to identify tolerant genotypes to water deficit. This perspective should be taken with caution since a loss of ABA sensitivity could increase water sensitivity in some species. As this review points out, some knowledge is already gathered on such actors; however, the complete picture is not yet drawn, and future studies will enable us to implement a model describing carbon fluxes in P. sativum and create new varieties more resilient to environmental changes. This could be reached through agronomic practices, as well as conventional breeding approaches, promoting beneficial microorganisms, or quantitative trait loci (QTLs) analyses (Nadeem et al., 2019).

Other promising ways to improve pea genotypes are biotechnol-ogy and functional genomics. New gene-editing technologies, such as CRISPR/Cas9, represent promising tools for legume improvement (Bhowmik et al., 2021; Nadeem et al., 2019). For instance, CRISPR/
Cas9 has successfully been developed in chickpeas with associated drought tolerance (Badhan et al., 2021). Targeting sugar transport system, such as vacuolar components maintaining cell turgescence in response to water deficit (Slawinski et al., 2021), may also be a hopeful opportunity to develop climate-resilient crops. In conclusion, plant membrane transporters represent a key target to increase crop yield and quality, as well as to improve the sustainable production of nutritious foods (Banasiak et al., 2021; Schroeder et al., 2013).

AUTHOR CONTRIBUTIONS
All authors contributed to proofreading the manuscript. Amélie Morin contributed to the part on germination and reproductive development, designed Figures 1, 2 and 3, and Table 1. Laurence Maurousset contributed to the part on abiotic stresses during vegetative and reproductive stages. Cécile Vriet contributed to the part on biotic interactions during vegetative and reproductive development. Rémi Lemoine contributed to the part on leaf development. Joan Doidy contributed to the part on biotic interactions during vegetative and reproductive development, the introduction and the abstract. Nathalie Pourtau contributed to the part on root development, abiotic stresses during the vegetative stage and edited the manuscript.

ACKNOWLEDGEMENT
This work was supported by the University of Poitiers, the Centre National de la Recherche Scientifique (CNRS) and by a doctoral grant from Grand Poitiers awarded to AM.

DATA AVAILABILITY STATEMENT
Data sharing is not applicable to this article as no new data were created or analyzed in this study.

ORCID
Amélie Morin https://orcid.org/0000-0003-3318-8778
Cécile Vriet https://orcid.org/0000-0001-6625-268X
Joan Doidy https://orcid.org/0000-0003-3966-3218
Nathalie Pourtau https://orcid.org/0000-0003-1200-4968

REFERENCES
Alcalde, J.A., Wheeler, T.R. & Summerfield, R.J. (2000) Genetic characterization of flowering of diverse cultivars of pea. Agronomy Journal, 92, 772–779.
Ali-Khan, S.T. & Snoad, B. (1977) Root and shoot development in peas. The Annals of Applied Biology, 85, 131–136.
Al-Quraan, N.A., Al-Ajlouni, Z.I. & Qawasma, N.F. (2021) Physiological and biochemical characterization of the GABA shunt pathway in pea (Pisum sativum L.) seedlings under drought stress. Horticulture, 7, 125. Alves-Carvalho, S., Aubert, G., Carrère, S., Cruaud, C., Brochet, A.-L., Jacquin, F. et al. (2015) Full-length de novo assembly of RNA-seq data in pea (Pisum sativum L.) provides a gene expression atlas and gives insights into root nodulation in this species. The Plant Journal, 84, 1–19.
An, J., Zeng, T., Ji, C., de Graaf, S., Zheng, Z., Xiao, T.T. et al. (2019) A Medicago truncatula SWEET transporter implicated in arbuscular mycorrhizal symbiosis. The New Phytologist, 224, 396–408.
Andrés, F., Kinoshita, A., Kalluri, N., Fernández, V., Falavigna, V.S., Cruz, T. M.D. et al. (2020) The sugar transporter SWEET10 acts downstream of FLOWERING LOCUS T during floral transition of Arabidopsis thaliana. BMC Plant Biology, 20, 53.
Armstrong, E.L., Pate, J.S. & Tennant, D. (1994) The field pea crop in South Western Australia – patterns of water use and root growth in genotypes of contrasting morphology and growth habit. Functional Plant Biology, 21, 517–532.
Ayre, B.G. (2011) Membrane-transport Systems for Sucrose in relation to whole-plant carbon partitioning. Molecular Plant, 4, 377–394.
Badhan, S., Ball, A.S. & Mantri, N. (2021) First report of CRISPR/Cas9 mediated DNA-free editing of 4CL and RVE7 genes in chickpea protoplasts. International Journal of Molecular Sciences, 22, E396.
Baigorri, H., Antolín, M.C. & Sánchez-Díaz, M. (1999) Reproductive response of two morphologically different pea cultivars to drought. European Journal of Agronomy, 10, 119–128.
Baldwin, L., Domon, J.-M., Klimek, J.F., Fournet, F., Sellier, H., Gillet, F. et al. (2014) Structural alteration of cell wall pectins accompanies pea development in response to cold. Phytochemistry, 104, 37–47.
Banasiak, J., Jamruszka, T., Murray, J.D. & Jasinski, M. (2021) A roadmap of plant membrane transporters in arbuscular mycorrhizal and legume–rhizobium symbioses. Plant Physiology, 187, 2071–2091.
Barbier, F., Péron, T., Lecerf, M.-P., Perez-García, M.-D., Barrière, Q., Rolčík, J. et al. (2015) Sucrose is an early modulator of the key hormonal mechanisms controlling bud outgrowth in Rosa hybrida. Journal of Experimental Botany, 66, 2569–2582.
Barbier, F.F., Cao, D., Fichtner, F., Weiste, C., Perez-García, M., Caradeuc, M. et al. (2021) HEXOKINASE1 signalling promotes shoot branching and interacts with cytokinin and strigolactone pathways. The New Phytologist, 231, 1088–1104.
Bastianelli, D., Crosjean, F., Peyronnet, C., Duparque, M. & Régnier, M. (1998) Feeding value of pea (Pisum sativum, L.) 1. Chemical composition of different categories of pea. Animal Science, 67, 609–619.
Bénézet, M., Blamé, V. & Jeufroy, M.-H. (2017) Impact of climate and diseases on pea yields: what perspectives with climate change? OCL, 24, D103.
Bertheloot, J., Barbier, F., Boudon, F., Perez-García, M.D., Péron, T., Citerne, S. et al. (2020) Sugar availability suppresses the auxin-induced strigolactone pathway to promote bud outgrowth. The New Phytologist, 225, 866–879.
Bertholdsson, N.-O. (1989) Rotutveckling och odlingssakerhet hos arter. Sveriges Utsadesforsknings Tidskrift, 99, 195–203.
Bhattacharyya, M.K., Smith, A.M., Ellis, T.H., Hedley, C. & Martin, C. (1990) The wrinkled-seed character of pea described by Mendel is caused by a transposon-like insertion in a gene encoding starch-branching enzyme. Cell, 60, 115–122.
Bhowmik, P., Konkin, D., Polowick, P., Hodgins, C.L., Subedi, M., Xiang, D. et al. (2021) CRISPR/Cas9 gene editing in legume crops: opportunities and challenges. Legume Sci., 3, e96.
Blicharz, S., Beemster, G.T.S., Ragni, L., De Diego, N., Spichal, L., Hernández, A.E. et al. (2021) Phloem exudate metabolic content reflects the response to water-deficit stress in pea plants (Pisum sativum L.). The Plant Journal, 106, 1338–1355.
Blöchl, A., Peterbauer, T. & Richter, A. (2007) Inhibition of raffinose oligosaccharide breakdown delays germination of pea seeds. Journal of Plant Physiology, 164, 1093–1096.
Bourion, V., Lejeune-Hénaut, I., Munier-Jolain, N. & Salon, C. (2003) Cold acclimation of winter and spring peas: carbon partitioning as affected by light intensity. European Journal of Agronomy, 19, 535–548.
Bretag, T.W., Keane, P.J. & Price, T.V. (1995) Effect of Ascochyta blight on the grain yield of field peas (*Pisum sativum* L.) grown in southern Australia. *Australian Journal of Experimental Agriculture, 35*, 531–536.

Chandran, D. (2015) Co-option of developmentally regulated plant SWEET transporters for pathogen nutrition and abiotic stress tolerance. *IUBMB Life, 67*, 461–471.

Chen, C., Miller, P., Muehlbauer, F., Neill, K., Wichman, D. & McPhee, K. (2006) Winter pea and lentil response to seeding date and micro- and macro-environments. *Agronomy Journal, 98*, 1655–1663.

Chen, L.-Q. (2014) SWEET sugar transporters for phloem transport and pathogen nutrition. *The New Phytologist, 201*, 1150–1155.

Chen, L.-Q., Hou, B.-H., Lalonde, S., Takanaga, H., Hartung, M.L., Qu, X.-Q. et al. (2010) Sugar transporters for intercellular exchange and nutrition of pathogens. *Nature, 468*, 527–532.

Chen, L.-Q., Lin, I.W., Qu, X.-Q., Sosso, D., McFarlane, H.E., Londóno, A. et al. (2015) A cascade of sequentially expressed sucrose transporters in the seed coat and endosperm provides nutrition for the Arabidopsis embryo. *Plant Cell, 27*, 607–619.

Chen, L.-Q., Qu, X.-Q., Hou, B.-H., Sosso, D., Osorio, S., Fernie, A.R. et al. (2012) Sucrose efflux mediated by SWEET proteins as a key step for phloem transport. *Science, 335*, 207–211.

Chen, W., Yao, Q., Patil, G.B., Agarwal, G., Deshmukh, R.K., Lin, L. et al. (2016) Identification and comparative analysis of differential gene expression in soybean leaf tissue under drought and flooding stress revealed by RNA-Seq. *Frontiers in Plant Science, 7*, 1044.

Cho, L.-H., Pasrlga, R., Yoon, J., Jeon, J.-S. & An, G. (2018) Roles of sugars in controlling flowering time. *Journal of Plant Biology, 61*, 121–130.

Côté, R., Geraud, J.M., Peterson, C.A. & Grodzinski, B. (1992a) Sink to source transition in tendrils of a Semileafless mutant, *Pisum sativum* cv curly. *Plant Physiology, 100*, 1640–1648.

Côté, R., Thompson, R.G. & Grodzinski, B. (1992b) Photosynthetic oxygen production facilitates translocation of 11C-labelled Photoassimilates from tendrils and leaflets of *Pisum sativum* var. *Journal of Experimental Botany, 43*, 819–829.

Couchoud, M., Salon, C., Girotet, S., Jeydy, C., Vermoud, V. & Prudent, M. (2020) Pea efficiency of post-drought recovery relies on the strategy to fine-tune nitrogen nutrition. *Frontiers in Plant Science, 11*, 204.

De Jong, A., Koerselman-Kooij, J.W., Schuurmans, J.A.M.J. & Borstlap, A.C. (1997) The mechanism of amino acid efflux from seed coats of developing pea seeds as revealed by uptake experiments. *Plant Physiology, 114*, 731–736.

de Sousa-Majer, M.J., Turner, N.C., Hardie, D.C., Morton, R.L., Lamont, B. & TJV, H. (2004) Response to water deficit and high temperature on transgenic *Pisum sativum* (cv. *Pisum sativum*) containing a seed-specific amylose inhibitor and the subsequent effects on pea weevil (*Bruchus pisorum*) survival. *Journal of Experimental Botany, 55*, 497–505.

Deng, L., Zhao, S., Yang, G., Zhu, S., Tian, J. & Wang, X. (2021) Soybean GmSUT1 transporter participates in sucrose transport to nodules during rhizobial symbiosis. *Plant Growth Regulation, 96*, 119–129. https://doi.org/10.1007/s10725-021-00764-y

Desrut, A., Moumen, B., Thibault, F., Le Hir, R., Coutos-Thévenot, P. & Vriet, C. (2020) Beneficial rhizobacteria *pseudomonas simiae* WCS417 induce major transcriptional changes in plant sugar transport. *Journal of Experimental Botany, 71*, 7301–7315.

Desrut, A., Thibault, F., Mercado-Blanco, J., Coutos-Thévenot, P. & Vriet, C. (2021) Transcriptional regulation of plant sugar transporter genes by beneficial rhizobacteria. *Journal of Plant Interactions*, 16, 443–451.

Devi, R., Munjal, N., Gupta, A.K. & Kaur, N. (2007) Cadmium induced changes in carbohydrate status and enzymes of carbohydrate metabolism, glycolysis and pentose phosphate pathway in pea. *Environmental and Experimental Botany, 61*, 167–174.

Dhandapani, P., Song, J., Novak, O. & Jameson, P.E. (2017) Infection by *Rhodococcus fascians* maintains cotyledons as a sink tissue for the pathogen. *Annals of Botany, 119*, 841–852.

Dhankher, O.P. & Foyer, C.H. (2018) Climate resilient crops for improving global food security and safety. *Plant, Cell & Environment, 41*, 877–884.

Dinant, S. & Lemoine, R. (2010) The phloem pathway: new issues and old debates. *Comptes Rendus Biologies, 333*, 307–319.

Doidy, J., Grace, E., Kühn, C., Simon-Plas, F., Casieri, L. & Wipf, D. (2012) Sugar transporters in plants and in their interactions with fungi. Trends in *Plant Science, 17*, 413–422.

Doidy, J., Vidal, U. & Lemoine, R. (2019) Sugar transporters in Fabaceae, featuring SUT MST and SWEET families of the model plant *Medicago trunculata* and the agricultural crop *Pisum sativum*. *PLoS One, 14*, e0223173.

Dong, S. & Beckles, D.M. (2019) Dynamic changes in the starch-sugar interconversion within plant source and sink tissues promote a better abiotic stress response. *Journal of Plant Physiology, 234–235*, 80–93.

Doré, T. (1994) Influence sur l’évolution du nombre de ramifications et de tiges chez le pois. *Agrophysiologie du pois protéagineux, UNIP - ITCF - INRA, mai*, 145–154.

Du, Y., Zhao, Q., Chen, L., Yao, X., Zhang, H., Wu, J. et al. (2020) Effect of drought stress during soybean R2–R6 growth stages on sucrose metabolism in leaf and seed. *International Journal of Molecular Sciences, 21*, 618.

Durand, M., Mainson, D., Porcheron, B., Mauroosset, L., Lemoine, R. & Pourtau, N. (2018) Carbon source-sink relationship in *Arabidopsis thaliana*: the role of sucrose transporters. *Planta, 247*, 587–611.

Durand, M., Porcheron, B., Hennion, N., Mauroosset, L., Lemoine, R. & Pourtau, N. (2016) Water deficit enhances C export to the roots in *Arabidopsis thaliana* plants with contribution of sucrose transporters in both shoot and roots. *Plant Physiology, 170*, 1460–1479.

Duthion, C. & Pigeaire, A. (1991) Seed lengths corresponding to the final stage in seed abortion of three grain legumes. *Crop Science, 31*, 1579–1583.

Fahad, S., Bajwa, A.A., Nazir, U., Anjum, S.A., Farooq, A., Zohaib, A. et al. (2017) Crop production under drought and heat stress: plant responses and management options. *Frontiers in Plant Science, 8*, 1147.

FAOSTAT 2021: http://www.fao.org/faostat/en/#data

Fichter, N. & Lunn, J.E. (2021) The role of Trehalose 6-phosphate (Tre6P) in plant metabolism and development. *Annual Review of Plant Biology, 72*, 737–760.

Figuerola, C.M. & Lunn, J.E. (2016) A tale of two sugars: Trehalose 6-phosphate and sucrose. *Plant Physiology, 172*, 7–27.

Fougerous, J.-A., Doré, T., Ladonne, F. & Fleury, A. (1997) Water stress during reproductive stages affects seed quality and yield of pea (*Pisum sativum*) L. *Crop Science, 37*, 1247–1252.

Foyer, C.H., Lam, H.-M., Nguyen, H.T., Siddique, K.H.M., Varshney, R.K., Colmer, T.D. et al. (2016) Neglecting legumes has compromised human health and sustainable food production. *Nat Plants, 2*, 1612.

Garry, G., Tivoli, B., Jeuffroy, M.H. & Citharel, J. (1996) Effects of drought stress on contrasting maize genotypes under water logging stress. *Crop Science, 37*, 1648–1655.

Gebyeuyu, S., Wiese, H. & Schubert, S. (2011) Effects of drought stress on seed sink strength and leaf protein patterns of common bean genotypes. *African Crop Science Journal, 18*, 75–88.

Geneva, M., Zehirov, G., Djonova, E., Kaloyanova, N., Georgiev, G. & Stancheva, I. (2011) The effect of inoculation of pea plants with mycorrhizal fungi and rhizobium on nitrogen and phosphorus assimilation. *Plant, Soil and Environment, 52*, 435–440.

Goyal, K., Kaur, K. & Kaur, G. (2020) Foliar treatment of potassium nitrate improves the fermentative and sucrose metabolizing pathways in contrasting maize genotypes under water logging stress. *Physiology and Molecular Biology of Plants, 26*, 899–906.

Gu, J., Zeng, Z., Wang, Y. & Lyu, Y. (2020) Transcriptome analysis of carbohydrate metabolism genes and molecular regulation of sucrose transport gene LoSUT on the flowering process of developing oriental
hybrid lily ‘Sorbonne’ bulb. International Journal of Molecular Sciences, 21, 3092.

Guilloni, L., Wéry, J. & Lecoeur, J. (2003) High temperature and water deficit may reduce seed number in field pea purely by decreasing plant growth rate. Functional Plant Biology, 30, 1151–1164.

Guilloni, L., Wéry, J. & Tardieu, F. (1997) Heat stress-induced abortion of buds and flowers in pea: is sensitivity linked to organ age or to relations between reproductive organs? Annals of Botany, 80, 159–168.

Guo, W., Chen, L., Herrera-Estrella, L., Cao, D. & Tran, L.-S.P. (2020) Altering plant architecture to improve performance and resistance. Trends in Plant Science, 25, 1154–1170.

Gupta, M., Dubey, S., Jain, D. & Chandran, D. (2021a) The Medicago truncatula sugar transport protein 13 and its Lr67res-like variant confer powdery mildew resistance in legumes via defense modulation. Plant & Cell Physiology, 62, 650–667.

Gupta, P.K., Balyan, H.S. & Gautam, T. (2021b) SWEET genes and TAL effectors for disease resistance in plants: present status and future prospects. Molecular Plant Pathology, 22(8), 1014–1026.

Hageman, A. & Van Voorden, E. (2021) Sink strength maintenance underlies drought tolerance in common bean. Plants (Basel), 10(3), 489.

Harvey, D.M. (1974) The translocation of 14C-Photoassimilate from leaves to shoots of Petunia axillaris. SUT/SUC and SWEET sugar transporters during flower development. Physiologia Plantarum, 165, 44–57.

Hennion, N., Durand, M., Vriet, C., Doidy, J., Maurousett, L., Lemoine, R. et al. (2019) Sugars en route to the roots. Transport, metabolism and storage within plant roots and towards microorganisms of the rhizosphere. Physiologia Plantarum, 165, 1776–1789.

Ho, L.C. (1988) Metabolism and compartmentation of imported sugars in sink organs in relation to sink strength. Annual Review of Plant Physiology and Plant Molecular Biology, 39, 355–378.

Husčková, A., Humplik, J., Hybl, M., Spichal, L. & Lazar, D. (2019) Analysis of cold-developed vs. cold-acclimated leaves reveals various strategies of cold acclimation of field pea cultivars. Remote Sensing, 11, 2964.

Iftikhar, J., Lu, M., Liu, Z., Mehmood, N., Munir, M., Ahmed, M.A.A. et al. (2020) Sugar and hormone dynamics and the expression profiles of SUT/SUC and SWEET sugar transporters during flower development in Petunia axillaris. Plants, 9, 1770.

Jameson, P.E., Dhandapani, P., Novak, O. & Song, J. (2016) Cytokinin and expression of SWEET, SUT, CWINV and AAP genes increase as pea seeds germinate. International Journal of Molecular Sciences, 17, 1-13.

Jeffroy, M.-H. & Devienne, F. (1995) A simulation model for assimilate partitioning between pods in Pisum sativum L. during the period of seed set: validation in field conditions. Field Crops Res, 2, 79–89.

Jeffroy, M.-H. & Warembourg, F.R. (1991) Carbon transfer and partitioning between vegetative and reproductive organs in Pisum sativum L. Plant Physiology, 97, 440–448.

Kafle A (2018) Tripartite interactions of legumes with Arbuscular Mycorrhizal fungi and Rhizobial bacteria: insight into plant growth, seed yield, and resource exchange. Electron. Theses Dissertation.

Kaschuk, G., Kuyper, T.W., Leffelaar, P.A., Hungria, M. & Giller, K.E. (2009) Are the rates of photosynthesis stimulated by the carbon sink strength of rhizobial and arbuscular mycorrhizal symbioses? Soil Biology and Biochemistry, 41, 1233–1244.

Khan, T.N., Timmerman-Vaughan, G.M., Rubiales, D., Warkentin, T.D., Siddique, K.H.M., Enskine, W. et al. (2013) Didymelono pinodes and its management in field pea: challenges and opportunities. Field Crops Research, 148, 61–77.

Khanna, S.M., Taxak, P.C., Jain, P.K., Saini, R. & Srivivasan, R. (2014) Glycolytic enzyme activities and gene expression in Cicer arietinum exposed to water-deficit stress. Applied Biochemistry and Biotechnology, 173, 2241–2253.

Kreplak, J., Madoui, M.-A., Cápal, P., Novák, P., Labadie, K., Aubert, G. et al. (2019) A reference genome for pea provides insight into legume genome evolution. Nature Genetics, 51, 1411–1422.

Kryvoruchko, I.S., Sinharoy, S., Torres-Jerez, I., Sosso, D., Plišar, I., Guan, D. et al. (2016) MtSWEET11, a nodule-specific sucrose transporter of Medicago truncatula. Plant Physiology, 171, 554–565.

Kumar, J., Sen Gupta, D., Djalioc, I., Kumar, S. & Siddique, K.H.M. (2021) Root-omics for drought tolerance in cool-season grain legumes. Physiologia Plantarum, 172, 629–644.

Kuo, T.M., VanMiddlesworth, J.F. & Wolf, W.J. (1988) Content of raffinose oligosaccharides and sucrose in various plant seeds. Journal of Agricultural and Food Chemistry, 36, 32–36.

Lahuta, L.B., Pluskota, W.E., Stelmaszewska, J. & Szablinska, J. (2014) Dehydration induces expression of GALACTINOL SYNTHASE and RAFFINOSE SYNTHASE in seedlings of pea (Pisum sativum L.). Journal of Plant Physiology, 171, 1306–1314.

Lahuta, L.B., Szablinska-Piernik, J. & Horbowicz, M. (2022) Changes in metabolic profiles of pea (Pisum sativum L.) as a result of repeated short-term soil drought and subsequent re-watering. International Journal of Molecular Sciences, 23, 1704.

Lalonde, S., Tegeder, M., Throne-Holst, M., Frommer, W.B. & Patrick, J.W. (2003) Phloem loading and unloading of sugars and amino acids: phloem loading and unloading. Plant, Cell & Environment, 26, 37–56.

Lalonde, S., Wipt, D. & Frommer, W.B. (2004) Transport mechanisms for organic forms of carbon and nitrogen between source and sink. Annual Review of Plant Biology, 55, 341–372.

Lejeune-Hénaut, I., Bourion, V., Etévé, G., Cunot, E., Delhaye, K. & Desmyter, C. (1999) Floral initiation in field-grown forage peas is delayed to a greater extent by short photoperiods, than in other types of European varieties. Euphytica, 109, 201–211.

Lemoine, R., La Camera, S., Atanassova, R., Dégaldecamp, F., Allario, T., Pourtau, N. et al. (2013) Source-to-sink transport of sugar and regulation by environmental factors. Frontiers in Plant Science, 4, 272.

Lepinay, C., Rigaud, T., Salom, C., Lemanceau, P. & Mougel, C. (2012) Interaction between Medicago truncatula and Pseudomonas fluorescens: evaluation of costs and benefits across an elevated atmospheric CO2. PLoS One, 7, e45740.

Li, C., Liu, Y., Tian, J., Zhu, Y. & Fan, J. (2020) Changes in sucrose metabolism in maize varieties with different cadmium sensitivities under cadmium stress. PLoS One, 15, e0243835.

Li, X., Lawas, L.M.F., Malo, R., Glaubitz, U., Erban, A., Mauleon, R. et al. (2015) Metabolic and transcriptomic signatures of rice floral organs reveal sugar starvation as a factor in reproductive failure under heat and drought stress: Rice floral organs under drought and heat stress. Plant, Cell & Environment, 38, 2171–2192.

Lin, I.W., Sosso, D., Chen, L.-Q., Guse, K., Kim, S.-G., Kessler, D. et al. (2014) Nectar secretion requires sucrose phosphate synthases and the sugar transporter SWEETY. Nature, 508, 546–549.

Lin, Q., Yang, J., Wang, Q., Zhu, H., Chen, Z., Dao, Y. et al. (2019) Over-expression of the trehalose-6-phosphate phosphatase family gene AtTPPF improves the drought tolerance of Arabidopsis thaliana. BMC Plant Biology, 19, 381.

Liu, W., Peng, B., Song, A., Jiang, J. & Chen, F. (2020) Sugar transporter, CmSWEET17, promotes bud outgrowth in Chrysanthemum morifolium. Genes, 11, 26.

Lu, M.-Z., Snyder, R., Grant, J. & Tegeder, M. (2020) Manipulation of sucrose phloem and embryo loading affects pea leaf metabolism, carbon and nitrogen partitioning to sinks as well as seed storage pools. The Plant Journal: for Cell and Molecular Biology, 101, 217–236.
Macherel, D., Benamar, A., Avelange-Macherel, M.-H. & Tolleter, D. (2007) Function and stress tolerance of seed mitochondria. *Physiologia Plantarum*, 129, 233–241.

Mahieu, S., Germon, F., Aveline, A., Hauggaard-Nielsen, H., Ambus, P. & Jensen, E.S. (2009) The influence of water stress on biomass and N accumulation, N partitioning between above and below ground parts and on N rhizodeposition during reproductive growth of pea (*Pisum sativum* L.). *Soil Biology and Biochemistry*, 41, 380–387.

Martignago, D., Rico-Medina, A., Blasco-Escámez, D., Fontanet-Manzaneque, J.B. & Caño-Delgado, A.J. (2020) Drought resistance by engineering plant tissue-specific responses. *Frontiers in Plant Science*, 10, 1676.

Mason, M.G., Ross, J.J., Babst, B.A., Wienclaw, B.N. & Beveridge, C.A. (2014) Sugar demand, not auxin, is the initial regulator of apical dominance. *Proceedings of the National Academy of Sciences*, 111, 6092–6097.

McAdam, E.L., Meitzel, T., Quittenden, L.J., Davidson, S.E., Dalmais, M., Bendahmane, A., et al. (2017) Evidence that auxin is required for normal seed size and starch synthesis in pea. *The New Physiologist*, 216, 193–204.

Meitzel, T., Radchuk, R., McAdam, E.L., Thormählen, I., Feil, R., Munz, E., et al. (2021) Trehalose 6-phosphate promotes seed filling by activating auxin biosynthesis. *The New Phytologist*, 229, 1553–1565.

Melkus, G., Rolletschek, H., Radchuk, R., Fuchs, J., Rutten, T., Wobus, U., et al. (2009) The metabolic role of the legume endosperm: A noninvasive imaging study. *Plant Physiology*, 151, 1139–1154.

Mitchell, R.L. & Russell, W.J. (1971) Root development and rooting patterns of soybean (*Glycine max* [L.] Merrill) evaluated under field conditions. *Agronomy Journal*, 63, 313–316.

Mittal, S. & Sawhney, S.K. (1990) Influence of lead on enzymes of nitrogen metabolism in germinating pea seeds. *Plant Physiology and Biochemistry*, 17, 75–81.

Morin, A., Kadi, F., Porcheron, B., Vriet, C., Maurousset, B., Lemoine, R., et al. (2022) Genome-wide identification of invertases in Fabaceae, focusing on transcriptional regulation of *Pisum sativum* invertases in seed subjected to drought. *Physiologia Plantarum*, 174, e13673.

Munier-Jolain, N. & Salter, P.J. (2001) Can sucrose content in the phloem sap reaching field pea seeds (*Pisum sativum* L.) affect the growth of a plant? *Journal of Experimental Botany*, 52, 551–564.

Pampana, S., Masoni, A. & Ardini, I. (2016a) Response of cool-season grain legumes to waterlogging at flowering. *Canadian Journal of Plant Science*, 96, 597–603.

Pampana, S., Masoni, A. & Ardini, I. (2016b) Grain legumes differ in nitrogen accumulation and remobilisation during seed filling. *Acta Agric Scand Sect B – Soil. Plant Science*, 66, 127–132.

Patrick, J.W. (1997) PHLOEM UNLOADING: sieve element unloading and post-sieve element transport. *Annual Review of Plant Physiology and Plant Molecular Biology*, 48, 191–222.

Patrick, J.W. & Collyvas, K. (2014) Crop yield components - photo-assimilate supply- or utilisation limited-organ development? *Functional Plant Biology*, 41, 893–913.

Patrick, J.W. & Offler, C.E. (2001) Compartmentation of transport and transfer events in developing seeds. *Journal of Experimental Botany*, 52, 155–165.

Pampana, S., Masoni, A. & Ardini, I. (2016b) Grain legumes differ in nitrogen accumulation and remobilisation during seed filling. *Acta Agric Scand Sect B – Soil. Plant Science*, 66, 127–132.

Pampana, S., Masoni, A. & Ardini, I. (2016b) Grain legumes differ in nitrogen accumulation and remobilisation during seed filling. *Acta Agric Scand Sect B – Soil. Plant Science*, 66, 127–132.

Pampana, S., Masoni, A. & Ardini, I. (2016b) Grain legumes differ in nitrogen accumulation and remobilisation during seed filling. *Acta Agric Scand Sect B – Soil. Plant Science*, 66, 127–132.

Pampana, S., Masoni, A. & Ardini, I. (2016b) Grain legumes differ in nitrogen accumulation and remobilisation during seed filling. *Acta Agric Scand Sect B – Soil. Plant Science*, 66, 127–132.

Pampana, S., Masoni, A. & Ardini, I. (2016b) Grain legumes differ in nitrogen accumulation and remobilisation during seed filling. *Acta Agric Scand Sect B – Soil. Plant Science*, 66, 127–132.

Pampana, S., Masoni, A. & Ardini, I. (2016b) Grain legumes differ in nitrogen accumulation and remobilisation during seed filling. *Acta Agric Scand Sect B – Soil. Plant Science*, 66, 127–132.

Pampana, S., Masoni, A. & Ardini, I. (2016b) Grain legumes differ in nitrogen accumulation and remobilisation during seed filling. *Acta Agric Scand Sect B – Soil. Plant Science*, 66, 127–132.

Pampana, S., Masoni, A. & Ardini, I. (2016b) Grain legumes differ in nitrogen accumulation and remobilisation during seed filling. *Acta Agric Scand Sect B – Soil. Plant Science*, 66, 127–132.

Pampana, S., Masoni, A. & Ardini, I. (2016b) Grain legumes differ in nitrogen accumulation and remobilisation during seed filling. *Acta Agric Scand Sect B – Soil. Plant Science*, 66, 127–132.

Pampana, S., Masoni, A. & Ardini, I. (2016b) Grain legumes differ in nitrogen accumulation and remobilisation during seed filling. *Acta Agric Scand Sect B – Soil. Plant Science*, 66, 127–132.
(Pisum sativum L) subjected to water stress. *Field Crops Research*, 86, 81–90.

Sánchez, F.J., Manzanares, M., de Andres, E.F., Tenorio, J.L. & Ayerbe, L. (1998) Turgor maintenance, osmotic adjustment and soluble sugar and proline accumulation in 49 pea cultivars in response to water stress. *Field Crops Research*, 59, 225–235.

Sandalo, L.M., Dalurzo, H.C., Gómez, M., Romero-Puertas, M.C. & del Rio, L.A. (2001) Cadmium-induced changes in the growth and oxidative metabolism of pea plants. *Journal of Experimental Botany*, 52, 2115–2126.

Sandberg, A.-S. (2011) Developing functional ingredients: a case study of pea protein. In: *Functional foods: concept to product*, Amsterdam: Elsevier, pp. 358–382.

Schneider, S., Hulpke, S., Schulz, A., Yaron, I., Höll, J., Imlau, A. et al. (2012) Vacuoles release sucrose via tonoplast-localised SUC4-type transporters: SUC4-type sucrose transporters in the tonoplast. *Plant Biology*, 14, 325–336.

Schroeder, J.J., Delhalze, E., Frommer, W.B., Gueinnot, M.L., Harrison, M.J., Herrera-Estrella, L. et al. (2013) Using membrane transporters to improve crops for sustainable food production. *Nature*, 497, 60–66.

Schulz, A. (1994) Phloem transport and differential unloading in pea seedlings after source and sink manipulations. *Planta*, 192, 239–248.

Shinde, B.P. & Thakur, J. (2016) The effect of co-inoculation of pea plants with arbuscular mycorrhizal fungi and rhizobium on the nodulation, growth and productivity. *International Journal of Biosciences*, 5, 4954.

Sinha, R., Fritschi, F.B., Zandalinas, S.I. & Mittler, R. (2021) The impact of stress combination on reproductive processes in crops. *Plant Science*, 311, 111007.

Sivitz, A.B., Reinders, A., Johnson, M.E., Krentz, A.D., Grof, C.P.L., Perroux, J.M. et al. (2007) Arabidopsis sucrose transporter AtSUC5. High-affinity transport activity, intragenic control of expression, and early flowering mutant phenotype. *Plant Physiology*, 143, 188–198.

Slawinski, L., Israel, A., Artault, C., Thibault, F., Atanassova, R., Laloi, M. et al. (2021) Responsiveness of early response to dehydration six-like transporter genes to water deficit in Arabidopsis thaliana leaves. *Frontiers in Plant Science*, 12, 708876.

Smith, M.R., Rao, I.M. & Merchant, A. (2018) Source-sink relationships in crop plants and their influence on yield development and nutritional quality. *Frontiers in Plant Science*, 9, 1889.

Smitha Ninan, A., Shah, A., Song, J. & Jameson, P.E. (2017) Differential gene expression in the meristem and during early fruit growth of Pisum sativum L. identifies potential targets for breeding. *International Journal of Molecular Sciences*, 18, 428.

Sosso, D., Luo, D., Li, Q.-B., Sasse, J., Yang, J., Gendrot, G. et al. (2015) Seed filling in domesticated maize and rice depends on SWEET-mediated hexose transport. *Nature Genetics*, 47, 1489–1493.

Storr, T. & Hall, J.L. (1992) The effect of infection by Erysiphe pisi DC on acid and alkaline Invertase activities and aspects of starch biochemistry in leaves of Pisum sativum L. *The New Phytologist*, 121, 535–543.

Strydhorst, S., Olson, M.A., Vasanthan, T., McPhee, K.E., McKenzie, R.H., Henriquez, B. et al. (2015) Adaptability and quality of winter pea and lentil in Alberta. *Agronomy Journal*, 107, 2431–2448.

Sugiyma, A., Said, Y., Yoshimizu, M., Takanashi, K., Sosso, D., Frommer, W.B. et al. (2017) Molecular characterization of LjSWEET3, a sugar transporter in nodules of Lotus japonicus. *Plant & Cell Physiology*, 58, 298–306.

Tegeder, M., Wang, X.D., Frommer, W.B., Offler, C.E. & Patrick, J.W. (1999) Sucrose transport into developing seeds of Pisum sativum L. *The Plant Journal*: for Cell and Molecular Biology, 18, 151–161.

Thorup-Kristensen, K. (1998) Root growth of green pea (Pisum sativum L. Genotypes). *Crop Science*, 38, 1445–1451.

Tutetschek, R., Desalegn, G., Epplle, T., Kaul, H.-P. & Wienkoop, S. (2017) Key metabolic traits of Pisum sativum maintain cell vitality during *Didymella pinodes* infection: cultivar resistance and the microsymbionts’ influence. *Journal of Proteomics*, 169, 189–201.

Vacheron, J., Desbrosses, G., Bouffaud, M.-L., Touraine, B., Moënne-Loccoz, Y., Muller, D. et al. (2013) Plant growth-promoting rhizobacteria and root system functioning. *Frontiers in Plant Science*, 4, 356.

Van Bel, A.J.E. & Gamafeli, Y.V. (1992) Ecophysiology of phloem loading in source leaves. *Plant, Cell & Environment*, 15, 265–270.

Van Dongen, J.T., Ammerlaan, A.M.H., Wouterlood, M., Van Aelst, A.C. & Borstlap, A.C. (2003) Structure of the developing pea seed coat and the post-phloem transport pathway of nutrients. *Annals of Botany*, 91, 729–737.

Van Emden, H.F., Ball, S.L. & Rao, M.R. (1988) Pest, disease and weed problems in pea, lentil, faba bean and chickpea. In: Summerfield, R.J. (Ed.) *World crops cool seas food legum*. Glob. Perspect. Probl. Prospects Crop Improv. Pea Lentil Faba Bean Chickpea. Dordrecht: Springer Netherlands, pp. 519–534.

Vidal-Valverde, C., Frias, J., Sierra, I., Blazquez, I., Lambein, F. & Kuo, Y.-H. (2002) New functional legume foods by germination: effect on the nutritive value of beans, lentils and peas. *European Food Research and Technology*, 215, 472–477.

Vocanson, A., Jeuffroy, M.-H. & Roger-Estrade, J. (2006) Effect of sowing date and cultivar on root system development in pea (Pisum sativum L.). *Plant and Soil*, 283, 339–352.

Voegele, R.T., Struck, C., Hahn, M. & Mendgen, K. (2001) The role of haustoria in sugar supply during infection of broad bean by the rust fungus *Uromyces fabae*. Proceedings of the National Academy of Sciences of the United States of America, 98, 8133–8138.

Voisin, A.S. (2003) Root and nodule growth in Pisum sativum L. in relation to photosynthesis: analysis using 13C-labelling. *Annals of Botany*, 92, 557–563.

Wang, E., Wang, J., Zhu, X., Hao, W., Wang, L., Li, Q. et al. (2008) Control of rice grain-filling and yield by a gene with a potential signature of domestication. *Nature Genetics*, 40, 1370–1374.

Wang, S., Yokosho, K., Guo, R., Whelan, J., Ruan, Y.-L., Ma, J.F. et al. (2019) The soybean sugar transporter GmSWEET15 mediates sucrose export from endosperm to early embryo. *Plant Physiology*, 180, 2133–2141.

Wardlaw, I.F. (1990) Tansley review no. 27 the control of carbon partitioning in plants. *The New Phytologist*, 116, 341–381.

Weber, H., Borisjuk, L., Heim, U., Buchner, P. & Wobus, U. (1995) Seed coat: associated Invertases of fava bean control both unloading and storage functions: cloning of cDNAs and cell-type specific expression. *Plant Cell*, 7, 1835–1846.

Weber, H., Borisjuk, L., Heim, U., Sauer, N. & Wobus, U. (1997) A role for sugar transporters during seed development: molecular characterization of a hexose and a sucrose carrier in fava bean seeds. *Plant Cell*, 9, 895–908.

Weber, H., Borisjuk, L. & Wobus, U. (1996) Controlling seed development and seed size in *Vicia faba*: a role for seed coat-associated invertases and carbohydrate state. *The Plant Journal*, 10, 823–834.

Weber, H., Borisjuk, L. & Wobus, U. (1997b) Sugar import and metabolism during seed development. Trends in *Plant Science*, 2, 169–174.

Weber, H., Borisjuk, L. & Wobus, U. (2005) Molecular physiology of legume seed development. *Annual Review of Plant Biology*, 56, 253–279.

Weller, J.L. & Ortega, R. (2015) Genetic control of flowering time in legumes. *Front. Plant Science*, 6, 1-13.

Westgate, M.E., Passioura, J.B. & Munns, R. (1996) Water status and ABA content of floral organs in drought-stressed wheat. *Functional Plant Biology*, 23, 763–772.

Xavier, L.J.C. & Germida, J.J. (2003) Selective interactions between arbuscular mycorrhizal fungi and rhizobium leguminosarum bv. *Viceae*
enhance pea yield and nutrition. *Biology and Fertility of Soils*, 37, 261–267.

Yamada, K., Saijo, Y., Nakagami, H. & Takano, Y. (2016) Regulation of sugar transporter activity for antibacterial defense in Arabidopsis. *Science*, 354, 1427–1430.

Yang, P., Li, Z., Wu, C., Luo, Y., Li, J., Wang, P. et al. (2019) Identification of differentially expressed genes involved in the molecular mechanism of pericarp elongation and differences in sucrose and starch accumulation between vegetable and grain pea (*Pisum sativum* L.). *International Journal of Molecular Sciences*, 20, 1–17.

Yomo, H. & Varner, J. (1973) Control of the formation of amylases and proteases in the cotyledons of germinating peas. *Plant Physiology*, 51, 708–713.

Yoon, J., Cho, L.-H., Tun, W., Jeon, J.-S. & An, G. (2021) Sucrose signaling in higher plants. *Plant Science*, 302, 110703.

Yu, B., Xiang, D., Mahfuz, H., Patterson, N. & Bing, D. (2021) Understanding starch metabolism in pea seeds towards tailoring functionality for value-added utilization. *International Journal of Molecular Sciences*, 22, 8972.

Yu, F., Wan, W., Lv, M.-J., Zhang, J.-L. & Meng, L.-S. (2020) Molecular mechanism underlying the effect of the intraspecific alternation of seed size on plant drought tolerance. *Journal of Agricultural and Food Chemistry*, 68, 703–711.

Yu, J., Jiang, M. & Guo, C. (2019) Crop pollen development under drought: from the phenotype to the mechanism. *International Journal of Molecular Sciences*, 20, 1550.

Zander, P., Amjath-Babu, T.S., Preissel, S., Reckling, M., Bues, A., Schläfke, N. et al. (2016) Grain legume decline and potential recovery in European agriculture: a review. *Agronomy for Sustainable Development*, 36, 26.

Zhang, W.-H., Zhou, Y., Dibley, K.E., Tyerman, S.D., Furbank, R.T. & Patrick, J.W. (2007) Nutrient loading of developing seeds. *Functional Plant Biology*, 34, 314–331.

Zhou, Y., Chan, K., Wang, T.L., Hedley, C.L., Offler, C.E. & Patrick, J.W. (2009) Intracellular sucrose communicates metabolic demand to sucrose transporters in developing pea cotyledons. *Journal of Experimental Botany*, 60, 71–85.

Zhou, Y., Qu, H., Dibley, K.E., Offler, C.E. & Patrick, J.W. (2007) A suite of sucrose transporters expressed in coats of developing legume seeds includes novel pH-independent facilitators. *Plant J Cell Mol Biol*, 49, 750–764.

---

**How to cite this article:** Morin, A., Maurousset, L., Vriet, C., Lemoine, R., Doidy, J. & Pourtau, N. (2022) Carbon fluxes and environmental interactions during legume development, with a specific focus on *Pisum sativum*. *Physiologia Plantarum*, 174(3), e13729. Available from: https://doi.org/10.1111/ppl.13729