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Genes of the most conserved WOX clade in plants affect root and flower development in *Arabidopsis*

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Abstract

Background: The Wuschel related homeobox (WOX) family proteins are key regulators implicated in the determination of cell fate in plants by preventing cell differentiation. A recent WOX phylogeny, based on WOX homeodomains, showed that all of the *Physcomitrella patens* and *Selaginella moellendorffii* WOX proteins clustered into a single orthologous group. We hypothesized that members of this group might preferentially share a significant part of their function in phylogenetically distant organisms. Hence, we first validated the limits of the WOX13 orthologous group (WOX13 OG) using the occurrence of other clade specific signatures and conserved intron insertion sites. Secondly, a functional analysis using expression data and mutants was undertaken.

Results: The WOX13 OG contained the most conserved plant WOX proteins including the only WOX detected in the highly proliferating basal unicellular and photosynthetic organism *Ostreococcus tauri*. A large expansion of the WOX family was observed after the separation of mosses from other land plants and before monocots and dicots have arisen. In *Arabidopsis thaliana*, *AtWOX13* was dynamically expressed during primary and lateral root initiation and development, in gynoecium and during embryo development. *AtWOX13* appeared to affect the floral transition. An intriguing clade, represented by the functional *AtWOX14* gene inside the WOX13 OG, was only found in the Brassicaceae. Compared to *AtWOX13*, the gene expression profile of *AtWOX14* was restricted to the early stages of lateral root formation and specific to developing anthers. A mutational insertion upstream of the *AtWOX14* homeodomain sequence led to abnormal root development, a delay in the floral transition and premature anther differentiation.

Conclusion: Our data provide evidence in favor of the WOX13 OG as the clade containing the most conserved WOX genes and established a functional link to organ initiation and development in *Arabidopsis*, most likely by preventing premature differentiation. The future use of *Ostreococcus tauri* and *Physcomitrella patens* as biological models should allow us to obtain a better insight into the functional importance of WOX13 OG genes.

Background

Homeodomain (HD) containing transcription factors are key regulators implicated in the determination of cell fate and cell differentiation in both plants and animals. In Angiosperms, a gene called *WUSCHEL* (*WUS*) has been isolated from many different species. *WUS* was the first identified member of the *Wuschel*-related homeobox (*WOX*) subfamily [1,2] that is only found in plants.

The specific expression of the *WOX* genes in different plant organs and cell types [2-7] suggested an important role for them during organogenesis. The *WUS* gene is expressed in a restricted region of the meristem called the organizing centre located below the stem cells of the shoot apical meristem (SAM) and functions non-cell autonomously to control the stem cell fate [8]. Interestingly Gallois et al [3] showed recently that ectopic expression of *WUS* in the root establishes shoot stem cells and leaf development. Other members of the *WOX* family might also prevent premature cell differentiation in developing organs or tissues. Recently, *WOX5* was shown to be involved in stem cell maintenance signaling in the root [9]. Wu et al [10] have shown that *WOX9* (*Stimpy*/*STIP*) prevents premature cell differentiation during organ growth, in addition to contributing positively to *WUS* expression. This work also revealed that *stip* mutants can be rescued by sucrose, a modulator of cell proliferation through *cyclin D* induction [11,12]. Two other *WOX* genes, named Narrow Sheath 1 and 2, that are maize orthologues of *Arabidopsis thaliana* *AtWOX6* were shown to be involved in the recruitment of the lateral founder cells within the SAM prior to the primordia development [13]. Last, Haecker et al [1] analyzed the expression dynamics of the *WOX* genes during *A. thaliana* embryo development and showed that members of the *WOX* family mark cell fate decisions during embryonic patterning. Taken together, these results suggest that not only *WUS* but several other *WOX* genes play a role in regulating cell division and preventing cell differentiation.

Characterized genomes of Angiosperms contain more than 10 different *WOX*s and phylogenetic studies suggest that the *WOX* family existed in the last common ancestor of monocots and core eudicots [14,15]. A recent *WOX* phylogeny based on *WOX* HDs from different Angiosperms, the moss *Physcomitrella patens* and the lycophyte *Selaginella moellendorffii* showed that all of the *P. patens* and eight of the *S. moellendorffii* *WOX* genes clustered in the same clade [16]. Indeed, the phylogenetic analyses of a transcription factor family like *WOX* suffer from some limitations due to the relatively short length of the HD, which is the only conserved sequence between the *WOX* protein family members, and the relatively few land plant species with a completely characterized proteome. As a consequence, the patterns of phylogenetic trees

of large families of plant transcription factors are often weakly supported by statistics and may be affected by long branch attraction [17]. Nevertheless, clade specific gene features and protein domains were observed in transcription factors [18,19]. Thus, in this study, we used clade specific signatures beyond the HD and clade specific conservation of intron insertion sites, to provide additional data in support of the orthology links predicted by the *WOX* phylogeny. In order to carry out the latter analyses, complete genome sequences with accurate structural annotations are essential. Our results strongly support the existence of a *WOX13* orthologous group (*WOX13* OG) that, contrary to the two other *WOX* OGs, contains genes from many different members of the plant kingdom, including basal species, and a branch apparently specific to Brassicaceae. To obtain information on the function of the genes belonging to the *WOX13* clade, the cell specificity of *AtWOX13* and *AtWOX14* expression was established experimentally and phenotypes resulting from mutations within these two genes were characterized in the plant model *A. thaliana*.

Results

A *WOX* distance tree from model genomes

We carefully screened several completely sequenced eukaryote genomes for *WOX* genes, *i.e.*, two Angiosperms, *A. thaliana* and *O. sativa*, three algae, *O. tauri*, *O. lucimarinus* and *Chlamydomonas reinhardtii* and two yeasts, *Saccharomyces cerevisiae* and *Schizosaccharomyces pombe*. No *WOX* genes were identified outside of the plant kingdom and *Ostreococcus* revealed only one *WOX* gene per genome. Moreover, using the HMMsearch tool [20], we found only one HD gene in *C. reinhardtii*, 2 in *S. pombe*, 4 in *O. tauri* and 8 in *S. cerevisiae* compared to the 92 HD genes in *A. thaliana*.

Distance and phylogenetic trees were computed from the HD sequences of the *WOX* proteins from the five plant model genomes, *O. tauri*, *P. patens*, *S. moellendorffii*, *A. thaliana* and *O. sativa* (see method section). The clustering of the *WOX* proteins in 3 different OGs were consistent using either the Neighbor-Joining (Figure 1) or the Maximum likelihood (see additional file 1: Figure 1) methods and different outgroups. The topology of the trees clearly showed three different OGs namely the *WOX1*, *WOX8* and *WOX13* MOGs (for Model plant OG). Each OG contained rice and *Arabidopsis* proteins, but the single *WOX* protein of *O. tauri*, the three *P. patens* and the three *S. Moellendorffii* *WOX* proteins were all found in the same group, the *WOX13* MOG. The *WOX1* MOG contained *AtWOX1-7* and *AtWUS*, the *WOX8* OG associated *AtWOX8*, 9, 11 and 12 and the *WOX13* OG included *AtWOX10*, 13 and 14. Thus, the previous observation of the presence of the *P. patens* proteins only in the *WOX13* OG group was extended to *O. tauri* and *S. moellendorffii*. However, the

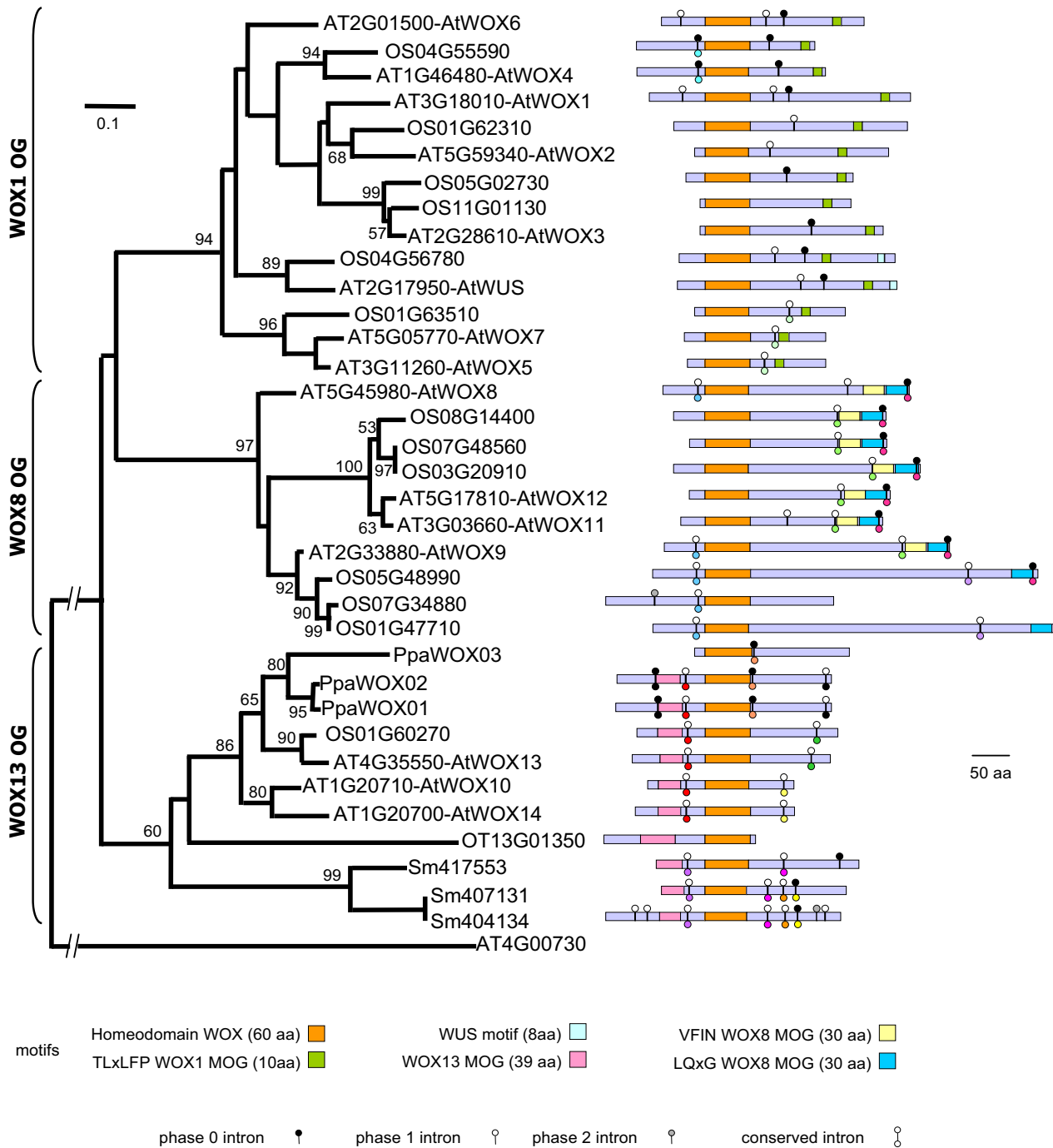


Figure 1
WOX genes from model plant genomes. The neighbor-joining tree of the WOX proteins from the 5 completely sequenced genomes of the green lineage: *Arabidopsis thaliana* (AT prefix), *Oryza sativa* (OS prefix), *Physcomitrella patens* (Ppa prefix), *Ostreococcus tauri* (OT prefix) and *Selaginella moellendorffii* (Sm prefix). The tree topology was inferred from Jones-Taylor-Thornton matrix distances. The significance of each node was tested using 1000 bootstrap replicates. Only bootstrap values above 50% are shown. The scale, at the top left of the tree, indicates 0.1 substitutions per site. Different structural features are represented by different signs and colors as described at the bottom of the figure.

introduction of OtWOX and the SmWOXs did not change the main branching of the trees nor significantly modified the bootstrap values at the internal node of the tree. The 14 available WOX sequences from different Gymnosperms were distributed between the three OGs (data not shown). Furthermore, the trees we obtained were not consistent with the phylogenetic tree of plant species as it is now known [21]. Thus the WOX trees we produced suggest probably several rounds of gene deletions in *O. tauri* and *P. patens* in order to explain the branching of the WOX13 OG. The finding that all the WOX sequences of *O. tauri* and *P. patens*, as well as those of *S. moellendorffii* were clustered in the WOX13 OG suggested to us that *AtWOX10*, *13*, *14* might represent the most interesting genes to study the functional diversification of the WOX genes although they are the WOX genes for which we have the least functional information (reviewed in [22]). This prompted us to look at the specificity of the WOX13 OG with regards to gene structure and conserved amino acid signatures other than the HD. Conservation in gene structure is an indication of a common ancestor while conserved amino acid signatures are linked to functions such as protein-protein interactions or regulatory sites.

WOX gene structures

The intron-exon organization of WOX genes from the model genomes showed different gene structures (see additional file 2: Figure 1). Alignments supporting the conservation of intron insertion sites are given in additional file 2: Figure 2. The organization of the gene structures along the phylogeny tree did not reveal any contradiction between gene structures and tree pattern. Intron site conservations were exclusively observed within each of the three OGs and never between them. In this context, it was interesting to observe that the insertion site of the first intron in *AtWOX13* is conserved in almost all the genes (from moss to Angiosperms) belonging to the WOX13 OG. The only exception was *PpaWOX03*, a possible pseudogene (see below). This observation cannot be extended to *O. tauri* as its genes do not have introns. *S. moellendorffii* genes have introns and the three full-length SmWOXs have one conserved intron insertion site downstream from the WOX13 motif. The latter intron of SmWOXs is in a similar position (relative to the coding region) to the first highly conserved intron of WOX13 OG genes in moss and Angiosperms. Nevertheless, there is no trace of amino acid sequence conservation in the corresponding protein region between *S. moellendorffii* and the other genes in the WOX13 OG (see additional file 3: Table 3). When taken together, our analyses of intron insertion sites support the boundaries of the WOX13 OG as defined by phylogeny. In addition to the highly conserved intron upstream of the HD coding sequence found in the WOX13 OG, *AtWOX10* and *AtWOX14* share another conserved intron insertion site downstream of the HD while

OS01G60270 and *AtWOX13* share a different intron insertion site in the 3' region of the gene. The tree pattern implies first a loss of the orthologues of *AtWOX10* and *14* in *O. sativa* and in *P. patens* and secondly that *AtWOX13* and *OS01G60270* are orthologous. Interestingly, the two latter genes share a specific intron insertion site.

Specific motifs in the WOX proteins

A search for specific motifs other than the HD used to build the distance and phylogeny trees led us to identify HMMs that were specific to each clade and located either within the N or the C terminal regions of the different proteins. These HMMs are illustrated by sequence logos (see additional file 2: Figure 3) and by alignments (see additional file 2: Figure 4). Only proteins of the WOX1 clade have a small motif of 10 amino acids (*i.e.*, the TLxLFP motif) in the C-terminal region, but at relatively variable distances from the HD. TLxLFP has previously been described as a characteristic motif of WUS proteins [1,23], but we show here that it is common to the entire WOX1 clade, *i.e.*, it is observed not only in WUS, but also in the *AtWOX1-7* proteins. However, the WUS proteins of *A. thaliana* (AT2G17950-*AtWUS*) and *O. sativa* (OS04G56780) share specific features not observed in any of the other proteins of the WOX1 MOG. Indeed, they display 1) the LELxL WUS motif [1] known as EAR-like or ERF-like motifs that are located in the outmost C-terminal region, similar to a motif found in potent transcriptional repressors in plants *i.e.*, Superman [24] and Aux/IAA [25]; 2) the same intron exon structure in the coding regions of each gene; and 3) an additional amino acid at a conserved position in the homeodomain.

Nearly all proteins of the WOX8 MOG contain two contiguous motifs of 30 amino acids situated near to their C-terminus (*i.e.*, VFIN-WOX8 and LQxG-WOX8 MOG motifs). The latter motifs are surrounded by two conserved introns. Remarkably, these introns are also found in OS05G48990 in which the VFIN-WOX8 MOG motif is replaced by a non-conserved sequence of a similar length.

All the proteins of the WOX13 MOG, except for *PpaWOX03*, share a 39 amino acid motif that is found upstream from the HD and named the WOX13 MOG motif. This observation provides further support for the validity of the WOX13 OG as defined by phylogeny and to the hypothesis of a specific function of the genes clustered within. For all of these reasons, it was decided to investigate further the WOX13 clade.

WOX13 OG distance tree

Thirty three full length WOX proteins were used to reconstruct the distance tree of the WOX13 clade (Figure 2), using OtWOX as the outgroup. The neighbor-joining tree showed two main branches. The first branch contained all

of the Gymnosperm and the moss WOXs. The second branch contained AtWOX13 together with many Angiosperm sequences. This branch was further separated into two branches, one contained AtWOX13 and many proteins from different Angiosperms while the second contained AtWOX10, AtWOX14 and a *Brassica rapa* sequence.

The preferential presence of amino acid motifs fully supported the separation of the different gene groups within the trees. Obviously, all proteins of the WOX13 clade displayed the WOX HD and the WOX13 motif. Moreover, we also identified specific motifs for the AtWOX10, 14 and for the AtWOX13 branches (see additional file 2: Figure 3). All Angiosperm proteins located inside the WOX13 OG shared two motifs, the YxDpl-WOX13 motif located between the WOX13 MOG motif and the HD, and the ESExE-WOX13 motif just downstream from the HD. The monocots of the AtWOX13 branch also share a specific motif called the monocot-WOX13 motif that was detected also in two other partial proteins of two monocotyledon species.

Proteins of the AtWOX10, 14 branch shared two specific motifs, the YFdPM-WOX10 and QAdDaAVTT-WOX10 motifs. They were located to the same region of the protein as the YxDpl-WOX13 and ESExE-WOX13 motifs, respectively. Hence, the similarity between the YFdPM-WOX10 OG and YxDpl-WOX13 motifs suggested a common origin. Partial protein sequences from *Brassica oleracea* and *Raphanus raphanistrum subsp. raphanistrum* (protein sequence derived from the two ESTs, gb|EX762111| and gb|EX766460|) confirmed the specificity of the motifs to Brassicaceae. Thus, the AtWOX14 containing branch appears to be Brassicaceae specific since: 1) all the sequences containing the two AtWOX10, 14 motifs are from Brassicaceae and 2) no species other than those belonging to the Brassicaceae displayed any of these two motifs.

The Gymnosperm and moss branch contained proteins that shared the ESExE-WOX13 OG motif with the Angiosperm WOX13 proteins, located at the same position relative to the HD. They also shared two other motifs specific to this branch: 1) the LxxGQ-gymnosperm/moss-WOX OG motif that was at the same position as the YxDpl-WOX13 OG or YFdPM-WOX10 OG motifs and 2) a supplementary motif in the C-terminal region of the proteins.

Functional WOX13 OG genes in the three model plants

We investigated the expression profiles of the WOX13 OG genes within 3 plant model species namely *O. tauri*, *P. patens* and *A. thaliana* because of their position within the green lineage. *O. tauri* is a highly proliferating unicellular

alga that has no known differentiated form. In addition to the WOX gene, we also identified a single copy of a *KNOX-like* gene (*Ot04g04460*) within the *O. tauri* genome. In Angiosperms, members of the class-1 KNOX subfamily act to prevent cell differentiation within the SAM [26]. Interestingly, the OtKNOX protein showed the characteristics of both the class-1 and class-2 KNOX conserved domains, suggesting that *O. tauri* had diverged from a common ancestor before the diversification of the KNOX gene into 2 classes.

In order to monitor the expression of WOX and KNOX genes during the progression of the cell cycle, we partially light-synchronized a cell suspension of *O. tauri* as described before in [27]. RT-PCR and quantitative PCR analyses of *Cyclin B* and *Histone 4* genes showed a peak of expression at 14 h, confirming the overlapping of the G2/M and S subpopulations (Figure 3A and see additional file 4: Figure 1). The *OtRH4* (*Ot07g03220*), and *OtRH21* (*Ot10g01950*) genes, respectively the orthologues of the Arabidopsis *DEAD-box RNA-Helicase 4* and 21 [28] were used as controls of constitutively expressed genes. They did not show any significant variation, and they were not cell cycle regulated (Figure 3A). More importantly, the WOX and the KNOX genes were found to be constitutively expressed and not cell cycle regulated (Figure 3A), with a higher expression level for WOX compared to KNOX (see additional file 4: Figure 1).

In *P. patens*, the initial gametophytic phase consists of an array of filamentous tubes that are formed after cell initials are produced from the main filament. Further development proceeds by the formation of buds which form the initial meristem from which originates the leafy adult gametophyte [29]. These processes rely on a tight control of cellular differentiation throughout the gametophyte development. An analysis of WOX expression in *P. patens* was carried out on 4-, 10- and 14-day-old plants corresponding respectively to the protonema, the bud and the gametophore stages under our growth conditions (data not shown). RT-PCR analyses showed that *PpaWOX01* and *PpaWOX02* were expressed throughout gametophyte development whereas *PpaWOX03* transcripts were not detected (Figure 3B), thus supporting the hypothesis that *PpaWOX3* could be a pseudogene.

An exhaustive data mining analysis of *A. thaliana* expression databases [30-33] indicated that, both *AtWOX13* and *AtWOX14* transcripts are present at different developmental stages *i. e.*, embryo development, plantlet stage, root formation, bolting stage, in flowers and in response to stress (see discussion). However, no *AtWOX10* transcripts were present in any of the databases analyzed. In addition, RT-PCR analyses of seedlings, even with a high PCR cycle number, did not allow us to detect the *AtWOX10* mRNA

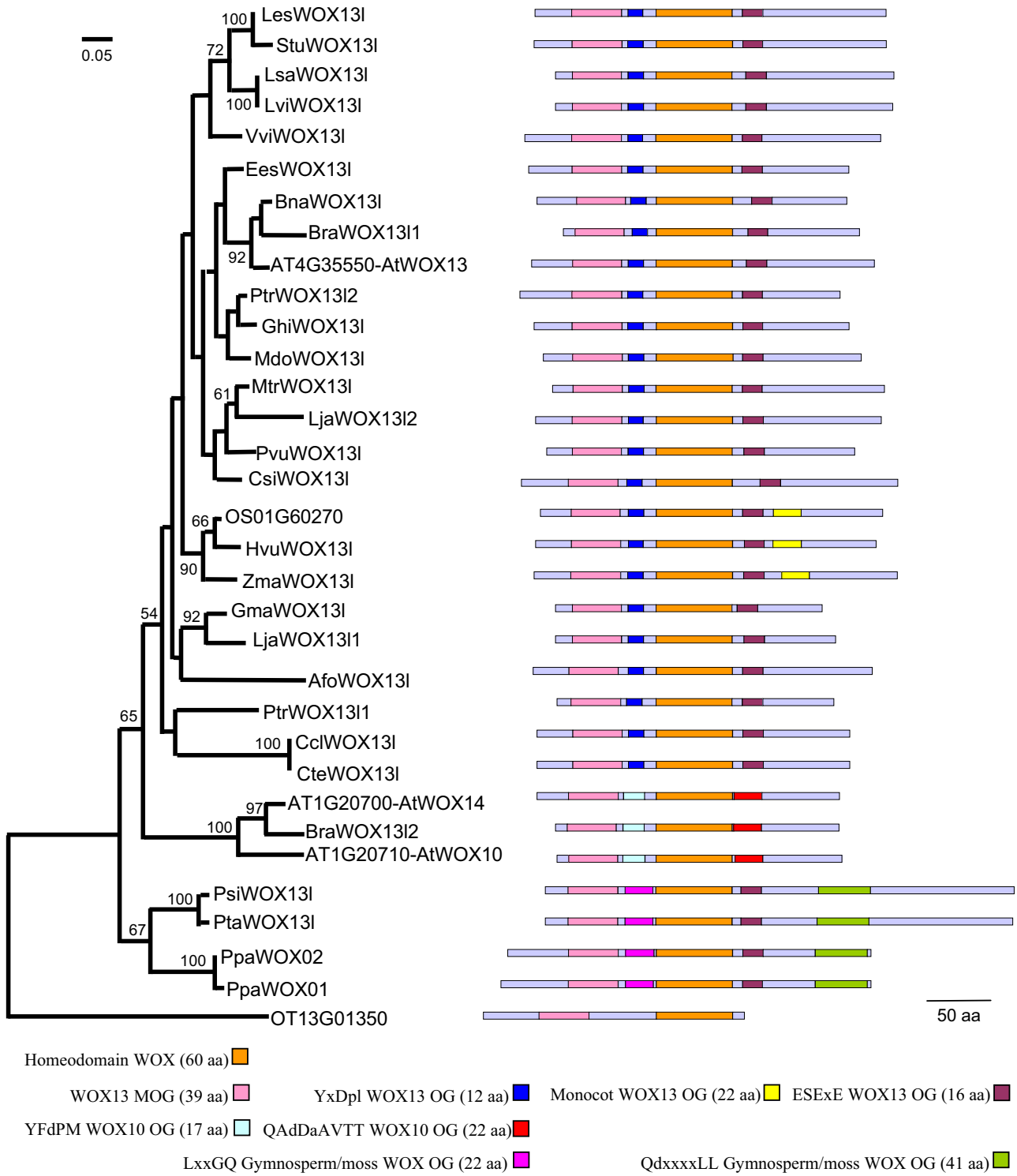


Figure 2 (see legend on next page)

Figure 2 (see previous page)

The WOX13 Orthology Group. The different elements of the figure are described from the left to the right. 1) The neighbor-joining tree of the WOX proteins from the WOX13 Orthology Group (see material and methods). The tree topology, the significance of the nodes, and the bootstrap values displayed are the same as in Figure 1. The scale at the top left corner indicates 0.05 substitutions per site. 2) The motif composition of each protein is denoted by colored boxes on a bar representing the length of the protein and aligned on the WOX homeodomain (displayed in orange). Species names, in alphabetic order, are: Afo: *Aquilegia formosa*; AT: *Arabidopsis thaliana*; Bna: *Brassica napus*; Bra: *Brassica rapa*; Ccl: *Citrus clementina*; Csi: *Citrus sinensis*; Cte: *Citrus temple*; Ees: *Euphorbia esula*; Ghi: *Gossypium hirsutum*; Gma: *Glycine max*; Gra: *Gossypium raimondii*; Han: *Helianthus annuus*; Hvu: *Hordeum vulgare*; Les: *Lycopersicon esculentum*; Lja: *Lotus japonicus*; Lsa: *Lactuca sativa*; Lvi: *Lactuca virosa*; Mdo: *Malus domestica*; Mtr: *Medicago truncatula*; OS: *Oryza sativa*; OT: *Ostreococcus tauri*; Ppa: *Physcomitrella patens*; Psi: *Picea sitchensis*; Pta: *Pinus taeda*; Ptr: *Populus tremula* × *Populus tremuloides*; Pvu: *Phaseolus vulgaris*; Stu: *Solanum tuberosum*; Vvi: *Vitis vinifera*; Zma: *Zea mays*. The color code of the different motifs is described at the bottom of the figure.

(Figure 3C). These results suggested that the recently duplicated *AtWOX10* gene, found only in the *A. thaliana* genome, might be a pseudogene. Based on these observations, this gene was not further characterized. We conclude that in the three plant model genomes studied at least one copy of the WOX13OG genes is expressed during development.

AtWOX13 shows a dynamic expression profile during *A. thaliana* development

Compared to other WOX proteins, the function of the *AtWOX13* and 14 proteins have received little attention to date (see review [22]). In order to carry out a more detailed analysis of the WOX13 OG genes in *A. thaliana*, we cloned the gene coding for the *GUS* reporter under the control of the *AtWOX13* and *AtWOX14* upstream regulatory sequences. Based on our data mining analyses (see above), we first analyzed the *GUS* expression during root development. *AtWOX13* promoter activity was detected during lateral root formation (Figure 4A, C, F and 4I). To validate the specificity of these data, *in situ* hybridization experiments were performed using respectively *AtWOX13* (Figure 4B, D, G and 4K), *GUS* (Figure 4J) and *CUC2* as probes (Figure 4E and 4H). The signal obtained using the *AtWOX13* and *GUS* probes matched perfectly with the above *GUS* staining, whereas the *CUC2* probe, as expected, did not give any specific signal. Hence, the *GUS* staining obtained reflects stable tissue specific mRNA expression and revealed that the *AtWOX13* promoter sequence, used to drive the reporter gene *GUS* expression was sufficient to mimic the tissue specific localization of the endogenous *AtWOX13* mRNA.

AtWOX13 expression was detected at the early stage of the lateral root (LR) formation of 10-days-old seedlings and it persisted during LR emergence (Figure 4A–D). The *GUS* staining was progressively restrained to the distal meristem (Figure 4E, and 4G) and it strongly accumulated in both the columella and the adjacent lateral root cap cells at a later developmental stage (Figure 4I–K). In mature LR

or primary roots of 10-day-old seedlings no staining was detected within the root tip.

The *AtWOX13* expression pattern was also investigated using the *AtWOX13::GUS* lines, during plant development. In 5-day-old seedlings, the promoter activity was strong in both cotyledon and root vasculatures, the pericycle and in the stem cells of the primary root meristem (Figure 4M and 4N). No *GUS* staining was detected in the division zone of the root. In the vegetative apex of 10 day-old seedlings, a strong staining was observed in the leaf primordia (Figure 4O). In the flower, the *AtWOX13* promoter was active in the vasculature, the stigma (Figure 4P) and in the gynoecium at the flower stage 13/14 (Figure 4Q, and 4R). At this developmental stage, when fertilization was ongoing and pollen grains were germinating at the tip of the gynoecium, ovules stained blue (Figure 4Q). Later on, the young embryos remained stained (Figure 4R), but no staining was observed in mature siliques indicating that the *AtWOX13* promoter was no longer active (Figure 4S), as observed before in all the other mature organs. During germination, soon after seed imbibition, *GUS* staining was also seen within the vasculature and the root tip, as observed in young seedlings (data not shown). Hence, *AtWOX13* expression was restricted to differentiating organs and therefore probably regulated by developmental signals.

AtWOX14 promoter specificity differs from that of AtWOX13

Compared to the *AtWOX13* expression patterns, the activity of *AtWOX14* was more restrictive during lateral root formation. As for *AtWOX13*, the *AtWOX14* promoter activity, revealed by *GUS* staining, was detected in the LR primordia, but decreased progressively as the LR emerged becoming restricted *in fine* to the vasculature up to the LR junction (Figure 5A–C). The *AtWOX13::GUS* gene continued to be expressed for a longer time (compare Figure 5B with Figure 4C). In 5-day-old seedlings, *AtWOX14* driven *GUS* activity was only found in the vascular system and in the pericycle of the primary root (Figure 5D). In the LR of

14-day-old seedlings (Figure 5E), GUS staining was again observed in the vasculature and the pericycle with *AtWOX14* expression being progressively induced in the pericycle and in the vasculature as the LR developed. Compared to the activity of the *AtWOX13* promoter,

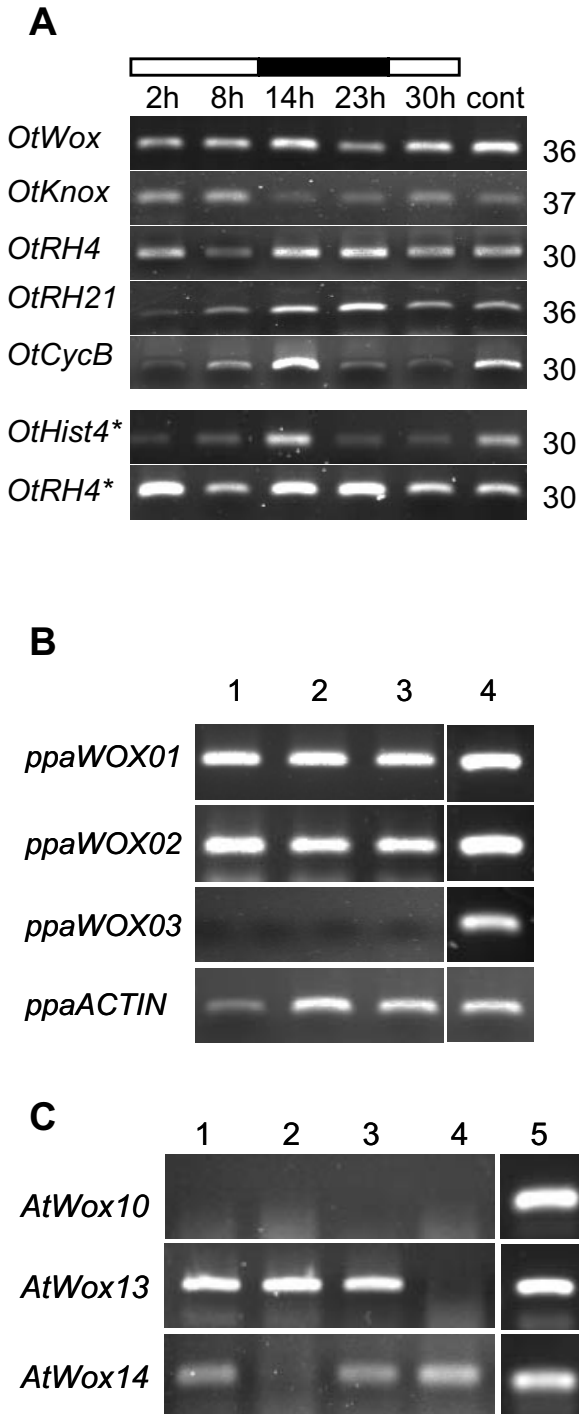


Figure 3

Figure 3

WOX13 OG gene expression in *O. tauri*, *P. patens* and *A. thaliana*.

(A) RT-PCR analysis of WOX gene expression in *Ostreococcus tauri*. 12 h light/dark synchronized or continuous light (cont) cell cultures of 10-day-old were used. Gene name (left side), number of PCR cycles done (right side), time of sampling (top side), light (blank bar) and dark (black bar) periods. The genes *OtRH4* and *OtRH21* were used as constitutive expression controls. RT were done using oligodT primers except in (*) where random primers were used **(B)** RT-PCR analysis of WOX gene expression during *Physcomitrella patens* development: (1) 4-day-old, (2) 10-day-old, (3) 14-day-old, (4) genomic DNA control. **(C)** RT-PCR analysis of the expression of WOX13 OG genes carried out on 7-day old seedlings of different *Arabidopsis thaliana* lines: (1) Wild type No ecotype, (2) the *pst13645 AtWOX14* mutant line, (3) Wild type WS ecotype, (4) the 193G10 *AtWOX13* mutant line, (5) Genomic DNA control.

AtWOX14::GUS expression stopped at a longer distance further up from the root tip (Figure 5E, black arrow), exactly at the position where the youngest LR primordia were initiated (see arrow in Figure 5F).

During flower development, *AtWOX14::GUS* expression was detected in the stamen at floral stage 11/12 (Figure 5G-I), when mitotic divisions of microspores are known to occur [34]. The staining decreased as the stamen matured, showing that the function of WOX14 is linked to the early stages of organ and tissue development. Hence, as for *AtWOX13*, *AtWOX14* promoter activity also seems to be regulated by developmental signals. The gene expression profile reported in the gene atlas [35] and the tissues and organs affected by the mutation of *AtWOX14* (see below) agree with the expression profile described above.

An *AtWOX13* gene truncated downstream of the homeodomain neither alters root nor flower development

The only available *wox13* mutant line, identified in GABI-Kat [36] and numbered 193G10, showed a T-DNA insertion upstream of the sequence coding for the characteristic WFQN motif located at the end of the WOX homeodomain (Figure 6A and see additional file 4: Figure 2A). RT-PCR experiments carried out using polyA⁺-RNA, prepared from the *wox13* 193G10 line, and primers matching each side of the insertion site, did not yield any PCR product (see Figure 3C). Because of the position of the insertion at the end of the WOX homeodomain, we also carried out an expression analysis using another couple of primers matching the 5'UTR of the *AtWOX13* mRNA and the left border of the T-DNA. The latter experiment led to the amplification of a 500 bp PCR product, the sequence of which coded for a putative chimeric protein of 148 resi-

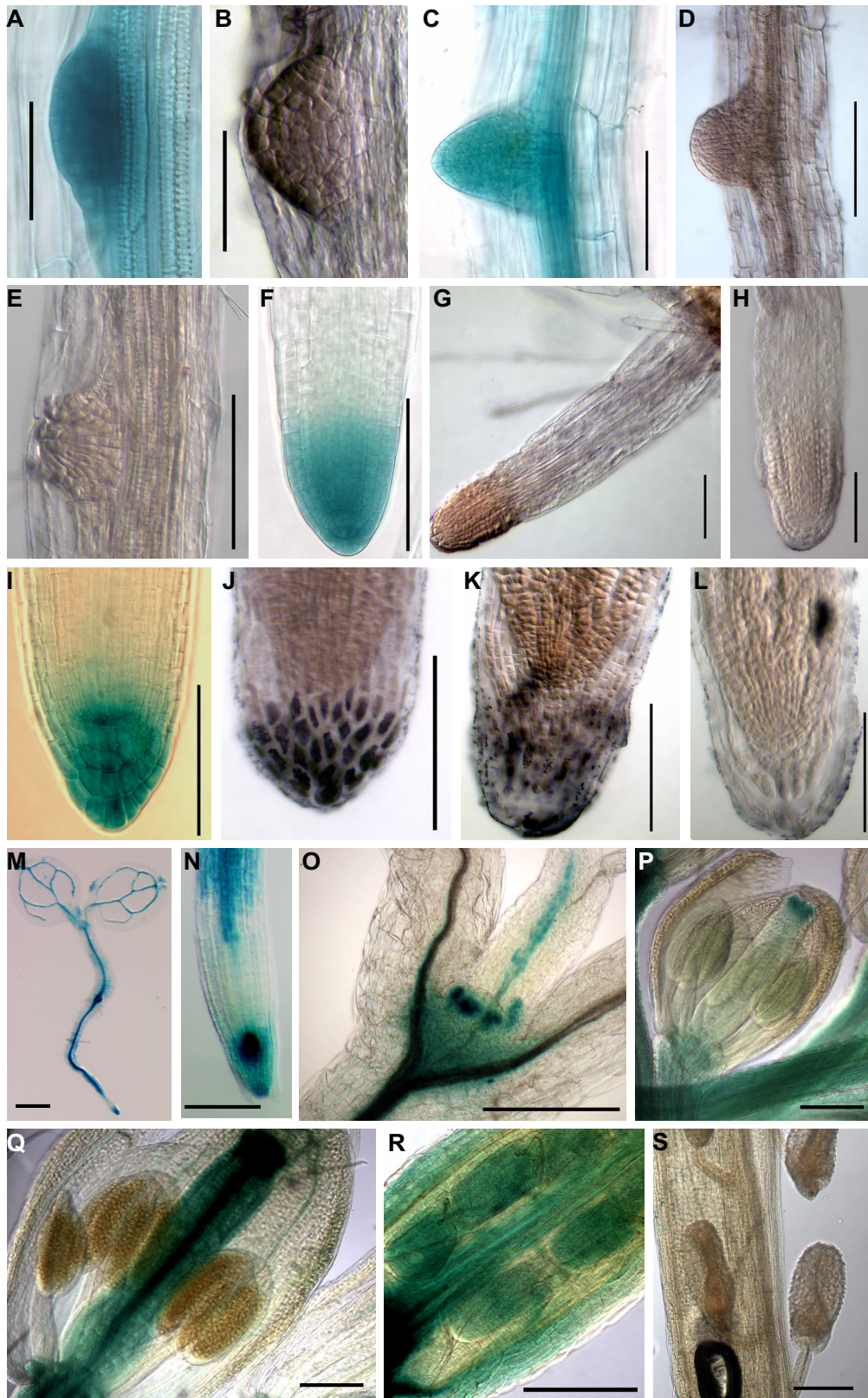


Figure 4 (see legend on next page)

Figure 4 (see previous page)

AtWOX13 gene expression during plant development. (A to L) Expression profile during the initiation and development of lateral roots obtained with 10-day-old *AtWOX13::GUS* seedlings. Independent lines were tested. (A, C, F and I) GUS staining. (B, D, and K) *In situ* hybridization with an *AtWOX13* antisense probe. (J) *In situ* hybridization with a GUS antisense probe used as a positive control. (E, H and L) *In situ* hybridization with a *CUC2* antisense probe used as a negative control. **(M to S)** Expression profile at different plant developmental stages with independent *AtWOX13::GUS* lines. (M) 5-day-old seedling with staining in the leaf, the hypocotyl, the root vasculature, the pericycle and the primary root tip. (N) Magnification of a 5-day-old seedling primary root tip showing high staining in the stem cell of the proximal meristem. (O) Expression within the shoot apex of 10-day-old seedlings. Note the staining in the leaf primordia. **(P to S)** Expression within the flower at different developmental stages where staining is seen in the stigmata and the vasculature (P) in the gynoecium (Q) at the floral stage 12–13 where the ovules (Q) are stained. (R) Young embryo, (S) Mature silique. Scale bars = 100 μm except for A, B, R and S (= 90 μm) and M (= 500 μm).

dues composed of the 143 amino acids of the *AtWOX13* N-terminal region plus an additional 5 amino acids specified by the T-DNA sequence.

Based on the expression profiles obtained using the GUS reporter gene constructs, an analysis of the primary root growth of the mutant lines was carried out after seed germination. Surprisingly, the above *WOX13* mutation neither affected primary root growth (Figure 6C), nor vegetative and flower development. The only phenotype appeared to be a much earlier floral transition (Figure 6E).

A *WOX14* knock-out mutant modifies lateral root and stamen development of *A. thaliana*

Conversely to the *wox13* mutant, a *wox14* insertion mutant line (*i.e.*, pst13645) with the entire *WOX* homeo-domain deleted (Figure 6B and see additional file 4: Figure 2B) displayed developmental defects in all the organs where GUS staining was observed. The primary root growth was retarded (Figure 6D) and after flowering, the pst13645 insertion line showed partial sterility (Figure 6M), with aborted and shorter siliques compared to the wild type (Figure 6N). The floral transition was also altered, however the phenotype observed was that of a late flowering mutant (Figure 6F).

The *wox14* mutant was further characterized during plant development. Whereas in the wild type Nossen a fully developed network of LR was observed (Figure 6G, H), the LR formation in *wox14* was strongly inhibited (Figure 6I, J). The pst13645 line displayed extra adventitious roots (see Figure 6K, L) whereas the wild type plants developed only one or two. At the reproductive stage, observations after tissue clearing showed that the shorter siliques of *wox14* contained many ovules that did not develop into seeds (Figure 6P), compared to the wild type (Figure 6O). Since the *AtWOX14* promoter was only active in the developing stamen, pollen grain viability was checked using Alexander staining. As shown in Figure 6Q and 6R, the anthers of both the wild type and the *wox14* mutant contained viable pollen grains when compared to the sterile

dmc1 RNAi line [37]. However, stamen development was severely affected in the *wox14* mutant. In open flowers of the wild type plants, the stamens were longer than the gynoecium allowing complete fertilization of the ovule (Figure 6T). In contrast, in the mutant line, the stamen did not fully develop and remained shorter than the gynoecium thereby preventing efficient fertilization and causing ovule abortion. In some *wox14* flowers, the phenotype was even more severe, with aborted stamens that developed to only half the size of the gynoecium (Figure 6V). These data suggested that *AtWOX14* prevents premature organ and tissue differentiation at the floral stage.

Discussion

To improve previously described and not completely congruent *WOX* phylogenies [14,16,38] we performed multi alignments on protein sequences (60–61 amino acids) longer than the usually defined HD (41–42 amino acids). Furthermore, we used gene or protein features other than the HD sequence to support the three OGs exhibited by our phylogenetic trees. Even if the *WOX* gene phylogeny remains difficult to reconcile with species phylogeny, we are confident about the gene distribution between the three main OGs. Only one *WOX* gene was detected in the basal unicellular and photosynthetic organism *O. tauri* and three *WOX* genes were identified in moss *P. patens*, including at least two functional paralogues. The *WOX13* OG is the clade containing the most conserved *WOX* genes as it is the only clade that contains the *O. tauri*, the *P. patens* and probably the eight *S. moellendorffii* genes. It also showed a recent and specific expansion in Brassicaceae. These observations point out the interest in the study of the *WOX13* OG with respect to its biological specificity and gene family evolution.

***WOX14* orthologues are only observed in Brassicaceae species**

We detected *AtWOX13* and *AtWOX14* mRNAs at different stages of plant development, in plantlets, roots, flowers and developing embryos. These expression profiles were in accordance with those described in a compendium of

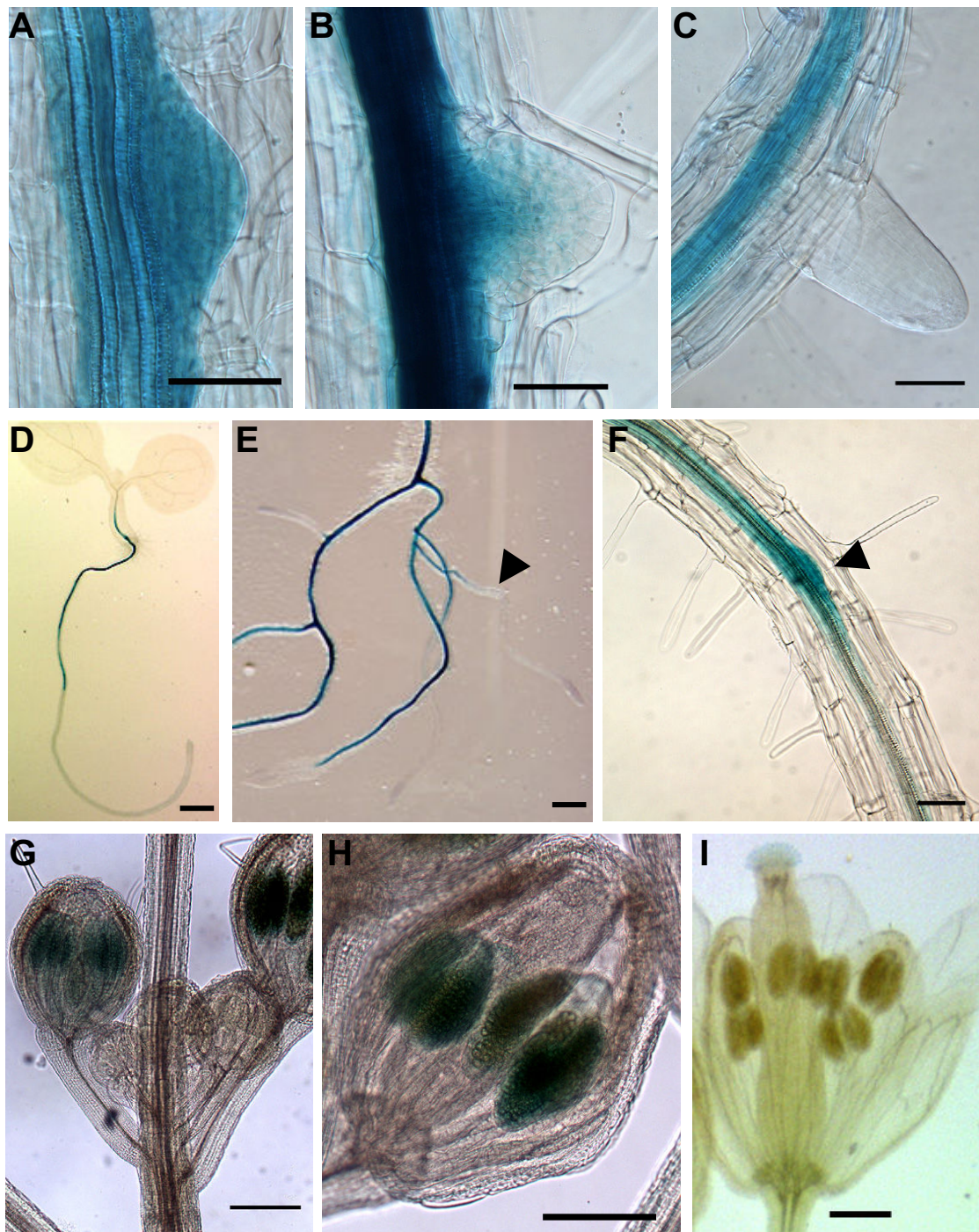


Figure 5
AtWOX14 gene expression during plant development. (A to C) Expression profile during the initiation and development of lateral roots obtained with 14-day-old *AtWOX14::GUS* seedlings. Independent lines were tested. (D) 5-day-old seedling showing staining in the lower part of the hypocotyl and the upper part of the primary root vasculature. (E) Staining profile in primary and lateral roots of 14-day-old seedlings with expression in the upper root vasculature and pericycle. (F) Magnification at the edge of the GUS coloration showing the presence of the youngest LR initial (black arrowhead). (G to I) Expression within the flower showing strong staining in the anther only at floral stage II–12 (G and H) before opening. Note the absence of staining in the mature anther (I). Scale bars = 100 μm except for C (= 200 μm) and D, E (= 500 μm).

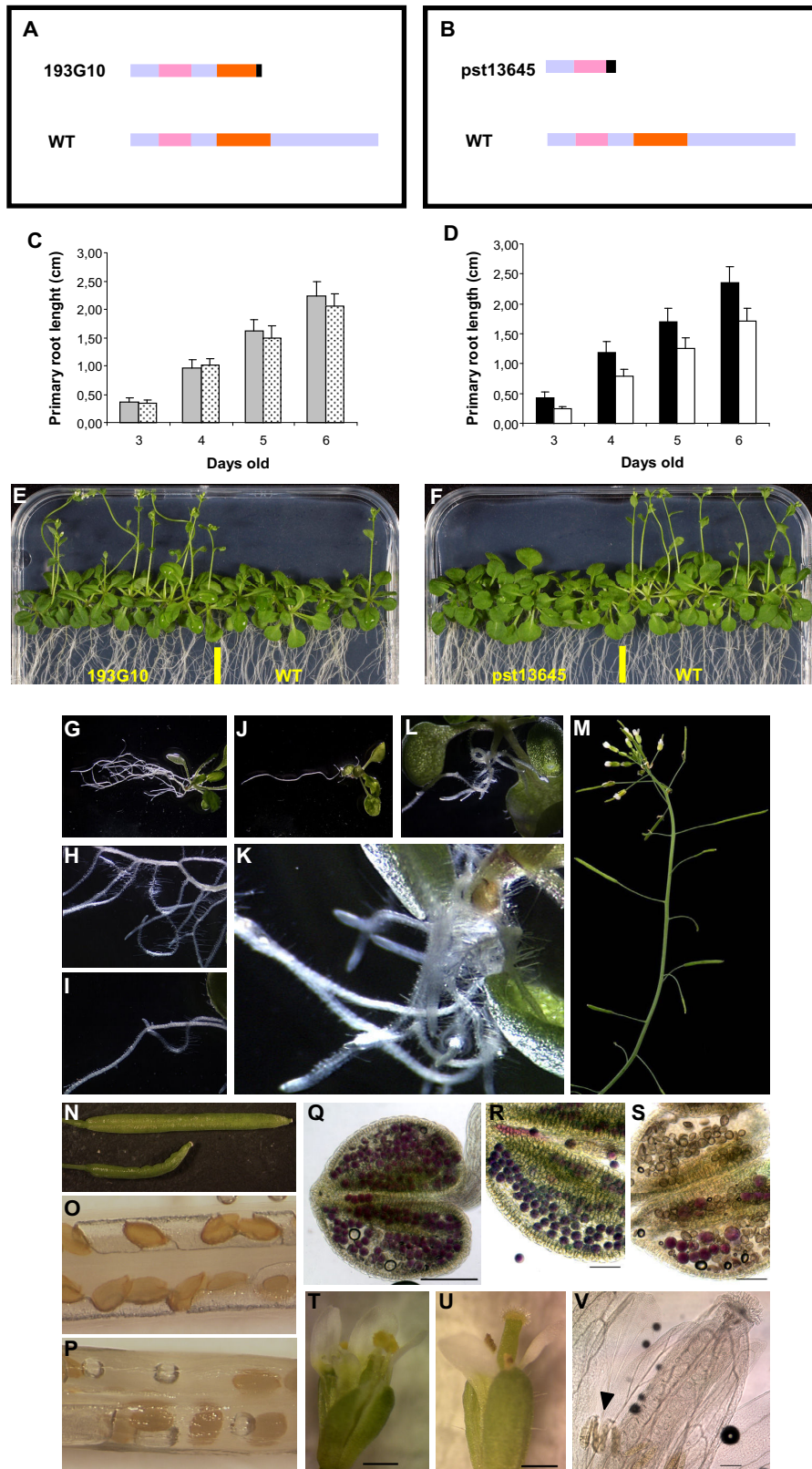


Figure 6 (see legend on next page)

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The effects of the WOX13 OG mutations on *Arabidopsis thaliana* development. (A) Diagram of the AtWOX13 protein of the *wox13* line 193G10 and the wild type plants (WT). (B) Diagram of the AtWOX14 protein of the *wox14* line pst13645 and the wild type plants (WT). *Legends:* WOX13MOG (pink), HD (orange). (C and D) Comparative primary root growth of mutant and wild type seedlings during the first 6 days of development. In (C), wild type WS ecotype (grey bar) and *wox13* line 193G10 (dot bar). In (D), wild type No ecotype (black bar) and *wox14* line pst13645 (white bar). (E) Floral transition of *wox13* and WS plants after a long day period of 4 weeks. (F) Floral transition of *wox14* and Nossen plants after a long day period of 4 weeks. (G to K) Root development of the wild type plant (G and H) compared to the *wox14* line pst13645 (I to K) that overproduces adventitious roots. (M to P) Flower development of *wox14* line pst13645 inflorescences that show partial sterility (M). Wild type (upper silique in N and O) and *wox14* semi-sterile silique (lower siliques in N and P). (N and P) Pictures obtained after clearing. (Q to S) Pollen viability tests with Alexander staining. (Q) Wild type, (R) *wox14* pollen. (S) *dmc1*/RNAi sterile mutant. (T to V). Mature flower development in wild type plants (T) and the *wox14* line (U and V). (V) Picture after clearing. Scale bars: Q, V (= 100 μ m), R, S (= 50 μ m) and T, U (= 500 μ m).

A. thaliana expression databases for the WOX13 OG genes [30-33]. For instance, in the gene expression map of the *Arabidopsis* root [39], the AtWOX13 mRNA profile shows a localized expression domain (LED) specific to the stele in the elongation zone of the mature primary root as observed in our study. These databases also revealed that the expression levels of both AtWOX14 and AtWOX13 were low compared to other WOX genes, suggesting either a tight control of their expression or a cell specific expression pattern. This has been confirmed by the expression data described in this paper.

The AtWOX14 branch within the WOX13 OG, has previously been suggested to be unique to *A. thaliana* [14]. Our study based on more species indicates that this branch also contains sequences from other Brassicaceae. Interestingly, differences in the expression profiles between AtWOX14 and AtWOX13 indicate that their 5' regulatory elements have diverged since gene duplication. In line with this conclusion, our data mining analyses showed that AtWOX13 expression, but not that of AtWOX14, was modulated in response to biotic and abiotic stresses. Hence, the AtWOX14 promoter appears to have diverged to presumably support specific developmental mechanisms found so far only in Brassicaceae. Analysis of the expression profile of a non Brassicaceae WOX13 gene should give us a better understanding of the evolutionary mechanisms giving rise to this particular branch.

WOX13 OG genes affect organogenesis and floral transition

The control of cell proliferation and differentiation during organ development are essential processes that require genes implicated in cell signaling, cell identity and cell cycle transitions. In higher plants, both the SAM and the root pericycle cells serve as initiation sites of organogenesis [40,41]. In this paper, we show that AtWOX13 and AtWOX14 are active in organ primordia and during early tissue differentiation. In the transcriptomic database CATdb [30], in which only the AtWOX13 probe is avail-

able, 5 out of 62 experiments associate AtWOX13 gene expression with cell proliferation and growth. Interestingly, the transcriptome profiling experiments, using protoplast cultures [42], showed that the AtWOX13 mRNA level followed an expression profile similar to that of the *proliferating cell nuclear antigen 1* gene (see additional file 4: Figure 3A), a key regulator of DNA metabolism and cell cycle progression [43]. Conversely, the homeobox gene *ATHB7*, a marker of cell differentiation and elongation [44], strongly decreased in the same protoplast culture. Compared to the cell cycle gene (see additional file 4: Figure 3B), AtWOX13 expression levels were maximal when the cells entered cell division. AtWOX13 expression was also up-regulated in the *tor* mutant or after a treatment with the IP3K inhibitor, wortmannin. The TOR protein and IP3K are regulators of cell division and growth. TOR has only been found in non-differentiated and rapidly dividing meristematic cells [45,46]. However, the dynamic expression of AtWOX13 during LR development described in this paper does not entirely overlap with the cell division zone, as AtWOX13 promoter activity became more restricted to the vasculature of the root.

Our data also showed that mutations of the AtWOX13 and 14 genes affected the floral transition. An increase of cell division in all the vegetative SAM was proposed to be a prerequisite for morphological changes at the floral transition in plants [47,48]. Furthermore, in *A. thaliana* stem axis growth is the result of cell elongation that occurs only in the rib zone and not in the upper part of the SAM [49]. Hence, the late flowering phenotype observed in the *wox14* mutant might reflect a default in the coordination of cell division and elongation in the transitional SAM. In contrast, the *wox13* mutant line showed a surprising early floral transition. However, a chimeric WOX13 mRNA coding for a putative protein containing the conserved domains was expressed in the *wox13* 193G10 line. We suggest that this line might act as a dominant negative mutant that induces early floral transition by activating precocious cell proliferation in the SAM. Alternatively, we

cannot exclude that other WOX family members could fulfill the function of the AtWOX13 protein, leading to a weak phenotype in the Arabidopsis mutant background. To date, the mechanism underlying gene regulation by the homeobox gene family is complex and remains largely unclear [50]. A monitoring of the promoter activity during the floral transition and an analysis of a mutant line with an insertion upstream of the HD or an RNAi strategy should give us better insights into the function of the WOX13 protein.

Finally, we found that the *wox14* null mutant showed root growth delay and early anther maturation. Interestingly, a similar phenotype was also observed in anthers when cyclin-dependent kinase inhibitors were ectopically expressed in plants [51-53]. Hence, it is tempting to speculate that the AtWOX14 protein is involved in the negative control of cell differentiation to allow for correct development.

In a similar way, the expression profile observed during the gametophytic development of *P. patens* is consistent with a putative function in cell differentiation control during the filamentous growth and the bud formation although *in situ* hybridization experiment are needed to validate this hypothesis. Last, the constitutive expression of the only WOX gene in *O. tauri* may account for its maintenance in an undifferentiated state. Future use of *O. tauri* and *P. patens* as biological models and ongoing complementation experiments of mutants using either full length or truncated proteins should allow us to obtain a better understanding of the functional importance of the WOX13 OG genes.

Conclusion

Our data provide evidence in favor of the WOX13 OG as the clade containing the most conserved WOX genes. The preferential presence of amino acid motifs fully supports the separation of the different gene groups in the phylogenetic trees. However, the latter is not consistent with the green plant phylogeny [21]. In this study, we linked the function of the WOX13OG Arabidopsis members to organ initiation and development, most likely by preventing premature differentiation as shown for other WOX proteins. The data also suggest that the WOX family has diverged both at the transcriptional and protein level to generate new features that control cell identity in specific domains of land plants. In line with such evolutionary events, the expression profile and mutant phenotype of the AtWOX14 gene suggest that the Brassicaceae might have developed specific mechanisms to control floral transition and pollen formation when compared to other plants.

Methods

Identification of WOX genes

The identification of WOX genes was undertaken using model plants for which the complete genome was available, *i.e.*, *A. thaliana* (a core eudicot), *O. sativa* (a monocot), *P. patens* (a moss) and *O. tauri* (a green alga). The WOX family was defined in *A. thaliana* and *O. sativa* genomes after a BLASTp [54] search in the FLAGdb++ database [55] using AtWUS (AT2G17950). AtWUS-like sequences were identified in the *P. patens* whole genome shotgun database ftp://ftp.ncbi.nih.gov/pub/TraceDB/physcomitrella_patens/, the *O. tauri* [56] and the *O. lucimarinus* [57] genomes after a BLASTp using the HD amino acid sequence. The limits of the sub-families were defined following the drop-in-Blast score method [58,59]. The structures of the identified genes were verified using all of the corresponding transcripts and when necessary the gene was reannotated. There was only one WOX gene, (OT13G01350), within the *O. tauri* and the *O. lucimarinus* genomes. Gene and protein sequences of the WOX families from completely sequenced genomes of model species are given (see additional file 3: Table 1). The positions of introns were obtained from nucleotide sequence alignments derived from the protein alignments. Only introns inserted into genomic regions coding for sufficiently similar protein sequences were used to study the conservation of intron insertion sites.

The proteome of *S. moellendorffii* was downloaded from the JGI ftp site ftp://ftp.jgi-psf.org/pub/JGI_data/Selaginella_moellendorffii/. Using the WOX-HD HMM (see additional file 3: Table 2), we found 8 WOX sequences, but only 3 were apparently full-length. Only the complete SmWOX sequences were used for the phylogeny reconstruction of the WOX family.

Distance trees and phylogeny reconstruction of the WOX family

Neighbor-Joining (NJ) [60], Maximum likelihood (ML) [61] trees, both based on Jones-Taylor-Thornton matrix [62], and Parsimony [63] trees were constructed using PHYLIP [64]. To assess support for the calculated relationships, 1000 bootstrap samples were generated [65]. To select appropriate outgroups for the WOX model plant phylogeny, we first established a tree using all of the HD proteins from *A. thaliana* and 3 outgroups (*i.e.*, At3G03260, At4G00730, At4G17710) were chosen from the three clades closest to the WOX clade. Trees were computed from 60 to 61 amino acid sequences containing the HD of the WOX proteins from the five plant model genomes, *O. tauri*, *P. patens*, *S. moellendorffii* (only the 3 available full-length proteins were used), *A. thaliana* and *O. sativa*. The clustering of the WOX proteins in three different OGs obtained from the genome models were consistent using both NJ (Figure 1) and ML (see additional

file 1: Figure 1) methods and the 3 above outgroups either independently or together (data not shown). We also built up phylogenetic tree from the same set of data using the Parsimony method. Different branching patterns were obtained depending upon which outgroup was used. With the multiple outgroup described above, *O. tauri* and *S. moellendorffii* were both independent branches at the base of the tree (see additional file 1: Figure 2).

The WOX13 OG trees has been established using the Neighbor-Joining [60] method based on Henikoffs evolutionary distances [66]. The trees obtained with the ML (see additional file 1: Figure 3) and Parsimony methods (see additional file 1: Figure 4) also gave the same main clusters of proteins.

Analysis of WOX protein sequences

In order to identify shared motifs among the protein sequences we used the MEME program, version 3.5.4 [67]. A gapped alignment using CLUSTALW was generated for each motif. Based on these multiple alignments a Hidden Markov Model (HMM) was built using HMMER 2.3.2 [20]. The HMM was used to search for motifs in the WOX sequences from the model genomes and in the whole proteome in GenBank NR (see additional file 3: Table 2). The sequence logos were made using Weblogo [68]. The specificity of the HMMs for the different WOX clades was tested using the two recently available *Vitis vinifera* genome sequences, PN40024 [69] and Pinot noir [70]. Using the WOX HD HMMs, an initial search in these two novel proteomes allowed us to specifically retrieve all members of the WOX gene family. A second search with the clade specific HMMs efficiently clustered the WOX genes into one of the three WOX clades (see additional file 3: Table 3). We found 13 different *Vitis* genes coding for WOX proteins, with 8 located in the WOX1 OG, 3 in the WOX13 OG, and 2 in the WOX8 OG. No *V. vinifera* proteins were located to the WOX14 clade.

Plant material

A naturally synchronized *O. tauri* strain OTTH0595 cell culture was used and the cell cycle analysed by flow cytometry as described by Farinas et al [27]. Using a 12 h light/dark condition, we obtained two subpopulations of synchronized cells at the light/dark transition, comprised of 12% cells in the G2/M phase and the remaining cells in the S phase of the cell-cycle. The moss *P. patens* was grown on BCD medium [71] overlaid with cellophane under 16 h light/8 h dark cycle at 65% hygrometry and at 20°C. The DS transposon tagged *A. thaliana* line *wox14*, i.e., pst13645, was obtained from the RIKEN BRC Japan. The T-DNA tagged *A. thaliana* line *wox13*, i.e., 193G10, was purchased from the INRA Versailles resource centre. The *wox13* and *wox14* mutant lines are in the Wassilewskija (Ws) and the Nossen (No) background, respectively. Wild

type plants of both ecotypes were used as background controls.

Root growth assays

Sterile seeds were placed on a solid growth medium (MS salt supplemented with 20 g/L sucrose and agar) at 4°C for 2 days, and grown under 16 h light/8 h dark cycles (20°C during illumination, 18°C in the dark) at 65% hygrometry. Root length was measured from day 3 to day 6 using the ImageJ free software (NIH, USA) after scanning.

Characterization of the mutant lines

The pst13645 and 193G10 lines were screened using media containing either 30 mg/L hygromycin or 30 mg/L kanamycin, respectively. In order to verify the mapping of the insertion site, F2 or F3 plants were genotyped at the *AtWOX13*, *AtWOX14* loci. Specific genomic DNA primers and either T-DNA or the Ds-element border-specific primers were designed based on the annotated insertion site of each mutant line (see additional file 4: Table 1). The resulting PCR products were separated on a 2% agarose gel and their size was verified. 341 bp and 163 bp fragments were expected for the *AtWOX13* wild type and mutated allele, respectively and 498 bp and 216 bp fragments for the *AtWOX14* wild type and mutated allele, respectively. Moreover, the entire *AtWOX14-AtWOX10* DNA region was genotyped to check for the absence of chromosome rearrangements of the duplicates and to confer that the phenotype was only due to the disruption of *AtWOX14*. Phenotypes were observed only in homozygous plants for each mutation and therefore they were used in all further experiments.

Reverse transcriptase PCR

Total RNA was extracted from the different plants using the Qiagen plant RNeasy extraction kit and DNaseI treated following the manufacturer's instructions. The RNA preparations were reverse transcribed using 1 to 5 µg of RNA. RT-PCR experiments were performed in order to verify the expression of the different WOX and control genes (see additional file 4: Table 1) within the different plant model species and the transgenic Arabidopsis plants. Primer efficiency was tested using genomic DNA and eventual genomic contamination was always checked by PCR amplification of the RNA samples using the equivalent amount of cDNA used for the RT-PCR. Quantitative real-time RT-PCR was also performed on *O. tauri* mRNA as described previously [72] using a Roche Light Cycler® 480 with the SYBER® green PCR mix from either Roche or Applied Biosystem. For each gene tested, PCR products from genomic DNA cloned into the PGEMT® vector and the *NotI* digested empty plasmid were used as standards to quantify the corresponding RNA copy number. These raw

copy numbers were then normalized to the amount of equivalent RNA used for the quantification.

Construction of the *WOX::GUS* reporter genes

To clone the *AtWOX13* and *AtWOX14* promoters, the entire intergenic regions upstream of the ATG initiators were PCR-amplified using gene specific primers containing either an *Xba*I or a *Bam*HI restriction site (5'-TCTC-GAGTGGAGCTTTTGCAGGTCTCT-3'; 5'-AGGATCCTCAG AATTCGCTCAGAAGATT-3') for *AtWOX13* and containing either a *Not*I or a *Spe*I restriction site (5'-GACAGCGGCCGCGGGTGTGAGTCCTATTGC-3'; 5'-GACACTAGTTGAACAAGACAATGAGAAAGTGAA-3') for *AtWOX14*. DNA fragments (of 3.6 kb and 1 kb for the *AtWOX13* and *AtWOX14* promoters, respectively) were subcloned into the pGEMT vector (Promega) to give the prom *WOX13* and prom *WOX14* plasmids. For the *WOX::GUS* reporter constructs, prom *WOX13* *Xho*I/*Bam*HI and prom *WOX14* *Spe*I fragments were subcloned into a *Sall*/*Bam*HI and an *Xba*I digested pPR97 binary vector, respectively [73]. Constructs were introduced into the *Agrabacterium tumefaciens* strain GV3101, then into Arabidopsis plants ecotype Ws as described in [74]. Transformed plants were selected by adding 30 mg/L kanamycin to the growth medium. Resistant plants were transferred to soil and grown in a growth chamber.

Staining and observations

Beta-glucuronidase (GUS) staining was performed for 24 h on T1 and T2 seedlings as previously described [75], except for the lateral roots where the incubation time was limited to 4 h. After fixing and clearing, samples were mounted on slides in HCG (80% Chloral hydrate, 10% Glycerol, 10% water) and observed under a Zeiss light microscope or a Zeiss binocular. Cytological observations of Alexander-stained anthers were carried out as previously described [76].

In situ hybridization

In situ hybridizations were conducted as previously described [77] using antisense probes. The *AtWOX13* antisense probe was synthesized in vitro using a PCR product obtained with the following primers: (CCTGCAGATGAT-GGAATGGGATAATCAGC and TGTAATACGACTCACTATAGGGCACTGCTTATGACT-GACTACCAAATCC). The CUC2 probe has been described previously [77] and the GUS probe was provided by the laboratory of P. Laufs (INRA Versailles).

Authors' contributions

The project was conducted by YD, MK and AL. The bioinformatics analyses were done by CTN, VT and AL; YD and GC carried out all the experimental works with the assistance of H. Morin (in conducting the *in situ* hybridization) and H. Moreau, the *O. tauri* experiments. YD, MK and AL

wrote this manuscript; PL revised several versions of this manuscript. All authors read and commented on drafts of the manuscript and approved the final manuscript.

Additional material

Additional file 1

Phylogenetic trees. Figure 1 : Maximum-likelihood tree of the *WOX* proteins from the 5 completely sequenced genomes of the green lineage: Arabidopsis thaliana (*AT* prefix), Oryza sativa (*OS* prefix), Physcomitrella patens (*Ppa* prefix), Ostreococcus tauri (*OT* prefix) and the three full-length proteins of Selaginella moellendorffii (*Sm* prefix). The significance of each node was tested using 1000 bootstrap replicates. Only bootstrap values above 50% are shown. The scale, at the right of the tree, indicates 0.2 substitution per site. **Figure 2 : Parsimony tree of the *WOX* proteins from the 5 completely sequenced genomes of the green lineage:** Arabidopsis thaliana (*AT* prefix), Oryza sativa (*OS* prefix), Physcomitrella patens (*Ppa* prefix), Ostreococcus tauri (*OT* prefix) and the three full-length proteins of Selaginella moellendorffii (*Sm* prefix). The significance of each node was tested using 1000 bootstrap replicates. Only bootstrap values above 50% are shown. The scale, at the right of the tree, indicates 0.1 substitution per site. Species names, in alphabetic order, are: *Afo*: Aquilegia formosa; *AT*: Arabidopsis thaliana; *Bna*: Brassica napus; *Bra*: Brassica rapa; *Ccl*: Citrus clementina; *Csi*: Citrus sinensis; *Cte*: Citrus temple; *Ees*: Euphorbia esula; *Ghi*: Gossypium hirsutum; *Gma*: Glycine max; *Gra*: Gossypium raimondii; *Han*: Helianthus annuus; *Hvu*: Hordeum vulgare; *Les*: Lycopersicon esculentum; *Lja*: Lotus japonicus; *Lsa*: Lactuca sativa; *Lvi*: Lactuca virosa; *Mdo*: Malus domestica; *Mtr*: Medicago truncatula; *OS*: Oryza sativa; *OT*: Ostreococcus tauri; *Ppa*: Physcomitrella patens; *Psi*: Picea sitchensis; *Pta*: Pinus taeda; *Ptr*: Populus tremula × Populus tremuloides; *Pvu*: Phaseolus vulgaris; *Stu*: Solanum tuberosum; *Vvi*: Vitis vinifera; *Zma*: Zea mays. **Figure 3 : Maximum-likelihood tree of the *WOX* proteins from the *WOX13* Orthology Group.** The significance of each node was tested using 1000 bootstrap replicates. Only bootstrap values above 50% are shown. The scale, at the right of the tree, indicates 0.1 substitution per site. Species names, in alphabetic order, are: *Afo*: Aquilegia formosa; *AT*: Arabidopsis thaliana; *Bna*: Brassica napus; *Bra*: Brassica rapa; *Ccl*: Citrus clementina; *Csi*: Citrus sinensis; *Cte*: Citrus temple; *Ees*: Euphorbia esula; *Ghi*: Gossypium hirsutum; *Gma*: Glycine max; *Gra*: Gossypium raimondii; *Han*: Helianthus annuus; *Hvu*: Hordeum vulgare; *Les*: Lycopersicon esculentum; *Lja*: Lotus japonicus; *Lsa*: Lactuca sativa; *Lvi*: Lactuca virosa; *Mdo*: Malus domestica; *Mtr*: Medicago truncatula; *OS*: Oryza sativa; *OT*: Ostreococcus tauri; *Ppa*: Physcomitrella patens; *Psi*: Picea sitchensis; *Pta*: Pinus taeda; *Ptr*: Populus tremula × Populus tremuloides; *Pvu*: Phaseolus vulgaris; *Stu*: Solanum tuberosum; *Vvi*: Vitis vinifera; *Zma*: Zea mays. Click here for file [http://www.biomedcentral.com/content/supplementary/1471-2148-8-291-S1.pdf]

Additional file 2

WOX gene structures and protein motifs. Figure 1 : The structure of WOX genes from model genomes. The gene structures from 4 model genomes (Arabidopsis thaliana, AT; Oryza sativa, OS; Physcomitrella patens, Ppa; Ostreococcus tauri, OT) and three Selaginella moellendorffii (Sm) genes are displayed following the order of the Neighbour-joining tree in Figure 1 and aligned on their translation start. Rectangles stand for exons while lines denote introns. Blue is for exonic regions in UTRs and light grey is for translated exons. Five different gene structures are given for PpaWOX02 due to the 5' UTR variants supported by transcripts. Figure 2 : WOX proteins: Sequence alignments around the virtual intron insertion sites. For each WOX OG, 15 amino acid sequences both upstream and downstream introns are given. The intron insertion sites are indicated by stars. Numbers on the left of sequences give the intron length. Digits between stars stand for the phase of the intron insertion site. The intron phase is defined by the position of its insertion into a codon. An intron can be located between two codons (phase 0) or within a codon, lying either after the first or after the second base pair (phase 1 and phase 2 respectively). Boxes indicate conserved introns with the same color code than in Figure 1. Figure 3 : Sequence logos of protein motifs in the WOX family. Each logo <http://weblogo.berkeley.edu> consists on stack of letters, one stack for each position in the sequence. The overall height of the stack indicates the sequence conservation at that position, while the letter's height within the stack indicates the relative information content of each amino acid at that position. The amino acid color code is dark blue for large polar, light blue for basic, black for small polar, green for nonpolar, orange for cysteine and purple for ambivalent amino acids. Figure 4 : WOX motifs: alignments for motifs without Logo. Motifs defined by less than 6 sequences cannot be shown as logo since the sequence information at each position is too weak. Thus multiple alignments performed with Clustalw with a majority consensus are shown here for these motifs. Amino acid color code is the same than in Figure 2. Click here for file [<http://www.biomedcentral.com/content/supplementary/1471-2148-8-291-S2.pdf>]

Additional file 3

WOX sequences and HMM motifs. Table 1: Gene and protein sequences of the WOX family in the model genomes A. thaliana, O. sativa, P. patens and O. tauri. Table 2: Specificity and sensitivity of HMM for WOX motifs. Scores can give indications to distinguish potential family -or subfamily-members from false positive. A clear drop in the score could be detected in most cases, indicating that sequence below this threshold did not fulfill the family model as well as above [21]. HMMER bit scores were obtained for each motif by querying the HMM against three protein sequence sets: WOX13 OG (33 sequences), model genome WOX (32 sequences) and GenBank NR (Release 157). The score drops above 50 and indicating the motifs exhibiting the highest specificities with a given database are in bold. a: no hit was found in the set. b: no specific hit was found in the GenBank NR database. Note that Physcomitrella sequences from genome model WOX set are not present in GenBank NR. c: best hits are not belonging to the WOX family and no score drop was found. Table 3: Vitis vinifera WOX protein prediction with HMM WOX motifs. HMM search scores above a drop give indication to assign novel WOX proteins to the different WOX OGs. AM(...) IDs are contigs from Pinot noir V. vinifera while contig(...) IDs are from PN40024 V. vinifera. Dash indicates that no score was found above the drop. a: no hit was found with this HMM profile. (*): 17.4 score is due to an undefined C or T nucleotide at one position leading to a stop in the CDS. After fixing the nucleotide to C, the score is 27.7. Click here for file [<http://www.biomedcentral.com/content/supplementary/1471-2148-8-291-S3.pdf>]

Additional file 4

Expression analyses. Figure 1 : Real-time PCR analysis of Ostreococcus tauri gene expression in a 12-hour light/12-hour dark synchronized cell culture. Copy number of WOX, KNOX, Histone4 and CyclinB RNA was quantified using total RNA sampling from the starting of the light period (0 h) or from continuous light grown cell culture. Figure 2 : Alignments of predicted sequences of AtWOX13 (A) and AtWOX14 (B) chimeric proteins with the full length ones. The homeodomain are indicated in bold and the amino acid changes at the end of the chimeric proteins indicated in lower case. Figure 3 : AtWOX13 expression is correlated with the induction of cell proliferation. Data mining analysis of the expression profiles of a set of genes in protoplast culture of Arabidopsis thaliana. (A) AtWOX13 (-♦-), ATHB-7 (■), PCNA1 (▲); (B) Expression profile of a cluster of cell cycle genes including AT4G37490 (CYC1); AT1G44110 (CYCA1;1 (■)); AT5G11300 (CYC3B); AT1G47210 (CYCA3;2); AT3G05330 (CDK); AT2G26760 (CYCB1;4); AT4G35620 (CYCB2;2); AT2G26430 (RCY1); AT2G31400 (ATPDNA Binding). Table 1: Primer sequences designed for expression analysis, genotyping (g primers) and cDNA cloning (2 last row) using primer3 software. Click here for file [<http://www.biomedcentral.com/content/supplementary/1471-2148-8-291-S4.pdf>]

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