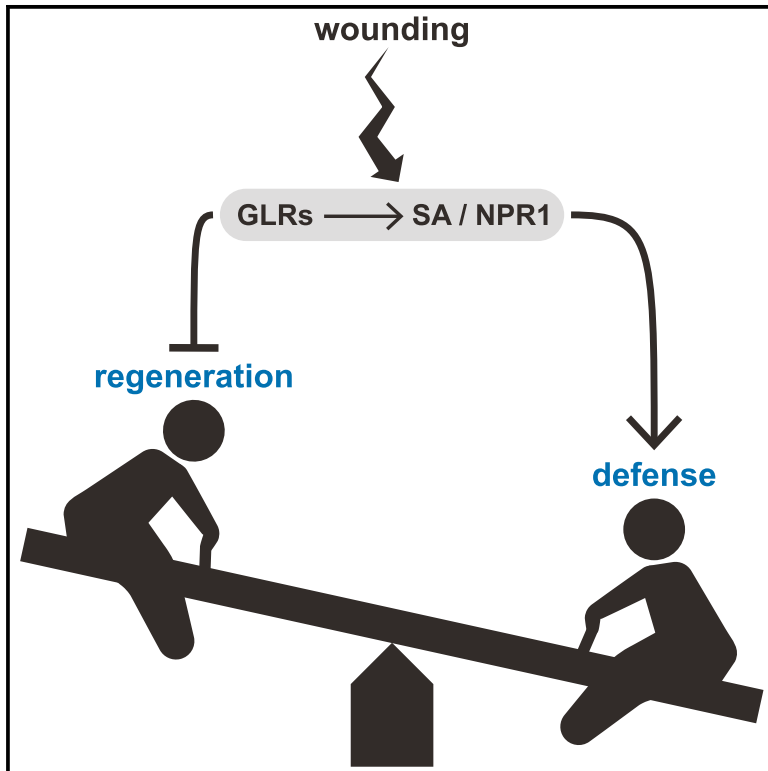


Developmental Cell

Plant glutamate receptors mediate a bet-hedging strategy between regeneration and defense

Graphical abstract



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In brief

When attacked, plants initiate defense responses and can also regenerate damaged body parts. It is not known whether the two responses are linked or represent trade-offs. Hernández-Coronado et al. show that plant glutamate receptors mediate a trade-off between defense and regeneration, favoring defense responses in an apparent “bet-hedging” strategy.

Highlights

- Perturbation of plant glutamate receptors improves regeneration in monocots and dicots
- The role of glutamate receptors in regeneration is mediated by salicylic acid signaling
- Increased salicylic acid response in older tissue is anticorrelated with regeneration
- Perturbing glutamate receptors improves regeneration of recalcitrant maize B73 lines

Article

Plant glutamate receptors mediate a bet-hedging strategy between regeneration and defense

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SUMMARY

Wounding is a trigger for both regeneration and defense in plants, but it is not clear whether the two responses are linked by common activation or regulated as trade-offs. Although plant glutamate-receptor-like proteins (GLRs) are known to mediate defense responses, here, we implicate GLRs in regeneration through dynamic changes in chromatin and transcription in reprogramming cells near wound sites. We show that genetic and pharmacological inhibition of GLR activity increases regeneration efficiency in multiple organ repair systems in *Arabidopsis* and maize. We show that the GLRs work through salicylic acid (SA) signaling in their effects on regeneration, and mutants in the SA receptor *NPR1* are hyper-regenerative and partially resistant to GLR perturbation. These findings reveal a conserved mechanism that regulates a trade-off between defense and regeneration, and they also offer a strategy to improve regeneration in agriculture and conservation.

INTRODUCTION

Plants have evolved intricate coping mechanisms to respond to biotic and abiotic insults. Two hallmarks of plant's response to attacks are the rapid initiation of defense responses and the replacement of damaged or lost body parts by whole organ regeneration (Ikeuchi et al., 2019; León et al., 2001). The first steps in both regeneration and defense pathways depend on the stress signals instigated by wounding (Florentin et al., 2013; Savatin et al., 2014), but it is not clear whether their circuitry is wired for differential regulation or their positive activation is linked. For example, the regulation of regenerative processes could require defense responses or rely on the same signals for activation.

There is a well-developed school of thought that growth and defense are trade-offs that emerge from limited resources (Huot et al., 2014; Züst and Agrawal, 2017; Guo et al., 2018). For example, regeneration calls upon many of the same processes as meristematic growth, such as cell division and organogenesis. However, defense and regenerative responses are tightly linked in large part through hormone and calcium signaling and their rapid activation after wounding (Hander et al., 2019; Hoermayer and Friml, 2019; Moeder et al., 2019; Shanmukhan et al., 2020).

In defense responses, the plant hormone jasmonic acid (JA) is a critical signal in the response to necrotrophic pathogens, but

JA also has roles in abiotic stress and development (Howe et al., 2018). The primary JA receptor is *CORONATINE-INSENSITIVE 1 (COI1)*, a key regulator of JA signaling (Katsir et al., 2008). Jasmonate promotes an interaction between COI1 (an E3 ligase complex) and JAZ repressor proteins, degrading the repressors and permitting downstream transcriptional responses (Howe et al., 2018). In addition, JA is also a necessary signal for root regeneration in multiple contexts, promoting cell division during repair (Zhang et al., 2019; Zhou et al., 2019).

Another primary defense-response hormone is salicylic acid (SA), which has roles in defense against biotrophic pathogens, mediating both independent and overlapping responses with JA (Yang, 2015). The primary receptor for SA is *NONEXPRESSOR OF PR GENES1 (NPR1)*, with *NPR3/NPR4* paralogs acting as repressors of *NPR1* (Ding et al., 2018). According to current models, NPR1 is part of a transcription complex interacting with DNA-binding transcriptional co-activators (TGAs) and the mediator/polymerase II complex with SA somehow promoting NPR1 activity to initiate broad transcriptional responses (Peng et al., 2021). Although other pathways mediate defense and regeneration (Ikeuchi et al., 2019; Marhavý et al., 2019), the prominent role of JA and SA in defense and wound responses make them good candidates to regulate interactions between defense and regeneration.

Plant glutamate receptor-like proteins (GLRs) also have a well-documented role in defense downstream of JA (Mousavi et al.,

2013; Nguyen et al., 2018). GLRs are non-specific ion channels, with a permeability for calcium Michard et al. (2011), Ortiz-Ramírez et al. (2017). In defense, GLRs have been implicated in calcium signaling that activates JA-dependent systemic signaling to prepare distant organs for a potentially imminent attack (Mousavi et al., 2013; Nguyen et al., 2018). In development, GLRs have been shown to signal through calcium fluxes associated with male gamete function during fertilization (Michard et al., 2011; Ortiz-Ramírez et al., 2017; Wudick et al., 2018b). GLR mutants also show defects in defense against biotrophic pathogens, suggesting that they may also have a role in the SA pathway (Forde and Roberts, 2014). Thus, GLRs mediate some of the plant's most rapid responses to environmental and developmental inputs. Their role in defense responses has been well established, but their role in regenerative responses remains unclear.

Exploring the regulatory wiring between defense and regeneration has practical implications for human nutrition, as many staple crops, such as potato, cassava, and yam, are grown via regenerative (clonal) propagation (Chin and Tan, 2018; Ikeuchi et al., 2019; Nishimura et al., 2005). Moreover, modern biotechnology approaches commonly employ genetic transformation and regrowth of a regenerative cell mass in plants known as callus (Ikeuchi et al., 2019; Yadava et al., 2016). However, many advanced cultivars in some of the world's most important crops, such as maize and wheat, are recalcitrant to regeneration (Altpeter et al., 2016; Delporte et al., 2012; Ikeuchi et al., 2019).

Current methods to increase regenerative capacity in crops rely on overexpression of plant stem cell/meristem initiation factors, such as *Baby Boom* and *Wuschel* (Lowe et al., 2016; Masters et al., 2020;). However, overexpression of these early growth and meristem regulators can lead to severe developmental effects, and they are impractical for wide use in a variety of plant species. Any mechanisms that could retune the balance between defense and regeneration—preferably limited to the regenerative phases—could be used to improve regeneration for biotechnology, conservation, and propagation of staple food crops.

Here, we uncover mechanisms that regulate defense and regeneration in an antagonistic manner. The suppression of *GLR* function significantly increased regeneration in *Arabidopsis* while downregulating defense responses. Transient block experiments with chemical inhibitors of *GLR* signaling showed that suppression of *GLR* activity at the time of injury and for several additional hours improved regeneration, implying an immediate and extended role for GLRs in wound responses that mediate regeneration. Although GLRs have been shown to signal through JA, a mutant in the JA receptor, *COI1*, shows lower regeneration rates in most assays, suggesting the *GLR* mutant phenotype was not mediated by JA. On the other hand, a mutant in the SA receptor, *NPR1*, showed dramatically higher regeneration efficiency than wild type. We also found that native SA markers anticorrelate with regeneration efficiency, suggesting that declining regeneration efficiency with age of tissue could be due, in part, to increasing SA levels. Finally, we show that GLRs have a conserved role in maize, as the *GLR* antagonist CNQX (6-Cyano-7-nitroquinoxaline-2,3-dione) and an SA inhibitor were effective in increasing callus formation in leaf and embryo explants of the recalcitrant B73 maize line. Overall, we

conclude that the GLRs regulate the balance between regeneration and defense in an array of contexts and different plant species, favoring defense responses. They now provide a “druggable target” for enhancing plant regeneration and clonal propagation.

RESULTS

Chromatin remodeling is a key step in cellular reprogramming during plant regeneration (Ikeuchi et al., 2019; Rymen et al., 2019), highlighting processes involved in the dramatic alteration of cell fates. To characterize chromatin dynamics during cellular reprogramming, we targeted cells in the microdomain of the root tip undergoing cell-fate transitions in the first 24 h post-injury in *Arabidopsis*. In these experiments, we used a root regeneration system in which we cut off most of the root meristem, including stem cells and cap, all of which are rapidly replaced within a day during a re-establishment of the proximo-distal (lengthwise) and radial (widthwise) axes (Figure 1A; Efroni et al., 2016; Sena et al., 2009). The expression domain of the functional marker of the root stem cell niche, *MONOPTEROS* (*MP*, *MP::MP-GFP*, Cole et al., 2009), is largely removed by root tip excision but then rapidly reappears in the regenerating stem cell niche region within 2 h after the cuts. We used this highly specific marker to collect reprogramming cells (hereafter, the reprogramming domain) at 2.5, 14, and 16 h after cuts via fluorescence-activated cell sorting (FACS) after rapid digestion of cell walls, followed by isolation of protoplast nuclei (Figure 1B).

The chromatin accessibility landscape changes rapidly in reprogramming cells

Up to 2,000 nuclei from the highly localized reprogramming domain were isolated from protoplasts and subjected to small-sample assay for transposase accessible chromatin (ATAC)-seq protocols (Buenrostro et al., 2013, 2015, STAR Methods). As a control for protoplast effects, open chromatin regions (ATAC-seq peaks) from nuclei of whole, uncut root tips were compared with nuclei taken from the same region but subjected to cell wall digestion before harvest, showing minimal differences. We then used MACS2 (Zhang et al., 2008) to call peaks and then DESeq2 (Love et al., 2014) to find regions of differential chromatin accessibility between control cell replicates and each of the post-cut time points (Figure 1B; STAR Methods).

Replicates for the 14-h cut time points had the strongest signals, showing about 750 promoter and 500 coding regions with a significant decrease in accessibility, whereas nearly 7,000 peaks in the promoter and 3,000 peaks in coding regions gained accessibility (examples in Figure 1C). A gene ontology analysis of loci associated with significantly changing ATAC peaks at 14 h revealed processes, such as auxin and wounding response (Figure 1D), providing validation that chromatin accessibility reflected categories of genes known to be upregulated in root regeneration.

Interestingly, among the highly significant gene ontology (GO) terms were calcium ion transport and calcium channel activity ($p < 10^{-3}$ and $<10^{-2}$, respectively, Figure 1D), which included genes from several *GLR* ion channels. In particular, the majority of the 20 annotated GLRs in *Arabidopsis* showed significant changes in open chromatin in their promoter coding regions,

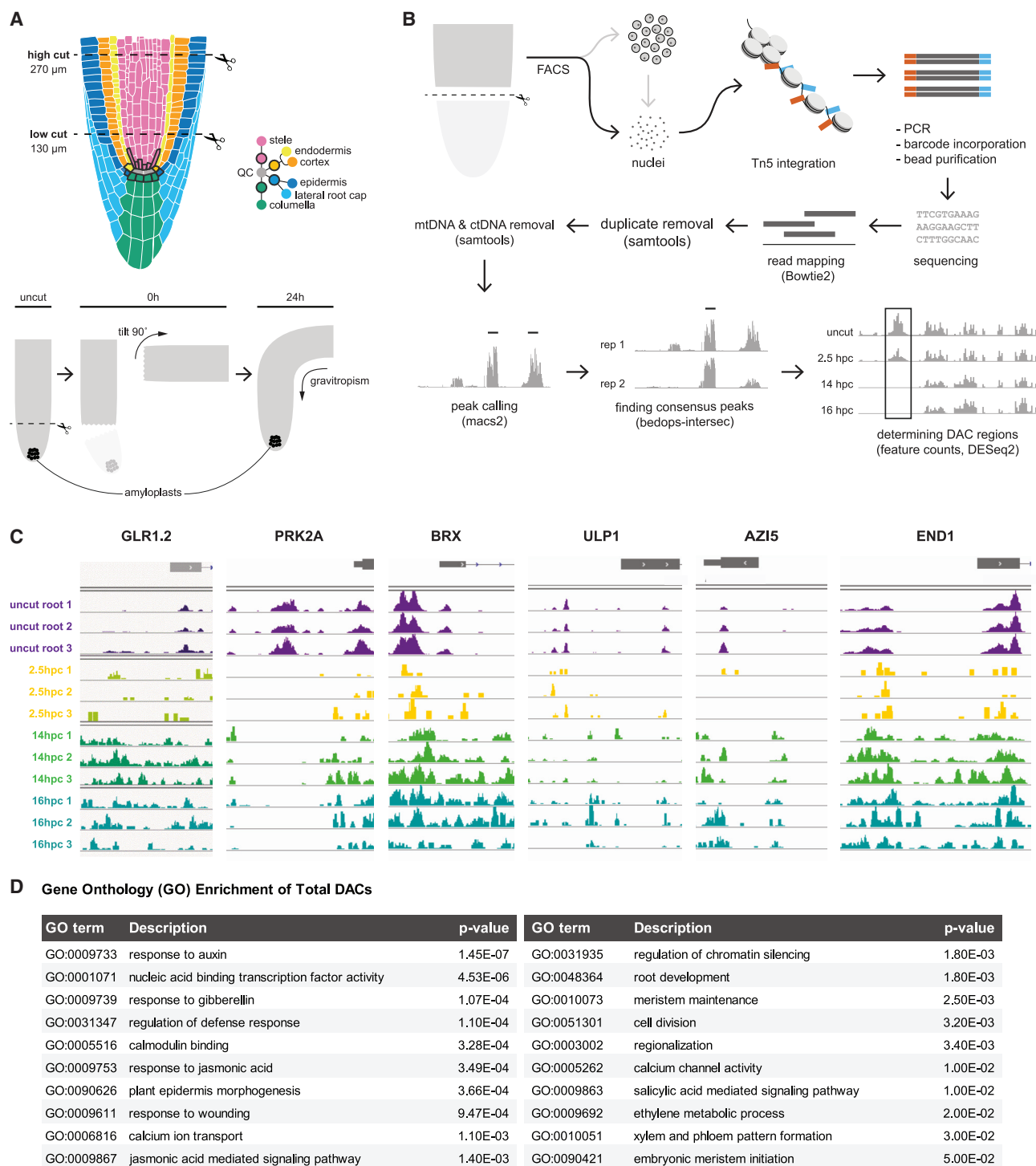


Figure 1. The chromatin accessibility landscape changes rapidly in reprogramming cells

(A) Schematics of the root anatomy and the root regeneration system in which the meristem (top) is excised. Regeneration can be scored by the re-specification of gravity-sensing amyloplasts in newly formed columella cells (below), showing, from left to right, uncut and cut root followed by regeneration of the root tip.

(B) Protocol for collecting nuclei and analyzing open chromatin regions in specific regenerating cells (DAC = differentially accessible chromatin) with arrows indicating workflow.

(C) Examples of changes in chromatin accessibility of the same region before and after cutting at various time points, showing both the gain and loss of chromatin accessibility in regulatory regions and gene bodies. Samples are taken in uncut roots or different hpc (hours post cut).

(D) GO terms associated with genome-wide changes in chromatin accessibility (either opening or closing) during regeneration. GO term analysis was performed using all genes associated with DAC regions versus all genes in the genome, with pvals indicating degree of over-representation by a Fisher's exact test.

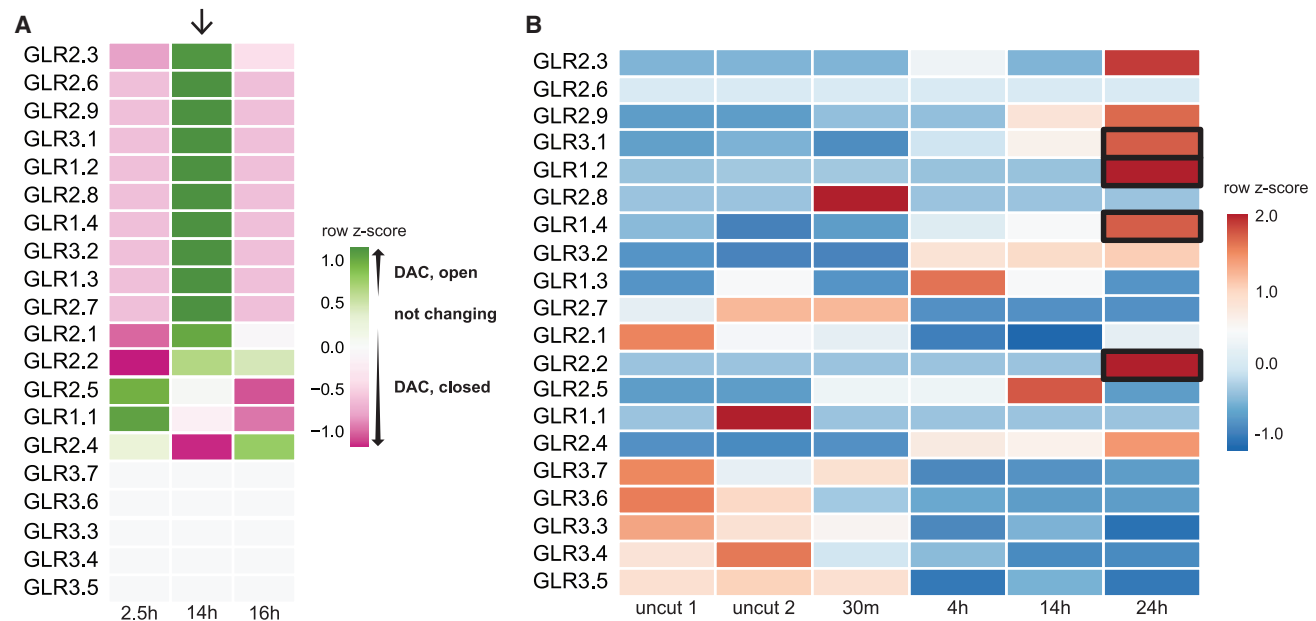


Figure 2. The GLR family undergoes rapid chromatin modification and transcriptional regulation

(A) Heatmap showing trends in chromatin accessibility among GLRs during regeneration at three time points representing hours post cut. Color represents the relative accessibility (per ATAC-seq analysis in Figure 1B) as a row Z score, which is standard deviations from the row mean. Arrow points to a trend of opening chromatin at 14 h among a subset of GLRs.

(B) Heatmap depicting GLR transcriptional analysis from RNA-seq profiles in a time course of uncut and regenerating root stumps at four time points after cutting (row Z score followed by log₂ normalization). Boxes highlight several GLRs that increase in expression during the first 24 h of regeneration and whose chromatin becomes more accessible.

with many showing increases in accessibility at 14 h (Figure 2A). At the transcriptional level, many GLRs were upregulated during the early stages of regeneration potentially influenced by changes in chromatin accessibility (Figure 2B). For example, *GLR1.2* showed a significant increase in chromatin accessibility in its promoter region at 14 h (Figure 2A) and an increase in mRNA abundance at later time points in the RNA-seq time course analysis of regenerating roots (Figure 2B). The potential involvement of the GLRs was intriguing because they could mediate early signals that trigger regenerative processes. Although chromatin accessibility was not always correlated with gene expression, we focused on GLRs that showed a trend of increased accessibility during regeneration together with either increasing expression during regeneration or high expression levels in the reprogramming zone (e.g., *GLR1.2*, *1.4*, *2.2*, *3.1*, *3.2*, *3.3*; Figures 2A and 2B).

Perturbation of GLR signaling improves regeneration

We initially hypothesized that GLRs, along with their well-characterized role in defense, could mediate ion signals that trigger cellular dedifferentiation and regeneration. In the root regeneration system (hereafter root-from-stump), regeneration can be screened quantitatively by the reprogramming of vascular cells to columella cap cells, which sense gravity and quickly restore downward growth (Sena et al., 2009). However, single mutants in several candidate genes, including *GLR1.2* (At5g48400), *GLR1.4* (At3g07520), and *GLR3.3* (At1g42540; Table S1), showed no difference in regeneration compared with wild type.

To address potential redundancy, we focused on the higher order *glr1.2/1.4/2.2/3.3* mutant, which included several of our

candidates and was previously shown to be severely impaired in calcium flux in pollen tubes (Wudick et al., 2018b). However, counter to expectations, the quadruple *glr1.2/1.4/2.2/3.3* mutant showed twice the regeneration efficiency as wild type in cuts in the more differentiated regions of the root (270 μ m from the tip, Figure 3A). In addition, other higher order *GLR* mutants also showed improvement in regeneration, although not as dramatic as the *glr1.2/1.4/2.2/3.3* combination (Figure 3A). None of the mutants, including the quadruple combinations, showed root growth phenotypes, with root length and meristem size indistinguishable from wild-type Columbia-0 (Col-0) plants (Figures S1A and S1B), providing evidence that enhanced regeneration was not trivially due to a larger meristem before cutting. We note that none of the mutants showed a regeneration phenotype (increased or decreased regeneration) in the more competent, younger region of the meristem (within $\sim$ 130 μ m from the tip, Figure S2A). This age effect of *GLR* mutations on regeneration was a consistent phenomenon in our subsequent experiments that we pursue further on.

To explore the effect of the strong mutant combination, we examined the expression domains of the *GLR* using an atlas of tissue-specific expression (Li et al., 2016), observing that GLRs *1.2*, *1.4*, *2.2*, and *3.3* had largely complementary patterns in root tissues (ground tissue, pericycle, epidermis, and vascular tissue, respectively, Figure S2B). This indicates that our mutant combination covered almost all domains of the meristem. We surmise the *glr1.2/1.4/2.2/3.3* combination had an additive effect in which the regeneration phenotype was not detectable until signaling from a critical number of tissues was blocked.

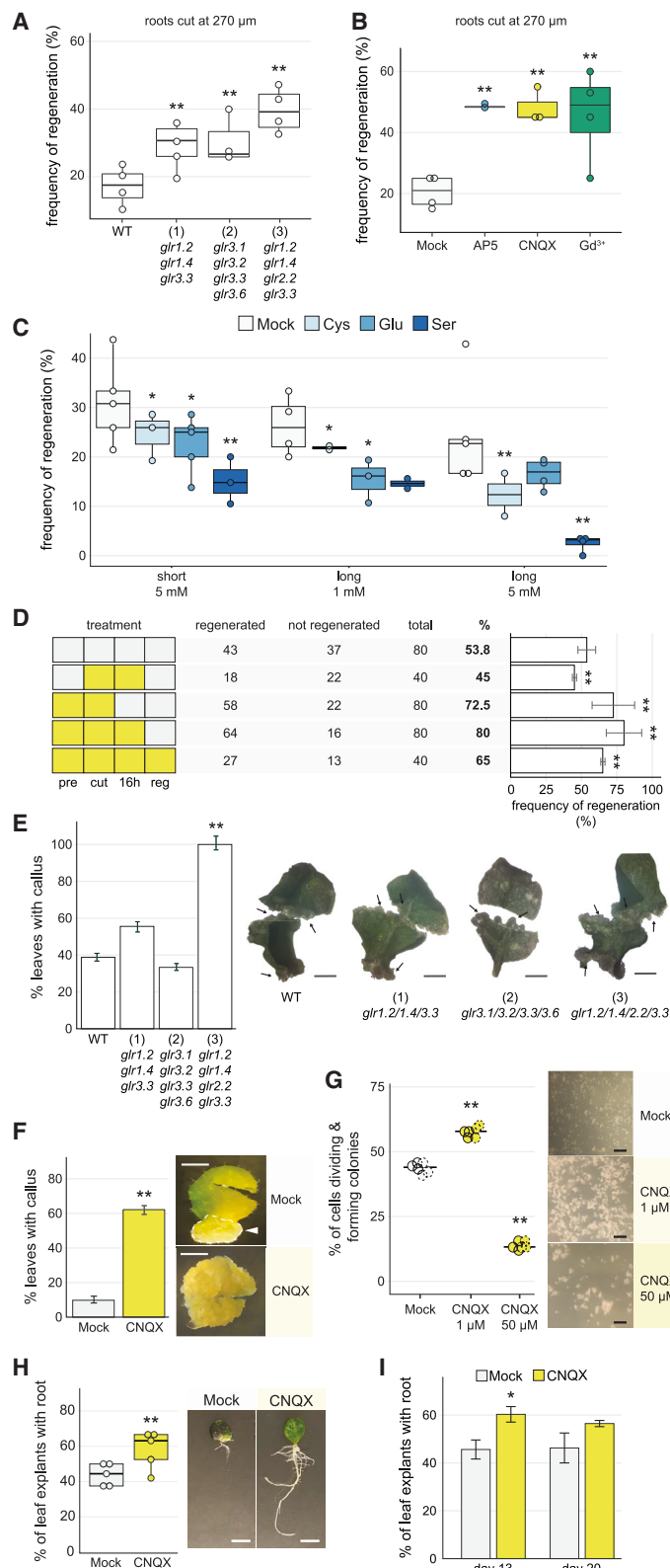


Figure 3. Perturbation of GLR signaling improves regeneration

(A) Regeneration frequency of wild-type plants (Col-0) compared with triple and quadruple *glr* mutants: (1) *glr1.2/1.4/3.3*, (2) *glr3.1/3.2/3.3/3.6*, and (3) *glr1.2/1.4/2.2/3.3* (chi-square test, Fisher's combined probability test; $n > 19$ for each genotype in each of 4 trials; $**p < 10^{-7}$).

(B) Constitutive post-injury treatment (cut on treatment plates and kept on treatment plates for the duration of the experiment) of cut roots on various calcium signaling antagonists, showing enhanced regeneration on all treatments. Roots were grown on normal media, cut, and transferred to treatment plates (chi-square test, Fisher's combined probability test; $n > 20$ for each treatment in each of 4 trials, Gd^{3+} $n > 12$ one trial, $**p < 10^{-11}$).

(C) Regeneration frequency in Col-0 plants treated with different amino acid agonists for GLRs. Regeneration frequencies were diminished in all treatments (chi-square test, Fisher's combined probability test; $n > 25$ for each agonist treatment in each condition, 6 trials short treatment, 4 trials long treatment, $*p < 0.02$, $**p < 10^{-6}$). Roots were grown on standard media, cut, and transferred to agonists. For "short" treatment, plants were exposed to 5 mM of L-glutamate (Glu), L-cysteine (Cys), or L-serine (Ser) for < 2 min and transferred back to $0.5 \times$ MS plates to assess regeneration. For "long" treatment, plants were maintained on plates with treatment until regeneration was assessed.

(D) CNQX transient "window" treatments, showing that inhibiting GLRs at the time of injury is critical for enhancing regeneration. Pattern at left shows treatment scheme with white boxes indicating mock and yellow boxes indicating CNQX treatment. Pre, 1 h pretreatment; Cut, cutting plate; 16 h, begins immediately after cutting and ends 16 h later; Reg, begins 16 h after cutting and continues for the duration of the experiment. (Chi-square, $n \geq 40$ for each treatment, $**p < 0.0008$).

(E) The *glr* quadruple mutant (3) shows enhanced callus formation, with bar graph showing percent of leaves that formed callus for each genotype, scored at 2 weeks post callus initiation. Right, representative images of callus formation in leaves from wild type, triple and quadruple GLR mutants, showing more extensive callus formation in mutants versus. Wild-type leaves that did form callus. In all cases, formation of callus occurred only in some regions of the leaf (partial). The error bars are proportional to the standard error of the pooled percentage computed using binomial distribution (chi-square test; $n \geq 32$ for each genotype; $**p < 10^{-9}$). Arrows indicate regions of callus formation. Scale bars, 0.5 cm.

(F) Callus formation in leaves treated with the GLR antagonist CNQX is enhanced, showing percent of leaves with callus. Images show roots in both conditions that formed callus. Arrow indicates region of callus formation in mock, while the treated sample is completely transformed to callus, indicating accelerated callus formation. The error bars are proportional to the standard error of the pooled percentage (chi-square test; $n > 48$ for each treatment; $*p < 10^{-29}$). Scale bars, 0.5 cm.

(G) Callus microcolony formation from single cells is enhanced by addition of $1\text{-}\mu\text{M}$ CNQX to the liquid media (chi-square test; $n = 8.0 \times 10^4$ cells, where frequency of callus formation was estimated from hemacytometer readings of 3 samples from each of two biological replicates and extrapolated; $**p < 10^{-30}$). At right, representative images of microcolony formation. Scale bar, 2 mm.

(H) Root-from-leaf regeneration under CNQX treatment, showing more leaves with roots in CNQX treatment. (Chi-square test; Fisher's combined probability test, $n \geq 18$ for each treatment in each of 6 trials, trial 3, $n = 12$; $**p < 0.0005$) Image shows more extensive root system of CNQX-treated leaves. Scale bars, 65 mm.

(I) Root-from-leaf regeneration under CNQX treatment assessed at 13 and 20 days. The difference in *de novo* root formation between treated and untreated leaves diminished over 20-day incubation (chi-square test; $n = 40$ for each treatment at each time point; $*p < 0.03$).

To test whether the GLRs function pre or post-wounding, we took advantage of *GLR* channel inhibitors used in animals and plants by conditionally blocking *GLR*-mediated signaling after wounding (Krystal et al., 1999; Michard et al., 2011; Wudick et al., 2018a). Gadolinium (Gd^{3+}) is a broad inhibitor of calcium channels, whereas AP5 and CNQX were originally described as antagonist of mammalian NMDA- and AMPA-type glutamate receptors, respectively (Krystal et al., 1999; Michard et al., 2011) and later found to efficiently inhibit various *GLR* members in *Physcomitrium* and *Arabidopsis* (Michard et al., 2011; Ortiz-Ramírez et al., 2017; Wudick et al., 2018a; Mou et al., 2020).

Using these inhibitors, plants were grown on standard plates, their meristems excised, and then transferred to plates containing calcium channel antagonists or to standard media (mock) for the remainder of the experiment. All three pharmacological treatments dramatically enhanced regeneration efficiency (Figure 3B), reaching a nearly 3-fold increase in regeneration in the challenging proximal wound cuts and exceeding the effects of the *glr* quadruple mutant (Figure 3A versus 3B). The fact that the glutamate receptor antagonists had the same effect as the broad calcium signaling antagonist Gd^{3+} suggests that GLRs—and not other calcium channels—mediate the effect on regeneration. We surmise that chemical treatments had a more dramatic effect on regeneration than mutants due to even higher order redundancy among the GLRs, many of which are expressed in the root (Figure S2B).

In addition, even brief post-injury treatment of roots with the canonical *GLR* agonists L-cysteine, L-serine, and L-glutamate, which has been shown to induce defense responses (Goto et al., 2020), resulted in the opposite effect, dramatically inhibiting regeneration (Figure 3C). Thus, *GLR* signaling strength has a direct effect on regeneration both above and below native levels.

Using the *GLR* inhibitor CNQX, we tested the time window in which glutamate receptor signaling affects regeneration. A 1-h pretreatment alone followed by transfer of cut roots onto mock plates improved regeneration significantly (Figure 3D), suggesting that the GLRs function, in large part, immediately after wounding. We could not observe enhancement of regeneration with a 16- or a 4-h CNQX treatment post-cutting alone without pretreatment (Figure 1D, compare rows 1 and 2, Figure S2C). However, when a pretreatment was combined with a 16-h incubation on CNQX post-cutting, regeneration increased to even higher levels than the maximal treatment (1-h pretreatment + transfer to CNQX plates for 3 days, Figure 3D). Thus, CNQX also improves regeneration in the hours after injury. Taken together, these results suggest that blocking the function of a redundant set of *GLR* acts immediately upon injury and continuously for several hours to suppress regeneration after wounding.

Plants can regenerate many different organs in varied developmental contexts (Ikeuchi et al., 2019; Sinnott, 1960). Thus, to ask if the GLRs function broadly in regeneration, we tested *GLR* mutants and chemical treatments in other regeneration systems in *Arabidopsis*. Many types of regeneration occur through callus, a root-like mass of cells analogous to blastema in animals, which gives rise to shoots and roots (Birbaum and Sánchez Alvarado, 2008; Sugimoto et al., 2010, 2011). To test the role of *GLR*-mediated signaling in callus formation (hereafter, callus-from-leaves), early-life-stage (8 days post germination, dpG) or later-life-stage (15 dpG) leaves were excised at the

base of the petiole, cut in half, and cultured on callus inducing media (CIM) plates for 2 weeks. Although no difference in callus formation efficiency was observed in early-stage leaves (Figure S2D), the *glr1.2/1.4/2.2/3.3* mutant showed an increase in callus formation in older leaves, with more than a 2-fold higher efficiency than wild type for the quadruple mutant (Figure 3E). We again draw attention to the fact that the more dramatic effect on later-stage leaves is reminiscent of the *glr* quadruple mutant's more pronounced effect on older cells in the root meristem (previous section)—a repeat of the so-called the age effect.

We repeated the callus-from-leaves experiments using chemical treatments, plating young (8 dpG) and later-stage (15 dpG) leaves on CIM plates containing CNQX or mock treatments (STAR Methods). Consistently, *GLR* inhibitors increased callus formation in normally older recalcitrant leaves (Figure 3F). After 15 days on CIM, untreated leaves formed callus at a frequency of 10%, whereas callus formation in CNQX-treated leaves was up to six times higher, reaching about ~60% regeneration in repeated trials (Figure 3F). In other regeneration contexts, CNQX improved the formation of microcalli from single cells (protoplasts, Figure 3G) and also improved root formation from the base of *Arabidopsis* leaves (hereafter, root-from-leaf, Chen et al., 2014; Figure 3H). The difference in root formation between treated and untreated leaves in the root-from-leaf system diminished over longer-term incubation (20 days, Figure 3I). However, we note that the effects of treatment were still critical for overall regeneration success in other systems (e.g., Figure 3B). Overall, the results show that GLRs inhibit the regeneration response in a broad set of regeneration contexts in *Arabidopsis*.

CNQX affects regeneration through the salicylic acid pathway

To examine the downstream effects of blocking *GLR* signaling after wounding, we generated an RNA-seq time course of control versus CNQX-treated roots after cutting (uncut [2×], 30 min, 4 h, 14 h, and 24 h). We dissected root tissues to harvest only the distal root tip or regenerating root stumps containing the reprogramming domain. We filtered the dataset for the top 10% most variable genes and performed a differential time series analysis on control versus CNQX-treated samples using ImpulseDE (Sander et al., 2017), identifying about 1,400 differentially expressed (DE) transcripts categorized into 11 different patterns (STAR Methods).

Most DE genes fell into a pattern of repression by CNQX throughout the time points as illustrated for cluster 9 (Figures 4A and S3A). These clusters were overwhelmingly represented by stress- and defense-responsive GO terms (response to abiotic stimulus $p < 10^{-8}$, response to stress $p < 10^{-19}$, response to other organisms $p < 10^{-7}$, Table S2). We then specifically examined genes annotated with a role in “defense.” Many of these genes were induced in mock but failed to show upregulation in the CNQX or in the *glr* quadruple mutant at 24 h post cut (hpc). (Figures 4B and 4C; Table S3). Thus, CNQX treatment appears to downregulate many stress responses in reprogramming cells (Figure 4B), showing that the GLRs mediate a response to root meristem damage next to the wound site.

Given that JA is involved in both root regeneration (Zhang et al., 2019; Zhou et al., 2019) and *GLR*-mediated defense (Mousavi et al., 2013; Nguyen et al., 2018), we tested a fertile

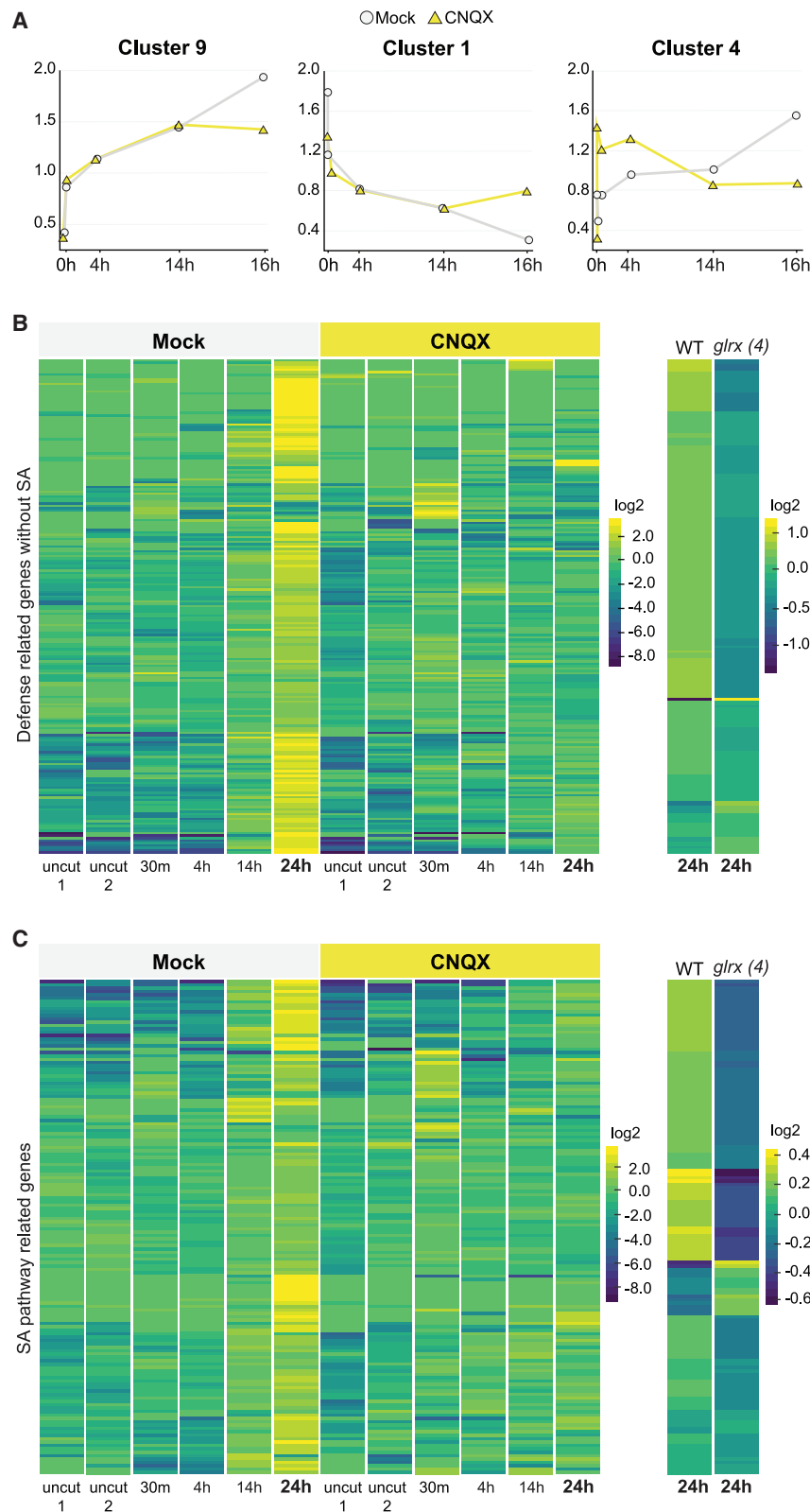


Figure 4. A time course shows dramatic changes in stress responses with CNQX treatment

(A) Plots summarizing 3 of the 11 different patterns found among approximately 1,400 differentially regulated transcripts along a regeneration time course (0 min, 30 min, 4 h, 14 h, and 24 h), using ImpulseDE (see experimental model and subject details).

(B and C) Heatmaps of defense-annotated genes (B) and SA responsive genes (C) that are upregulated in regeneration at 24 h in untreated roots, showing a failure to upregulate the same genes in CNQX treatments and in the *glrx4* mutant (*glr1.2/1.4/2.2/3.3*). For treatments, plants were grown on standard plates and then transferred to mock or CNQX plates (50 μ M). Read counts were row normalized and $\log_2$ transformed. Col-0 versus *glr1.2/1.4/2.2/3.3* mutant plants were profiled at 24 h after injury only and normalized separately. Genes in heatmaps are listed in Table S3.

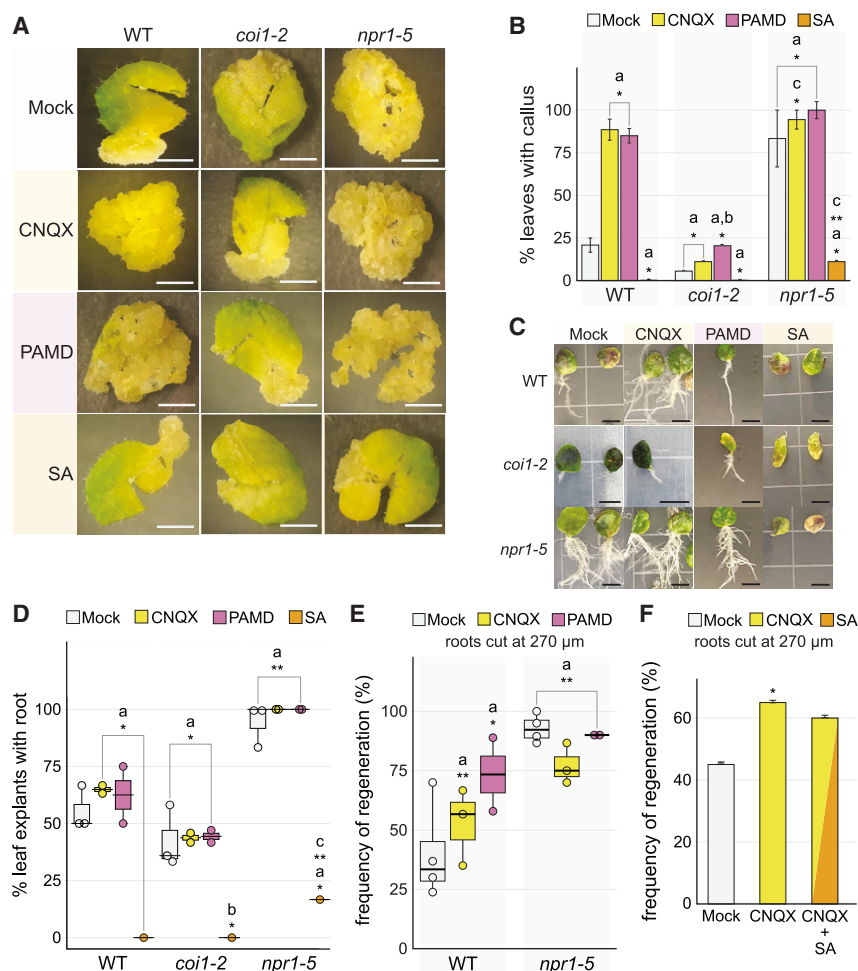


Figure 5. CNQX affects regeneration through the SA pathway

(A) Representative images of callus formation in leaves of wild type (Col-0), *coi1-2*, and *npr1-5* on Mock, CNQX, PAMD, or SA, all under constitutive post-injury treatment. Productive callus appears as clusters of amorphous, pale (yellow) cell masses. Scale bars, 0.5 cm.

(B) Quantification of callus formation shown in (A) (chi-square test; $n \geq 18$ for each genotype in each treatment; ^asignificant difference with respect to Col-0 Mock, $*p < 10^{-6}$, $**p < 10^{-11}$; ^bsignificant difference with respect to *coi1-2* Mock, $*p = 0.003$; ^csignificant difference with respect to *npr1* Mock, $*p < 0.04$, $**p < 10^{-9}$).

(C) Root-from-leaf regeneration in wild type (Col-0), *coi1-2*, and *npr1-5* treated on Mock, CNQX, SA or PAMD under constitutive post-injury treatments. Representative images show differences in responses based on genotype and treatment. Scale bars, 65 mm.

(D) Quantification of root-from-leaf regeneration shown in (C) (chi-square test, Fisher's combined probability test; $n \geq 12$ for each treatment in each genotype in each of 3 trials; ^asignificant difference with respect to Col-0 Mock, $*p < 0.03$, $**p < 10^{-7}$; ^bsignificant difference with respect to *coi1-2* Mock, $*p < 0.02$; ^csignificant difference with respect to *npr1* Mock, $*p < 10^{-20}$).

(E) Regeneration frequency of Col-0 and *npr1-5* mutant on different constitutive post-injury treatments. The error bars are proportional to the standard error of the pooled percentage computed using binomial distribution (chi-square test, Fisher's combined probability test; $n \geq 18$ for each treatment in each genotype in each of 4 trials; ^asignificant difference with respect to Col-0 Mock, $*p < 0.05$, $**p < 10^{-6}$).

(F) Root regeneration frequency under combined constitutive post-injury treatment of CNQX and SA, showing that SA addition does not fully counteract GLR blockage by CNQX (chi-square test; $n = 40$; $*p < 0.02$).

hypomorph mutant of *COI1* (Katsir et al., 2008). If the GLR-mediated ion signaling inhibits regeneration through JA signaling, we reasoned that loss-of-function mutants in *COI1* would phenocopy GLR mutants.

However, the *coi1-2* hypomorph showed lower regeneration in callus formation (Figures 5A and 5B) and root-from-leaf regeneration (Figures 5C and 5D), the latter in agreement with previous reports (Zhang et al., 2019). In the root-from-stump assay, the *coi1-2* allele did not show consistent results (Figures S3B and S3C). Nonetheless, overall, the largely lower regeneration efficiency of the *coi1* mutant fails to implicate JA in mediating the GLR effect on regeneration.

On the other hand, the RNA-seq time course showed that many SA responsive genes were upregulated in regeneration under mock conditions but failed to show that upregulation on CNQX treatment and in *glr1.2/1.4/2.2/3.3* mutant (Figure 4C). In addition, we observed that SA responsive markers increased along the longitudinal axis of the root as cells matured away from the stem cell niche (Figure S4). The increase in SA response is inversely correlated with the competence of cells to regenerate

along the longitudinal axis of the root (Sena et al., 2009). The observation is analogous to a previous report showing that callus formation in garlic leaf decreased as leaf tissues matured and SA levels rose (Mostafa et al., 2020). These trends provided an intriguing connection between SA and the “age effect” we observed in several contexts in which perturbing GLRs improved regeneration only in older tissues. This led to the hypothesis that SA mediates GLR signaling in older tissues that normally inhibits regeneration.

To test the role of SA in GLR-mediated regeneration response, we assayed several mutants of the SA signal transduction pathway for regeneration phenotypes, with the receptor, *NPR1*, showing the most dramatic effect (Ding et al., 2018; Figures S5A–S5D). Compared with wild type, callus formation in the *npr1-5* mutant showed more than a 3-fold increase, including more complete reprogramming of explants to callus (Figures 5A and 5B). In root-from-leaf regeneration, the *npr1* mutant showed more than a 2-fold increase in efficiency (Figures 5C and 5D) and, similarly, in the root-from-stump assay, *npr1* roots showed an approximate 2-fold increase in regeneration in high cuts

(Figure 5E). Consistently, transient treatment with the SA inhibitor 4-phenyl-2-[[3-(trifluoromethyl)-aniline]methylidene]cyclohexane-1,3-dione (PAMD) showed higher regeneration than mock-treated wild-type roots in all three regeneration systems, although PAMD has a strong inhibitory effect on long-term growth (Figures 5A–5E; Seo et al., 2012). In contrast, treatment of roots with SA greatly inhibited regeneration in all the systems tested (Figures 5A–5D, S5E, and S5F). These strong effects with transient treatments support a role for SA signaling in the inhibition of regeneration in the post-injury environment.

Examining epistatic effects between the GLR and SA pathway, we observed at least partial insensitivity of *npr1* to regeneration enhancement by CNQX treatment in most assays (Figures 5A–5D). The insensitivity of *npr1* to CNQX was most apparent in the more challenging regeneration assays, such as high cuts in root-from-stump regeneration (Figures S5A and S5D). In addition, although wild-type plants showed increased regeneration in SA inhibitor PAMD, the same treatment had no effect on regeneration in the *glr1.2/1.4/2.2/3.3* mutant combination, showing resistance to the SA inhibitor (Figure S5E). The results are consistent with GLR perturbations already suppressing downstream SA signaling. Simultaneous treatment with CNQX and SA in wild-type plants showed only a small, non-significant decrease in root regeneration levels compared with CNQX treatment alone (Figure 5F). However, treatment of the *glr1.2/1.4/2.2/3.3* mutant with SA still inhibits regeneration as it does in wild type, a partial rescue of the wild-type response by apparent restoration of downstream SA signaling (Figure S4E). These results support a role for SA downstream of GLR signaling. Together, the results suggest that the GLR-mediated effects on regeneration work in large part through the SA signaling pathway.

Blocking GLRs severely disrupts ion flux, inhibits defense, and augments repair

The involvement of the GLR calcium channels in our regeneration assays implies a role for rapid ion signaling near the injury site. To test whether GLRs mediate ion signaling in our root-from-stump assay, we used a calcium-specific vibrating probe to measure extracellular calcium fluxes in wild type and *glr1.2/1.4/2.2/3.3* root stumps for several minutes after wounding. Without wounding, *glr1.2/1.4/2.2/3.3* roots did not show a significant difference in calcium flux when compared with wild type (Figure S6A). Upon wounding, wild-type roots showed high Ca^{2+} influx at the injury site for 10 min after which the influx decreased over the recovery time (Figure 6A, gray markers and gray trend line). In contrast, the *glr* quadruple mutant best-fit curve showed lower Ca^{2+} influx immediately after wounding and a faster recovery to stable levels (Figure 6A, blue markers and blue trend line).

The steeper decline in influx over time in the mutant curve amounts to a higher recovery constant compared with wild type ($K = 2.041$ versus $K = 0.485$, $p < 0.0001$), exhibiting lower levels of calcium influx within the first 5 min after wounding. At recovery phase, however, the *glr* quadruple mutant plateaued at an influx rate one order of magnitude higher than wild type (920.2 versus 74.6 $\text{pmol}/\text{cm}^2/\text{s}$, $p = 0.0002$; Figure 6A, compare fitted curves), indicating that a larger calcium influx is sustained in the mutant.

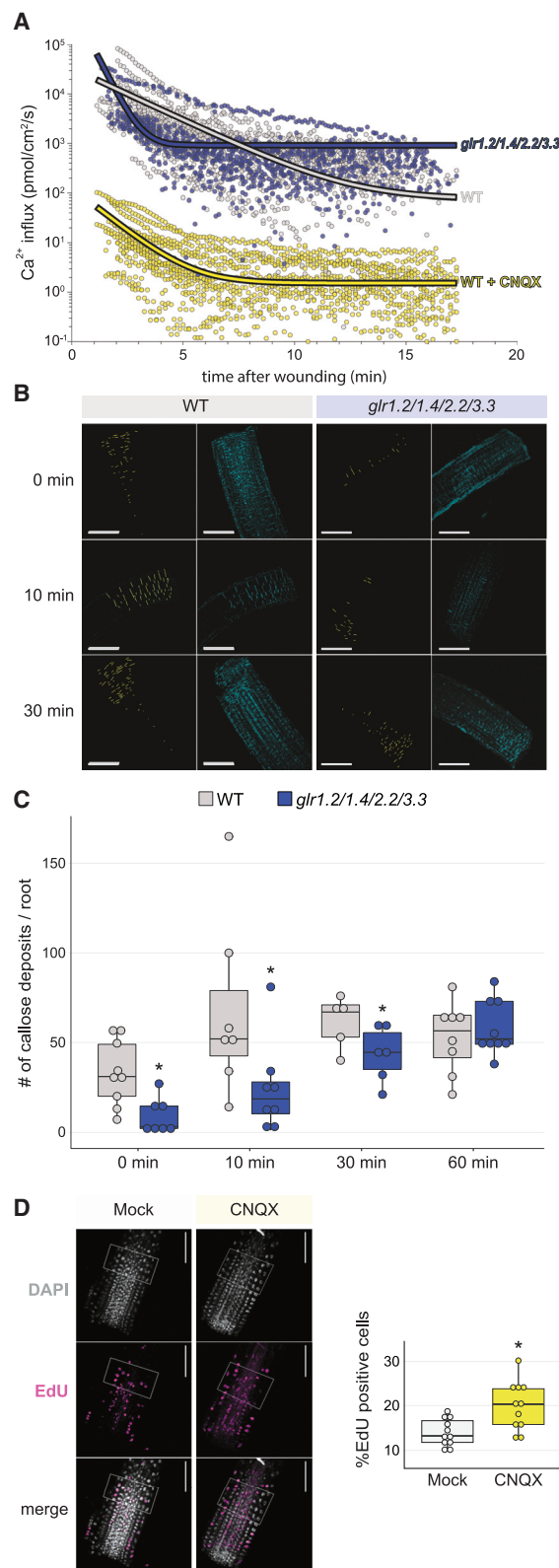
Lower calcium influx observed in the mutant immediately after injury could result from a direct contribution of the GLRs or

an indirect consequence of an altered cellular physiology in the mutant. To distinguish these possibilities, we performed the same measurements on wild-type roots treated with CNQX after wounding. Upon the chemical treatment, calcium influx is severely affected decreasing by four orders of magnitude. Nonetheless, similar dynamics were observed with the lowest calcium influx occurring at 6 min after wounding ($K = 0.8218$) followed by sustained calcium levels in the recovery phase (Figure 6A, yellow markers and yellow trend line). The comparable dynamics of *glr* mutants and the transient CNQX-blocking of GLR function confirms an overall impairment in the kinetics of homeostatic re-establishment of calcium after wounding further supporting that GLR channels mediate Ca^{2+} influxes in response to injury.

Earlier studies of injury in soybean showed that callose synthase, which deposits callose on the cell cortex in response to injury (Ellinger and Voigt, 2014), is directly activated by a calcium flux (Kauss, 1985; Wissemeier and Horst, 1995). Using this evidence, we hypothesized that GLRs activate callose synthase in response to wounding through their calcium permeability. In this scenario, callose buildup then seals the injury helping to re-establish normal homeostasis. The model implies that perturbation of GLRs, which we showed earlier as decreasing calcium fluxes upon wounding, should prevent callose buildup.

To test the model, we measured callose buildup in cells at 0, 10, 30, and 60 min after wounding by quantitative confocal imaging of aniline blue stained roots in wild type and the *glr1.2/1.4/2.2/3.3* mutant. Indeed, the *glr* mutants had significantly less callose deposition (Figures 6B and 6C), showing that perturbation of GLR signaling reduced a defense response in the immediate vicinity of the wound. Thus, the *glr* mutant's impaired Ca^{2+} influx response immediately after wounding might not suffice to activate callose deposition to the same extent as in wild type. However, after 10 min, when influxes plateau, the sustained higher Ca^{2+} influx seems to compensate for the early low influx, and the differences in callose deposition become less evident until no significant difference was observed at 60 min (Figures 6C and S6B).

Focusing on regenerative responses, we also asked whether perturbation of GLRs lead to cellular changes that could enhance regeneration. We first examined cell division rates, a well-documented regeneration response in the root injury model (Heyman et al., 2016). To measure cell-cycle activity, we used the nucleoside analog 5-ethynyl-2'-deoxyuridine (EdU), which is incorporated into DNA during synthesis and labels cells in S phase. Quantitative measurements of EdU-stained roots showed an increase in the frequency of S-phase-labeled cells in the root stump of wild-type CNQX-treated plants where cells reprogram to reform the distal meristem, suggesting an increase in division rates (Figure 6D). In addition, a quantitative index of cell identity (ICI, Efroni et al., 2015; Efroni et al., 2016) showed that CNQX-treated cells had a more “youthful” stem cell-like character at 24 h after cutting compared with the mock cells (Figure S6C). Together, these results show the physiological consequences of GLR signaling near the wound site favor defense responses in wild type, such as the buildup of callose, while inhibiting regenerative responses, such as cell division and cellular reprogramming. Thus, the GLRs manage a regeneration versus defense balance in the aftermath of wounding.



GLR-mediated signaling affects regeneration in flowering plants broadly

To determine if the GLRs' role in regeneration is conserved among plants, we took advantage of GLR antagonists to conditionally perturb GLR signaling in other species. Maize is a monocot crop species separated from Arabidopsis, a dicot, by an estimated 150 million years (Chaw et al., 2004). Due to the recalcitrance of callus formation from vegetative tissues in maize, biotechnology protocols employ a callus from embryo protocol for transformation (Green and Phillips, 1975). We cultured maize embryo cells on CIM adjusted for maize using the regeneration competent Hi-II line, which is commonly used for transformation, and the recalcitrant B73 line, which is a parental line used in advanced hybrid seed production, scoring callus formation after 15 days (STAR Methods).

Both lines showed an increase in callus formation when incubated on CNQX after embryo isolation (Figure 7A). In the recalcitrant B73 line, CNQX treatment increased the formation of productive calli (>5 mm) by 50% (Figure 7A). In addition, the productive calli were 30% larger in CNQX treatment (Figure 7B). CNQX-treated B73 maize embryos even reached similar callus formation rates of Hi-II on mock treatments (Figure 7A). Moreover, B73 embryos incubated on CNQX had smaller, cytoplasmically clear cells that typically give rise to regenerative organs (Figures 7B and S7A).

To determine whether calcium signaling antagonists could enhance regeneration in highly recalcitrant tissue, we incubated the first leaves and the sheath (coleoptile) of B73 on mock or CNQX plates for 15 days. Callus formed on around 4.4% of mock-treated B73 leaf explants, whereas callus formation on CNQX-treated B73 leaf explants more than tripled to about 15% (Figure 7C). Callus microcolony formation from Hi-II protoplasts also increased on CNQX (Figure S7B). In addition, treatment with PAMD increased coleoptile to callus formation rate and size to similar levels as CNQX for the coleoptile (Figure 7D). The result suggests that the role of GLRs and the SA pathway in regeneration is conserved in maize. Thus, GLR-mediated signaling inhibits regeneration in a range of contexts across flowering plant species.

Figure 6. Blocking GLRs severely disrupts ion flux, inhibits defense, and augments repair

(A) Vibrating probe measurements over time after root tip cutting show an increase in the rate of positive ion (Ca²⁺) flux in the *glr1.2/1.4/2.2/3.3* quadruple mutant compared with wild type (top), showing a severe disruption in ion flux (wild type, *n* = 13; *glr4*, *n* = 11). In wild type (bottom), treatment with CNQX decreases calcium influx 4-fold overall, showing a similar pattern of calcium influx over time compared with the mutant trend above.

(B and C) Representative images (B) and quantification (C) of callose deposition in the minutes after injury, showing a decrease in callose accumulation in the *glr4* mutant compared with wild type (t test; *n* ≥ 5 for each genotype at each time point; **p* = 0.003 for 0 min, which is effectively 3–5 min after wounding due to experimental limitations, **p* = 0.035 for 10 min, **p* = 0.03 for 30 min & n.s. for 60 min) with images over time. Callose deposition was analyzed using the DAPI filter (cyan) after aniline blue staining and particle analysis (e.g., yellow pixels in first and third column of B). Scale bars, 100 μm. (D) Representative images and quantification of EdU staining for cells that have entered or undergone S phase, showing increased division rates in Col-0 roots treated with CNQX (t test; mock, *n* = 18; CNQX, *n* = 16; **p* = 0.003).

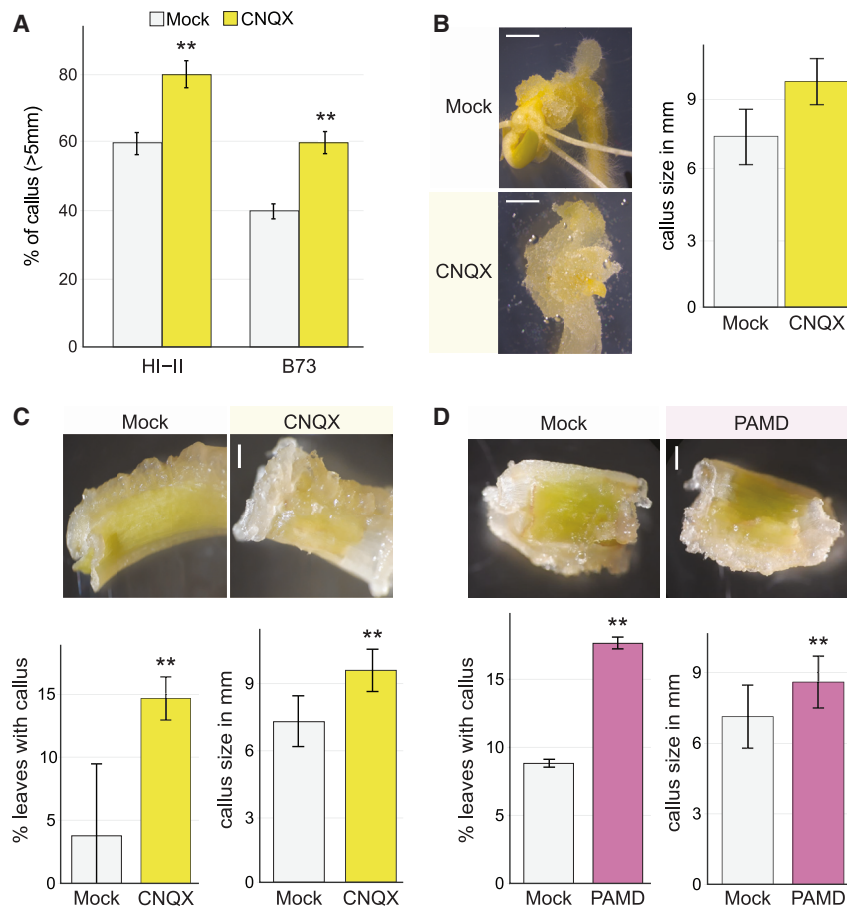


Figure 7. Blocking calcium channels improves regeneration in the maize recalcitrant line B73

(A) CNQX constitutive post embryo harvest treatment increases the percent of callus formation from embryos in the regeneration competent HI-II line (chi-square test; $n = 48$; $**p < 0.0001$) and the regeneration recalcitrant B73 line (chi-square test; $n = 48$ for each genotype in each treatment; $**p < 0.0001$).

(B) Representative images and quantification of callus formation from embryo in the B73 line and quantification. Under CNQX constitutive post embryo harvest treatment, a larger mass of callus was obtained (t test; $n = 24$ for each treatment; $**p < 10^{-8}$). Scale bars, 2 mm.

(C) Regeneration of the maize seedling's first leaf and coleoptile in the recalcitrant B73 line comparing mock and CNQX constitutive post-leaf harvest treated samples with quantitative analysis showing more than 3× higher frequency of callus formation in CNQX-treated cells (chi-square test; $n \geq 34$ for each treatment; $**p < 0.0001$) and improvement in callus size (t-test; $n = 24$; $**p < 0.0001$). Scale bars, 1 mm.

(D) Callus formation from B73 leaves with PAMD constitutive post leaf harvest treatment, showing an improvement in callus formation frequency (chi-square test; $n \geq 34$ for each treatment; $**p < 0.0001$) and callus size due to treatment (t test; $n = 34$ for each treatment; $**p < 0.001$). Scale bars, 1 mm.

GLR signaling acts mainly through the SA pathway

The concept of a trade-off between growth and defense is well established

(Huot et al., 2014; van Butselar and Van den Ackerveken, 2020; Züst and Agrawal, 2017, but also see Kliebenstein [2016]). Both SA and JA have been implicated in boosting defense while slowing growth (Huot et al., 2014; van Butselar and Van den Ackerveken, 2020); therefore, the increase in regeneration we observed by blocking GLRs or the SA pathway may reflect a growth versus defense trade-off. However, perturbations of defense pathways did not always affect regeneration and growth in the same way. For example, the *coi1-2* mutant also shows increased growth but lower regeneration in most of the assays. In addition, the quadruple *glr* mutant, which showed increased regeneration, did not show enhanced root growth. (In fact, other groups found that a *glr3.6* knockdown had root growth defects, Singh et al., 2016.) Furthermore, our conditional treatments showed that the effects of GLR signaling on regeneration and defense responses could be entirely recapitulated by blocking the channels transiently upon wounding, and many of the effects of the *glr* mutants, such as decreased callose deposition, were apparent only after wounding. Thus, we posit that GLR-mediated signaling represents a different type of trade-off—not in balancing growth versus defense—but routing the complex responses to an injury. The local GLR-mediated response appears to allow a “bet-hedging” strategy between defense and regeneration.

Both SA and JA have been shown previously to mediate GLR responses in wounding (Forde and Roberts, 2014).

DISCUSSION

Upon wounding, GLRs balance defense and regeneration

The plant's two primary strategies to defend against attacks—defense and regeneration—share many of the same rapid wound responsive triggers, such as calcium, reactive oxygen species, and electrical signals (Chen et al., 2016; Efroni et al., 2016; Florentin et al., 2013; Hoermayer and Friml, 2019; Moeder et al., 2019; Savatin et al., 2014). The key finding here is that GLR-mediated signals negatively regulate regeneration but positively regulate defense response. The result shows that the activation of the two processes can be decoupled. GLR-mediated signaling tunes responses in favor of defense, enhancing callose deposition and the expression of defense-related genes. At the same time, GLR signaling dampens the regeneration response, slowing division rates, cellular reprogramming, and meristem recovery. The effects all appear to take place in the immediate vicinity of the wound without long-distance systemic signaling, as conditional GLR inhibition improved regeneration in isolated explants of leaves or embryos where no systemic signaling is possible. Thus, the plant's wound response appears to be, at least to some degree, a trade-off between efficient repair and a robust defense response and the plant does not necessarily optimize wound responses for either process.

Therefore, why were the regeneration effects mediated by SA and not JA? Although both hormone pathways have overlapping function, they also mediate response to different types of pathogens independently (Yang, 2015). Still, it is not apparent how those independent roles would lead to their differential effects on regeneration. Another explanation centers on JA's competing roles in wound response. That is, in addition to its well-documented roles in defense (Howe et al., 2018), recent reports have implicated JA with a positive role in regeneration (Zhang et al., 2019; Zhou et al., 2019). Thus, in disturbing JA signaling, any gain in regenerative capacity resulting from a down regulation of defense is offset by also inhibiting the developmental responses needed for repair. Indeed, whereas our transcriptional analysis showed that the GLRs activated SA responses, the effect on JA response was more complex, with both up- and downregulation of JA-responsive genes. Overall, SA may provide a less “conflicted” pathway to downregulate regeneration and upregulate defense. Thus, SA may be a better pathway to mediate the trade-off between the two.

Calcium fluxes are severely disturbed around the wound in GLR mutants

GLR channels are known to mediate calcium signaling (Michard et al., 2017; Wudick et al., 2018a), whereas other work has documented the complexity of ion-mediated signaling during wounding in plants (Meyer and Weisenseel, 1997). In the wounding experiments, maize roots wounded by deep incisions showed very fast membrane depolarizations, followed by periods of gradual re- or hyper-polarization. Similarly, our results show high Ca^{2+} influxes at the injury site in Arabidopsis wild-type roots followed by gradual recovery of fluxes. Maize root wounding also showed that the observed re-polarization was composed of major effluxes of potassium and chloride and smaller influxes of calcium and protons, which could be expected given the concentration gradients of these metabolites and ions.

Our experiments with the *glr* quadruple mutant demonstrate an impairment of calcium influxes that are reduced even further to almost background levels by pretreatment with CNQX. These results suggest that, without Ca^{2+} signaling, the default pathway reverts to favor regeneration. Although we cannot rule out that the observed effects mediated by the GLRs were caused by ion fluxes other than calcium, the equivalent calcium flux dynamics obtained from CNQX-exposed roots and the observed phenotype in callose deposition further support a role in calcium signaling. The lower levels of calcium influx observed in the mutant 2–8 min after wounding could account for a lower callose synthase activity, resulting in less callose deposition (observed at time 0 after wounding). Following the observations of the maize results described earlier, the subsequent disturbances we observed in Ca^{2+} flux levels could be due to lower callose deposits that allow a leakage of ions and other metabolites, such as charged macromolecules.

In this model, the GLRs represent a rapid response mechanism to injury; therefore, a key question is when after wounding do they exert their effects? Ion signaling acts fast in most contexts, which leads to the expectation that GLR signaling affects regeneration in the seconds or minutes after wounding. Indeed, our experiments showed that a 1-h pretreatment ending right

after injury led to a significant increase in regeneration. This shows that GLRs mediate rapid signaling after wounding. However, we could increase regeneration further with prolonged treatment on CNQX, showing the GLRs also mediate a longer-term response. One explanation is that certain responses, such as callose deposition, represent early outcomes of GLR signaling, whereas GLRs also mediate responses that act on a longer time frame, such as the SA responses we observed in our expression analysis around 24 h after wounding. This leads to a scenario in which defense responses can inhibit regeneration for some time after wounding. To augment regeneration, defense responses need to be inhibited continually for some time after wounding, such as the clinical use of fever suppressants to prevent temperature spiking during an infection. In the context of normal function, the GLRs may signal over an extended time to sustain the defense response and inhibit regeneration.

SA contributes to age-dependent regeneration

Interestingly, the involvement of the SA pathway in regeneration may also provide a clue to the longstanding observation that regeneration efficiency decreases with tissue age (Binding, 1975; Chen et al., 2014; Hoque and Mansfield, 2004; Prakash and Gurumurthi, 2010; Sinnott, 1960; Vasil and Vasil, 1974). SA levels were previously shown to be anticorrelated with callus formation in garlic leaves (Mostafa et al., 2020), and our own analysis showed that SA-response markers were anticorrelated with zones of regeneration efficiency in roots. Furthermore, our data showed that perturbing GLRs, which signal to SA, had stronger effects in enhancing regeneration in older tissues. A consistent explanation to these observations is that mature tissues exhibit higher levels of SA signaling that prime them for defense responses at the cost of regenerative processes. The model implies that inhibiting SA response in these older tissues reverts regenerative competence—in part—to a younger developmental state.

Finally, plants deal with a wide array of insults and attacks, including pathogens, herbivores, and mechanical damage. In many contexts, the strategic response is a combination of defense and regeneration. However, neither response is apparently optimized. Early screens established the prevalence of mutants for enhanced defense responses (Glazebrook et al., 1996), whereas we show here perturbations that can enhance regeneration. Inhibition of the GLR-SA pathway provides evidence of a trade-off between the two responses and exposes one mechanism that the plant uses to balance its complex response to the environment.

Limitations of the study

Regeneration can vary by ambient conditions, time of year, and subtle growth conditions; therefore, mock control or WT plants are needed for each experimental trial. Nonetheless, although starting regeneration rates may vary by trial, effects are generally consistent over multiple trials. Repeats of the same experiment need to be treated as independent trials. Here, we used Fisher's combined p value method. The net result is that absolute regeneration levels for a given experimental set should be considered in a range rather than an absolute value.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.devcel.2022.01.013>.

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AUTHOR CONTRIBUTIONS

Conceptualization, M.H.C., P.C.D.A., and K.D.B.; methodology, M.H.C., P.C.D.A., C.O.N., J.A.F., and K.D.B.; materials and expertise, M.M.W. and M.A.L.; experiments, M.H.C., P.C.D.A., P.-L.I., C.O.N., R.R., and M.A.L.; data analysis, M.H.C., P.C.D.A., C.O.N., J.A.F., and K.D.B.; writing, M.H.C. and K.D.B.

DECLARATION OF INTERESTS

A patent (application #63120640) has been filed related to this work on compositions and methods to enhance plant regeneration.

INCLUSION AND DIVERSITY

One or more of the authors of this paper self-identifies as an underrepresented ethnic minority in science. One or more of the authors of this paper self-identifies as a member of the LGBTQ+ community. The author list of this paper includes contributors from the location where the research was conducted who participated in the data collection, design, analysis, and/or interpretation of the work. While citing references scientifically relevant for this work, we also actively worked to promote gender balance in our reference list.

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STAR★METHODS

KEY RESOURCES TABLE

Reagent or resource	Source	Identifier
Biological samples		
None	N/A	N/A
Chemicals, peptides, and recombinant proteins		
(2E)-4-phenyl-2-amino]methylidene] cyclohexane-1,3-dione	Aurora Fine Chemicals	180.584.762
SULFO-CYANINE5 AZIDE 1MG	LUMIPROBE	A3330
CNQX	Santa Cruz	sc-203003A
Gadolinium(III) chloride	Santa Cruz	sc-224004
D(-)-2-Amino-5-phosphonovaleric acid (D-AP5)	Santa Cruz	sc-20043
ABA	Sigma	862169
Aniline Blue	Sigma	415049
L-ASCORBIC ACID	Sigma	A4544
6-Benzylaminopurine	Sigma	B3408
Copper (II) Sulfate	Sigma	451657
DAPI	Sigma	D9542
2,4-Dichlorophenoxyacetic acid	Sigma	D7299
L-Glutamic acid potassium salt monohydrate	Sigma	G1501
L-Cysteine (Cys)	Sigma	C7352
L-Serine (Ser)	Sigma	S4311
Inositol	Sigma	I3011
1-Naphthaleneacetic Acid	Sigma	N0640
Nicotinic acid	Sigma	N0671
Salicylic Acid	Sigma	247588
Vitamin B1	Sigma	T4625
Vitamin B6	Sigma	P5669
ZEATIN	Sigma	Z0164
Critical commercial assays		
Nugen OvationRNA Amplification System V2	Tecan Genomics	3100-A01
Nugen Ovation Ultralow DR Multiplex System	Tecan Genomics	0344NB-08
Deposited data		
Raw and processed ATAC-seq data from root reprogramming cells	This paper	GSE168381
Raw and processed RNA-seq data from root reprogramming cells	This paper	GSE168385
Experimental models: Organisms/strains		
<i>Arabidopsis thaliana</i> Columbia ecotype	ABRC	N/A
<i>Arabidopsis thaliana</i> <i>glr1.2</i> mutant	ABRC	SALK_114822
<i>Arabidopsis thaliana</i> <i>glr3.3-1</i> mutant	ABRC	SALK_077608
<i>Arabidopsis thaliana</i> <i>glr3.3-2</i> mutant	ABRC	SALK_082194
<i>Arabidopsis thaliana</i> <i>glr1.2/1.4-1/3.3-1</i> mutant	Wudick et al., 2018b	N/A

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Continued

Reagent or resource	Source	Identifier
<i>Arabidopsis thaliana</i> glr1.2/1.4-1/2.2/3.3-1 mutant	Wudick et al., 2018b	N/A
<i>Arabidopsis thaliana</i> glr3.1/3.2/3.3/3.6 mutant	Mou et al., 2020	N/A
<i>Arabidopsis thaliana</i> coi1-2 mutant	Xu et al., 2002	N/A
<i>Arabidopsis thaliana</i> npr1-5 mutant	Zipfel et al., 2004	N/A
<i>Arabidopsis thaliana</i> cbp60g-1 mutant	ABRC	SALK_023199C
<i>Arabidopsis thaliana</i> cbp60a-1/cbp60g-1 mutant	ABRC	CS72190
<i>Arabidopsis thaliana</i> npr3-2/npr4-2 mutant	ABRC	CS72351
<i>Arabidopsis thaliana</i> cpk11-2/cpk5/cpk6 mutant	ABRC	CS69906
<i>Arabidopsis thaliana</i> cpk6(a) mutant	ABRC	CS65905
<i>Arabidopsis thaliana</i> cpk6(b) mutant	ABRC	SALK_025460
<i>Zea mays</i> B73	Jackson Lab CSHL	N/A
<i>Zea mays</i> Hill	Jackson Lab CSHL	N/A
Oligonucleotides		
Ad1_noMX	Buenrostro et al., 2013	AATGATACGGCGACCAC CGAGATCTACACTCGTCGGCA GCGTCAGATGTG
Ad2.1_TAAGGCGA	Buenrostro et al., 2013	CAAGCAGAAGACGGC ATACGAGATTCGCCTTAGTCTC GTGGGCTCGGAGATGT
Ad2.2_CGTACTAG	Buenrostro et al., 2013	CAAGCAGAAGACGG CATACGAGATCTAGTACGGTCTCG TGGGCTCGGAGATGT
Ad2.3_AGGCAGAA	Buenrostro et al., 2013	CAAGCAGAAGACGGC ATACGAGATTCTGCCTGTCTCG TGGGCTCGGAGATG
Ad2.4_TCCTGAGC	Buenrostro et al., 2015	CAAGCAGAAGACGG CATACGAGATGCTCAGGAG TCTCGTGGGCTCGGAGATGT
Software and algorithms		
Bowtie2	Langmead and Salzberg, 2012	http://bowtie-bio.sourceforge.net/bowtie2/index.shtml
macs2 version 2.1.1	Zhang et al., 2008	https://pypi.org/project/MACS2/2.1.1.20160309/
DESeq2	Love et al., 2014	https://bioconductor.org/packages/release/bioc/html/DESeq2.html
HISAT2	Kim, et al., 2015	http://daehwankimlab.github.io/hisat2/manual/
Image J software Fiji version 1.52p	Schindelin et al., 2012	http://imagej.nih.gov/ij/
VirtualPlant 1.3	Katari et al., 2010	http://virtualplant.bio.nyu.edu/cgi-bin/vpweb/

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be made available upon reasonable request by the Lead Contact, Kenneth D. Birnbaum, ken.birnbaum@nyu.edu.

Materials availability

Arabidopsis mutant lines used in this study are available from the authors or from the sources cited in the text or methods. Chemical reagents can be obtained by suppliers, as listed in the reagent table below.

Data and code availability

- Raw and processed data files have been submitted to GEO and are publicly available as of the date of publication under accession numbers GSE168381 (ATAC-seq data) and GSE168385 (RNA-seq data).
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

The ATACseq and RNAseq analysis, as well as all wild type controls, were performed using Arabidopsis Columbia (Col-0) ecotype. The mutant lines used in this study are: *glr1.2* (SALK_114822), *glr3.3-1* (SALK_077608), *glr3.3-2* (SALK_082194), *glr1.2/1.4-1/3.3-1*, *glr1.2/1.4-1/2.2/3.3-1* (Wudick et al., 2018b), *coi1-2* (Xu et al., 2002) and *npr1-5* (Zipfel et al., 2004), *glr3.1/3.2/3.3/3.6* (Mou et al., 2020). Other SA pathway related mutants tested were: *cbp60g-1* (SALK_023199C), *cbp60a-1/cbp60g-1* (CS72190), *npr3-2/npr4-2* (CS72351), *cpk11-2/cpk5/cpk6* (CS69906), *cpk6(a)* (CS65905), *cpk6(b)* (SALK_025460). All in the Col-0 genetic background. All plants were grown on regular agar plates (0.8% agar, ½ MS, 0.5% sucrose) in Percival chambers at 23°C under 16 hr light/8 hr darkness.

METHOD DETAILS

Root stump regeneration assay

Seeds were stratified at 4°C for two days, placed on regular agar plates, and then grown vertically for 7 days. Plants were transferred to a “cutting board” (3% agar, ½ MS, 0.5% sucrose), upon which root meristems were cut at ~130µm (low cuts) or 270µm (high cuts), which removed most of the meristem. Cuts were made by using a 30G sterile dental needle (Exellnt) by hand under a dissecting microscope. Plates were turned 90° degrees immediately after cutting. Root regeneration was scored for gravitropism as in Figure 1A at 1, 2, and 3 days post cut (dpc). A clear gravitropic response determined regeneration, where regenerated roots were also checked under a dissecting microscope to ensure the primary root attained morphological characteristics of a regenerated meristem.

ATAC-seq analysis

Seeds of the *pMP::MP-GFP* background were stratified and grown in standard growth conditions on standard plates for 7 days. Plants were then transferred to cutting board plates, as above, and cut at ~130µm. The excised roots were then harvested at 2.5h, 14h, or 16 h after cutting, with triplicate samples collected. *pMP::MP-GFP* is typically induced within 2 h, and localized mainly in the nucleus but also in the cytoplasm. However, due to the low signal observed in the nucleus at early time points, the excised roots were subjected to rapid enzymatic digestion of cell walls as described for maize (Ortiz-Ramirez et al., 2017), with a shorter enzymatic incubation of 1 hr. GFP+ cells were collected by FACS and protoplasts were lysed in deionized water. For the time points after root tip cutting, about 1000-2000 GFP-expressing protoplasts were used per sample. After protoplasts were lysed, the low-input Tn5 tagmentation protocol was used, decreasing the concentration of both Tn5 and primers (Buenrostro et al., 2015). A final library purification was done using Ampure XP beads to selectively avoid fragments of less than 100 bp that correspond to primers and primer dimers. For uncut controls, about 50K protoplasts from distal meristem dissections were used. Nuclei were either mechanically released or obtained after root protoplasting, stained with DAPI (1µM) and isolated by FACS. The standard Tn5 tagmentation protocol was used according to the described protocols (Buenrostro et al., 2015; Lu et al., 2017). In brief, a purification step of the transposed DNA fragments was done using Qiagen MinElute PCR Purification Kit prior library amplification (see key resources table for primers used). The appropriate number of cycles for library amplification were experimentally determined by qPCR.

All sequenced libraries were aligned using Bowtie2 and the peaks corresponding to accessible chromatin regions were identified with MACS2. Consensus peaks among replicates were obtained and the counts were determined. The Differential Accessible Chromatin (DAC) regions between samples were analyzed for significance using DESeq2 comparing each time point vs. uncut roots. To eliminate false-positives and to increase the confidence in our results, we 1) only considered accessible regions present in all replicates, and 2) used a padj value below 0.01 in our DAC analysis.

Regeneration assays in calcium antagonist treatments or SA treatments

For treatments, seeds were grown and meristems were cut as detailed above for standard growth on standard plates. For the different treatments, plates were prepared as followed: mock plates (0.8% agar, ½ MS, 0.5% sucrose, water); mock cutting board (3% agar, ½ MS, 0.5% sucrose, water); CNQX, GD³⁺ or AP5 plates (0.8% agar, ½ MS, 0.5% sucrose and either 50µM CNQX, 1µM GD³⁺ or 200µM AP5); CNQX, GD³⁺ or AP5 cutting board (3% agar, ½ MS, 0.5% sucrose and 50µM CNQX, 1µM GD³⁺ or 200µM AP5 respectively); Mock plates for SA treatments (0.8% agar, ½ MS, 0.5% sucrose, DMSO); mock cutting board for SA treatments (3% agar, ½ MS, 0.5% sucrose, DMSO); SA or PAMD plates (0.8% agar, ½ MS, 0.5% sucrose and either 1µM SA or 1µM PAMD); CNQX and SA plates (0.8% agar, ½ MS, 0.5% sucrose, 50µM CNQX, 1µM SA). Seven days after plating, seeds from standard plates were transferred to either the CNQX or mock cutting board for rapid resection and then immediately placed back on either CNQX plates or control plates. In the experiment shown in the main Figure, plants were pre-incubated for 1 h on either mock or CNQX plates to ensure cells were exposed to CNQX at the time of cutting.

Regeneration assays in GLR agonist treatments

Arabidopsis Col-0 seeds were grown and meristems were cut as detailed above for standard growth on standard plates. For the different concentrations used in the agonist treatments, regular plates (0.8% agar, ½ MS, 0.5% sucrose) were supplemented with 1mM or 5mM of either L-Glutamate (Glu), L-Cysteine (Cys) or L-Serine (Ser). In short treatment plants were exposed to the amino acid tested (5mM) for <2 min during root tip cut and transfer to half strength MS plates to assess regeneration. For long treatment, plants were cut and maintained on plates with the treatment until regeneration was assessed.

Root stump regeneration assays with transient CNQX window treatments

For transient treatments, seeds were grown and meristems were cut as detailed for CNQX treatments with regeneration. For the short early window treatments, plants were cut on the CNQX cutting board and transferred to CNQX plates for 4 h then transferred to mock plates for the remainder of the period. For the later treatment, plants were transferred to the mock cutting board for resection, transferred again to a mock plate after cutting for 4 h and then transferred to a CNQX plate for the remainder of the experiment. For the 16 h window treatments, plants were exposed to a 1h pretreatment by placing them on mock or CNQX plates and then transferred to the cutting board without the treatment. After root tip excision, plants were placed on or off CNQX for 16 h followed by a second transfer to either mock or CNQX plates for regeneration. Regeneration was scored as above. Controls included all the same transfers where one set of plates was transferred on and off mock plates (mock) and a second set of plates was transferred on and off only CNQX plates (positive control).

RNA-seq time course

Plants were grown on standard plates in standard conditions and transferred to CNQX or mock cutting board plates at 7 days after plating. Immediately after transfer, plants were cut at 270 µm (high cuts) and then transferred to CNQX plates or mock plates. Approximately 20 stump tips consisting of about 100 µm of the stump were collected at each time point (30 min, 4 h, 14 h, 24 h). For controls, two separate sets of uncut roots were transferred to CNQX plates or mock plates and incubated for 14 h before collecting root tip tissue, which also consisted of about 100 µm of the distal root tip. RNA was extracted using the Qiagen MicroRNAeasy Kit. We performed cDNA construction and library amplification with Nugen OvationRNA Amplification System V2. Libraries were made using the Nugen Ovation Ultralow DR Multiplex System (Tecan). DNA fragmenting was performed using the Covaris S220. Short read sequences were generated on an Illumina NextSeq 500 and were aligned using HISAT2. Counts were normalized between samples using DESeq2's median of ratios method before analyzing for differentially expressed genes.

Analysis of RNA-seq time course

The time series included six time points for mock and CNQX treated plants (2 uncut controls, 30 min post cut, 4h post cut, 14 h post cut, 24 h post cut). First, of the 32,833 annotated transcripts, the top 10th percentile most variable genes were selected [(genevarfilter(data,'Percentile',90)) in Matlab], resulting in a list of 3,283 genes. Any gene with a zero value in any sample was removed (leaving 3131 genes). The resulting matrix was then subject to time series analysis using *ImpulseDE* version 1.12.0 in R (Chechik and Koller, 2009; Sander et al., 2017). All default values were used (n_iter = 100, n_randoms = 50000, Q_value = 0.01) in a case vs. control analysis with the 6 mock samples vs. 6 CNQX-treated samples used for comparative differential expression. Genes classified as differentially expressed 'DE' in 'at least 2 TPs for case vs. control' were selected for further analysis (1436 genes). *Impulse* categorized the DE genes into 19 different clusters. Each cluster was then subjected to GO term analysis using the *biomaps* function in *VirtualPlant* 1.3 (Katari et al., 2010). Overall trends for each cluster (control vs. treat) were derived by taking the row sum and normalizing by the row mean.

Extracellular Ca²⁺ flux measurements after wounding

The ion-selective vibrating probe was used to estimate extracellular calcium flux at the wounding site (as per Wudick et al., 2018b) between 1–2 min after the cut due to the constraints in the technique. *Arabidopsis* roots were hydrated approximately 1 h before the measurements to equilibrate with the Basic Measuring Solution (BMS): 0.5 mM KCl, 0.1 mM CaCl₂, pH 5.5 (unbuffered) (Shabala et al., 2005) or BMS+CNQX (50 µM). All the roots were cut and measured in BMS. The vibrating probe was moved with an excursion of 10µm, completing the cycle of acquisition in 10 s. Each of the roots was measured until the fluxes stabilized at approximately 10–15 min after starting the measures. All the points were plotted, and an exponential decay fit was employed to extract the asymptote (plateau) and the decay constant (k) that were then compared with the extra sum of squares F-test. The statistics and fitting of the data were run on Graphpad Prism.

EdU staining

Cell proliferation analysis was performed with the EdU click reaction containing 2Mm CuSO₄·5H₂O (Sigma), 8 µM Sulfo-Cyanine5 azide (Lumiprobe) and 20mg/ml Ascorbic acid (Sigma). For EdU incorporation, Col-0 plants were grown under standard conditions described above, and, at 7 days after plating, were transferred to ½ MS media supplemented with CNQX (50 µM) or mock. After 60 min pre-incubation, plants were placed on CNQX or mock cutting board, cut at 270 µm (high cuts) and immediately transferred to ½ MS medium containing CNQX (50 µM)/ 5 µM EdU or mock/5 µM EdU for 30min. Plants were fixed and permeabilized according to published protocols (Kazda et al., 2016). The next day, samples were washed with PBS twice and incubated in the EdU click reaction cocktail for 30 min in the dark. After two washes with PBS, plants were placed on 0.1mM DAPI in PBS and incubated for 30 min

at 22 °C in the dark. Two final washes with PBS were performed. Samples were observed under an inverted fluorescence microscope. Images were analyzed with Image J software Fiji 1.52p version (<http://imagej.nih.gov/ij/>). Using images from a median optical section, labeled cells were quantified for EdU and DAPI in the meristematic zone. The boundary between the meristematic and the elongation zones is defined by a twofold or more increase in the distance between nuclei in the cortical cell layer.

Callose staining

Plants grown under standard conditions on standard plates, as above, were transferred to CNQX or mock cutting board 7 days after plating and cut at 270 μ m (high cuts). Plants were fixed on a 1:3 acetic acid/ethanol solution overnight after harvesting either immediately (0 min), after 10 min, or after 30 min. Note that 0 min is effectively 3–5 min after wounding accounting for the time it takes to fix the roots. The next day plants were stained in aniline blue solution (750 μ l of 67 mM K_3PO_4 pH 9.5, 240 μ l of distilled water, 1 μ l of Silwet-77, and 10 μ l of 1% aniline blue) in the dark for 1 h prior to imaging as described previously (Cui and Lee, 2016). Images were analyzed with Image J software Fiji 1.52p version (<http://imagej.nih.gov/ij/>).

Arabidopsis tissue culture and callus induction

Plants were grown under standard conditions in standard plates for each genotype shown in the main text. Detached leaf explants from 8-day-old (young leaves) and 15-day-old (old leaves) seedlings were cultured on media for callus induction: $\frac{1}{2}$ MS + 3% sucrose + 1% Agar + 2 mg/L 2,4 D + 0.5 mg/l AIA + 0.5 mg/L Zeatin, pH 5.7. For callus induction, two different calcium channel antagonists were tested in the media: Gd^{3+} 1 μ M or CNQX at 1 μ M or 50 μ M. SA treatment was tested using 50 μ M SA or 10 μ M PAMD on CIM. Sterile water was used instead of calcium channel antagonists as a mock control. Detached leaves were cut in half. Each plate contained either nine 15-day-old leaves or sixteen 8-day-old leaves. The plates were left at 23 °C under dark photoperiod. Callus formation was scored at day 15 after plating.

Root from leaf regeneration

For root-from-leaf regeneration, seeds were stratified at 4°C for two days, placed on regular agar plates (0.7% agar, $\frac{1}{2}$ MS, 0.5% sucrose) and grown vertically in Percival chambers at 23°C under 16 hr light/8 hr darkness for 15 days. Different chemicals were tested to improve root regeneration, supplementing either with 50 μ M CNQX, 50 μ M SA or 10 μ M PAMD. Detached leaf explants from 15-day seedlings were cultured on regular agar plates grown under the above conditions. Each plate contained twelve leaves. Root from leaf regeneration was scored at day 15 after plating.

Maize tissue culture and callus induction

Mature dry seeds of maize inbred lines HI-II and B73 were surface sterilized in 50ml falcon tube with 70% ethanol for 10 min and then in 4% sodium hypochlorite (NaOCl) supplemented with 0.1% Tween20 for 30 min, washed 3 times in sterile distilled water and incubated in 4% chloramine for 20min under shaking. The seeds were rinsed 4 times with sterilized water and plated on $\frac{1}{2}$ MS basal media with 0.7% agar and pH 5.7, and incubated at 28 °C in darkness. Immature embryos were excised to use as explants after 3 days on $\frac{1}{2}$ MS medium. The swollen mature embryos were removed from seeds with a scalpel and cultured, scutellum side up, on induction medium. The first leaf and coleoptile were taken for explants at 6 days after germination on $\frac{1}{2}$ MS medium. The coleoptiles were longitudinally split with a sharp scalpel to expose shoot meristem and simultaneously cultured on callus induction media with the split side facing the media. Callus induction media contained N6 basal media supplemented with 3% sucrose, 2 mg/l glycine, 1 mg/l vitamin B1, 0.5 mg/l vitamin B6, 0.5 nicotinic acid, 120 mg/l inositol, 7 g/l powdered agar, 2 mg/l 2,4-dichlorophenoxyacetic acid (2,4-D), 1 mg/l 1-naphthaleneacetic acid (NAA), CNQX (50 μ M for treated samples) and pH 5.7 (Shen et al., 2012). The embryos and first leaf/coleoptile were incubated for 15 days in darkness at 28 °C. After 3 weeks of culture on induction medium, the percentages and size of primary calli were determined.

Protoplast culture

Plants were grown under standard conditions and approximately 0.6 g of the aerial parts of 1-week-old sterile plantlets was soaked in 5 mL of maceration medium (MGG) in a Petri dish. Five milliliters of MGG were added, bringing the total volume to 10 mL. Maceration was performed in the dark, at 24°C, overnight (12 h). After cell wall digestion, 20 mL of protoplast suspension in MGG was filtered through a sterile 80-mm mesh filter into 10 mL washing solution (2.5% KCl and 0.2% $CaCl_2$) already added to a 30-mL tube. After centrifugation (70g, 6 min), protoplast pellets were gently resuspended in 25 mL washing solution and centrifuged again (70g, 6 min). Washing was performed two more times. Protoplast numbers were estimated before the last centrifugation. After a 1 h incubation at 4°C, the protoplast suspension was diluted in protoplast induction media (PIM) to a concentration of 8.3×10^5 protoplasts per milliliter. Protoplasts were diluted 1:10 in PIM. The effect of CNQX on cell division was evaluated by adding 1 μ M or 50 μ M CNQX to the liquid media. The plates were subsequently placed in large plastic containers to limit evaporation and cell suspensions were cultured in the dark at 20°C (Chupeau et al, 2013). On day 11, cell colonies formed in a liquid culture dish were visualized and counted.

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical data of all experiments, as the statistical tests used, p values and the number of individuals tested (n) can be found in the figure legends and in the results section. All detailed raw data on phenotypes, their statistical analysis, and multiple testing results are

contained in [Table S4](#). For the analysis of mutants and treatments for regeneration, standard Chi-square (e.g., regeneration proportions) or t tests (e.g., root length) were used for statistical analysis of phenotypes and traits. In many cases, experiments were repeated up to 4 times for rigor and/or to test more subtle differences between phenotypic classes. If repeat experiments were performed under the same condition at the approximate same time, the results were pooled for statistical tests. In some cases, experiments were performed months or years apart and batch effects were apparent. For example, background control (WT or mock) regeneration rates can vary by season and ambient humidity levels. In these cases, we combined separate tests for a given hypothesis using Fisher's Method for Combining P Values, using the formula $\text{pchisq}((\sum(\log(p)) \cdot 2), \text{df}=\text{length}(p) \cdot 2, \text{lower.tail}=\text{F})$ in R, where p is a row vector of all probability values for a given hypothesis. The multiple regeneration systems tested led to 127 tests in total, so, to account for multiple testing, we applied the Benjamini-Hochberg correction for all pvals reported for each test as the false discovery rate, FDR. None of the p values reported as significant lost their significance (fell above the significance cutoff of 5% FDR).