Review

Highlighting reactive oxygen species as multitaskers in root development

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Summary

Reactive oxygen species (ROS) are naturally produced by several redox reactions during plant regular metabolism such as photosynthesis and respiration. Due to their chemical properties and high reactivity, ROS were initially described as detrimental for cells during oxidative stress. However, they have been further recognized as key players in numerous developmental and physiological processes throughout the plant life cycle. Recent studies report the important role of ROS as growth regulators during plant root developmental processes such as in meristem maintenance, in root elongation, and in lateral root, root hair, endodermis, and vascular tissue differentiation. All involve multifaceted interplays between steady-state levels of ROS with transcriptional regulators, phytohormones, and nutrients. In this review, we attempt to summarize recent findings about how ROS are involved in multiple stages of plant root development during cell proliferation, elongation, and differentiation.

Introduction

Reactive oxygen species

The emergence of complex multicellular life was interconnected with the buildup of oxygen on Earth. Oxygen accumulation resulted from the emergence of oxygenic photosynthesis and consequently enabled the evolution of aerobic life. This evolving era is characterized by high-energy yielding metabolic pathways (Ward et al., 2019) such as aerobic respiration and oxidative phosphorylation. These pathways, where oxygen serves as the final acceptor of electrons, are more thermodynamically efficient in the evolving complexity than other nonaerobic pathways with nonoxygenic oxidants. This denotes oxygen as the most energetic oxidant in biological life (Reinhard et al., 2016). The reduction of oxygen by the ATP-producing electron transport chains, in addition to other metabolic pathways in which molecular oxygen is the ultimate oxidizing agent, can generate various levels of reactive oxygen species (ROS) (see Box 1). Therefore, ROS are constitutive products of aerobic metabolism (Singh et al., 2016; Rabinovitch et al., 2017), and it is definite that the emergence and permanence of aerobic life was never “ROS-free” (Mittler, 2017).

ROS homeostasis

Earlier on, our understanding to ROS was restricted to the lethal manifestations of the disparity between ROS production and scavenging (Sies, 1986, 1997). This disparity leads to oxidative damage to different molecules and cellular components as nucleic acids, proteins, and membranes and cell death (Das and Roychoudhury, 2014; Anjum et al., 2015). Therefore, as a part of their evolutionary adaptations, plants have developed important ROS-detoxifying machineries (see Box 1). In planta, a large battery of antioxidative enzymes was identified in addition to various antioxidants that form together a double-armed ROS-scavenging mechanism (Das and Roychoudhury, 2014). Together, they forage ROS and counterattack the elevations in their concentrations, preventing oxidative bursts and the consequent detrimental manifestations (Mittler et al., 2004) and maintaining a threshold boundary between redox potential and cytotoxicity, referred to as ROS homeostasis (Saini et al., 2018). In Arabidopsis thaliana, there exists a highly dynamic and complementary network of more than 150 genes that have been identified and correlated to ROS homeostasis (Miller et al., 2009). Few examples related to root development are listed in Table 1. On another side, ROS have been lately recognized for playing fundamental roles as signaling molecules engaged in plant development. They are currently considered as key actors that control cell proliferation, patterning, and differentiation during plant growth (Schippers et al., 2016; Tsukagoshi, 2016; Tognetti et al., 2017). They are also involved in the perception of environmental signals, their transduction, and their
translation into physiological responses. In fact, changes in ROS homeostasis are signals that can trigger growth modifications (Miller et al., 2010; Das and Roychoudhury, 2014; Saini et al., 2018). These modifications derive from the association between ROS homeostasis and subsequent adjustments of gene expression. Hence, being fundamental in plant growth and development, ROS detection in time and space is required to better understand their roles and regulations. This triggered the development of several dyes, probes, and, recently, genetically encoded sensors to detect ROS, mostly as H$_2$O$_2$ (see Box 2).

**Box 1. ROS homeostasis concepts**

**ROS homeostasis box**

- What are ROS? ROS are highly reactive molecules that originate from oxygen (O$_2$) where each has its distinct lifetime (space and chemical properties (Ozcan and Ogun, 2015).

- ROS molecules: ROS are in various forms among which are the radicals having free electrons such as superoxide (O$_2^-$), hydroxyl radical (OH$^\cdot$), peroxyl (RO$_2^-$), and alkoxyl (RO$^\cdot$); others are nonradicals as hypochlorous acid (HOCI), ozone (O$_3$), singlet oxygen ($^1$O$_2$), and hydrogen peroxide (H$_2$O$_2$) (Mhamdi and Van Breusegem, 2018b). Each type of ROS molecule has distinct chemical properties: singlet oxygen can oxidize lipids, proteins, and guanidine residues of DNA; superoxide like singlet oxygen has a half-life time of 1-4 μs and reacts with Fe-S proteins; and hydroxyl radicals are extremely reactive and unstable with a half-life time of 1 ns (Mittler, 2017; Waszczak et al., 2018). In contrast, hydrogen peroxide is the most stable ROS (more than 1 ms) and, therefore, is the predominant ROS involved in cellular signaling.

- ROS homeostasis is controlled by the delicate balance between ROS production and scavenging, maintaining a threshold boundary between redox potential and cytotoxicity (Saini et al., 2018).

- ROS in the organelles. ROS are produced in the mitochondria by mitochondrial respiration, in the chloroplasts during photosynthesis, in the peroxisome-localized photorespiratory reactions, and in the apoplast by NADPH oxidases such as the respiratory burst oxidase homologs (RBOHs), peroxidases, and other oxidases.

- ROS in the apoplast. Two enzymatic families contribute to the apoplastic ROS production. In addition, other families of enzymes are also able to modify the ROS homeostasis in the apoplast such as oxalate oxidases, polyamine oxidases, quinone oxidases, lipoygenases, etc.

  1. The NADPH oxidases also called respiratory burst oxidative homologs (RBOHs) (Orman-Ligeza et al., 2016) consist of transmembrane proteins which produce superoxide O$_2^-$. In Arabidopsis thaliana, ten genes have been asserted to encode RBOHs (RBOHA-RBOHJ) (Marino et al., 2012).

  2. The class III peroxidase family (PRX) consists of apoplastic proteins that oxidize various substrates by using H$_2$O$_2$ as an electron acceptor but can also regulate the local concentration of H$_2$O$_2$ or produce ROS (Francoz et al., 2015). In Arabidopsis thaliana, 73 members of this family have been found (Tognolli et al., 2002), but only few of them have been characterized.

- Apoplastic ROS can be channeled into the cytoplasm by multiple aquaporin of the PIP-type (Tian et al., 2016).

- ROS scavenging is a two-armed mechanism:

  1. The first arm consists of the antioxidative enzymes as the superoxide dismutase (SOD), catalase (CAT), ascorbate peroxidase (APX), monodehydroascorbate reductase (MDHAR), dehydroascorbate reductase (DHAR), glutathione reductase (GR), glutathione peroxidase (GPX), and class III peroxidase (PRX) (Das and Roychoudhury, 2014; Choudhary et al., 2020).

  2. The second arm comprises nonenzymatic antioxidants such as ascorbic acid (ASC), reduced glutathione (GSH), carotenoids, flavonoids, and proline (Eltayeb et al., 2007).

See also Table 1 for more details.

**ROS production and scavenging**

In plants, ROS are constitutively produced during different biochemical reactions located in various cellular compartments, i.e., chloroplast, mitochondria, and the peroxisome in addition to the apoplast. In the chloroplasts, ROS production is associated with the light-dependent reactions and is due to the leakage of electrons (Ivanov et al., 2018). Nevertheless, in the mitochondria, ROS production is relatively lower and is related to the respiratory chain (Marchi et al., 2012; Kröller-Schön et al., 2014). In addition, peroxisomes are also sites for hydrogen peroxide production during different processes as the glycolate oxidase...
Table 1. Selected examples of components that regulate ROS homeostasis: ROS scavengers, ROS producers, and an ROS sensor involved in root development processes

| Gene name/protein | Subcellular localization | Functional evidence, mutant analysis, phenotypes | References |
|-------------------|--------------------------|--------------------------------------------------|------------|
| **ROS scavengers** |                          |                                                  |            |
| Superoxide dismutase (SOD) | Chloroplast, mitochondria, peroxisome, and cytosol | Defense against oxidative damage. It eradicates O₂⁻ by producing O₂ and H₂O₂ | (Saini et al., 2018) |
| Catalase (CAT) CAT1, CAT2, CAT3 | Peroxisome and mitochondria in pollen and seeds; photosynthetic tissues and roots | Removes H₂O₂ and transforms it into H₂O and O₂. | (Das and Roychoudhury, 2014) |
| Ascorbate peroxidase (APX) | Chloroplast and cytosol | Redundant to CAT. Transforms H₂O₂ into H₂O and dehydroascorbate (DHA) | (Sharma and Dubey, 2004) |
| Monodehydroascorbate reductase (MDHAR) | Colocalized with APX in the peroxisome | Replenishes the ascorbic acid (AA) reservoir | (Das and Roychoudhury, 2014) |
| Dehydroascorbate reductase (DHAR) | Intracellular and in the apoplast | Reduces DHA to AA using glutathione (GSH) as a reductant | (Eltayeb et al., 2007) |
| Glutathione reductase (GR) | Chloroplast, mitochondria and cytosol | Maintain a high cellular GSH:GSSG ratio, reducing glutathione disulfide (GSSG) into glutathione (GSH). | (Das and Roychoudhury, 2014) |
| Guaiacol peroxidase (GP) | Active both intracellular and extracellular | Remove of H₂O₂ during either regular metabolism or oxidative burst. | (Saini et al., 2018) |
| **RBOHs** |                          |                                                  |            |
| RBOH (AT1G64060) | Roots: endodermis, lateral roots and vascular tissues; stems, seedlings, inflorescences, leaves and guard cells. RBOHF is phosphorylated by CIPK26, binding Ca²⁺. Besides interacts with SRC2. It is upregulated by aba. RBOHD can interacts with BIK1 and PBL13 Atbrohf and Atbrohd control ROI spatial production | Are required for CS formation, producing H₂O₂. The number and density of LR increase in the loss-of-function mutants of rbohd and rbohf, also show lower H₂O₂ and O₂⁻ concentration in the root cap, MZ and EZ as compared to the WT. But has and increase of O₂⁻ in the DZ increasing LR density in rbohd and rbohf. RBOHD and RBOHF negatively regulate LRP formation and LR emergence by downregulating the peroxidase activity. | (Lee et al., 2013; Li et al., 2014; Franke, 2015; Pedreira et al., 2011) |
| RBOHC (AT5G51060) | In roots, is expressed in the epidermis, TZ, EZ, DZ and in elongating root hairs. RBOHC actively produces OH⁻ in the apoplast at the tip of the root hair. | It is required for the production of reactive oxygen species that regulate cell expansion through the activation of Ca²⁺ channels. His expression is increased in potassium starvation. Participates in root hair elongation. rbohc mutants shows shorter root hairs | (Mhamdi and Van Breusegem, 2018a, 2018b; Foreman et al., 2003) |
| RBOHE (AT1G19230) | Expressed in roots: lateral root; inflorescences, leaves and stems. GUS reporter shows accumulation of RBOHE at the starting site of the lateral root | Loss-of-function mutant rbohe delays the development of the lateral root | (Chapman et al., 2019) |

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### Table 1. Continued

| Gene name/protein | Subcellular localization | Functional evidence, mutant analysis, phenotypes | References |
|-------------------|--------------------------|-----------------------------------------------|------------|
| **Apoplastic type-III peroxidases (PRXs)** |
| PRX72 (AT5G66390) | Roots endodermis and xylem | PRX72 expression is upregulated when the xylem vessel elements are actively forming, playing a role in xylem formation. In prx72 mutants, lignin levels were 40% less compared to the WT. It has been observed that PRX72 expression is increased in the lac4-2 lac11 lac17 triple mutants, implying a possible functional linkage between PRXs and LACs | (Herrero et al., 2013; Zhao et al., 2013) |
| PRX17 (AT2G22420) | Root endodermis and xylem. Is regulated by the MADS-box transcription factor AGL15 and interacts with GRF3 3SS:PRX17 over accumulates xylem compared to WT | Participates in lignin polymerization of TEs or indirectly via ROS signaling. Loss-of-function mutant, prx17, has lower lignin content in most organs | (Cosio et al., 2017; Herrero et al., 2013; Hoffmann et al., 2020) |
| PRX64 (AT5G42180) | Root endodermis Upregulated by cold | Uses the H2O2 for monolignols oxidation, catalyzing lignin formation | (Lee et al., 2013; Franke, 2015) |
| PRX37 (AT4G08770) | Root endodermis and vascular bundles. Interacts with RGS1 and NAC3 | Participates in phenolic “cross-linking” activity during lignin deposition in the cell wall. Overexpression of prx37 shows a dwarf phenotype with smaller plants and delayed development, and a increase in the amount of esterified phenolic products | (Pedreira et al., 2011) |
| PRX01 (AT1G05240) | Root hair. Coexpressed with other genes as extensins and glycoproteins in the cell wall | Participation in cell wall expansion in root hairs and ROS homeostasis. PRX01 is involved in the lignification of cells walls. Double mutants of t px44 px73, px01 px44, and px01 px73, as well as the triple null mutant, px01 px44 px73 showed shorter root hairs. PRX1 and PRX73 are upregulated in the overexpressors of RSL4 and downregulated in rsl4 KO mutants | (Mangano et al., 2017; Marzol et al., 2020) |
| **PRX57 (AT5G17820)** | Lateral roots. Root hair PRX57: Positively light-regulated during the first stage of development, upregulated by cold treatment, and downregulated by salicylic acid | PRX7 participates in root hair elongation, is target of RSL4. PRX7 and PRX57 were detected to be auxin-independent. KO mutants px7 and px57 have lower LR density and number due to changes of ROS balance in the initiation stage | (Manzano et al., 2014; Vijayakumar et al., 2016) |

(Continued on next page)
| Gene name/protein                      | Subcellular localization | Functional evidence, mutant analysis, phenotypes                                                                                                                                                                                                 | References                                      |
|----------------------------------------|--------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------|
| Glutathione peroxidase (GPX) family    | Lateral root             | gpx mutants develop larger lateral root primordia, indicating that the GPX family plays a role in regulating root architecture, especially GPX1 and GPX7, having a critical role in development and formation of the lateral root.                                                                                           | (Choudhary et al., 2020)                        |
| ROS sensor                             |                          |                                                                                          |                                                 |
| HYDROGEN PEROXIDE-INDUCED Ca\(^{2+}\) INCREASES 1 (HPCA1) (AT5G49760) | HPCA1 is localized to the plasma membrane and is activated by H\(_2\)O\(_2\) via covalent modification of extracellular cysteine residues, which leads to autophosphorylation of HPCA1.                                                                                 | (Wu et al., 2020)                               |
| ROS regulators                         |                          |                                                                                          |                                                 |
| Retarded root growth (RRG) (AT1G69380) | Root meristem, primary and lateral root tips. Also present in leaves and pollen. In root tips, is preferentially expressed in QC cells, cortex/endodermis stem cells and endodermal and stele cells of root meristems | Promote cellular division in the RAM. KO mutants, rrg-1 and rrg-2 revealed a short-root phenotype compared to the WT. Required for the maintenance of mitochondrial structure. Positive regulator of cell division and but negative regulator of cell expansion in the postembryonic root meristem. | (Zhou et al., 2011)                             |
| Prohibitin3 (PHB3) (AT5G40770)         | Expressed in proliferative tissues, vasculature, stems, leaves, and roots. Preserves redox homeostasis in the mitochondria                                                                                                                                  | Inhibits the cell cycle. Maintains the stem cell niche in QC and activate the cellular division in the proximal meristem. ROS production is induced and subsequently inhibits PHB3 expression. In phb3 mutants, genes implicated in ROS production showed higher expression (i.e. AOX1 and NA | (Kong et al., 2018)                             |
| Caspian strip membrane domain proteins (CASPs) CASP1 (AT2G36100) | Root endodermis. Interacts with CASP2, CASP3, CASP4, and CASP5. Along with CASP2, it is required for the localization of ESB1                                                                                                                          | CASPs coordinate the proper functions of the different components in the lignin-polymerizing machinery. Their interaction with PRX64 and RBOHF and determine their precise localization at the CsS development site. CASP1 guides this localization by interacting with SGN1 and SGN3 | (Alassimone et al., 2016; Li et al., 2018a) |
| Gene name/protein | Subcellular localization | Functional evidence, mutant analysis, phenotypes | References |
|------------------|--------------------------|-------------------------------------------------|------------|
| MYB domain protein 36 (MYB36) (AT5G57620) | Leaves, roots endodermis, lateral root and seedlings. Directly upregulated by SCR | It promotes differentiation of the endodermis during root development. Regulates the spatial expression of genes involved in positioning and build of CSs in the root endodermis (i.e., CASPs, PRX64, and ESB1). Loss-of-function mutants, myb36, show deficit CS, with an abnormal distribution of lignin in endodermal cells. The mutant alleles myb36-5 and myb36-2 had less LRs verse the WT | (Cui et al., 2014; Kamiya et al., 2015; Liberman et al., 2015; Fernández-Marcos et al., 2017) |
| Like sex four 2 (LSF2) (AT3G10940) | Widely expressed, especially in root hair | lsf2-1 KO mutants showed lower O2-generation rate but higher levels of H2O2 upon oxidative stress treatments. LSF2 interacts with MPK8, which regulates the cytosolic redox balance | (Zhao et al., 2016) |
| Feronia (FER) (AT3G51550) | Widely expressed, especially in root hairs. It is induced by brassinosteroids. | FER acts as an upstream regulator for the Rac/Rop-signaling pathway that controls ROS-mediated root hair development. fer and rhd2 mutants shows short root hairs, and a decreased in ROS accumulation. FER interacts with ROP2, promoting radical hair growth upstream of RBOHC, interacts with RopGEF1, RALF1, ROP2 and BRL3 | (Chapman et al., 2019) |
| Root hair defective six like4 (RSL4) (AT1G27740) | Root hair, leaves, and flowers. Promotes post mitotic cell growth in root hair cells. Induced by jasmonate (JA) and low-phosphate conditions | RSL4 is involved in the regulation of root hair elongation. KO rsl4 plants have no root hair. RSL4 is controlled by ARFs, inducing localized ROS synthesis. Has targets genes that promotes root hair growth, like GBF2, EXO70A1, CPK11 and SAC1. RSL4 is a direct transcripational target of RHD6. RSL4 can bind to the RHE in the regulatory region of PRX1, PRX44, PRX60, and PRX73 | (Mangano et al., 2017; Chapman et al., 2019) |
| Hypoxia responsive ERF genes (HRE1 and HRE2) Ethylene response factor genes (ERF and RAP2) | Lateral roots | KO mutants rap2, hre1, hre2, erf have more LRs compared to the WT, these mutants affect the delayed emergence in LR development | (Shukla et al., 2019) |
reaction, fatty acid \(\beta\)-oxidation by acyl Co-A oxidase, and an enzymatic reaction of flavin oxidases. Furthermore, with reduced CO\(_2\) availability during photosynthesis, RUBISCO binds O\(_2\) and oxygenates ribulose-1,5-bisphosphate into glycolate. The latter is subsequently oxidized to glyoxylate in the peroxisome with concomitant production of hydrogen peroxide. Likewise, superoxide radicals are produced by xanthine oxidase in the peroxisomal matrix and by NADPH:ferri cyanide reductase in the peroxisomal membrane (Del Ríoa and López-Huertas, 2016). Moreover, two enzymatic families contribute to the apoplastic ROS production in a controlled manner: the respiratory burst oxidative homologs (RBOHs) (Orman-Ligeza et al., 2016) and the class III peroxidases (PRXs) (Francoz et al., 2015).

ROS released in the chloroplast during photosynthesis represent the biggest pool of ROS among the different compartments. These ROS are implicated in signaling processes such as the regulation programmed cell death (PCD) (Shapiguzov et al., 2012). In addition, ROS produced in mitochondria during oxidative phosphorylation are associated to roles in redox signaling, retrograde signaling, plant hormone action, and PCD during pathogen attacks (Huang et al., 2016). First, these ROS may possibly adjust protein functions in the mitochondrial matrix by oxidizing specific protein thiols as for alternative oxidases (AOXs) (Yoshida et al., 2013) and for several tricarboxylic acid cycle (TCA) enzymes (Daloso et al., 2015). Second, they are probably involved in the mitochondrial retrograde regulation (MRR) where the mitochondria communicate back their functional status to the nucleus. The observation of MRR responses shows remarkable overlaps with transcriptional responses to different ROS triggers and abiotic stress, and the induction of MRR with complex III inhibitor antimycin A increased ROS production (Van Aken et al., 2016). Moreover, the interaction between hormonal signaling that regulates all layers of plant development and cellular functions and mitochondrial signaling, including mitochondrial ROS signaling, has been recently reviewed (Berkowitz et al., 2016). Lastly, these ROS trigger the execution of PCD during pathogen attacks, where various mutants in mitochondrial functions have been reported to have diverse PCD or pathogen defense responses (Huang et al., 2016). To exemplify, Arabidopsis mosaic death 1, MOD1, codes for an enoyl-acyl carrier protein (ACP) reductase essential for fatty acid biosynthesis in the chloroplast and negatively regulates PCD in Arabidopsis. Recently, a study demonstrated that mitochondrial complex I is responsible for the ROS production that leads to PCD in the mod1 mutant (Wu et al., 2015). Furthermore, a comprehensive chloroplast-mitochondria communication pathway that regulates the generation of mitochondrial ROS and triggers PCD has been elucidated (Zhao et al., 2018). In turn, apoplastic ROS are engaged in plant development and responses to extrinsic signals. These ROS regulate plant cell division and cell elongation rate, and hence the whole plant growth (Podgórska et al., 2017); they are also implicated in regulating stomatal closure, cell wall lignification, and callose deposition in response to external signals (Podgórska et al., 2017; Qi et al., 2017; Janku et al., 2019; Farvardin et al., 2020). Obviously, ROS signaling encompasses several pathways within and between various organelles and cellular compartments. An elaborated understanding of all these pathways and their interconnections is still lacking, and their investigation remains a big challenge for plant biology research.

To sum up, in plants, ROS production is an inevitable part of the basal metabolism and occurs in various cellular compartments. Remarkably, ROS production may be suddenly enhanced in response to

| Gene name/protein | Subcellular localization | Functional evidence, mutant analysis, phenotypes | References |
|-------------------|--------------------------|-----------------------------------------------|------------|
| UPBEAT1 (UPB1) (AT2G47270) | Expressed in the root vascular tissue and in root hairs and lateral root caps. J55:UPB1-YFP, showed reduced root length and a decrease in cortex cell number in the MZ | UPB1 manipulates the ROS homeostasis, regulating cellular division and differentiation. upb1 KO mutants had more and longer emerging LRs compared to WT, another mutant upb1-1, shows significant increase in the number of cortical cells, indicating enlargement of the meristem. UPB1 controls the expression of several peroxidases (PRX39, 40, and 57) | (Tsukagoshi et al., 2010; Manzano et al., 2014) |
environmental cues; this is defined as the "oxidative burst" (Choudhary et al., 2020). The oxidative burst is one of the earliest and most important physiological responses to stresses (Saini et al., 2018); it fires various signaling pathways leading to transcriptional changes in the stress-related genes, resulting in a defensive response (Baxter et al., 2014; Sewelam et al., 2016; Zhang et al., 2016). ROS overproduction may lead to an irreversible oxidative damage as the degradation of biomolecules such as lipids, proteins, nucleic acids, and carbohydrates (Chapman et al., 2017). This may lead to physiological disruption or even cellular death (Anjum et al., 2015; Choudhary et al., 2020). Therefore, the normal cellular homeostasis requires maintaining the delicate balance between ROS production and scavenging (Mittler, 2017). ROS scavenging is a two-armed mechanism (Eltayeb et al., 2007) that shields the various cellular compartments against any provisional oxidative damage. Furthermore, many actors in ROS scavenging are also implicated in plant growth and development by taking part in processes such as mitosis, elongation, senescence, and PCD (Noctor et al., 2018).

ROS sensing

Understanding how plants sense ROS and translate their signals into physiological responses was thoroughly investigated. One mechanism to sense ROS is the oxidative post-translational modification (Ox-PTM) of proteins, which was recently reviewed (Waszczak et al., 2015). Numerous proteins with Cys residues have been identified and reviewed as targets of Ox-PTMs such as enzymes of the Calvin cycle or signal transduction enzymes such as transcription factors, kinases, phosphatases, and RNA-binding proteins (Dietz, 2014; Waszczak et al., 2015; Mittler, 2017; Sies et al., 2017). Apoplastic H$_2$O$_2$ is sufficiently stable to penetrate into the cytoplasm via aquaporin membrane proteins where it covalently modifies proteins. However, a recent breakthrough discovery characterized a cell surface ROS sensor, the HPCA1 (HYDROGEN PEROXIDE-INDUCED Ca$^{2+}$ INCREASES 1) identified as the first known cell-surface-specific sensor for H$_2$O$_2$ in plants (Wu et al., 2020). HPCA1 encodes a leucine-rich-repeat receptor kinase (LRR-RK) that contains two special pairs of Cys residues in its extracellular domain. The thiol groups of Cys residues are targets for oxidation to sulfenic acids by H$_2$O$_2$, and these sites are the place for extracellular

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**Box 2. Tool examples for ROS detection in plant cells**

**Tools for ROS detection**
- Diaminobenzidine (DAB) and nitro blue tetrazolium (NBT) staining. Both staining methods are used to visualize H$_2$O$_2$ and superoxide radicals. DAB and NBT are not specific or direct measurements of both ROS, and color formation does not always reflect the levels of the desired ROS. NBT staining can reflect the presence of ascorbate or the activity of dehydrogenases, while DAB browning color accumulates in the presence of higher peroxidase activity. Dihydroethidium (DHE) is also used to detect O$_2^-$.  
- Dichlorofluorescin (DCF)-derived fluorescent dyes. DCF is widely used for quantification of H$_2$O$_2$ although it quantifies total ROS. This method is not very reliable and is not specific. Dichlorofluorescin (DCFH) does not react with H$_2$O$_2$ or other ROS directly. ROS imaging can be further complicated by the presence of endogenous autofluorescent compounds. Other fluorescent dyes exist such as BES-H$_2$O$_2$-Ac to detect H$_2$O$_2$ (Tsukagoshi et al., 2010).
- Amplex Red (AR) and Amplex Ultra Red (AUR) fluorescent dyes. AR is used for extracellular-apoplastic H$_2$O$_2$ since it cannot go through the plasma membrane. Both, AU/AUR, undergoes one electron oxidation and yields the fluorescent product resorufin, a reaction with 1:1 (AR/AUR: H$_2$O$_2$) stoichiometry, in which concentration of horseradish peroxidase and pH play important roles. These two dyes have been used to track H$_2$O$_2$ levels in several plant tissues (Rhee et al., 2010; Deng et al., 2011; Sang et al., 2012; Chakraborty et al., 2016; Tian et al., 2016).
- Genetically encoded probes. Genetically encoded sensors have been engineered to measure H$_2$O$_2$ which have several advantages over the dyes because they are noninvasive, can be targeted in time and space (at tissues and cellular level), and are stable over time. To name some of the most widely used in plant cells are roGFP1 (Schwarzländer et al., 2008), roGFP-GSH (Bratt et al., 2016), GRX1-roGFP2 (Gutschert et al., 2008; Schwarzländer et al., 2008), roGFP-Orp1 (Dagda et al., 2017), and HyPer (Costa et al., 2010; Hernández-Barrera et al., 2015; Mullineaux et al., 2018).

Several tools have been developed for the detection and visualization of ROS in plant tissues. For more details on limitation and controls to be included for their correct use, please see Noctor and Foyer, 2016; Mhamdi and Van Breusegem, 2018a; Fichman et al., 2019.
H$_2$O$_2$ sensing. More importantly, this covalent modification leads to the activation of HPCA1 intracellular kinase activity by its autophosphorylation, which subsequently triggers through unknown mechanisms Ca$^{2+}$-channel gating and Ca$^{2+}$ influx, followed by activation of downstream signaling pathways (Wu et al., 2020). Remarkably, the extracellular H$_2$O$_2$ sensing mechanism by HPCA1 does not resemble any H$_2$O$_2$ receptors or sensors previously reported for other eukaryotic organisms, highlighting that plants have evolved a unique manner to deal with ROS. Since H$_2$O$_2$ is also produced in several organelles, other H$_2$O$_2$ sensors may act together with HPCA1 to transmit organelle-specific redox messages to the nucleus, along with messages arising from the apoplastic face of the plasma membrane. On another hand, HPCA1 was recently identified to be involved in quinone sensing which triggers defense-related gene expression by elevating cytoplasmic Ca$^{2+}$ and activating MAPks (mitogen-activated protein kinase) signaling pathway named as CARD1 (CANNOT RESPOND TO DMBQ 1) (Laohavisit et al., 2020). The recognition of quinone molecules by CARD1 is also based on Cys residues in its apoplastic ectodomain. Yet, it is still unclear how extracellular quinone and H$_2$O$_2$ sensing are related to each other in the HPCA1/CARD1 proteins.

**ROS recap**

During the past years, several reviews recapped the recent findings about ROS as signaling molecules during natural metabolism or in response to exogenous factors. In 2004, the mechanisms of generation and removal of ROS during development and under biotic and abiotic stresses were reported, in addition to new insights into their complexity and roles from genetic analyses of ROS-detoxifying and ROS-signaling mutants (Apel and Hirt, 2004). Moreover, the important role of ROS waves as signaling molecules in and across different cells, the dynamics and specificity of these signals, and their cross talk with other networks were thoroughly interpreted (Mittler et al., 2011). Furthermore, upon zooming into the ROS gene network, the RBOHs were identified as a key protein at the center of this ROS signaling network. In fact, RBOHs demonstrated important functions in the cellular biological processes as integrating calcium signaling and protein phosphorylation necessary for controlled ROS production (Suzuki et al., 2011). In 2012, the influence of apoplastic and chloroplastic ROS on nuclear gene regulation and transcriptional reprogramming was pinpointed (Shapiguzov et al., 2012). Also, the oxidative burst after pathogen recognition and the consequent metabolic and proteomic fluxes during plant defense were described (O’Brien et al., 2012). Soon after came the findings that stress responses in plants are mediated by a spatiotemporal coordination between ROS and other signals to confer systemic acquired resistance or systemic acquired acclimation (Baxter et al., 2014). Additionally, numerous updates conferred the role of ROS during abiotic stresses, such as drought (Noctor et al., 2014), salinity (Bose et al., 2014), and anoxia (Gonzali et al., 2015). Further advancement in understanding the signaling aspect of ROS has been made in the past five years (Dietz et al., 2016; Sewelam et al., 2016; Mittler, 2017; Qi et al., 2017; Waszczak et al., 2018). Recently, the regulatory actions of ROS during plant development have also been extensively studied, yet many are still not well known. In fact, ROS have been shown to be involved in cell proliferation and differentiation, PCD, seed germination, organ growth, flower and pollen tube development, and leaf senescence (Singh et al., 2016; Mhamdi and Van Breusegem, 2018a).

**ROS & roots**

However, less is known about the involvement of ROS in roots growth and development despite the knowledge of their fundamentality. As roots grow into dark heterogeneous soils, their investigation by researchers is indeed challenging and limited, although these obstacles are being surpassed, thanks to the developing root imaging technologies (Mooney et al., 2012; Mairhofer et al., 2013; Rellán-Alvarez et al., 2015). In A. thaliana, the plant root architecture is established by the cross talk between two key players: the genetic background and the environmental signals (Wachsman et al., 2015). Similar to the aerial part of the plant, the primary development and formatting follow genetically programmed intrinsic patterns (Novoplansky, 2019), but the fate of root morphology is strongly adjusted by extrinsic environmental signals. These adjustments by exogenous cues reflect the phenotypic plasticity of plants, which allow them to cope with the fact that they are sessile organisms existing in a continuously changing environment (Simard, 2018). In A. thaliana, the root consists of three major zones: the meristematic zone (MZ), the elongation zone (EZ), and the differentiation zone (DZ). The root morphology is set up by highly orchestrated and dynamic balances between these zones. In fact, during this establishment of root architecture, ROS are involved as signaling molecules that choreograph root structures and anatomies (Tsukagoshi, 2016) through various developmental processes (Czarnecka and Karpíš, 2018). Several other molecules have been characterized for their role in establishing and maintaining such balances as phytohormones and nutrients (Majda and Robert, 2018; Mangano et al., 2018; Noctor et al., 2018; Choudhary et al., 2018a).
Additionally, roots represent a perfect model for studying plant growth since all the developmental stages that a cell goes through are present. In fact, the stem cell phase is present at the root tip where cell proliferation occurs. Stem cells and their derived cells divide and shift away from a mitotically inactive quiescent center (QC). Afterward, the proliferative phase ceases and cells start to elongate and differentiate until the fully differentiated phase in the DZ (Street et al., 2016). Therefore, this review aims to address the multiple roles of ROS during plant root development, elongation, and differentiation, focusing on the recently uncovered molecular mechanisms.

**Role of ROS balances in the meristematic and elongation root zones during proliferation and elongation**

**ROS & progression of cell cycle**

Root growth and development result from the constant generation of cells in its MZ (De Veylder, Beeckman and Inze, 2007). Afterward, these dividing cells acquire specialized functions during differentiation where they undergo orchestrated modifications. A cell cycle controls the number of dividing cells and the duration of these divisions which determines the growth rate of the root. A proper cell cycle is hence highly important for a proper root growth because it maintains the MZ size and prevents the premature differentiation (González-García et al., 2011). In fact, the molecular mechanisms that coordinate the phase-to-phase transitions in the cell cycle have been thoroughly studied. The roles of cyclins (CYCs) and cyclin-dependent kinases (CDKs) as key actors in these transitions have been highlighted (Hamdoun et al., 2016; Tamirisa et al., 2017). In addition, ROS have been found to influence these actors and govern the interphasic transition in the cell cycle (Mhamdi and Van Breusegem, 2018b). First, the cell cycle stops owing to changes in the redox homeostasis. CYCs expression is controlled by the transcription factor TEOSINTE BRANCHED1-CYCLOIDEA-PROLIFERATING CELL FACTOR1 (TCP), which contains a redox-sensitive cysteine residue that determines its activity. When ROS levels increase, a disulfide bond is formed, preventing the binding of TCP on the CYC promoters (Viola et al., 2016). Moreover, CYCs and CDKs expression patterns have been correlated with the ascorbate-glutathione (ASC:GSH) cycle and their redox status (Figure 1). Several glutathione-deficit mutants revealed a cell cycle arrest as in root meristem less 1 (rml1) (Schnaubelt et al., 2015). The concentrations of ROS, glutathione, and ascorbate in various organelles were found synchronized with transition events in the cell cycle. Accordingly, ascorbate-deficit mutants had slower cell cycles than the wild-type (WT) ones (Tognetti et al., 2017; Ortiz-Espín et al., 2017). In addition, further studies reveal imperative roles of ROS in the cell plate synthesis during cytokinesis, the last stage of the cell cycle. During this process, the key to a proper patterning of the plant cell wall (PCW) is held by a dynamic network of filamentous structures, i.e., the cytoskeleton (Szymanski and Staiger, 2018). ROS have been shown as actors during this morphogenesis. Modification of ROS homeostasis disturbed the cytoskeleton structure and formed many atypical tubulin polymers (Livanos et al., 2012). This aberrant formation negatively affected the cytokinesis and the cell plate synthesis. Similarly, Arabidopsis mutated for RBOHs, ROS-producing enzymes, develop disorganized cytoskeletal tubulin and hence the cytokinesis is disturbed (Foreman et al., 2003; Livanos et al., 2017). Therefore, ROS are essential regulators of the cell cycle, hence contributing to the regulation of MZ size and progressive root growth.

**ROS & maintenance of stem cell niche**

ROS are involved in maintaining the stem cell identity in the root apical meristem (RAM) and regulating the organogenesis (Wany et al., 2018). For this aim, they form extensive networks with other signaling molecules as nitric oxide (NO) in addition to several phytohormones (auxin, cytokinin, ethylene, etc.) and their associated signaling pathways (Wany et al., 2018). As well, ROS fine-tunes the stem cell fate in the RAM (Yang et al., 2018). PROHIBITIN3 (PHB3) maintains the root stem cell niche maintenance. In phb3 mutants, the transcript of the cell cycle inhibitor retinoblastoma RB is higher than the WT. However, both KNOLLE, a member of SYG11 syntaxin gene family, and cyclin-dependent protein kinase (CYCB1) which are markers of the cell cycle are downregulated. These results confirm the role of PHB3 in the RAM size maintenance. Moreover, phb3 mutants show an increased division rate in the quiescent center (QC) (>80%) compared with the WT (~5%) which indicates the role of PHB3 in the QC identity maintenance. In fact, this PHB3 regulation is in cross talk with the redox status since hydrogen peroxide and superoxide overaccumulate in the phb3 plants (Kong et al., 2018). In the RAM, ROS production is regulated according to the nutrient availability in the soil. Under nitrate deficiency, ROS production is induced and subsequently inhibits PHB3 expression. This triggers ROS-responsive factors (ERF109, ERF114, and ERF115) leading to cellular proliferation and differentiation (Figure 1). On the contrary, nitrate sufficiency promotes the production...
of glutathione antioxidant which reduces ROS concentrations. Therefore, PHB3 becomes active and inhibits the underlying ETHYLENE RESPONSE FACTOR 115 (ERF115), thus maintaining the pluripotency of the QC (Figure 1) (Kong et al., 2018). Simultaneously, ETHYLENE RESPONSE FACTOR 109 and 114 (ERF109 and ERF114) are induced to promote cellular elongation and differentiation in the elongation and differentiation zones, respectively (Wany et al., 2018). Moreover, PHB3 plays a role in preserving redox homeostasis in the mitochondria. In Arabidopsis phb3 mutants, genes implicated in ROS production and signaling such as alternative oxidases (AOXs), ALTERNATIVE NAD(P)H DEHYDROGENASE 1 (NDA1), and NAD(P)H DEHYDROGENASE B2 (NDB2) showed higher expression (Kong et al., 2018). These studies suggest a vital role for ROS in conserving the stem cell niche, which enables a continued growth of the root.

**ROS & cell elongation**

The dividing cells enter a new phase of cellular elongation in the EZ. This elongation is mostly dependent on the structure and plasticity of the primary cell walls (PCWs). The dynamics of the PCW are associated with the properties of its components, polysaccharides and proteins (Schmidt et al., 2016). In our scope,
numerous studies reported the role of apoplastic compounds in controlling this cellular expansion in the EZ. Cellular expansion rates are tightly related to the balance between cell wall stiffening and loosening, where both are controlled by ROS homeostasis (Francoz et al., 2015). Primarily, during their peroxidative cycle, PRXs oxidize several PCW components and consequently enhance PCW stiffening by directly affecting its components’ cross-links (Dunand et al., 2007; Marzol et al., 2020). PRXs indeed catalyze the formation of cross-links between phenolics and extensins. In this sense, the cell wall stiffens, and hence the cellular elongation capacity is reduced (Francoz et al., 2015). Thus, on the one hand, elevated levels of $H_2O_2$ in the apoplast were found to increase cell wall rigidity. However, in their hydroxyl cycle, PRXs produce ROS which enhance the nonenzymatic PCW loosening and thus promote cellular elongation. In this sense, auxin-induced RBOHs produce $O_2^-$ into the PCW. There, these short-living molecules are rapidly dismutated into $H_2O_2$. Then, $H_2O_2$ is utilized by PRXs to produce $OH^-$ (Majda and Robert, 2018; Marzol et al., 2020). In fact, the produced $OH^-$ catalyzes the oxidative cleavage of xyloglucan and other pectins, which loosens the PCW and promotes cellular expansion (Schmidt et al., 2016). These contrasted findings suggest that ROS have a dual role in determining the cell wall status and reveal the effects of hydroxyl-peroxide ($OH^:-H_2O_2$) balance in the apoplast (Schmidt et al., 2016). Yet, it is unclear how these opposite effects on cell wall polymers are coordinated during cell elongation (Lee et al., 2018; Francoz et al., 2019).

In addition, a transcription factor of the MYB family, MYB30, which regulates the expression of several genes involved in the transport of very-long-chain fatty acids (VLCFAs), has been identified to regulate cellular expansion in the root in response to ROS (Mabuchi et al., 2018). The transfer DNA insertion line myb30-1, which has reduced levels of MYB30, showed a higher root elongation rate than the WT upon $H_2O_2$ treatment. By counting the number and measuring the length of cortical cells in the MZ and EZ, it was proved that this increase in elongation rate is due to longer cells in the EZ and not to an increase in cells number (Mabuchi et al., 2018). Also, the overexpression of MYB30 in myb30 mutants resulted in shorter roots under control conditions. These results suggested that MYB30 is sufficient to inhibit root elongation. The observation of MYB30 expression by transcriptional and translational fusions revealed that MYB30 is induced by $H_2O_2$ and expressed in the MZ and EZ (Mabuchi et al., 2018).

**ROS at the interface between proliferation and differentiation**

An additional complexity in the redox regulation of the developmental processes is presented at the transition zone (TZ) of the primary root at the interface between the MZ and the EZ (Singh et al., 2016). At the root TZ, UPBEAT1 (UPB1), a bHLH TF, regulates the ROS homeostasis between two ROS species, i.e., superoxide ($O_2^-$) and peroxide ($H_2O_2$), to direct growth either toward proliferation in the MZ or toward differentiation in the upper zones (Figure 1). It was detected that a high ($O_2^:-H_2O_2$) ratio stimulates cellular division that requires advanced $O_2$ concentrations. However, a low ($O_2^:+H_2O_2$) ratio gives preference to cellular differentiation apart from the MZ (Tsukagoshi et al., 2010). The upb1 mutant shows a significant increase in the number of cortical cells, indicating enlargement of the meristem. An overexpression of UPB1 showed reduced root length and a decrease in the cortex cell number in the MZ (Tsukagoshi et al., 2010). UPB1 acts as a regulator of root growth through modulation of the transition from cell proliferation to elongation as well as plays a role in controlling cell size. UPB1 has 166 putative direct target genes, among which are several TFs. It also inhibits expression of several peroxidase genes (PRX39, PRX40, and PRX57), where PRX57 is of higher importance in this context as it is colocalized at the TZ (Figure 1). This inhibition of PRX expression leads to an accumulation of $H_2O_2$, changing the redox homeostasis in favor of differentiation rather than proliferation. Hence, upb1 mutants develop longer roots than the WT (Tsukagoshi et al., 2010).

**ROS & differential elongation**

ROS are required to establish the gravitropic curvature response of plant roots. This response is an asymmetrical cellular elongation, where gravity stimulates the arrest of elongation at the lower flank of the root (Singh et al., 2016). In fact, gravitropism is a versatile and multistep process. First, gravity sensing begins in specialized columella cells containing statoliths, i.e., plastids rich in starch, in the root cap (Blancaflor et al., 1998; Leitz et al., 2009). Afterward, an auxin-dependent complex signaling pathway (Zhang and Friml, 2020), which also involves calcium ($Ca^{2+}$), translocates this message into the EZ, where the curvature response occurs (Su et al., 2017). ROS are required to establish the root elongation in response to gravitropism (Swarup et al., 2013). In Zea mays roots, gravistimulation induces higher ROS production in the apex and the lower cortex than in the upper zones. A prolonged stimulation leads to consequent ROS accumulation in the upper zones, particularly in the EZ. This asymmetric distribution of ROS suggests that ROS are actors in developing the gravitropic response of roots. In fact, the gravity-dependent redistribution of auxin triggers the root bending responses by stimulating ROS generation (Joo et al., 2001). Auxin via
phosphatidylinositol 3-kinase (PI3K)-dependent pathway activates transmembrane RBOHs and triggers subsequent ROS production at the bending region (Krieger et al., 2016). The produced ROS inhibit cellular elongation by modulating the cell wall stiffening.

**Role of ROS during root differentiation**

**ROS & tracheary elements**

ROS are implicated in the differentiation of the tracheary elements, particularly in their lignification process. In *A. thaliana*, vasculature is arranged into a central stele, where xylem and phloem are interspersed with undifferentiated procambial cells and surrounded by the pericycle. These longitudinal vessels transport water and nutrients through the roots to the upper plant parts. Accordingly, they develop reinforcing and lignified secondary wall thickenings to cope with the negative water potential and prevent cellular collapse (Seagull, 2016). Lignin formation has been extensively studied as part of xylem differentiation. Lignin is a poly phenolic polymer generated by the radical coupling of aromatic monolignol, which is oxidized by ROS-dependent PRXs (PRX72 and PRX17) and laccases (LAC4) (Tsukagoshi, 2016; Fujita et al., 2020; Hoffmann et al., 2020). In addition, a putative role of ROS as a lignification regulator is derived from the roles of these PRXs acting during xylem development. The cationic peroxidase from *Zinnia elegans* (ZePRX) catalyzes the last step of lignification in *Zinnia elegans*. The ortholog in *A. thaliana*, PRX72, is upregulated in the xylem vessel elements, suggesting a role in xylem formation. In *prx72* mutants, lignin levels were 40% less than in the WT. It has been observed that PRX72 expression is increased in the *lac4 lac11 lac17* triple mutants, implying a possible functional linkage between PRXs and LACs during monolignol coupling (Zhao et al., 2013). However, increased PRX72 expression is not sufficient to rescue the lignin defect in the triple mutant, indicating that LACs and PRXs can still have distinct or complementary roles during monolignol assembly (Rojas-Murcia et al., 2020). Moreover, PRX17 is also involved in lignified tissue formation (Figure 2) and, more precisely, in regulating the kinetics of lignified wall formation (Herrero et al., 2013; Hoffmann et al., 2020). Its corresponding loss-of-function mutant, *prx17*, has lower lignin content in most organs, whereas xylans were overaccumulated in the *PRX17* overexressor compared with the WT. Additionally, three MYBPLANT motifs have been localized in the *PRX17* promoter; MYBPLANT-binding sites specify the expression of genes in lignifying cells (Cosio et al., 2017). However, it remains puzzling whether PRX17 acts directly in vessel lignification or indirectly via ROS signaling.

**ROS & root endodermis**

Root endodermis is the innermost cortical layer that surrounds the vascular stele in the root that resembles a polarized epithelium and functions in nutrient uptake and stress resistance. The development and differentiation of this layer are triggered by both intrinsic and extrinsic cues such as abiotic stresses, nutritional signals, and abscisic acid (ABA)-dependent metabolism (Lee et al., 2013). Its primary function is to form a selective barrier that filters the passage of the soil-absorbed water, solutes, and nutrients into the plant’s transporting vessels. This vital role is acknowledged to the hydrophobic coating of lignified Casparian strips (CSs) and its suberin lamellae (Cohen et al., 2020). CSs have belt-like structures middling the anticlinal cell walls of the endodermal cells. They seal the extracellular spaces by tying the cell walls between adjacent cells (Franke, 2015). CS formation is tightly regulated by ROS and their mediated enzymes RBOHs and PRXs (Figure 3). Nevertheless, the regulatory mechanisms underlying the endodermal selectivity are still under investigation.

In *Arabidopsis*, CS formation sites are determined by the localization of the transmembrane protein family, CS membrane domain proteins (CASPs). CASPs are explicitly expressed in the endodermis (Li et al., 2018a, 2018b) and localize specifically at the Casparian strip membrane domain (CSD) (Barbosa et al., 2019), where CASP1 guides CS-specific localization by interacting with two receptor-like kinases SCHENGEN1 (SGN1) and SCHENGEN3 (SGN3) (Allassimone et al., 2016). CASPs act as a transmembrane scaffold (Barbosa et al., 2019) and coordinate the proper localization of the different actors of the lignin-polymerizing machine: RBOHF, PRX64, and ENHANCED SUBERIN 1 (ESB1) at the CSD (Li et al., 2018a). RBOHFs function downstream the formation of the localized CASP scaffold. It generates superoxide (O2·−) which readily mutates into H2O2. Subsequently, PRX64 utilizes the produced H2O2 for monolignol oxidation, hence catalyzing lignin formation. Additionally, ESB1 forms a part of this assembly; it stabilizes the CASP protein scaffold and, contrariwise, requires the CASP complex for its CS localization (Lee et al., 2013; Franke, 2015).
Moreover, MYB36 has been revealed to regulate the expression of CASPs, PRX64, and ESB1 (Kamiya et al., 2015). Loss-of-function mutants, myb36, show deficit CS and hence lose the apoplastic barrier in this region of the root. Beforehand, during the endodermal differentiation, MYB36 expression is driven by the interaction of two upstream TFs, SHORTROOT (SHR) and SCARECROW (SCR). SHR and SCR are responsible for the initial specification of endodermal cell identity (Liberman et al., 2015; Li et al., 2018b). Apart from PRXs, MYB36 may also target the endodermis-enriched LACs. LACs are glycosylated, multi-copper enzymes that catalyze the oxidation of various phenolic substrates using O2 as final electron acceptor, not requiring H2O2 (Rojas-Murcia et al., 2020). Previously, the role of LACs in lignification in the vascular tissue was reported as

Figure 2. RBOHs and PRXs are important molecular players in several ROS-mediated developmental processes in roots
(A) RBOHE accumulates at the initiation site of the LR, while LR development in the loss-of-function mutant rbohe is delayed. GPXs are also expressed in the LR. gpx mutants develop larger lateral root primordia, indicating that the GPX family plays a role in regulating root architecture, especially GPX1 and GPX7, having a critical role in the development and formation of the lateral root. PRX07 and PRX75 control LR growth by maintaining ROS balance during the initiation stage in an auxin-independent manner. prx07 and prx75 have lower LR density and number. DZ: differentiation zone; EZ: elongation zone; LR: lateral root; MZ: meristematic zone; RC: root cap; RH: root hair.
(B) Localization of expression of various peroxidases. PRX17 is expressed in the xylem and root endodermis and participates in the lignin polymerization. prx17 mutants have lower lignin content in most organs. PRX37 is expressed in the root endodermis, and it participates in the phenolic cross-linking activity during lignin deposition in the cell wall. The overexpression of PRX17 results in a dwarf phenotype with small plants and delayed development. PRX64 is expressed in the root endodermis; it utilizes peroxide for a monolignol oxidation during lignin formation. PRX72 is expressed in the root endodermis and the xylem. Its expression is upregulated when the xylem is actively forming. The lignin levels in prx72 mutants are 40% less compared with those in the WT.
(C) RBOHC and PRX01, PRX07, PRX44, and PRX73 are expressed in the trichoblast cells in the epidermis. They are key enzymes responsible for root hair elongation by controlling the ROS homeostasis at the tip of the root hair cell. RBOHD and RBOHF are active in the mature root region especially in the vascular cylinder and LRP. RBOHD is expressed in the endodermis and procambium; the number and density of LR is higher in rbohd mutants than in the WT. RBOHF is required during CS formation in the endodermis, where its localization is coordinated by several CASPs and participates in lignin polymerization.

Moreover, MYB36 has been revealed to regulate the expression of CASPs, PRX64, and ESB1 (Kamiya et al., 2015). Loss-of-function mutants, myb36, show deficit CS and hence lose the apoplastic barrier in this region of the root. Beforehand, during the endodermal differentiation, MYB36 expression is driven by the interaction of two upstream TFs, SHORTROOT (SHR) and SCARECROW (SCR). SHR and SCR are responsible for the initial specification of endodermal cell identity (Liberman et al., 2015; Li et al., 2018b). Apart from PRXs, MYB36 may also target the endodermis-enriched LACs. LACs are glycosylated, multi-copper enzymes that catalyze the oxidation of various phenolic substrates using O2 as final electron acceptor, not requiring H2O2 (Rojas-Murcia et al., 2020). Previously, the role of LACs in lignification in the vascular tissue was reported as
the overall lignin content was reduced in both lac and prx mutants (Pfister et al., 2014; von Wangenheim et al., 2017; Wunderling et al., 2018). However, the role of LACs in the endodermis and whether some cell types preferentially use LACs and others PRXs were addressed recently (Rojas-Murcia et al., 2020). In fact, the CS was devoid in the quintuple mutant prx03 prx09 prx39 prx72 prx64, while it showed no difference in the lac13578912136 nonuple mutant compared with the WT. Together, these findings suggest that PRXs play very specific roles in various contexts, and LACs are redundant for lignification of primary cell walls but are required for lignification of secondary cell walls (Rojas-Murcia et al., 2020). Additionally, other RBOH and PRX enzymes are also known to be involved in ROS homeostasis and cell wall remodeling in the endodermis, including apoplastic RBOHD, PRX37, and PRX72. PRX37 participates in phenolic “cross-linking” activity during lignin deposition (Pedreira et al., 2011), and PRX72 is involved in lignin biosynthesis when plant growth has ceased (Figure 2) (Kim et al., 2010; Fernández-Pérez et al., 2015; Tognetti et al., 2017; Fujita et al., 2020).

**ROS & root hair**

Presence of root hairs marks the DZ of the primary root and is among its exhaustively studied features. In fact, root hairs are tube-like extensions projecting from single specialized epidermal cells. They massively increase the surface area of the root and hence improve its absorption potential of water and soil-borne nutrients. ROS play a critical role in root hair development (Sundaravelpandian et al., 2013; Zhao et al., 2016; Mhamdi and Van Breusegem, 2018b; Choudhary et al., 2020). In Arabidopsis, RBOHC is the key enzyme responsible for root hair elongation under ROS regulation although RBOHH and RBOHJ also contribute to a much lower extent (Mangano et al., 2017). RBOHC actively produces superoxide, and the hydroxyl radicals formed downstream of superoxide are the specific ROS that stimulates root hair

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**Figure 3. Selected examples of ROS-mediated root developmental process**

(A) In lateral roots (LRs), UPBEAT1, a basic helix-loop-helix TF expressed at the TZ, directly regulates peroxidases that alter the distribution of ROS. Cell proliferation requires high levels of O2, whereas high H2O2 induces differentiation. UPB1 is transcriptionally enriched during the early stages of lateral root primordia (LRP) when ROS significantly accumulate in the emerging LR. UPB1 overexpressors have more LRPs. upb1 has more emerging LRs than WT. UPB1 controls the expression of a subset of PRXs genes that manipulate the ROS balance. Besides, MYB36 acts in stages V and VI of LR development, by manipulating the ROS balance via the regulation of at least PRX09 and PRX64. (B) In growing root hairs, RBOHC is the key enzyme that produces ROS in the apoplast. RBOHC-mediated ROS leads to the activation of a MAPK cascade through OX11 during cell elongation of root hairs. At the transcriptional level, RSL4, a bHLH transcription factor, regulates the expression of several class III PRXs (PRX01, PRX44, and PRX73), which participate in root hair growth and ROS homeostasis. In addition, PRX07 is also required for root hair elongation. (C) In inner root tissues, SHORTROOT (ShR) transcription factor (TF) moves from the stelo to the endodermis where it interacts with another TF SCARECROW (SCR), and then the SHR:SCR complex specifies the endodermal cell identity and activates the transcription of MYB36. MYB36 regulates the biosynthetic apparatus for Casparian strips (CSs). It also plays role in the lateral root primordia differentiation and lateral root initiation. MYB36 regulates the expression of several underlying genes such as CASPS, PRX64, and ES81 to trigger the proper endodermis development. CASP is responsible for the proper localization at the CS development site by the interaction with two receptor-like kinases SCHENGEN1 (SGN1) and SCHENGEN3 (SGN3). ES81 stabilizes the CAPS proteins scaffold. RBOHF generates superoxide that mutates into peroxide. PRX64 and LACs oxidize monolignols and catalyze lignin formation.
elongation (Chapman et al., 2019), and expectedly, rbohc mutants have much shorter root hairs than the WT (Foreman et al., 2003). ROS stimulate ROS-dependent calcium gates, creating a Ca\textsuperscript{2+} gradient from the tip inward. This gradient drives the apical elongation of the root hair (Mhamdi and Van Breusegem, 2018b). RBOHC trafficking, transport to its correct subcellular compartment, and localization are complex processes regulated by the receptor-like kinase FERONIA (FER) and members of the Rho family of plant GTPases (ROP) (Chapman et al., 2019). The FER extracellular domain binds to the pectin components of the cell wall and activates Ca\textsuperscript{2+} channels that maintain the integrity of the cell wall during growth. fer and rbohc mutants possess similar phenotypes, with short root hairs and decreased ROS accumulation. Pull-down assays that map protein-protein interactions suggest that FER interacts with ROP2, promoting radical hair growth upstream of RBOHC (Chapman et al., 2019). Another actor on the ROS homeostasis during root hair development is the LIKE SEX FOUR 2 (LSF2) gene encoding a phosphatase. This gene was identified for its novel role in redox regulation (Zhao et al., 2016). Isf2 Arabidopsis mutants showed lower superoxide generation rate but higher levels of hydrogen peroxide upon oxidative stress treatments. LSF2 interacts with the mitogen-activated protein kinase 8 (MPK8) which regulates the cytosolic redox balance (Zhao et al., 2016). Besides RBOHC, another component of ROS signaling in growing root hair is the OXIDATIVE BURST INDUCIBLE1 (OXI1) kinase. OXI1 is a Ser-Thr kinase, which plays a role in elongation of root hairs. It was proposed that RBOHC-mediated ROS lead to the activation of a MAPK cascade through OXI1 during cell elongation of root hairs (Figure 3) (Rentel et al., 2004).

Moreover, auxin triggers a higher ROS level in the root hairs and enhances their growth by ROOT HAIR DEFECTIVE SIX LIKE4 (RSL4) activation (Mangano et al., 2017). RSL4 encoding a bHLH TF has been identified as a vital regulator of root hair development (Yi et al., 2010; Mangano et al., 2017; Pires et al., 2013). Furthermore, it is known that the expression of RSL4 is controlled by several auxin response factors (ARFs), including ARF5, ARF7, ARF8, and ARF19; this ARF-RSL network can induce localized ROS synthesis (Figure 3), promoting the development of root hairs (Mangano et al., 2017, Mangano et al., 2018; Chapman et al., 2019). Moreover, in addition to RBOHC, the key enzyme involved in root hair growth RBOHH and RBOHJ have been identified as important enzymes in this process. These RBOHs have root hair cis-elements (RHE) in their genes’ promoters highly similar to the RSL4 response elements (RSL-RE). It was suggested that RSL4 binds at least one RHE region in the RBOHC and RBOHJ promoters and hence directly regulates their expression (Mangano et al., 2017). In parallel, among the thirty-four putative targets of RSL4, PRX07 has been identified which codes the PRXs required for root hair elongation (Vijayakumar et al., 2016). PRX regulatory regions include several putative RHE motifs. Chromatin immunoprecipitation (ChIP) experiment indicated that RSL4 can bind to the RHE in the regulatory region of PRX01, PRX44, PRX60, and PRX73. Consistently, PRX01 and PRX73 were upregulated in the overexpressors of RSL4 and downregulated in rsl4 mutants (Mangano et al., 2017). It is proposed that the phenotypes observed for peroxidase null mutants, especially pxr01, pxr44, and pxr73, are due to their participation in root hair growth and ROS homeostasis (Figure 3) (Mangano et al., 2017). These three genes in turn are highly coexpressed with other root hair-specific genes such as extensins and glycoproteins, which jointly participate in the expansion of the cell wall (Mangano et al., 2018; Marzol et al., 2020). Double mutants pxr44 pxr73, pxr01 pxr44, and pxr01 pxr73, as well as the triple null mutant, pxr01 pxr44 pxr73, showed shorter root hairs, possibly due to the high accumulation of H\textsubscript{2}O\textsubscript{2} in the apoplast, affecting ROS homeostasis, the root hair growth, and the thickness of the cell wall (Marzol et al., 2020). ROOT HAIR DEFECTIVE SIX LIKE 2 (RSL2), a close relative of RSL4, is also involved in root hair growth in an ROS-dependent manner involving PRXs and RBOHC enzymes, suggesting that RSL2-RSL4 may act together to regulate ROS homeostasis (Mangano et al., 2017, 2018). In addition, the subunit of the mediator complex PHYTOCHROME AND FLOWERING TIME1 (PFT1)/MED25 triggers H\textsubscript{2}O\textsubscript{2} production by the upregulation of a subset of PRX expression and regulates the equilibrium between H\textsubscript{2}O\textsubscript{2} and during root hair differentiation and cell elongation (Sundaravelpandian et al., 2013).

**ROS & lateral roots**

Plants have the capacity of the de novo development of new organs in which new cells are enrolled in the formation of primordia organs (Fernández-Marcos et al., 2017). Lateral roots (LRs) serve as an example of this feature and choreograph the dynamic architecture of the root system. LRs enhance the absorption of water and nutrients from the neighboring soil environment. They also contribute to the root acclimation to various biotic and abiotic constraints. LRs are intrinsically diverse as they have contrasted morphologies and anatomies, such as having different lengths and orientations (Muller et al., 2019). LRs are initiated when pericycle founder cells undergo several rounds of anticlinal divisions to create one-layered primordia of about ten equal cells; this is termed stage I. Next, the cells divide periclinally, forming an inner and an...
outer layer denoting stage II. Further anticlinal and periclinal divisions create dome-shaped primordia (stages III–VII) that eventually emerge at stage VIII from the parental root (Malamy and Benfey, 1997; Péret et al., 2009; Banda et al., 2019). In fact, the establishment of lateral root primordia (LRP) from the root pericycle is auxin dependent. These LRP further develop into LRs by orchestrated divisions of the lateral root founder cells (Fernández-Marcos et al., 2017).

Redox signaling is notably involved in LR development by a complex genetic network that has been thoroughly studied (Manzano et al., 2014; Reyt et al., 2015; Orman-Ligeza et al., 2016). First, SKP2B (S-phase kinase-associated protein2), a marker for LR development, is coexpressed with many ROS-related genes. UPB1 is transcriptionally enriched during the early stages of LRP, and ROS significantly accumulate in the emerging LR. In addition, plants overexpressing UPB1 have more LRP in stages IV and V. However, in stage VIII, when LRP emerge from the epidermis, their number was significantly reduced in the UPB1 overexpressors. Coherently, upb1 mutants had more emerging LRs than WT (Manzano et al., 2014). UPB1 controls the expression of a subset of PRX genes that manipulate the ROS balance (Figure 3). Altogether, this suggests that ROS play a role in the early development of LRP. Moreover, a potassium cyanide treatment, which inhibits PRX activity, reduced LR emergence. Diaminobenzidine and nitro blue tetrazolium staining revealed, respectively, that H₂O₂ and superoxide are present in all the developmental stages of LRs (Manzano et al., 2014).

Additionally, MYB36, a previously identified advocate in CS formation during endodermal differentiation (Kamiya et al., 2015; Liberman et al., 2015), was also identified as an actor in LR development (Fernández-Marcos et al., 2017). Initially, MYB36 protein was revealed to accumulate in the root endodermis. Its overexpression leads to shorter primary roots, whereas three allelic myb36 mutants develop longer roots (Fernández-Marcos et al., 2017). This phenotype results from the increase in the number of cells in the meristem, where ROS balance is known to regulate the meristematic boundaries (Kamiya et al., 2015; Liberman et al., 2015). The new role of MYB36 in LR development was first discovered in 2017 (Fernández-Marcos et al., 2017). Two allelic mutants of myb36 had less LRs than the WT. A spatiotemporal tracking of the expression pattern of this gene showed that it was not expressed in the endodermal cells adjacent to the LRP. However, it was cell-autonomously enriched in the lateral root primordia boundary (LRPB) at stage V of the LR development. LRP in the two mutants were flat, and they failed to form into domes and hence failed to penetrate the overlying cellular layers. Further spatiotemporal analyses of the expression pattern of
MYB36 and its corresponding PRX targets confirmed that MYB36 acts in stages V and VI of LR development, by manipulating the ROS balance via the regulation of at least PRX09 and PRX64 (Fernández-Marcos et al., 2017). Finally, supplementary forward genetics approaches identified another ROS-related genetic basis affecting the LR phenotype. PRX07 and PRX57 were explicitly identified as the actors in LR growth among other peroxidases (Figure 2) since prx07 and prx57 mutants have lower LR density owing to changes of ROS balance in the initiation stage. Interestingly, despite that LRP establishment depends on auxin, PRX07 and PRX57 were detected to be auxin-independent (Manzano et al., 2014). Together, all these studies highlight the importance of ROS homeostasis in shaping and growing LRs.

Decoding ROS signal

As demonstrated above, ROS production and scavenging naturally takes place in various cellular compartments and is surveilled by the profound ROS gene network to maintain their homeostasis (Das and Roychoudhury, 2014; Singh et al., 2016; Mittler, 2017). It is worth noting that ROS do not occur in isolation but rather as part of a large and complex redox signaling that involves many components. However, the different ROS levels in the different cellular compartments may be contemplated as ROS signatures in resemblance to the famous Ca²⁺ signature. In fact, Ca²⁺ signaling is implicated in countless cellular activities, where varying spatiotemporal distribution, vibration amplitude, and frequency of Ca²⁺ determine a distinct signature. This specific signature, triggered by a particular stimulus, results in a precise adaptive response (Yuan et al., 2017). As for ROS, different natures and intensities of stimuli lead to the formation of ROS signatures, which are decoded by ROS sensors to generate a stimulus-specific signal, and consequently to customize an acclimation response in the plant cell. This signature varies in compliance with the cell type, developmental stage, and environmental status (Choudhury et al., 2017), and its decoding can alter protein structures or functions via PTMs. For instance, these alterations can affect TFs leading to modifications in their transcription (Dietz et al., 2016).

This concept is exceptionally important, as it explains how plants are able to decrypt different ROS signals provoked by, chemically, the same molecules, and then elaborate a specific response to each particular signature (Choudhury et al., 2017). It is notable how these ROS-producing or ROS-scavenging machineries are copresent simultaneously within cells or even organelles in widely ranging proportions and activities. What turns them on or off remains an intriguing question. They are surely dependent on the ambient O₂ concentration and on the exogenous environmental cues. Furthermore, they are also dependent on the cell type, localization, and status, which is subsequently depending on the gene expression pattern. Thus, an optimal pattern of oxidants and antioxidants exists for a given physiological condition (Sies et al., 2017).

Conclusion

Recent studies investigated the vital roles of ROS in plant root developmental processes. In summary, during cell proliferation, ROS concentrations regulate the cell cycle since oxidative bursts can block cyclin expression. In the MZ, ROS act to conserve the stem cell niche in the QC, hence allowing a sustained growth of the root. In the EZ, apoplastic ROS balance determine the elongation rates of cells in the root EZ. In fact, apoplastic ROS can promote either PCW stiffening or loosening, by catalyzing either the cross-links between PCW components or their oxidative cleavage, respectively. At the transition zone, ROS balances give preference to either proliferation or differentiation according to UPB1 regulation. These aforementioned mechanisms reveal a vital role of ROS during the plant root primary growth. Moreover, ROS are involved in modulating the differentiation stage of development at multiple levels. In the tracheary elements, ROS regulate xylem lignification by the functioning of the ROS-producing enzymes, PRX72 and PRX17. Similarly, PRX64 and RBOHF are essential for the proper formation of the CS in the root endodermis. In addition, in the epidermis, ROS trigger cell growth in root hairs. Last but not least, numerous ROS-related genes are implicated in LR formation. ROS production and scavenging are two co-occurring processes that are minutely adjusted to maintain the manifold “ROS homeostasis” (Mittler et al., 2004; Suzuki et al., 2012; Mittler, 2017; Mhamdi and Van Breusegem, 2018b). This homeostasis is essential for regulating the multiple stages of plant root development and hence choreographing the final root architecture. Furthermore, this redox balance is central to life since redox practices encompass all fundamental processes, from bioenergetics to metabolism and life functions (Sies et al., 2017). In our scope, ROS homeostasis has been shown as a requisite for proper root growth (Tsukagoshi, 2016; Choudhury et al., 2020), development (Mhamdi and Van Breusegem, 2018b), and response to stimuli (Miller et al., 2010;
Liu and He, 2016; Zhang et al., 2016). In view of the above, ROS of underground plant cells are multitaskers in root development.

Future key issues

a. Apoplastic ROS-sensing mechanism by cell surface receptors has been recently proposed to be mediated by HPCA1 (Laohavisit et al., 2020; Wu et al., 2020). The downstream post-translational modifications relative to the kinase domain of CARD1/HPCA1 were also described. However, the further detailed downstream signaling components still to be identified.

b. ROS fluctuations are tightly connected with Ca^{2+} signals (e.g. polar growing cells like root hairs). However, it is not clear how these two signals are interconnected. In this context, it is known that Ca^{2+} activates RBOHs, but which Ca^{2+} channels are activated by ROS (e.g. H_{2}O_{2}) remain unknown. In addition, it is unknown if this process is direct and not mediated by other components (e.g. CARD1/HPCA1 or other kinases).

c. How apoplastic ROS pool is coordinated with the “cytoplasmic” and “organellar” ROS pools in order to maintain a coherent ROS homeostasis in the plant cells along growth and development? The use of multiple genetically encoded ROS sensors targeted to different subcellular zones will allow us to get a high-resolution spatial map distribution of ROS species.

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Author contributions

A.E. analyzed the bibliography, wrote the paper, and helped on the figures’ design. Y.d.C.R.G. analyzed the bibliography and helped on the figures’ design. A.E, J.M.E., and Ch.D. analyzed the bibliography, supervised the project, and wrote the paper. This manuscript has not been published and is not under consideration for publication elsewhere. All the authors have read the manuscript and have approved this submission.

Declaration of interests

The authors declare no competing interests.

References

Allassimone, J., Fujita, S., Doblás, V.G., van Dop, M., Barberon, M., Kalmbach, L., Vermeer, J.E., Rojas-Murcia, N., Santuari, L., Hardtke, C.S., et al. (2016). Polarity localized kinase SGN1 is required for Casparian strip integrity and positioning. Nat. Plants 2, 1–10.

Anjum, N.A., Szfo, A., Scopa, A., Roychoudhury, A., Gill, S.S., Iqbal, M., Lukatin, A.S., Pereira, E., Duarte, A.C., and Ahmad, J. (2015). Lipids and proteins—major targets of oxidative modifications in abiotic stressed plants. Environ. Sci. Pollut. Res. 22, 4099–4121.

Apel, K., and Hirt, H. (2004). Reactive oxygen species: metabolism, oxidative stress, and signal transduction. Annu. Rev. Plant Biol. 55, 373–399.

Banda, J., Bellande, K., von Wangenheim, D., Goh, T., Guymarch’, S., Laplaze, L., and Bennett, M.J. (2019). Lateral root formation in arabidopsis: a well-ordered LReXit. Trends Plant Sci. 24, 826–839.

Barbosa, I.C.R., Rojas-Murcia, N., and Geldner, N. (2019). The Casparian strip—one ring to bring cell biology to lignification? Curr. Opin. Biotechnol. 56, 121–129.

Baxter, A., Mittler, R., and Suzuki, N. (2014). ROS as key players in plant stress signalling. J. Exp. Bot. 65, 1229–1240.

Berkowitz, O., De Clercq, I., Van Breusegem, F., and Whelan, J. (2016). Interaction between hormonal and mitochondrial signalling during growth, development and in plant defence responses. Plant Cell Environ. 39, 1127–1139.

Blancaflor, E.B., Fasano, J.M., and Gilroy, S. (1998). Mapping the functional roles of cap cells in the response of arabidopsis primary roots to gravity. Plant Physiol. 116, 213–222.

Bose, J., Rodrigo-Moreno, A., and Shabala, S. (2014). ROS homeostasis in halophytes in the context of salinity stress tolerance. J. Exp. Bot. 65, 1241–1257.

Bratt, A., Rosenwasser, S., Meyer, A., and Fluhr, R. (2016). Organelle redox autonomy during environmental stress. Plant Cell Environ. 39, 1909–1919.

Chakraborty, K., Bish, S.K., Goswami, N., Singh, A.L., and Zala, P.V. (2016). Differential fine-regulation of enzyme driven ROS detoxification network imparts salt tolerance in contrasting peanut genotypes. Environ. Exp. Bot. 128, 79–90.

Champion, J.M., Muhlemann, J.K., Gayomba, S.R., and Muday, G.K. (2017). RBOH-dependent ROS synthesis and ROS scavenging by plant specialized Metabolites to modulate plant development and stress responses. Physiol. Behav. 176, 139–148.

Champion, J.M., Muhlemann, J.K., Gayomba, S.R., and Muday, G.K. (2019). RBOH-dependent ROS synthesis and ROS scavenging by plant specialized Metabolites to modulate plant development and stress responses. Chem. Res. Toxicol. 32, 370–396.
Choudhary, A., Kumar, A., and Kaur, N. (2020). ROS and oxidative burst: roots in plant development. Plant Divers. 42, 33–43.

Choudhary, F.K., Rivero, R.M., Blumwald, E., and Mittler, R. (2017). Reactive oxygen species, abiotic stress and stress combination. Plant J. 90, 856–867.

Cohen, H., Fedyuk, V., Wang, C., Wu, S., and Aharoni, A. (2020). SUBERMAN regulates developmental suberization of the Arabidopsis root endodermis. Plant J. 102, 431–447.

Cosio, C., Ranocha, P., Francoz, E., Burlat, V., Zheng, Y., Perry, S.E., Ripoll, J.J., Yanofsky, M., and Dunand, C. (2017). The class III peroxidase PXR1 is a direct target of the MADS-box transcription factor AGAMOUS-LIKE15 (AGL15) and participates in lignified tissue formation. New Phytol. 213, 250–263.

Cosio, C., Vuillemin, L., De Meyer, M., Kevers, C., Penel, C., and Dunand, C. (2009). An anionic class III peroxidase from zucchini may regulate hypocotyl elongation through its auxin oxidase activity. Planta 229, 823–836.

costa, A., Dragi, I., Behera, S., Zottini, M., Pizzo, P., Schrower, J.I., Pozzan, T., and Lo Schiavo, F. (2010). H2O2 in plant peroxisomes: an in vivo analysis uncovers a Ca2+-dependent scavenging system. Plant J. 62, 760–772.

Cui, H., Kong, D., Wei, P., Hao, Y., Torri, K.U., Lee, J.S., and Li, J. (2014). SPINDLY, ERECTA, and its ligand STOMAGEN have a role in redox-mediated cortex proliferation in the arabidopsis root. Mol Plant 7, 1727–1739.

Czarnocka, W., and Karpinski, S. (2018). Friend or foe? Reactive oxygen species production, scavenging and signaling in plant response to environmental stresses. Free Radic. Biol. Med. 122, 4–20.

Dagda, R.K., Rice, M., Merrill, R.A., Filipo, K.H., and Strack, S. (2017). Techniques to investigate mitochondrial function in neurons. Neurmeothods 123, 1–27.

Daloso, D.M., Müller, K., Obata, T., Florian, A., Tohge, T., Bottcher, A., Riondet, C., Bariat, L., Carrari, F., Nunes-Nesi, A., et al. (2015). Thioredoxin, a master regulator of the tricarboxylic acid cycle in plant mitochondria. Proc. Natl. Acad. Sci. U S A 112, E1392–E1400.

Das, K., and Roychoudhury, A. (2014). Reactive oxygen species (ROS) and response of antioxidant s as ROS-scavengers during environmental stress in plants. Front. Environ. Sci. 2, 1–13.

Deng, B., Deng, S., Sun, F., Zhang, S., and Dong, H. (2011). Down-regulation of free riboflavin content induces hydrogen peroxide and a pathogen defense in Arabidopsis. Plant Mol. Biol. 77, 185–201.

Dietz, K.J. (2014). Redox regulation of transcription factors in plant stress acclimation and development. Antioxid. Redox Signal. 21, 1356–1372.

Dietz, K.J., Mittler, R., and Noctor, G. (2016). Recent progress in understanding the role of reactive oxygen species in plant cell signaling. Plant Physiol. 171, 1535–1539.

Dunand, C., Crèvecoeur, M., and Penel, C. (2007). Distribution of superoxide and hydrogen peroxide in Arabidopsis root and their influence on root development: possible interaction with peroxidases. New Phytol. 174, 332–341.

Eltaeye, A.E., Kawano, N., Badawi, G.H., Kaminaka, H., Sanekata, T., Shibahara, T., Inanaga, S., and Tanaka, K. (2007). Overexpression of monodehydroascorbate reductase in transgenic tobacco confers enhanced tolerance to ozone, salt and polyethylene glycol stresses. Planta 225, 1255–1264.

Farvardin, A., Gonzalez-hernandez, A.I., Llorens, E., Garcia-agustin, P., Scalschi, L., and Vicedo, B. (2020). The apoplastic: a key player in plant survival. Antioxidants 9, 1–26.

Fernández-Marcos, M., Descovsky, B., Manzano, C., Uberman, C., del Pozo, J.C., and Gutierrez, C. (2017). Control of Arabidopsis lateral root primordium boundaries by MYB36. New Phytol. 213, 105–112.

Fernández-Pérez, F., Pomar, F., Pedreño, M.A., and Novo-Uzal, E. (2015). The suppression of AtPmS2 affects fibers but not xylem lignification in Arabidopsis by altering the proportion of syringyl units. Physiologia Plantarum 155, 395–406.

Fichman, Y., Miller, G., and Mittler, R. (2019). Whole-plant live imaging of reactive oxygen species. Mol Plant 12, 1203–1210.

Foreman, J., Demidchik, V., Bothwell, J.H., Mylona, P., Medema, H., Torres, M.A., Linstead, P., Costa, S., Brownlee, C., Jones, J.D., et al. (2003). Reactive oxygen species produced by NADPH oxidase regulate plant cell growth. , Nat. Cell Biol. 5, 442–446.

Franke, R.B. (2015). ‘Caspary’s conductor’. Proc. Natl. Acad. Sci. U S A 112, 10084–10085.

Franke, E., Fonseca, E., Le Ru, A., Martinez, Y., Fourquaux, I., Jaunee, A., Dunand, C., and Burlat, V. (2019). Fectin demethylcitification generates platforms that anchor peroxidases to remodel plant cell walls. Dev. Cell 48, 261–276.e8.

Francoz, E., Ranocha, P., Nguyen-Kim, H., Jamet, E., Burlat, V., and Dunand, C. (2015). Roles of cell wall peroxidases in plant development. Phytochemistry 112, 15–21.

Franke, R.B. (2015). ‘Caspary’s conductor’. Proc. Natl. Acad. Sci. U S A 112, 10084–10085.

Fujita, S., De Bellis, D., Edel, K.H., Ko¨ ster, P., Andersen, T.G., Schmid-Siegert, E., De ´ nervaud Janku, M., Luhová, L., and Petrivalsky ´, M. (2019). Hyper, a hydrogen peroxide sensor, indicates the sensitivity of the Arabidopsis root elongation zone to aluminum treatment. Sensors (Switzerland) 15, 855–867.

Herrero, J., Fernández-Pérez, F., Yebra, T., Novo-Uzal, E., Pomar, F., Pedreño, M.A., Cuello, J., Guér, A., Esteban-Carrasco, A., and Zapata, J.M. (2013). Bioinformatic and functional characterization of the basic peroxidase 72 from Arabidopsis thaliana involved in lignin biosynthesis. Plants 237, 1599–1612.

Hoffmann, N., Benske, A., Betz, H., Schuentz, M., and Samuels, A.L. (2020). Laccases and peroxidases co-localize in lignified secondary cell walls throughout stem development. Plant Physiol. 184, 806–822.

Huang, S., Yen Aken, O., Schwä zlar, M., Belt, K., and Millar, A.H. (2016). The roles of mitochondrial reactive oxygen species in cellular signaling and stress response in plants. Plant Physiol. 171, 1551–1559.

Ivanov, B.N., Borisova-Mubarakhshina, M.M., and Kozuleva, M.A. (2018). Formation mechanisms of superoxide radical and hydrogen peroxide in chloroplasts, and factors determining the signalling by hydrogen peroxide. Funct. Plant Biol. 45, 102–110.

Janku, M., Luhová, L., and Petřivalský, M. (2019). On the origin and fate of reactive oxygen species in plant cell compartments. Antioxidants 8, 105.

Joo, J.H., Bae, Y.S., and Lee, J.S. (2001). Role of auxin-induced reactive oxygen species in plant growth. Physiologia Plantarum 126, 1055–1060.

Kamiya, T., Borghi, M., Wang, P., Danku, J.M.C., Kalmbach, L., Hosmani, P.S., Naseer, S., Fujiwara, T., Geldner, N., and Salt, D.E. (2015). The MYB36 transcription factor orchestrates Casparian strip formation. Proc. Natl. Acad. Sci. U S A 112, 10533–10538.

Kim, M.J., Ciani, S., and Schachtman, D.P. (2010). A peroxidase contributes to ros production during Arabidopsis root response to potassium deficiency. Mol. Plant 3, 420–427.

Kong, X., Tian, H., Yu, Q., Zhang, F., Wang, R., Gao, S., Xu, W., Li, J., Shani, E., Fu, C., et al. (2018). PHB3 maintains root stem cell niche identity through ROS-responsive AP2/ERF transcription factors in arabidopsis. Cell Rep. 22, 1350–1363.
Krieger, G., Shkolkov, D., Miller, G., and Fromm, H. (2016). Reactive oxygen species tune root tropic responses. Plant Physiol. 172, 1209–1220.

Kroiler-Schön, S., Steven, S., Kossmann, S., Scholz, A., Daub, S., Oelze, M., Xia, N., Hausding, M., Mikhed, Y., Zinnius, E., et al. (2014). Molecular mechanisms of the crosstalk between mitochondria and NADPH oxidase through reactive oxygen species - studies in white blood cells and in animal models. Antioxid. Redox Signal. 20, 247–266.

Laohavivat, A., Wakatake, T., Ishihama, N., Mulvey, H., Takiwara, K., Suzuki, T., and Shirasu, K. (2020). Quinone perception in plants via leucine-rich-repeat receptor-like kinases. Nature 587, 92–97.

Lee, Y., Rubio, M.C., Alassimone, J., and Geldner, N. (2013). A mechanism for localized lignin deposition in the endodermis. Cell 153, 402–412.

Lee, Y., Yoon, T.H., Lee, J., Jeon, S.Y., Lee, J.H., Lee, M.K., Chen, H., Yun, J., Oh, S.Y., Wen, X., et al. (2018). A lignin molecular brace controls precision processing of cell walls critical for surface integrity in Arabidopsis. Cell 173, 1468–1480.e9.

Leitz, G., Kang, B-H., and Schoenwaelder, M.E.A. (2009). Statolith sedimentation kinetics and force responses: learning from AtRBOHD. Plant Cell 21, 843–860.

Li, N., Sun, L., Zhang, L., Song, Y., Hu, P., Li, C., and Hao, F.S. (2014). AtRbohD and AtRbohF negatively regulate lateral root development by changing the localized accumulation of superoxide in primary roots of Arabidopsis. Planta 241, 591–602.

Li, P., Yu, Q., Gu, X., Xu, C., Qi, S., Wang, H., Zhong, F., Baxin, T.I., Rahman, A., and Wu, S. (2018a). Construction of a functional casparian strip in non-endodermal lineages is orchestrated by two parallel signaling systems in Arabidopsis thaliana. Curr. Biol. 28, 2777–2786.e2.

Li, P., Yang, M., Chang, J., Wu, J., Zhong, F., Rahman, A., Qin, H., and Wu, S. (2018b). Spatial expression and functional analysis of casparian strip regulatory genes in endodermis reveals the conserved mechanism in tomato. Front. Plant Sci. 9, 832.

Liberman, L.M., Sparks, E.E., Moreno-Ruano, M.A., Petricka, J.J., and Benfey, P.N. (2015). MYB36 regulates the transition from proliferation to differentiation in the Arabidopsis root. Proc. Natl. Acad. Sci. U S A 112, 12099–12104.

Liu, Y., and He, C. (2016). Regulation of plant reactive oxygen species (ROS) in stress responses: learning from AtRBOHD. Plant Cell Rep. 35, 995–1007.

Livanos, P., Galatis, B., Quader, H., and Apostolakos, P. (2012). Disturbance of reactive oxygen species homeostasis induces atypical tubulin polymer formation and affects mitosis in root-tip cells of Triticum turgidum and Arabidopsis thaliana. Cytoskeleton 69, 1–21.

Livanos, P., Galatis, B., Quader, H., and Apostolakos, P. (2017). ROS homeostasis as a prerequisite for the accomplishment of plant cytokinesis. Protoplasma 254, 569–586.
Patricka, J.J., Winter, C.M., and Benfey, P.N. (2012). Control of arabidopsis root development. Annu. Rev. Plant Biol. 63, 563–590.

Pfister, A., Barberon, M., Alassium, J., Kalmbach, L., Lee, Y., Vermeer, J.E., Yamazaki, M., Li, G., Maurel, C., Takano, J., et al. (2014). A receptor-like kinase mutant with absent endodermal diffusion barrier displays selective nutrient homeostasis defects. Elife 3, e03113.

Pires, N.D., Yi, K., Breuninger, H., Catarino, B., Menard, B., and Dolan, L. (2013). Recruitment and remodeling of an ancient gene regulatory network during land plant evolution. Proc. Natl. Acad. Sci. U S A 110, 9571–9576.

Podgór ska, A., Burian, M., and Szal, B. (2017). GLO-Roots: an imaging platform enabling multidimensional characterization of soil-grown root systems. Elife 4, e07597.

Qi, J., Wang, J., Gong, Z., and Zhou, J.M. (2017). Mitochondrial and chloroplast stress responses are modulated in distinct touch and chemotaxis. Front. Plant Sci. 8, 1–15.

Rentel, M.C., Del Río, L.A., and López-Huertas, E. (2016). ROS balance is critical for multiple stages of root hair development in Arabidopsis. Plant Signaling Behav. 8, 10–12.

Suzuki, N., Kousukevitzky, S., Mittler, R., and Miller, G. (2012). ROS and redox signalling in the response of plants to abiotic stress. Plant Cell Environ. 35, 259–270.

Szymanski, D., and Staiger, C.J. (2018). The actin cytoskeleton: functional arrays for cytoplasmic organization and cell shape control. Plant Physiol. 176, 106–118.

Wardenburg, J., Winter, C.M., and Benfey, P.N. (2011). Methods for detection and measurement of hydrogen peroxide inside and outside of cells. Anal. Cell Environ. 37, 266–279.

Shukla, V., Lombardi, L., Iacopino, S., Pencik, A., Shulaev, V., Torres, M.A., and Mittler, R. (2011). Respiratory burst oxidases: the engines of ROS signaling. Curr. Opin. Plant Biol. 14, 691–699.

Swarup, R., Wells, D.M., and Bennett, M.J. (2013). Root gravitropism. In Plant Roots: The Hidden Half, Fourth Edition, pp. 265–278.

Tognetti, V.B., Bielach, A., and Hrtyan, M. (2017). Endogenous hypoxia in lateral root primordia controls root architecture by antagonizing auxin signaling in arabidopsis. Mol. Plant 12, 538–551.

Tognetti, V.B., Bielach, A., and Hrtyan, M. (2017). Mycorrhizal networks facilitate tree communication, learning, and memory. Memory and Learning in Plants (Cham: Springer), pp. 191–213.

Tsuchakoshi, H. (2016). Control of root growth and development by reactive oxygen species. Curr. Opin. Plant Biol. 29, 57–63.

Tsuchakoshi, H., Busch, W., and Benfey, P.N. (2010). Transcriptional regulation of ROS controls transition from proliferation to differentiation in the root. Cell 143, 606–616.

De Veylder, L., Beeckman, T., and Inzé, D. (2007). The ins and outs of the plant cell cycle. Nat. Rev. Mol. Cell Biol. 8, 655–665.

van Aken, O., de Clercq, I., Ivanova, A., Law, S.R., van Breusegem, F., Millar, A.H., and Whelan, J. (2016). Mitochondrial and chloroplast stress responses are modulated in distinct touch and chemical inhibition phases. Plant Physiol. 171, 2150–2165.

Street, I.H., Mathevo, D.E., Yamburkenko, M.V., Sorososhadze, A., John, R.T., Swaprup, N., Bennett, M.J., Kieber, J.J., and Schaller, G.E. (2016). Cytokinin acts through the auxin influx carrier AUX1 to regulate cell elongation in the root. Development (Cambridge) 143, 3982–3993.

Su, S.H., Gibbs, N.M., Janecewicz, A.L., and Masson, P.H. (2017). Molecular mechanisms of root gravitropism. Curr. Biol. 27, R964–R972.

Sundaravelpandian, K., Chandrika, N.P.T., Tsai, Y.H., and Schmidt, W. (2013). FPT1-controlled ROS balance is critical for multiple stages of root hair development in Arabidopsis. Plant Signaling Behav. 8, 1–15.

Suzuki, N., Miller, G., Morales, J., Shulaev, V., Torres, M.A., and Mittler, R. (2011). Respiratory burst oxidases: the engines of ROS signaling. Curr. Opin. Plant Biol. 14, 691–699.

Swarup, R., Wells, D.M., and Bennett, M.J. (2013). Root gravitropism. In Plant Roots: The Hidden Half, Fourth Edition, pp. 265–278.

De Veylder, L., Beeckman, T., and Inzé, D. (2007). The ins and outs of the plant cell cycle. Nat. Rev. Mol. Cell Biol. 8, 655–665.

van Aken, O., de Clercq, I., Ivanova, A., Law, S.R., van Breusegem, F., Millar, A.H., and Whelan, J. (2016). Mitochondrial and chloroplast stress responses are modulated in distinct touch and chemical inhibition phases. Plant Physiol. 171, 2150–2165.

Street, I.H., Mathevo, D.E., Yamburkenko, M.V., Sorososhadze, A., John, R.T., Swaprup, N., Bennett, M.J., Kieber, J.J., and Schaller, G.E. (2016). Cytokinin acts through the auxin influx carrier AUX1 to regulate cell elongation in the root. Development (Cambridge) 143, 3982–3993.

Su, S.H., Gibbs, N.M., Janecewicz, A.L., and Masson, P.H. (2017). Molecular mechanisms of root gravitropism. Curr. Biol. 27, R964–R972.
Vijayakumar, P., Datta, S., and Dolan, L. (2016). Root Hair Defective SIX-Like4 (RSL4) promotes root hair elongation by transcriptionally regulating the expression of genes required for cell growth. New Phytol. 212, 944–953.

Viola, I.L., Camoirano, A., and Gonzalez, D.H. (2016). Redox-dependent modulation of anthocyanin biosynthesis by the TCP transcription factor TCP15 during exposure to high light intensity conditions in Arabidopsis. Plant Physiol. 170, 74–85.

don Wangenheim, D., Goh, T., Dietrich, D., and Bennett, M.J. (2017). Plant biology: building barriers in roots. Curr. Biol. 27, R172–R174.

Wachsman, G., Sparks, E.E., and Benfey, P.N. (2015). Genes and networks regulating root anatomy and architecture. New Phytol. 208, 26–38.

Wany, A., Foyer, C.H., and Gupta, K.J. (2018). Nitrate, NO and ROS signaling in stem cell homeostasis. Trends Plant Sci. 23, 1041–1044.

Yi, K., Menand, B., Bell, E., and Dolan, L. (2010). A basic helix-loop-helix transcription factor controls cell growth and size in root hairs. Nat. Genet. 42, 264–267.

Zhao, Q., Nakashima, J., Chen, F., Yin, Y., Fu, C., Yun, J., Shao, H., Wang, X., Wang, Z.Y., and Dixon, R.A. (2013). LACCASE is necessary and nonredundant with PEROXIDASE for lignin polymerization during vascular development in Arabidopsis. Plant Cell 25, 3976–3987.

Zhao, Y., Luo, L., Xu, J., Xin, P., Guo, H., Wu, J., Bai, L., Wang, G., Chu, J., Zuo, J., et al. (2018). Malate transported from chloroplast to mitochondrion triggers production of ROS and PCD in Arabidopsis thaliana. Cell Res. 28, 448–461.

Zhao, P., Sokolov, L.N., Ye, J., Tang, C.Y., Shi, J., Zhen, Y., Lan, W., Hong, Z., Qi, J., Lu, G.H., et al. (2016). The LIKE SEX FOUR2 regulates root development by modulating reactive oxygen species homeostasis in Arabidopsis. Scientific Rep. 6, 1–11.

Zhou, X., Li, Q., Chen, X., Liu, J., Zhang, Q., Liu, Y., Liu, K., and Xu, J. (2011). The Arabidopsis RETaRDED ROOT GROWTH gene encodes a mitochondria-localized protein that is required for cell division in the root meristem. Plant Physiol. 157, 1793–1804.