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# Genes & Development

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## RESEARCH COMMUNICATION

# WUSCHEL induces shoot stem cell activity and developmental plasticity in the root meristem

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**Most of the plant shoot originates from a small group of stem cells, which in *Arabidopsis* are specified by *WUSCHEL* (*WUS*). It is unknown whether these cells have an intrinsic potential to generate shoot tissues, or whether differentiation is guided by signals from more mature tissues. Here we show that *WUS* expression in the root induced shoot stem cell identity and leaf development (without additional cues), floral development (together with *LEAFY*), or embryogenesis (in response to increased auxin). Thus, *WUS* establishes stem cells with intrinsic shoot identity and responsive to developmental inputs that normally do not change root identity.**

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Plants generate new organs and tissues reiteratively at the meristems, which are groups of undifferentiated, actively dividing cells present in the growing apices and in axillary buds (Weigel and Jürgens 2002). Within the shoot apical meristem (SAM), a small group of centrally located, slowly dividing stem cells is the ultimate source of all meristem cells and therefore all new shoot organs (Stewart and Dermien 1970). In the SAM, stem cells reside in the meristem central zone (CZ), which regularly provides new cells to replenish the peripheral zone (PZ), where new organs are initiated (Gross-Hardt and Laux 2003).

In *Arabidopsis*, maintenance of the stem cells in the CZ requires *WUSCHEL* (*WUS*), which encodes a homeodomain protein (Laux et al. 1996; Mayer et al. 1998). In strong *wus* mutants, SAM development during embryogenesis is defective, with the CZ occupied by cells that are larger and more vacuolated than the normal meristem cells. When the seedling germinates, one or two leaf primordia emerge at the shoot apex, indicating that PZ activity is present, but the pool of undifferentiated

cells is not replenished and organogenesis stops. New leaves are eventually initiated, presumably by a process related to the establishment of axillary meristems, but organogenesis again terminates prematurely. Some mutant plants finally form flowers that lack the innermost organs (stamens and carpels), showing that *WUS* is also required to sustain organogenesis in the floral meristem.

*WUS* is first expressed in the 16-cell embryo, preceding meristem development, within the region that originates the embryonic shoot (Mayer et al. 1998). Although no clear function has been attributed to this early expression, ectopic *WUS* expression induced somatic embryogenesis, suggesting that *WUS* promotes embryonic identity (Zuo et al. 2002). Subsequent expression of *WUS* in the vegetative and reproductive meristems is confined to a small group of cells below the CZ. Because of its expression beneath the SAM stem cells, it has been proposed that *WUS* acts through an intercellular signal that maintains the stem cells (Mayer et al. 1998), but the signal has not yet been identified. The maintenance of stem cells by signals from specialized cells is also seen in the root meristem, and is in fact a common feature of stem cells in plants and animals (Spradling et al. 2001; Weigel and Jürgens 2002; Laux 2003; Sabatini et al. 2003).

One general question in stem cell biology is the extent to which stem cells can be directed to alternative fates by signals from surrounding tissues (transdifferentiation), or whether they have an intrinsically limited range of fates (Weissman et al. 2001). In plants, laser ablation experiments and genetic evidence suggest that the differentiation of meristem cells can be directed by signals from more mature tissues (van den Berg et al. 1995; Stuurman et al. 2002). Thus *WUS* could act by establishing naïve cells that subsequently differentiate as shoot cell types in response to signals from surrounding shoot tissues. If *WUS* protects stem cell identity by antagonizing differentiation signals that emanate from surrounding tissues, the question arises whether this antagonism is specialized for signals that promote shoot cell fates, or whether *WUS* could have a general role in blocking differentiation. Alternatively, *WUS* could establish stem cells that give rise to shoot tissues, regardless of the differentiating cells surrounding them. To address these questions, we studied the effects of ectopic expression of *WUS* outside shoots.

## Results and Discussion

To be able to induce the non-cell-autonomous effects of *WUS* (Mayer et al. 1998), we expressed *WUS* in roots using a Cre-*loxP*-based mosaic expression system (Gallois et al. 2002). The plants contained heat shock-inducible Cre recombinase, which catalyzed excision of a  $\beta$ -glucuronidase (*GUS*) reporter gene to activate *WUS* expression from the widely expressed 35S promoter (for simplicity, this genotype will be called "*WUSMOS*").

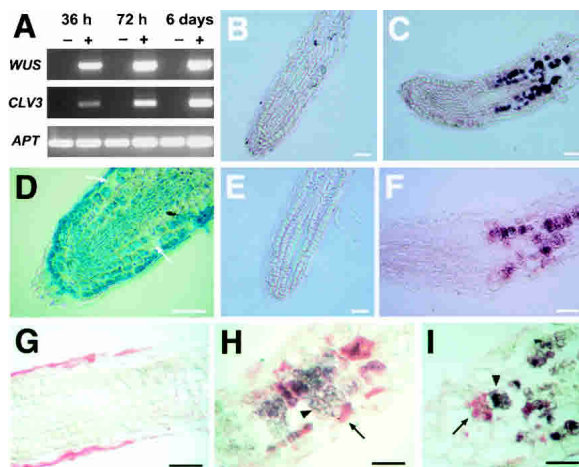
RT-PCR confirmed that *WUS* expression was activated by heat shock in *WUSMOS* roots, whereas *WUS* mRNA was undetectable in non-heat-shocked controls (Fig. 1A) or heat-shocked roots that lacked the *WUSMOS* construct (data not shown). RNA in situ hybridization confirmed that *WUS* was activated in a mosaic pattern near the root tips (Fig. 1B,C); *GUS* staining revealed a

[Keywords: *Arabidopsis*; *WUSCHEL*; auxin; *LEAFY*; pluripotency; embryogenesis]

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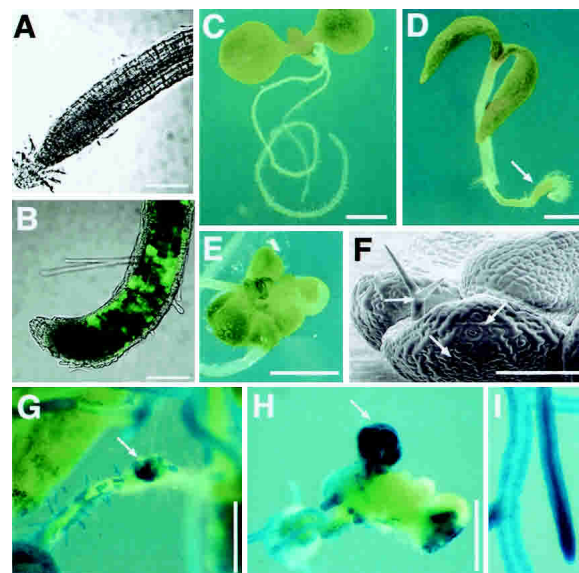
**Figure 1.** Ectopic WUS activated a shoot stem cell marker in roots. (A) RT-PCR detection of *WUS*, *CLV3*, and *APT* (constitutive control) mRNAs in roots from *WUSMOS* seedlings at different times after heat shock (+) or in non-heat-shocked controls (-). (B,C) RNA in situ hybridization on longitudinal sections of *WUSMOS* root tips, 3 d after heat shock (C) or non-heat-shocked control (B; same genotype as in C); the dark signal in C reveals *WUS*-expressing cells. (D) Longitudinal section of *WUSMOS* root tips, stained for GUS 3 d after heat shock; arrows indicate GUS-negative cells. (E,F) As in B,C, but hybridized with *CLV3* antisense probe. (G-I) Double-labeling RNA in situ hybridization, with *WUS* signal in red (arrows) and *CLV3* signal in blue (arrowheads). (G) Non-heat-shocked control. (H,I) Fixed 3 d after heat shock. Bar, 40  $\mu$ m.

complementary pattern, with scattered GUS-negative cells in the vicinity of the root meristem (Fig. 1D). The lack of *WUS*-expressing, GUS-negative cells very close to the root tip, 3 d after heat shock, suggests that either Cre activation was inefficient within the root meristem, or that *WUS*-expressing cells were left behind as the root tip continued to grow. The region of the root meristem that did not express *WUS* showed disorganized cell division (Supplemental Material); we do not know whether this was the direct effect of a *WUS*-induced intercellular signal, or an indirect consequence of the changes in the cells adjacent to the root tip (see following).

In the shoot apex, *WUS* activates *CLAVATA3* (*CLV3*; Schoof et al. 2000; Brand et al. 2002), which functions in a negative feedback loop that antagonizes *WUS* activity to control the size of the stem cell population (Fletcher et al. 1999; Brand et al. 2000; Schoof et al. 2000). *CLV3* expression specifically marks the shoot stem cells (Laux 2003). We saw that ectopic *WUS* activated *CLV3* in roots, detectable 36 h after heat shock and maintained for at least 6 d (Fig. 1A). In situ hybridization showed that the *CLV3*-expressing cells were present in the same region of the root tip where *WUS* was activated (Fig. 1E,F). Double-labeling in situ hybridization, however, revealed that the expression patterns of *WUS* and *CLV3* did not coincide. Separate expression of *CLV3* and *WUS* was seen in adjacent cells (Fig. 1G-I), showing that, as in the shoot apex, *WUS* expression in the root was able to activate *CLV3* non-cell-autonomously.

The *CLV3* expression suggested that ectopic *WUS* was sufficient to induce shoot stem cell identity in the root tips. If the subsequent fate of ectopic shoot stem cells

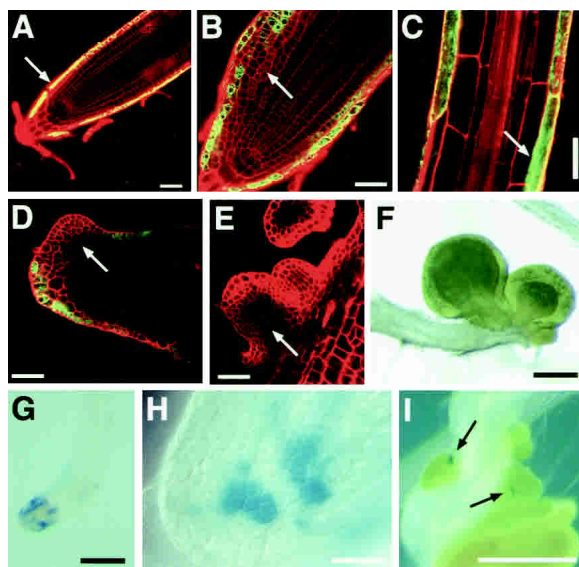
was determined by input from neighboring tissues, the cells should eventually reacquire root identity. Instead, we saw that the *WUS*- and *CLV3*-expressing root tip soon developed shoot features. Three to four days after heat shock, the root tips expressed green fluorescent protein (GFP) directed by the promoter from *AINTEGUMENTA* (*ANT*, a marker for shoot organ primordia; Elliott et al. 1996; Fig. 2A,B). Activation of endogenous *ANT* was also confirmed by RT-PCR (data not shown). Six days after heat shock, the root tip contained green tissues (Fig. 2C,D), which either replaced the root tip (Supplemental Material) or were left behind as the tip continued to grow. Between 2 and 3 wk after heat shock, the primary root had developed leaf-like organs in 50% of the plants ( $n = 318$ ), with characteristic leaf cell types such as guard cells and trichomes (Fig. 2E,F). In other cases (24%), the root tip formed a green callus containing leaf cell types such as guard cells; in 4% of the plants, the root tips formed embryo-like structures similar to those described previously (Zuo et al. 2002), whereas the remaining 22% of the plants had no visible green tissues in the primary root (data not shown). The ectopic leaves formed in heat-shocked *WUSMOS* root tips were made entirely or partially of GUS-positive cells (Fig. 2G,H). As root cells that



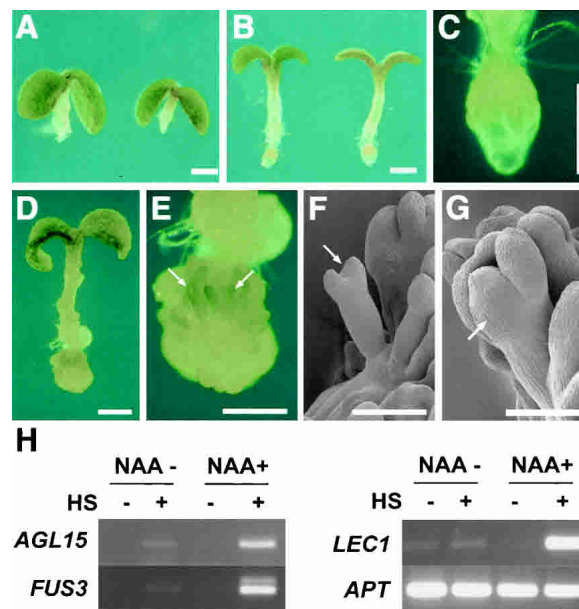
**Figure 2.** *WUS* induced the development of shoot tissues from roots. (A,B) Optical sections (combined bright field and GFP channels) of *WUSMOS*, *ANT:GFP* root tips. (A) Non-heat-shocked control. (B) Four days after heat shock. (C,D) Eight-day-old *WUSMOS* plants. (C) Non-heat-shocked control. (D) Six days after heat shock, with green tissue near the root tip (arrow). (E) Ectopic leaves on root, 18 d after heat shock. (F) Electromicrograph of ectopic leaves on *WUSMOS* root, 21 d after heat shock; arrows indicate leaf cell types such as interdigitated epidermal cells, guard cells, and a trichome. (G-I) GUS staining of *WUSMOS* roots of 20-day-old plants, 18 d after heat shock (G,H) or in non-heat-shocked control (I). (G) Mosaic GUS expression on the primary root and ectopic shoot tissue developing on a lateral root tip (arrow). (H) Ectopic shoot tissue on the primary root tip, 18 d after heat shock, with a mixture of GUS-positive and GUS-negative tissues; the arrow points to a GUS-positive ectopic leaf. Bars: A,B, 200  $\mu$ m; C-E, 1 mm; F, 100  $\mu$ m; G-I, 500  $\mu$ m.

had not expressed WUS were converted to shoot identity, this response to WUS in the roots was non-cell-autonomous. When the roots formed disorganized green tissues, these were made of variable proportions of GUS-positive and GUS-negative cells (Fig. 2G,H; Supplemental Material), suggesting that the WUS-expressing cells also proliferated in WUSMOS root tips.

The induction of ectopic shoot tissues in the roots was confirmed using a gene trap line (*J2301*), in which genes containing the *UAS* sequence are activated by GAL4-VP16 in the lateral root cap and in the atrichoblasts of the root epidermis (Fig. 3A). Shortly after germination, the root tips of *J2301*, *UAS:GFP*, *UAS:WUS* seedlings showed aberrant cell divisions both in GFP-negative epidermal cells and in adjacent cells that expressed GFP (and presumably also WUS; Fig. 3B). Subsequently, GFP-negative cells formed outgrowths resembling leaf primordia (Fig. 3D,E), which eventually gave rise to ectopic leaves (Fig. 3F). As in *WUSMOS* roots, development of ectopic shoot tissues was preceded by *CLV3* expression (this time revealed by a *CLV3:GUS* reporter gene; Brand et al. 2002). Small groups of *CLV3:GUS*-expressing cells were seen at the root tip shortly after germination (Fig. 3G,H) and later associated with the ectopic primordia (Fig. 3I), although *CLV3:GUS* expression eventually dis-



**Figure 3.** Expression of WUS in the lateral root cap induced ectopic leaf development. (A–C) Optical sections of *J2301* roots [GFP and propidium iodide channels combined]. (A) *J2301*, *UAS:GFP* root tip, 4 d after germination; the arrow indicates GFP expression in the lateral root cap. (B) *J2301*, *UAS:GFP*, *UAS::WUS* root tip, 7 d after germination; the arrow indicates abnormal cell proliferation. (C) Mature section of *J2301*, *UAS:GFP*, *UAS::WUS* root, 7 d after germination; the arrow indicates GFP expression in atrichoblasts. (D–E) Ectopic leaf primordia (arrows) on secondary root tips of *J2301*, *UAS:GFP*, *UAS::WUS* plants, 21 d after germination. (F) Ectopic leaves on the primary root tip of *J2301*, *UAS:GFP*, *UAS:WUS* plant, 21 d after germination. (G–I) GUS staining of *J2301*, *UAS:GFP*, *UAS:WUS*, *CLV3:GUS* root tips. (G,H) Primary root tip, 7 d after germination. (I) Secondary root tip, 21 d after germination; arrows indicate *CLV3:GUS* expression (blue signal) associated with ectopic leaf primordia. Bars: A–E,H, 40  $\mu$ m; F,G,I, 200  $\mu$ m.



**Figure 4.** WUS expression combined with auxin induced somatic embryogenesis in roots. (A–C) Eight-day-old *WUSMOS* seedlings grown on medium with 5  $\mu$ M NAA, without heat shock (A) or 6 d after heat shock (B); C is a higher-magnification view of the root tip of a seedling equivalent to B. (D,E) Twenty-day-old *WUSMOS* seedling grown with 5  $\mu$ M NAA, 18 d after heat shock; the root tip of D is shown at higher magnification in E, with arrows indicating somatic embryos. (F,G) Cryo-scanning electron micrographs of 2-week-old heat shocked *WUSMOS* plants grown on medium containing 5  $\mu$ M NAA. Arrows indicate somatic embryos. (H) RT-PCR detection of embryo marker mRNAs and constitutive control (APT) in roots dissected from *WUSMOS* seedlings, not heat shocked (HS–) or 2 wk after heat shock (HS+), grown in GM medium (NAA–) or medium supplemented with NAA 5  $\mu$ M (NAA+). Bars: A–E, 1 mm; F,G, 100  $\mu$ m.

appeared as the leaves grew (data not shown). These results are compatible with the idea that descendants of *CLV3*-expressing cells formed the ectopic shoot tissues, although definitive proof of this will require clonal analysis.

In both the *WUSMOS* and in the *UAS:WUS* experiments, phenotypic effects were only seen in the primary and lateral root tips, despite the fact that heat shock-induced Cre catalyzed GUS excision throughout the roots (Fig. 2G), and that GFP (and presumably WUS) was expressed in the atrichoblasts of more mature root regions (Fig. 3C). As the response to WUS expression in roots was restricted to the meristematic regions, we aimed to expand WUS action, using external application of auxin to induce larger numbers of lateral root meristems (King et al. 1995). Unexpectedly, these experiments showed a change in the fate of the cells responding to WUS. When heat-shocked *WUSMOS* seedlings were plated on medium containing 5  $\mu$ M of the synthetic auxin  $\alpha$ -naphthalene acetic acid (NAA), none of the roots formed ectopic leaves, and instead virtually all developed structures resembling embryos, mostly oriented with their shoot poles away from the root tip (Fig. 4). We also noticed that the inhibition of hypocotyl elongation by auxin was somewhat suppressed in seedlings where

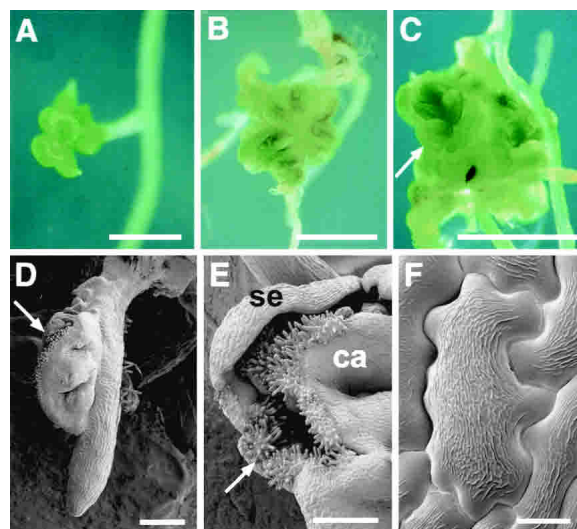
WUS was activated (Fig. 4, cf. A and B); we do not know whether this is an effect of ectopic WUS in the hypocotyl or in the root. Embryonic identity was confirmed by activation of embryo marker genes such as *LEC1*, *FUS3*, and *AGL15* (Fig. 4H; Heck et al. 1995; Rounsley et al. 1995; Lotan et al. 1998; Luerssen et al. 1998). In the absence of external auxin, heat-shocked *WUSMOS* roots expressed these embryonic markers at a low level, yet higher than in the roots of non-heat-shocked plants. The low levels of embryonic gene expression in *WUSMOS* plants without added auxin are consistent with the embryo-like structures that developed on the root tips of these plants at a low frequency and may be due to the endogenous auxin that accumulates at the root tips (Sabatini et al. 1999).

The somatic embryos described earlier are in agreement with the previous report (Zuo et al. 2002) that expression of WUS in roots induced somatic embryogenesis. The effect of auxin is in accordance with the observation that it promotes somatic embryogenesis in tissue culture, is present at high levels in early embryos, and is essential for normal embryo patterning (Feher et al. 2003; Friml et al. 2003). Thus, it appeared that, depending on auxin levels, ectopic WUS could redirect root cells to at least two different developmental pathways: shoot organogenesis or somatic embryogenesis.

To test whether *WUSMOS* roots could be directed to another of the developmental pathways in which WUS normally functions, we combined *WUSMOS* with constitutive expression of *LEAFY* (*LFY*), which is a master regulator of floral development (Weigel et al. 1992). Expression of *LFY* using the *35S* promoter (*35S:LFY*) caused early flowering and conversion of the inflorescence meristem to floral meristem (Weigel and Nilsson 1995). *35S:LFY* alone, however, does not bypass the vegetative phase (when the shoot meristem produces leaves), does not alter embryogenesis, and has no effect on root development.

Heat shock had no effect on the development of *35S:LFY* plants. When *35S:LFY*, *WUSMOS* seedlings were heat shocked, floral organs and tissues developed from primary and lateral root tips (Fig. 5). These organs were not organized in the normal whorled arrangement seen in flowers, and their identity seemed random, with various combinations of sepals (Fig. 5A,E), stamens (Fig. 5C), and carpel tissue (such as the stigmatic papillae shown in Fig. 5D,E). We cannot discriminate whether WUS only acted to establish pluripotent cells, with *LFY* subsequently directing their development, or whether WUS also acted directly in combination with *LFY* to control genes involved in floral development. The latter possibility is based on the fact that during flower development, the organ identity gene *AGAMOUS* (*AG*) is directly activated by WUS combined with *LFY* (Lenhard et al. 2001; Lohmann et al. 2001). However, an indirect interaction between *LFY* and WUS is also evident during normal floral development: *LFY* confers floral identity throughout the floral meristem, including cells that do not express WUS, but whose maintenance requires WUS.

Our results showed that WUS expression made root cells developmentally flexible and able to be directed to embryo, leaf, or floral organ development, depending on additional cues. The ability to enter alternative developmental pathways, combined with expression of a stem



**Figure 5.** Floral tissues in the roots of 21-day-old *WUSMOS*, *35S:LFY* plants, 19 d after heat shock. (A) Sepals on lateral root tip. (B) Sepal and carpel-like organs on lateral root tip. (C) Arrow indicates anther originated from lateral root tip. (D–F) Cryo-scanning electron micrographs showing carpelloid tissue on primary root tip (D), sepaloid (se) and carpelloid (ca) tissues on lateral root tip, with arrow indicating stigmatic papillae (E), and an epidermal cell with the epicuticular ridges typically seen in floral organs (F). Bars: A–C, 1 mm; D,E, 200  $\mu$ m; F, 10  $\mu$ m.

cell marker (*CLV3*), indicates that expression of WUS in the root caused ectopic activation of stem cell functions. The ectopic organs and embryo-like structures, however, did not maintain a stable pool of stem cells, and their development eventually terminated. This may be due to the fact that maintenance of a stable stem cell population requires feedback regulation of WUS by the *CLAVATA* pathway (Brand et al. 2000; Schoof et al. 2000), which cannot operate on the heterologous promoters used here (Gallois et al. 2002). Alternatively, other genes required for meristem activity, such as *SHOOT MERISTEMLESS* (Long et al. 1996), may not have been activated.

It is striking that when WUS was expressed in the roots with no additional cues (i.e., not combined with *LFY* or external auxin), ectopic leaves developed. Ectopic shoot identity did not occur simply because WUS rendered the root cells responsive to light as a developmental input, because *ANT* was still activated by WUS in the root tips of seedlings that were heat shocked and grown in the dark (data not shown). Our experiments leave two possibilities open. One is that, unless root identity is actively maintained, shoot development occurs by default. This could reflect the evolutionary origin of roots as an addition to preexisting shoots (Kenrick and Crane 1997). The alternative is that WUS itself provided the cues that converted the cells to shoot identity. Either way, the results imply that WUS does not simply establish naïve cells that require input from surrounding tissues to develop as shoot cells, but instead establishes cells with intrinsic potential to generate shoot tissues. Although we cannot exclude that WUS could induce stem cells and shoot identity through parallel pathways, the most straightforward interpretation of our results is

that shoot identity is a property of the stem cells specified by WUS.

We also saw that WUS made cells within or in the vicinity of the root meristem responsive to inputs that normally do not redirect root cell identity (e.g., LFY for floral development, increased auxin for embryogenesis). In addition to supporting the proposed role of WUS in promoting pluripotency (Mayer et al. 1998), this developmental plasticity may have practical use. Although plant development is remarkably plastic and virtually all parts of plants can be regenerated in tissue culture from fragments of adult plants, the conditions defined in tissue culture cannot override developmental controls in whole plants. We have shown that a gene that controls stem cell identity can be used to redirect root cells of an intact plant to any of the other major sporophytic developmental pathways—leaf, floral, or embryo development.

## Materials and methods

### *Arabidopsis lines*

*hsp18.2:Cre* (gift from Leslie Sieburth, Ohio State University), 35S:*lox-uidA-lox-WUS* and 35S:*lox-uidA-lox-GFP*, and *CLV3:GUS* have already been described (Sieburth et al. 1998; Brand et al. 2002; Gallois et al. 2002). The gene trap line J2301 (from Jim Haseloff, Cambridge University) was obtained through the Nottingham *Arabidopsis* Stock Centre (<http://nasc.life.nott.ac.uk>). UAS:WUS was constructed with the UAS sequence (HindIII–BamHI, from the same vector used to create the gene trap lines), the WUS cDNA (BamHI–SpeI; Gallois et al. 2002) and the NOS terminator (XbaI–PstI, from pCGN18; Krizek and Meyerowitz 1996) inserted between the SmaI and PstI sites of pZP222 (Hajdukiewicz et al. 1994). The plasmid was transformed into *Arabidopsis thaliana* Landsberg-*erecta* (*L-er*) by the floral dip method (Clough and Bent 1998) and lines were selected that segregated a single UAS:WUS locus (based on gentamycin resistance). GAL4–VP16-directed WUS expression was analyzed in the progeny between homozygous UAS:WUS and gene trap lines, with crosses between *L-er* and the gene traps as controls. *ANT:GFP* was generated by inserting a 4.2-kb region upstream of the *ANT* initiation codon into the HindIII and BamHI sites of mGFP5-ER (Haseloff 1999) and transformed in *Arabidopsis thaliana* Columbia background. Expression in more than 20 independent lines was essentially as described by Schoof et al. (2000; Y. Mizukami, unpubl.).

For LEAFY (LFY) overexpression, *Arabidopsis thaliana* Columbia were transformed by the floral dip method with pDW151 (Weigel and Nilsson 1995). Expression of LFY mRNA in the roots of the transformed lines was checked by RT–PCR with primers 5′-GATTCCGGTACTCATCACGC-3′ and 5′-GGCTTGTAACAAGCCTGACGCCA-3′.

### *Growth conditions*

Seeds were surface-sterilized by chlorine gas by being kept 7 h in a desiccator with a mixture of 100 mL commercial bleach and 3 mL concentrated hydrochloric acid in a fume hood. Sterile seeds were plated on GM medium (Valvekens et al. 1988), stratified for 2 d at 4°C, and grown in 16 h light/8 h dark cycles (fluorescent lights at ~100 μmole photons m<sup>-2</sup>sec<sup>-1</sup>) at 18°C–20°C. For auxin treatment, GM was supplemented with 5 μM NAA solution (Sigma).

### *Activation of Cre recombinase*

Wild-type (*L-er*) or 35S:*LFY* plants were emasculated and fertilized 2 d later with 35S:*lox-uidA-lox-WUS*; *hsp18.2:Cre* pollen. For heat shock, seeds were plated on GM medium and sealed plates were incubated for 30 min at 38°C.

### *Microscopy*

mRNA in situ hybridization on tissue sections with digoxigenin (DIG)-labeled WUS or CLV3 cDNA was as described (Fobert et al. 1996). Double labeling was as described (Fobert et al. 1996), with the WUS probe labeled with fluorescein, developed with Fast Red TR/Naphthol AS-MX (Sigma)

and DIG-labeled CLV3 developed with BCIP/NIBT. For whole-mount GUS detection, tissues were fixed for 10 min in ice-cold 90% acetone and stained for GUS as described (Sieburth et al. 1998). For GUS detection in sectioned tissues, roots were first stained for GUS for 90 min at 30°C, followed by fixation with and sectioning as for in situ hybridization. For root confocal imaging, cell outlines were marked by staining with 50 μM propidium iodide. A Leica TCS SP microscope was used, with excitation set at 488 nm; emission was filtered to 500–550 nm (GFP) or 600–660 nm (propidium iodide), or was not filtered, for bright field. GFP-negative controls gave no signal in the GFP channel with the settings used. For cryo-scanning electron microscopy, seedlings were frozen in nitrogen slush at –190°C. Ice was sublimated at –90°C; the specimen was sputter coated and examined on a Philips XL 30 FEG SEM fitted with a cold stage. Images were processed (color balance, contrast, cropping, orientation) using Adobe Photoshop 5.0.

### *RNA extraction and RT–PCR*

RNA was extracted using TRI reagent (Sigma). RT–PCR were carried out with Superscript Rnase H<sup>-</sup> reverse transcriptase (GIBCO-BRL) on 1 μg total RNA according to the supplier's instructions. One-twentieth of the RT product was used for each subsequent PCR amplification. Amplification was initiated by adding *Taq* polymerase at 94°C, followed by 25–35 cycles of 30 sec at 94°C, 30 sec at 55°C, and 60 sec at 72°C. The primers used were as follows: LEC1, 5′-ACCAGCTCAGTCATAGTAGCCG-3′ and 5′-CTTATACTGACCATAATGGTCAAA-3′; FUS3, 5′-GAATGCAAGGAAGGGATTCTCA-3′ and 5′-CCCAAACCATCGAA TGTCCGAAC-3′; AGL15, 5′-ATCGAGATAAAGAGGATCGAGA-3′ and 5′-GAGAAAGCTCCTCAGTTCTTG-3′; WUS, 5′-AGTCGGATC CAACACACATGGAGCCGCCAC-3′ and 5′-CGGCTCTAGAGCTAG TCCAGACGTAGCTCA-3′; CLV3, 5′-CTTTGGATCCAAAAATGGAT TCTAAAAGCTTTG-3′ and 5′-ATAATCTAGAGCAACAAGAGAT TAGGTCAAG-3′. The APT cDNA (Moffatt et al. 1994) was amplified as a control with oligonucleotides 5′-CCTTCCCTTAAGCTCTG-3′ and 5′-TCCCAGAATCGCTAAGATTGCC-3′.

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