Title
An evolutionarily young defense metabolite influences the root growth of plants via the ancient TOR signaling pathway.

Permalink
https://escholarship.org/uc/item/273791wd

Authors
Malinovsky, Frederikke Gro
Thomsen, Marie-Louise F
Nintemann, Sebastian J
et al.

Publication Date
2017-12-12

DOI
10.7554/elife.29353

Peer reviewed
An evolutionarily young defense metabolite influences the root growth of plants via the ancient TOR signaling pathway

Frederikke Gro Malinovsky¹, Marie-Louise F Thomsen¹, Sebastian J Nintemann¹, Lea Møller Jagd¹, Baptiste Bourgine†, Meike Burow¹, Daniel J Kliebenstein¹,²*

¹DynaMo Center, Copenhagen Plant Science Center, Department of Plant and Environmental Sciences, University of Copenhagen, Copenhagen, Denmark; ²Department of Plant Sciences, University of California, Davis, Davis, United States

Abstract To optimize fitness a plant should monitor its metabolism to appropriately control growth and defense. Primary metabolism can be measured by the universally conserved TOR (Target of Rapamycin) pathway to balance growth and development with the available energy and nutrients. Recent work suggests that plants may measure defense metabolites to potentially provide a strategy ensuring fast reallocation of resources to coordinate plant growth and defense. There is little understanding of mechanisms enabling defense metabolite signaling. To identify mechanisms of defense metabolite signaling, we used glucosinolates, an important class of plant defense metabolites. We report novel signaling properties specific to one distinct glucosinolate, 3-hydroxypropylglucosinolate across plants and fungi. This defense metabolite, or derived compounds, reversibly inhibits root growth and development. 3-hydroxypropylglucosinolate signaling functions via genes in the ancient TOR pathway. If this event is not unique, this raises the possibility that other evolutionarily new plant metabolites may link to ancient signaling pathways.

DOI: https://doi.org/10.7554/eLife.29353.001

Introduction

Herbivory, pathogen attacks and weather fluctuations are just some of the factors that constantly fluctuate within a plants environment. To optimize fitness under this wide range of conditions, plants utilize numerous internal and external signals and associated signaling networks to plastically control metabolism and development (Henriques et al., 2014; Smith and Stitt, 2007; Rexin et al., 2015). This metabolic and developmental plasticity begins at seed germination, where early seedling growth is maintained by heterotrophic metabolism relying solely on nutrients and energy stored in the seed including the embryo. Upon reaching light, the seedling transitions to autotrophy by shifting metabolism to initiate photosynthesis and alters development to maximize photosynthetic capacity (Rexin et al., 2015; Galli et al., 2014). Until light is available, it is vital for the plant to prioritize usage from the maternal energy pool, to ensure the shoot will breach the soil before resources are depleted. Because the time to obtaining light is unpredictable, seedlings that had the ability to measure and accordingly adjust their own metabolism would likely enjoy a selective advantage. In this model, energy availability would an essential cue controlling growth throughout a plant’s life and not solely at early life-stages. On a nearly continuous basis, photo-assimilates, such as glucose and sucrose, are monitored and their internal levels used to determine the growth potential by partitioning just the right amount of sugars between immediate use and storage (Smith and Stitt, 2007).
Illustrating the key nature of metabolite measurement within plants is that glucose is measured by two separate kinase systems that are oppositely repressed and activated to determine the potential growth capacity, SnRKs1 (sucrose non-fermenting 1 (SNF1)-related protein kinases 1) and the Target of Rapamycin (TOR) kinase (Sheen, 2014). SnRKs1s are evolutionarily conserved kinases that are activated when sugars are limiting (Lastdrager et al., 2014). Arabidopsis thaliana (Arabidopsis) has two catalytic SnRK1-subunits, KIN10 and KIN11 (SNF kinase homolog 10 and 11), that activate vast transcriptional responses to repress energy-consuming processes and promote catabolism (Baena-González, 2010; Crozet et al., 2014; Baena-González et al., 2007). This leads to enhanced survival during periods of energy starvation. Oppositely, the TOR kinase is a central developmental regulator, whose sugar-dependent activity controls a myriad of developmental processes including cell growth, cell-cycle, and cell-wall processes. The TOR pathway functions to modulate growth and metabolism by altering transcription, translation, primary and secondary metabolism, as well as autophagy (Sheen, 2014; Sablowski et al., 2014). The TOR kinase primarily functions in meristematic regions where it promotes meristem proliferation. Within these cell types, TOR measures the sugar content and if the tissue is low in sugar, TOR halts growth, even overruling hormone signals that would otherwise stimulate growth (Xiong et al., 2013). In plants, TOR functions within a conserved complex that includes RAPTOR (regulatory-associated protein of TOR) and LST8 (lethal with sec-13 protein 8) (Henriques et al., 2014). RAPTOR likely functions as an essential substrate-recruiting scaffold enabling TOR substrate phosphorylation (Rexin et al., 2015), and LST8 is a seven WD40 repeats protein with unclear function (Moreau et al., 2012). TOR complex (TORC) activity is
positively linked with growth (Rexin et al., 2015) as mutants in any component lead to qualitative or quantitative defects in growth and development and even embryo arrest in strong loss-of-function alleles (Menand et al., 2002; Deprost et al., 2005). Although the energy sensory kinases KIN10/11 and TOR sense opposite energy levels, they govern partially overlapping transcriptional networks, which are intimately connected to glucose-derived energy and metabolite signaling (Sheen, 2014; Baena-González et al., 2007). Having two systems to independently sense sugar shows the importance of measuring internal metabolism. A key pathway controlled by TOR in all eukaryotes is autophagy (Liu and Bassham, 2010; Shibutani and Yoshimori, 2014). In non-stressed conditions, continuous autophagy allows the removal of unwanted cell components like damaged, aggregated or misfolded proteins by vacuolar/lysosomal degradation (Inoue et al., 2006). Under low energy conditions, TORC inhibition leads to an induction of autophagy to free up energy and building blocks, through degradation of cytosolic macromolecules and organelles (Shibutani and Yoshimori, 2014). Autophagy-mediated degradation is facilitated by formation of autophagosomes; double membrane structures that enclose cytoplasmic cargo, and delivers it to the vacuole (Shibutani and Yoshimori, 2014; Feng et al., 2014; Le Bars et al., 2014; Zhuang et al., 2016).

In nature, plant plasticity is not only limited to responding to the internal energy status, but to an array of external environmental inputs. The multitude of abiotic and biotic factors that plants continuously face often require choices between contradictory responses, that requires integrating numerous signals across an array of regulatory levels to create the proper answer. For example; plant defense against biotic organisms requires coordination of metabolic flux to defense and development while in continuous interaction with other organisms and the potential for interaction with other organisms. A proper defense response is vital for the plant as a metabolic defense response to one organism can impart an ecological cost by making the plant more sensitive to a different organism (Züst et al., 2012). Therefore, a plant must choose the most appropriate defense response for each situation to optimize its fitness and properly coordinate its defense response with growth and development. A key defense mechanism intricately coordinated across development is the synthesis of specific bioactive metabolites that are often produced in discrete tissues at specific times. A current model is that developmental decisions hierarchically regulate defense metabolism with little to no feedback from defense metabolism to development. However, work on the glucosinolate and phenolic pathways is beginning to suggest that defense metabolites can equally modulate development (Katz et al., 2015; Bonawitz et al., 2012; Bonawitz et al., 2014; Kim et al., 2014; Francisco et al., 2016a; Francisco et al., 2016b), thus suggesting that development and defense metabolism can directly cross-talk.

To assess if and how defense metabolites can signal developmental changes, we chose to investigate the glucosinolate (GSL) defense metabolites. The evolution of the core of GSL biosynthesis is relatively young, and specifically modified GSL structures are even more recent (Sønderby et al., 2012). There are >120 known GSL structures limited to plants from the Brassicales order and some Euphorbiaceae family members, with Arabidopsis containing at least 40 structures (Sønderby et al., 2010). GSLs are amino acid derived defense metabolites that, after conversion to an array of bioactive compounds, provide resistance against a broad suite of biotic attackers (Sønderby et al., 2010; Lambrix et al., 2001; Kliebenstein et al., 2002). GSLs not only exhibit a wide structural diversity, but their composition varies depending on environmental stimuli, developmental stage and even across tissues. This, combined with the information-rich side chain, makes GSLs not only an adaptable defense system, but also prime candidates for having distinctive signaling functions. Previous work has suggested that there may be multiple signaling roles within the GSLs (Francisco et al., 2016a; Francisco et al., 2016b; Clay et al., 2009; Khokon et al., 2011; Bednarek et al., 2009). Tryptophan-derived indole GSLs or their breakdown products alter defense responses to non-host pathogens illustrated by biosynthetic mutants devoid of indole GSLs being unable to deposit PAMP-induced callose in cell walls via an unknown signal and pathway (Clay et al., 2009; Bednarek et al., 2009). Similarly, indole GSL activation products have the ability to directly alter auxin perception by interacting with the TIR1 auxin receptor (Katz et al., 2015). In contrast to indole GSLs, mutants in aliphatic GSL accumulation alter flowering time and circadian clock oscillations (Kerwin et al., 2011; Jensen et al., 2015). The aliphatic GSL activation product, allyl isothiocyanate can induce stomatal closure but it is unknown if this is specific to allylGSL or a broader GSL property (Khokon et al., 2011; Boller and Felix, 2009). AllylGSL (other names are 2-propenylGSL and sinigrin) can also alter plant biomass and metabolism in Arabidopsis (Francisco et al., 2016a; Francisco et al., 2016b).
While these studies have provided hints that the GSL may have signaling potential, there is little understanding of the underlying mechanism or the structural specificity of the signal. To explore whether built-in signaling properties are a common attribute of GSLs, we screened for altered plant growth and development in the presence of specific purified aliphatic GSLs. In particular, we were interested in identifying candidate signals whose activity could ensure fast repartitioning of resources between development and defense. Here we present a novel signaling capacity specific to the aliphatic 3-hydroxypropylglucosinolate (3OHPGSL). Our results suggest that 3OHPGSL signaling involves the universally conserved TOR pathway for growth and development, as mutants in TORC and autophagy pathways alter responsiveness to 3OHPGSL application.

Results

3ohpgsl inhibits root growth in arabidopsis

We reasoned that if a GSL can prompt changes in plant growth it is an indication of an inherent signaling capacity. Using purified compounds, we screened for endogenous signaling properties among short-chain methionine-derived aliphatic GSLs by testing their ability to induce visual phenotypic responses in Arabidopsis seedlings. We found that 3OHPGSL causes root meristem inhibition, at concentrations down to 1 \( \mu \)M (Figure 1A). The observed response is concentration-dependent (Figure 1A–B). All Arabidopsis accessions accumulate 3OHPGSL in the seeds, and this pool is maintained at early seedling stages (Chan et al., 2011; Petersen et al., 2002; Brown et al., 2003). Previous research has shown that GSLs in the seed are primarily deposited in the embryo, accumulating to about 3 mmol/g suggesting that we are working with concentrations within the endogenous physiological range (Fang et al., 2012; Kliebenstein et al., 2007).

We tested how exogenous 3OHPGSL exposure to the roots alters 3OHPGSL accumulation within the shoot, and how this compares to endogenously synthesized 3OHPGSL levels. We grew the Col-0 reference accession and the myb28-1 myb29-1 mutant that is devoid of endogenous aliphatic GSLs in the presence and absence of exogenous 3OHPGSL. At day 10, the foliar 3OHPGSL levels were analyzed. Col-0 without treatment had average foliar levels of 3OHPGSL of 3.2 \( \mu \)mol/g and grown on media containing 5 \( \mu \)M, 3OHPGSL contributed an additional 2.2 \( \mu \)mol/g raising the total 3OHPGSL to 5.4 \( \mu \)mol/g (Figure 1C). The myb28-1 myb29-1 mutant had no measurable 3OHPGSL on the control plates and accumulated ~1.4 \( \mu \)mol/g upon treatment (Figure 1D). In agreement with the lower foliar 3OHPGSL accumulation in the myb28-1 myb29-1 mutant background, this double mutant had a lower root growth response to exogenous 3OHPGSL (Figure 1—figure supplement 1). Importantly, this confirms that the level of 3OHPGSL application is within the physiological range.

We then tested if 3OHPGSL or potential activation products inhibit root growth because of cell death or toxicity. The first evidence against toxicity came from the observation that even during prolonged exposures, up to 14 days of length, Col-0 seedlings continued being vital and green (Figure 1E). If there was toxicity the seedlings would be expected to senesce and die. We next tested if the strong root growth inhibition by 50 \( \mu \)M 3OHPGSL is reversible. Importantly, root inhibition is reversible, as the 3OHPGSL-mediated root stunting could be switched on and off by transfer between control media and media containing 3OHPGSL (Figure 1E). Based on the toxicity and GSL assays, we conclude that the 3OHPGSL treatments are at reasonable levels compared to normal Arabidopsis physiology, and that the phenotypic responses we observed were not caused by flooding the system with 3OHPGSL or toxicity.

Root inhibition is specific to 3OHPGSL

To evaluate whether 3OHPGSL mediated root inhibition is a general GSL effect or if it is structurally specific to 3OHPGSL, we tested if aliphatic GSLs with similar side-chain lengths, but different chain modifications, would induce similar root growth effects. First, we assessed 3-methylsulfinyl-propyl (3MSP) GSL, the precursor of 3OHPGSL, and the alkenyl-modified three carbon glucosinolate allyl (Figure 2A). In contrast to 3OHPGSL, neither of these structurally related GSLs possessed similar root-inhibiting activities within the tested concentration range (Figure 2B–D). We also analyzed the potential root inhibition for the one carbon longer C4-GSLs 4-methylsulfinylbutyl (4MSB) and but-3-enyl (Figure 2E). Neither of these compounds could inhibit root growth at the tested concentrations (Figure 2F–H). There is no viable commercial, synthetic or natural source for the 4-hydroxybutyl GSL.
Figure 1. 3OHP reversibly inhibits root growth. (A) 7-d-old seedlings grown on MS medium supplemented with a concentration gradient of 3OHP. (B) Quantification of root lengths of 7-d-old. Results are averages ± SE (n = 3–7; p<0.001). (C) Accumulation of 3OHP in shoots/areal tissue of 10-d-old Col-0 wildtype seedlings grown on MS medium supplemented with 5 µM 3OHP. Results are least squared means ± SE over three independent experimental replicates with each experiment having an average eleven replicates of each condition (n = 31–33; ANOVA P<0.001). (D) Accumulation of 3OHP in shoots of 10-d-old myb28 myb29 seedlings (aliphatic GSL-free) grown on MS medium supplemented with 5 µM 3OHP. Results are least squared means ± SE over two independent experimental replicates with each experiment having an average of four independent biological replicates of each condition (n = 8–14; ANOVA P<0.001). (E) 14-d-old seedlings grown for 1 week with or without 3OHP as indicated. After one week of development, the plants were moved to the respective conditions showed in week 2.

DOI: https://doi.org/10.7554/eLife.29353.003
The following figure supplement is available for figure 1:

Figure supplement 1. Root inhibition is affected by endogenous GSL levels.
DOI: https://doi.org/10.7554/eLife.29353.004
which prevented us from testing this compound. To test if the presence of a hydroxyl is essential for this response, we tested if either the R or S form of 2-hydroxybut-3-enylGSL had similar effects. This showed that neither enantiomer was similar to 3OHPGSL and that both actually stimulated root growth (Figure 2—figure supplement 1) The fact that only 3OHPGSL inhibits root elongation suggests that the core GSL structure (comprising of a sulfate and thioglucose) does not cause the effect. Importantly, this indicates that 3OHPGSL root inhibition is not a generic result of providing extra sulfur or glucose from the GSL core structure to the plant, as these compounds would be equally contributed by the other GSLs. Furthermore, the results confirm that there is no general toxic activity when applying GSLs to Arabidopsis. This evidence argues that the 3OHPGSL root inhibition effect

**Figure 2.** Root growth is not inhibited by all aliphatic GSLs. (A) The aliphatic glucosinolate biosynthetic pathway, from the C3 3-methyl-sulphinyl-propyl (3MSP) to the secondary modified 3-hydroxyl-propyl (3OHP) and 2-propenyl (allyl/sinigrin). (B–C) Root lengths of 7-d-old Col-0 wildtype seedlings grown on MS medium supplemented with a concentration gradient of the indicated aliphatic C3-GSL. The left most point in each plot shows the root length grown in the absence of the specific GSL treatment. Results are least squared means ± SE over four independent experimental replicates with each experiment having an average of 21 replicates per condition (n_{3MSP} = 59–153; n_{Allyl} = 52–153). Significance was determined via two-way ANOVA combining all experiments. (D) 7-d-old seedlings grown on MS medium with or without 50 µM of the indicated GSL. (E) The aliphatic glucosinolate biosynthetic pathway from the C4 4-methyl-sulphinyl-buty1 (4MSB) to But-3-enyl. (F–G) Root lengths of 7-d-old Col-0 wildtype seedlings grown on MS medium supplemented with a concentration gradient of the indicated aliphatic C4-GSL. The left most point in each plot shows the root length grown in the absence of the specific GSL treatment. Least squared means ± SE over four independent experimental replicates with each experiment having an average of 22 replicates condition (n_{4MSB} = 38–153; n_{But-3-enyl} = 68–164). Significance was determined via two-way ANOVA combining all experiments. (H) 7-d-old seedlings grown on MS medium with or without 50 µM of the indicated GSL. DOI: https://doi.org/10.7554/eLife.29353.005

The following figure supplement is available for figure 2:

**Figure supplement 1.** Root lengths of 7 DAG Col-0 WT grown on MS media supplemented with 50 µM of the indicated GSL. DOI: https://doi.org/10.7554/eLife.29353.006
links to the specific 3OHP side chain structure, indicating the presence of a specific molecular target mediating the root inhibition response.

3OHPGSL responsiveness is wider spread in the plant kingdom than GSL biosynthesis

The evolution of the GSL defense system is a relatively young phylogenetic event that occurred within the last ~92 Ma and is largely limited to the Brassicales order (Edger et al., 2015). The aliphatic GSL pathway is younger still (~60 Ma) and is limited to the Brassicaceae family with the enzyme required for 3OHPGSL production, AOP3, being limited to Arabidopsis thaliana and Arabidopsis lyrata within the Arabidopsis lineage (Edger et al., 2015; Kliebenstein et al., 2001a). However, 3OHPGSL is also found in the vegetative tissue of the close relative Olmarabidopsis pumila (dwarf rocket) (Windsor et al., 2005), and in seeds of more distant Brassicaceae family members such as the hawkweed-leaved treacle mustard (Erysimum hieracifolium), virginia stock (Malcolmia maritima), shepherd’s cress (Teesdalia nudicaulis), and alpine pennycress (Thlaspi alpestre) (Daxenbichler et al., 1980; Daxenbichler et al., 1991). These species are evolutionarily isolated from each other, suggesting that they may have independently evolved the ability to make 3OHPGSL (Kliebenstein et al., 2001a; Windsor et al., 2005; Cni, 2016). As such, 3OHPGSL is an evolutionarily very young compound and we wanted to determine if the molecular pathway affected by 3OHPGSL is equally young, or whether 3OHPGSL affects an evolutionarily older, more conserved pathway.

First we tested for 3OHPGSL responsiveness in plant species belonging to the GSL-producing Brassicales order (Figure 3A). We found that four of the five tested Brassicales species responded to 5 µM 3OHPGSL with root growth inhibition regardless of their ability to synthesize 3OHPGSL (Figure 3B–F). This suggests that responsiveness to 3OHPGSL application does not link to the ability to make 3OHPGSL. We expanded the survey by including plants within the eudicot lineage that do not have the biosynthetic capacity to produce any GSLs (Figure 3G) and found that 5 µM 3OHPGSL can inhibit root growth in several of the non-Brassicales species tested (Figure 3H–L). The ability of 3OHPGSL to alter growth extended to Saccharomyces cerevisae where 3OHPGSL led to slower log phase growth than the untreated control (Figure 3—figure supplement 1). Allyl GSL in the media had no effect on S. cerevisae growth showing that this was a 3OHPGSL mediated process (Figure 3—figure supplement 1). The observation that 3OHPGSL responsiveness is evolutionarily older than the ability to synthesize 3OHPGSL suggests that the molecular target of

3OHP reduces root meristem and elongation zone sizes

We hypothesized that 3OHPGSL application may alter root cellular development to create the altered root elongation phenotype. A reduction in root growth can be caused by inadequate cell division in the root meristematic zone or by limited cell elongation in the elongation zones (Figure 4A) (Henriques et al., 2014). To investigate how 3OHPGSL affects the root cellular morphology, we used confocal microscopy of 4-d-old Arabidopsis seedlings grown vertically with or without 10 µM 3OHPGSL. We used propidium iodide stain to visualize the cell walls of individual cells, manually counted the meristematic cells, and measured the distance to the point of first root hair emergence. Root meristems of 3OHPGSL treated seedlings were significantly reduced in cell number compared to untreated controls (Figure 4B–C). Moreover, we also observed a premature initiation of the differentiation zone, as the first root hairs were closer to the root tip upon 3OHPGSL treatment (Figure 4D–E). In addition, we saw bulging and branching of the root hairs in 3OHPGSL treated roots (Figure 4F). There was no morphological evidence of cell death in any root supporting the argument that 3OHPGSL is not a toxin. These results indicate that 3OHPGSL leads to root growth inhibition by reducing the size of the meristematic zone within the developing Arabidopsis root.

TORC-associated mutants alter 3OHPGSL responsiveness

The observed response to 3OHPGSL suggests that the target of this compound is evolutionarily conserved and alters root growth but does not affect the patterning of the root meristem. This indicates that key root development genes like SHR and SCR are not the targets as they affect meristem patterning (Sabatini et al., 2003; Hao and Cui, 2012). Mutants in GSL biosynthetic genes can lead to
Figure 3. Conservation of 3OHP responsiveness suggests a evolutionally conserved target. (A) Stylized phylogeny showing the phylogenetic relationship of the selected plants from the Brassicales family, branch lengths are not drawn to scale. (B–F) plants from the Brassicales family, grown on MS medium supplemented with or without 5 μM 3OHP. (G) Stylized phylogeny showing the phylogenetic relationship of all the selected crop and model plants, branch lengths are not drawn to scale. (H–L) Root growth of plants from diverse eudicot lineages, grown on MS medium supplemented with or without 5 μM 3OHP. Results are least squared means ± SE for each species using the following number of experiments with the given biological replication. Camelina three independent experimental replicates (n_{ctrl} = 8 and n_{3OHP}=12). Rucola three independent experimental replicates (n_{ctrl} = 17 and n_{3OHP}=17). Cress; three independent experimental replicates (n_{ctrl} = 19 and n_{3OHP}=18). Rape; seed four independent experimental replicates (n_{ctrl} = 14 and n_{3OHP}=13). Broccoli; three independent experimental replicates (n_{ctrl} = 10 and n_{3OHP}=13). Lotus; three independent experimental replicates (n_{ctrl} = 10 and n_{3OHP}=10). Linseed; three independent experimental replicates (n_{ctrl} = 11 and n_{3OHP}=11). Dill; three independent experimental replicates (n_{ctrl} = 14 and n_{3OHP}=13). Oregano; four independent experimental replicates (n_{ctrl} = 40 and n_{3OHP}=39). Tomato; three independent experimental replicates (n_{ctrl} = 11 and n_{3OHP}=15). A significant effect of treatment on the various species was tested by two-way ANOVA combining all the experimental replicates in a single model with treatment as a fixed effect and experiment as a random effect. DOI: https://doi.org/10.7554/eLife.29353.007
Figure 3 continued on next page
auxin over-production phenotypes as indicated by the superroot (SUR) 1 and 2 loci (Mikkelsen et al., 2004; Boerjan et al., 1995). However, the SUR genes are not evolutionarily conserved and 3OHPGSL does not create a superroot phenotype, showing that the genes are not the targets. A remaining conserved root regulator that does not alter meristem formation, but still alters root growth, is the TOR pathway (Xiong et al., 2013). Thus, we proceeded to test if mutants in the TOR pathway alter sensitivity to 3OHPGSL. Because TORC activity is sugar responsive, we investigated whether 3OHPGSL application may alter the response to sugar in genotypes with altered TORC activity. We first used the TOR kinase overexpression line GK548 (TORox) because it was the only one of several published TOR overexpression lines (Deprost et al., 2007) that behaved as a TOR overexpressor within our conditions (Figure 5—figure supplement 1). The GK548 TORox line exhibits accelerated TORC signaling and consequently grows longer roots on media containing sucrose (Figure 5 and (Deprost et al., 2007)). In addition, GK548 TORox meristems are harder to arrest (Figure 5B). Applying 3OHPGSL to the GK548 TORox line showed that this genotype had an

---

**Figure 3 continued**

The following figure supplement is available for figure 3:

**Figure supplement 1.** Yeast response to 3OHP suggests a conserved target throughout eukaryotes.

DOI: https://doi.org/10.7554/eLife.29353.008

---

**Figure 4.** 3OHP reduces root zone sizes. (A) Diagrammatic organization of a root tip; the meristem zone from the QC to the first cell elongation; the elongation zone ends when first root hair appears (Dolan and Davies, 2004). (B) Meristem size of 4-d-old Arabidopsis seedlings grown on MS medium with sucrose ±10 µM 3OHP. Results are least squared means ± SE over three independent experimental replicates with each experiment having an average of three replicates per condition (n_{ctrl} = 6; n_{3OHP}=9). Significance was tested via two-way ANOVA with treatment as a fixed effect and experiment as a random effect. (C) Confocal images of 4-d-old propidium iodide stained seedlings grown with and without 3OHP. Meristematic cells are marked with white asterisks, elongated cells with blue asterisks. (D) Appearance of first root hair; measured from the root tip on 4-d-old seedlings grown on MS medium with sucrose ±10 µM 3OHP. Results are least squared means ± SE over two independent experimental replicates with each experiment having an average of nine replicates per condition (n_{ctrl} = 17; n_{3OHP}=20). Significance was tested via two-way ANOVA with treatment as a fixed effect and experiment as a random effect. (E) Confocal images of 4-d-old propidium iodide stained seedlings grown with and without 3OHP. Protruding root hairs are marked with white/black asterisks. (F) 3OHP induced root hair deformations, confocal images of 4-d-old propidium iodide stained seedlings grown with 3OHP.

DOI: https://doi.org/10.7554/eLife.29353.009

---

Malinsky et al. eLife 2017;6:e29353. DOI: https://doi.org/10.7554/eLife.29353 9 of 24

---

Genomics and Evolutionary Biology | Plant Biology
**Figure 5.** TOR over-activation amplifies 3OHP response. (A) Root growth for low light grown seedlings. The seedlings were grown on MS medium without sucrose for 3 days, then transferred to the indicated media (Suc; sucrose). Multi-factorial ANOVA was used to test the impact of Genotype (Col-0 v TORox), Treatment (Control v Sucrose) and their interaction on root length. All experiments were combined in the model and experiment treated as a random effect. The ANOVA results from each day are presented in the table. (B) The root lengths grown photo-constrained and without sucrose (from A) displayed at each time point as relative to the respective sucrose activated roots. Results least squared means ± SE over three independent experimental replicates with each experiment having an average of nine replicates per condition (n = 26–30). Multi-factorial ANOVA was used to test the impact of Genotype (Col-0 v TORox), Treatment (Sucrose v Sucrose/3OHP) and their interaction on root length. All experiments were combined in the model and experiment treated as a random effect. The ANOVA results from each day are presented in the table. (C) Root growth for low light grown seedlings. The seedlings were grown on MS medium without sucrose for 3 days, then transferred to the indicated media. (D) Photo-constrained root lengths in response to sucrose and 3OHP (from A) displayed at each time point as relative to the respective sucrose activated roots. Results are least squared means ± SE over two independent experimental replicates with each experiment having an average of six replicates per condition (n = 11–14). (E) Schematic model; over expression of the catalytic subunit TOR increases growth and the relative 3OHP response.

DOI: https://doi.org/10.7554/eLife.29353.010

The following figure supplements are available for figure 5:

**Figure supplement 1.** Published TORox lines that did not display the TORox phenotype under our conditions.

DOI: https://doi.org/10.7554/eLife.29353.011

**Figure supplement 2.** RAPTOR1 haplo-insufficiency does not affect 3OHP response.

DOI: https://doi.org/10.7554/eLife.29353.012

**Figure supplement 3.** Loss of one of the two substrate-binding TORC-subunits affect 3OHP response.

DOI: https://doi.org/10.7554/eLife.29353.013
elevated 3OHPGSL-mediated inhibition of meristem reactivation in comparison to the WT (Figure 5C–D). This suggests that TORC activity influences the response to 3OHPGSL (Figure 5E).

We next investigated how genetically disrupting additional components of TORC affects 3OHPGSL responsiveness. In addition to the catalytic TOR kinase subunit, TORC consists of the substrate binding RAPTOR (Henriques et al., 2014; Rexin et al., 2015), and LST8 (13) (Figure 5E). In Arabidopsis—there is one copy of TOR, and two copies of both RAPTOR and LST8 (RAPTOR1/RAPTOR2 and LST8-1/LST8-2) (Moreau et al., 2012; Deprost et al., 2005).

RAPTOR1 and TOR null mutants are lethal as homozygotes (Menand et al., 2002; Deprost et al., 2005), and heterozygous raptor1 mutants did not display a significant change in 3OHPGSL responsiveness (Figure 5—figure supplement 2). We therefore tested insertion mutants within the weaker homolog RAPTOR2, whose null mutant is viable, and in our conditions shows mildly reduced root length on sucrose-containing media (Figure 5—figure supplement 3A and C). We found that, for two independent insertion lines raptor2-1 (Deprost et al., 2005; Anderson et al., 2005) and raptor2-2 (Deprost et al., 2005) (Figure 5—figure supplement 3E), there was a statistically significant reduction in 3OHPGSL response (Figure 5—figure supplement 3A–C). This supports the hypothesis that 3OHPGSL-associated signaling proceeds through TORC and that RAPTOR2 may play a stronger role in 3OH perception than RAPTOR1.

3OHPGSL treatment inhibits sugar responses

A key function of TORC activity is to control meristem cell division and this can be measured by meristem reactivation assays (Xiong et al., 2013). Thus, to further test if TORC dependent responses are altered by 3OHPGSL, seedlings were germinated in sugar-free media and photosynthesis-constrained under low light conditions to induce root meristem arrest when the maternal glucose is depleted (three days after germination). The root meristems were reactivated by applying exogenous sucrose (Figure 6A–C). By treating arrested root meristems with sucrose alone or in combination with 3OHPGSL we found that 3OHPGSL could inhibit meristem reactivation of sugar-depleted and photosynthesis-constrained seedlings (Figure 6D–E). Further, this response was dependent upon the 3OHPGSL concentration utilized. A similar response was found when treating with a TOR inhibitor such as rapamycin (Xiong et al., 2013), providing additional support to the hypothesis that 3OHPGSL may reduce root growth by altering TORC activity.

3OHPGSL pharmacologically interacts with the TOR-inhibitor AZD-8055

To further examine the possibility that 3OHPGSL may be affecting the TOR pathway, we proceeded to compare the effect of 3OHPGSL to published chemical TOR inhibitors. The active site TOR inhibitors were originally developed for mammalian cells and inhibit root growth in various plant species (Montané and Menand, 2013). Similar to 3OHPGSL, the active-site TOR inhibitor AZD-8055 (AZD) induces a reversible concentration-dependent root meristem inhibition (Montané and Menand, 2013). By directly comparing 3OHPGSL treatment with known TOR chemical inhibitors in the same system, we can test for interactions between 3OHPGSL and the known TOR inhibitors. An interaction between 3OHPGSL application and a known TOR inhibitor, e.g. an antagonistic relationship, is an indication that the same target is affected. To assess whether interactions between 3OHPGSL and TOR signaling occur, we grew seedlings vertically on media with combinations of 3OHPGSL and AZD and root-phenotyped the plants to compare the effect on root morphology. This identified a significant antagonistic interaction between AZD and 3OHPGSL (3OH x AZD), both in terms of root length response (Figure 7A) and in initiation of the differentiation zone (Figure 7B). This antagonistic interaction is also supported by the appearance of first root hair (Figure 7B), as the premature initiation of the differentiation zone in the presence of 10 μM 3OHPGSL did not change further upon co-treatment (Figure 7B). Moreover, there was a vast overlap in the phenotypic response to both compounds (Figure 7C–D); notably the closer initiation of the root differentiation zone to the root tip (Figure 7B–C) and the decreased cell elongation (Figure 7C, right panel). Together, this suggests that the TOR inhibitor AZD and 3OHPGSL have a target in the same signaling pathway as no additive effect is observed. Supporting this is the observation that 3OHPGSL treatment is phenotypically similar to a range of TOR active site inhibitors, as well as an inhibitor of S6K1 (one of the direct targets of TOR) (Figure 7—figure supplement 1).
Interestingly, the short root hair phenotype induced by AZD showed a synergistic interaction between AZD and 3OHPGSL suggesting that they may target different components of the TORC pathway that interact (Figure 7C). Further, while there is strong phenotypic overlap between AZD and 3OHPGSL, there are also specific activities. AZD induced a rounding of the root tip (Dolan and Davies, 2004), but co-treatment with 3OHPGSL restored a wildtype-like tip phenotype (Figure 7D). The lack of root rounding and root hair inhibition suggest that AZD and 3OHPGSL both target the TOR pathway, but at different positions. Alternatively, the 3OHPGSL may be a more specific TOR inhibitor and the additional AZD phenotypes could be caused by the ATP-competitive inhibitor having alternative targets in plants. Together, these results suggest that 3OHPGSL directly or indirectly targets the same molecular pathway as known TOR inhibitors (Jia et al., 2009).
Figure 7. Pharmacological interaction of 3OHPGSL and the TOR inhibitor, AZD. (A) Root lengths of 7-d-old Col-0 wildtype seedlings grown on MS medium with sucrose ± combinations of AZD and different concentrations of 3OHP. Results are least squared means ± SE over three independent experimental replicates with each experiment having an average of nine replicates per condition (n = 18–58). Multi-factorial ANOVA was used to test Figure 7 continued on next page
Blocking parts of the autophagy machinery affects 3OHPGSL associated signaling

Activation or repression of the TOR pathway leads to regulatory shifts in numerous downstream pathways (Figure 8E) (Sheen, 2014; Sablowski et al., 2014). For example, active TOR negatively regulates autophagy across eukaryotic species including Arabidopsis (Liu and Bassham, 2010; Shibutani and Yoshimori, 2014). To test if pathways downstream of TORC are affected by, or involved in, 3OHPGSL signaling, we analyzed mutants of two key autophagic (ATG) components, atg2-1 (18) and atg5-1 (58). ATG2 is part of the ATG9 cycling system that is essential for autophagosome formation (Feng et al., 2014; Velikkakath et al., 2012; Ryabovol and Mini-bayeva, 2016). ATG9-containing vesicles are a suggested membrane source for the autophagosome, and vesicles containing ATG9 are cycled to-and-from the phagophore via the ATG9 cycling system (Feng et al., 2014; Ryabovol and Minibayeva, 2016). ATG5, is part of the dual ubiquitin-like conjugation systems responsible for ATG8 lipidation (Feng et al., 2014; Ryabovol and Minibayeva, 2016; Fujioka et al., 2008). There are nine ATG8 paralogues in Arabidopsis (Ryabovol and Minibayeva, 2016) and together with the single copy of ATG5, they are essential for autphagosome initiation, expansion, closure, and vacuolar fusion (Feng et al., 2014; Ryabovol and Minibayeva, 2016). After the first conjugation system has conjugated ATG8 to an E2-like enzyme, the E3 ligase-like activity of the second ATG5-containing system enables ATG8 lipidation at the autophagic membrane (Feng et al., 2014; Walczak and Martens, 2013; Kaufmann et al., 2014). We found that atg5-1 enhanced 3OHPGSL responsiveness (Figure 8A–B) while atg2-1 had a wild type response (Figure 8C–D). One possible explanation for this difference between the two mutants is that, apart from macro-autophagy, plants also have micro-autophagy (Ryabovol and Minibayeva, 2016), a process that, in animal systems, has been shown to be negatively regulated by TOR (Li et al., 2012). Micro-autophagy does not involve de novo assembly of autophagosomes, and ATG5 has been shown to be involved in several forms of micro-autophagy whereas the role of ATG2 is more elusive and may not be required (Li et al., 2012). Thus, the elevated 3OHPGSL response in the atg5-1 mutant supports the hypothesis that 3OHPGSL signaling proceeds through the TOR pathway, but also suggests that this response requires parts of the autophagic machinery as it was not observed for atg2-1.

Discussion

In this study we describe a novel signaling capacity associated with 3OHPGSL, a defense metabolite present in Arabidopsis, and provide evidence that the linked signal proceeds via the TOR pathway. Application of exogenous 3OHPGSL caused reversible root meristem inhibition by morphological reprogramming of the root zones, i.e. dramatically reduced the root meristem size and limited root cell elongation (Figure 4). This response occurred at levels within the endogenous range and there was no evidence of cell death in any treated root, suggesting that this is not a toxicity response (Figure 1). Additionally, these morphological responses were specific to 3OHPGSL and not caused by
any structurally or biosynthetically related GSL, suggesting that these responses were not because of generic properties shared by GSLs (Figure 2). Exposing a wide phylogenetic array of plants, including lineages that have never produced GSLs, to 3OHPGSL showed that application of this compound can inhibit growth broadly across the plant kingdom as well as in yeast (Figure 3,
Figure 3—figure supplement 1). This suggests conservation of the downstream signaling pathway across these diverse plant lineages. Equally, if the signaling compound is not 3OHPGSL itself, but a derivative, then the required biosynthetic processes must be conserved. This conservation largely rules out the specific GSL activation pathway controlled by Brassicales specific thioglucosidases, myrosinases (Barth and Jander, 2006; Nakano et al., 2017; Bones and Rossiter, 2006). The phylogenetic conservation of the 3OHPGSL response led us to search for a target pathway controlling growth and development that would be evolutionary well conserved between the tested species.

By comparing the root phenotype identified with 3OHPGSL application to the published literature, we hypothesized that 3OHPGSL treatment may affect TORC, a key primary metabolic sensor that controls growth and development, and is conserved back to the last common eukaryotic ancestor (Henriques et al., 2014). Active site TOR inhibitors inhibit root growth in numerous plant species similar to 3OHPGSL application (Montané and Menand, 2013), supporting the hypothesis that 3OHPGSL may function via TORC. A model with 3OHPGSL affecting TORC would explain how 3OHPGSL can alter root development across the plant kingdom (Figure 3). Mechanistic support for this hypothesis came from a number of avenues. First, 3OHPGSL can block the TOR-mediated sugar activation of arrested meristems (Figure 6). Second, the TORox mutant intensifies 3OHPGSL linked signaling (Figure 5), and correspondingly loss-of-function mutants of the substrate binding TORC component raptor2 diminish the 3OHPGSL effect (Figure 5—figure supplement 3). Additionally, there are clear phenotypic overlaps between the root phenotypes induced by known TOR inhibitors and 3OHPGSL, e.g. root inhibition, inhibition of cell elongation, and notably the dramatic reduction of the meristem sizes (Figure 7). Critically, 3OHPGSL and known small-molecule inhibitors of TOR were mutually antagonistic for a number of phenotypes. In pharmacology, the outcomes of a drug combination can either be antagonistic, additive or synergistic, depending on whether the effect is less than, equal to, or greater than the sum of the effects of the two drugs (Jia et al., 2009). Antagonistic interactions, as observed with 3OHPGSL and AZD, can occur if two drugs exhibit mutual interference against the same target site, or if their targets converge on the same regulatory hub (Jia et al., 2009). Together, these lines of evidence suggest that 3OHPGSL or a derived metabolite targets the TOR pathway to alter root meristem development within Arabidopsis and potentially other plant species.

Extending the analysis to pathways downstream of TORC, showed that loss of ATG5, a vital component of the autophagic machinery (Liu and Bassham, 2010; Shibutani and Yoshimori, 2014), intensifies the 3OHPGSL response (Figure 8A-DB). This supports the hypothesis that 3OHPGSL signaling proceeds through the TOR complex, but also suggest that this signal requires parts of the autophagic machinery. Loss of another autophagic component, ATG2, did not influence the 3OH response (Figure 8C-D). Together, this raises the possibility that 3OHPGSL influenced responses involve predominantly a micro-autophagy pathway, which is ATG5- but may not be ATG2-dependent, rather than the macro-autophagy pathway that depends upon both genes (Ryabovol and Minibayeva, 2016; Li et al., 2012). Micro-autophagy removes captured cytoplasmic components directly at the site of the vacuole via tonoplast invagination. The cargo to be degraded ranges from non-selective fractions of the cytoplasm to entire organelles, dependent on the type of micro-autophagy. The two ubiquitin-like conjugation systems, and thereby ATG5, have been shown to be involved in several forms of micro-autophagy, such as starvation-induced, non-selective, and glucose-induced selective autophagy (Li et al., 2012). Interestingly, micro-autophagy involves vacuolar movement of cargo, and the vacuole is considered the main storage site for glucosinolates (Mikkelsen et al., 2004). Thus, ATG5 may be responsible for enabling the movement of exogenously applied 3OHPGSL out of the cytoplasm where it or a derivative metabolite could interact with the TORC pathway and into the vacuole. This would decrease the concentration of the 3OHPGSL associated signal and could explain why the atg5-1 mutant is more sensitive to 3OHPGSL application. Further work is required to test if ATG5 is functioning to attenuate the 3OHPGSL associated signal.

A conundrum for defense signaling compounds to affect growth is the evolutionary age discrepancy; defense metabolites are typically evolutionarily very young, as they are often species or taxa specific, while growth regulatory pathways are highly conserved across broad sets of plant taxa. This raises the question of which mechanism(s) may allow this connection between young metabolites and old regulatory pathways. This suggests that plants may sense young metabolites using evolutionarily old signaling pathways. Similar evidence is coming from other secondary metabolite
systems suggesting that this may be a general phenomenon. For example, an indolic GSL activation product can interact with the conserved TIR1 auxin receptor to alter auxin sensitivity within Arabidopsis (Katz et al., 2015). Similarly, an unknown phenolic metabolite appears to affect regulation of growth and development by influencing the Mediator complex that is conserved across all eukaryotes (Bonawitz et al., 2012; Bonawitz et al., 2014; Kim et al., 2014); and the plant polyphenol resveratrol directly inhibits the mammalian TOR to induce autophagy (Park et al., 2016). Thus, young plant metabolites can influence evolutionarily conserved pathways. Interestingly, this strongly resembles the action of virulence-associated metabolites within plant pathogens. Pseudomonad bacteria produce the evolutionarily young coronatine that alters the plant defense response by interacting with the conserved JA-Ile receptor COI1 (Xie et al., 1998). In plant/pathogen interactions, this ability of pathogen-derived metabolites to alter plant defense signaling is evolutionarily beneficial because it boosts the pathogen’s virulence in planta. It is less clear if this selective pressure model also applies to plant defense compounds that interact with endogenous signaling pathways. Following the plant/pathogen derived model, it is tempting to assume that such plant defense metabolites have been co-selected on their ability to affect the biotic attacker and simultaneously provide information to the plant. However, an alternative hypothesis that is that these examples may simply be serendipitous cases, where the defense metabolites happened to interact with a pathway and are potentially of no evolutionary benefit. In this model, the plant might still be adapting to the evolution of this new regulatory linkage. In the particular case of 3OHPGSL, the AOP3 enzyme that makes this compound evolved prior to the split between A. thaliana and A. lyrata suggesting that these species have had at least several million years/generations of potential to adapt. However, the two hypotheses need to be empirically tested. Central to testing between the two hypotheses is to assess if the observed signaling effects have any fitness benefit for the plant suggesting that even if the connections arose by serendipity that they have been maintained by a selective benefit. This will require field testing the fitness of plants that contrast for the presence of these connections. An alternate way to test between these hypotheses would be to conduct a broad survey of plant metabolites to test how many can affect signaling within the plant. If a large fraction of metabolites have potential signaling function, it is unlikely that all of these are simply serendipitous cases that have not had sufficient time to be removed by natural selection. However; earlier studies have provided evidence that both allyl GSL and the GSL breakdown product indole-3-carbinol affect plant signaling and growth (Katz et al., 2015; Francisco et al., 2016a; Francisco et al., 2016b). In addition, R- and S-2-hydroxybut-3-enyl GSL promoted root growth (Figure 2—figure supplement 1), suggesting that dual effects of defense metabolites such as 3OHPGSL are possibly more general.

Within this report, we provided evidence that 3OHPGSL, or derived compounds, appears to function as a natural endogenous TORC inhibitor that can work across plant lineages. This creates a link whereby the plant’s endogenous defense metabolism can simultaneously coordinate with growth. Such a built-in signaling capacity would allow coordination between development and defense, as the plant could use the defense compound itself as a measure of the local progress of any defense response and readjust development and defense to optimize against the preeminent threat. Future work is required to identify the specific molecular interaction that allows this communication to occur, this will help to illuminate how and why plants measure their own defense metabolism to coordinate available resources more broadly with growth. Future work might also ascertain whether there is a broader class of plant produced TOR inhibitors. If this is true, they might be highly useful in understanding TOR function across kingdoms of life and possibly to reveal significant aspects of this universally conserved pathway that may have gone unnoticed in other eukaryotic models.

Materials and methods

Plant materials

The genetic background for the Arabidopsis (Arabidopsis thaliana) mutants and transgenic lines described in this study is the Col-0 accession. The following lines were described previously: myb28-1 myb29-1 (SonderbySonderby et al., 2007), atg2-1 (Inoue et al., 2006), atg5-1 (Thompson et al., 2005), raptor1-1 (Deprost et al., 2005; Anderson et al., 2005), raptor1-2 (Deprost et al., 2005), raptor2-2 (Deprost et al., 2005), raptor2-1 (Deprost et al., 2005; Anderson et al., 2005), and the TORox lines G548, G166, S784, and S7817 (Deprost et al., 2007).
All genotypes were obtained and validated both genetically and phenotypically as homozygous for the correct allele.

Plant growth media and in vitro root growth assays

Seeds were vapour sterilized for 2–3 hr, by exposure to a solution of 100 mL household bleach (Klorin Original, Colgate-Palmolive A/S) mixed with 5 mL hydrochloric acid (12M), and ventilated for 30 min to one hour. After plating, on ½ strength Murashige and Skoog (MS) medium (2.2 g/l MS + vitamins) (Duchefa) with 1% (w/v) sucrose (Nordic Sugar), and 0.8% (w/v) micro agar (Duchefa), pH adjusted to 5.8, the seeds were stratified for two days in the dark at 4°C. For root length assays at normal light (115-130 μE) Arabidopsis seedlings were grown vertically at 22 °C day 20°C night under a 16 hr photoperiod and 80% humidity (long day). Concentrated 3OHPGSL in water was added to the agar post-sterilization to create media with the described concentration for each assay. The same method and water was used to create the media for the testing of the other specific GSLs. For meristem reactivation assays plants were grown as described in (Xiong et al., 2013), except that in our conditions we needed to go to 25μE to obtain meristem inhibition. Daily root lengths were manually marked (from day 3) with a permanent marker pen on the backside of the plate. After photography of 7-d-old seedlings the root growth was quantified using the ImageJ software (Schneider et al., 2012). The least square means (lsmeans) for the genotypes in response to different treatments were calculated across experiments (in R, see statistics), and plotted in excel.

In vitro root growth assays for the species (seed plating and growth conditions)

To test 3OHPGSL perception in other plant orders, seeds were obtained as listed in Supplementary File 1. Except for Solanum lycopersicum, here San Marzano tomatoes were bought in a local supermarket and the seeds were harvested, fermented and dried. All seeds were vapour sterilized for three hours (as above). Before plating, and Lotus japonicus MG20 (Lotus) were emerged in water and kept at 4°C for 1–2 weeks. Seeds were plated on vertical ½MS plates as specified in Supplementary File 1, stratified for four days in the dark at 4°C before being transferred to a long day growth chamber. Root growth was measured approximately every 24 hr (as described above).

Yeast strain, media, and growth conditions

The yeast strain, NMY51 with pOST1-NubI and pDHB1-LargeT ((Stagljar et al., 1998; Möckli et al., 2007); DUALsystem Biotech), was grown in liquid YPD media (2% w/v bactopeptone (Duchefa Biochemie), 1% w/v yeast extract (Becton, Dickinson and Company), 2% w/v glucose) with or without added GSLs, at 30°C and 150 rpm shaking.

Yeast growth assay

On day one; a 5 ml overnight culture was started from cryostock. Day two; four new 4 ml cultures were inoculated with 1 ml overnight culture, and grown overnight. On day three; an OD600 0.4 and a 0.04 dilution was prepared from each of the four cultures. 500 μl of each of the four cultures, at both dilutions, were transferred to a 96-well culture plate containing 500 μl YPD liquid media with 3OHPGSL or Allyl GSL, to final OD600 0.2 and GSL concentrations of 50, 10, 5, 1 and 0 μM. The yeast growth was measured at 0, 4, 6, 8, 24 and 48 hr. For each growth measurement 100 μl culture was transferred to a 96-well Elisa-plate together with three wells of YPD liquid media for standardization. Growth was measured with a SpectraMAX 190 (Molecular Devices) and SoftMax Pro 6.2.2 software. Growth rates and statistical analysis was calculated using the R software. The linear growth range was determined, and a linear regression using the lm() function in R was carried out to determine OD600 increase per hour (slope) and the yeast doubling time was calculated.

Glucosinolate analysis

Glucosinolates were extracted from whole plant tissue of adult plants (for 3OHPGSL extraction), or from or 10-d-old seedlings (3OHPGSL uptake) (Kliebenstein et al., 2001a; Kliebenstein et al., 2001b; Kliebenstein et al., 2001c), and desulfo-glucosinolates were analysed by LC-MS/TQ as desulfo-GSLs as described in (Crocoll et al., 2016).
Statistics
The R software with the R studio interface was used for statistical analysis (R Core Team, 2017; RStudio Team, 2015). Significance was tested using the Anova function (aov), lsmeans were obtained using the ‘lsmeans’ package (version 2.17) (Lenth, 2016). The letter groupings (Tukey’s HSD Test) were obtained using the ‘agricolae’ package (version 1.2–3) (de Mendiburu, 2010).

Confocal microscopy
To examine the root tip zones, we used confocal laser-scanning microscopy of 4-d-old seedlings grown vertically with or without treatment (with 3OHPGSL and/or various inhibitors). Samples were mounted on microscopy slides in propidium iodide solution (40 μM, Sigma) and incubated for 15 min. Confocal laser scanning microscopy was carried out on a Leica SP5-X confocal microscope equipped with a HC PL FLUOTAR 10 DRY (0.3 numerical aperture, 10X magnification) or a HCX lambda blue PL APO 320 objective (0.7 numerical aperture, 20X magnification) for close-up pictures of the meristem. To visualize the cell walls of individual cells the propidium iodide stain was excited at 514 nm and emission was collected at 600 nm to 680 nm. To determine the size of the meristems the confocal pictures we manually inspected and the meristematic cells marked and counted (the meristem region is defined as in (Dolan and Davies, 2004; Perilli and Sabatini, 2010)). To measure the distance from the root tip to the point of first root hair emergence we used ImageJ (Schneider et al., 2012).

Chemicals
The AZD8055 (Chresta et al., 2010), Torin2 (Liu et al., 2011), KU-63794 (García-Martínez et al., 2009), and WYE-132 (Yu et al., 2010) were purchased from Selleckchem. PF-4708671 (Pearce et al., 2010) and allyl/sinigrin were purchased from Sigma-Aldrich. 4MSB and 3MSP GSLs were purchased at C2 Bioengineering. But-3-enyl GSL was purified from Brassica rapa seeds while 3OHPGSL was purified from the aerial parts of 4–5 weeks old greenhouse-grown plants of the Arabidopsis accession Landsberg erecta (Kliebenstein et al., 2001c; Crocoll et al., 2016). The concentration of 3OHP and but-3-enyl GSL was determined by LC-MS/TQ as desulfo-GSLs. All inhibitors were dissolved in DMSO and stored as 10 mM stocks at –20°C. For allyl, 3MSP, and 4MSB ~ 100 mM GSL stocks were made with H2O and the concentration of GSLs within these stocks was determined by LC-MS/TQ (see above).

Acknowledgements
We thank the excellent technical assistance of the PLEN Greenhouse staff, and the DynaMo student helpers. We thank Dr. Svend Roesen Madsen for providing Camelina, and rape seeds, and Dr. Camilla Knudsen Baden for giving us lotus seeds.

Additional information

Competing interests
Daniel J Kliebenstein: Reviewing editor, eLife. The other authors declare that no competing interests exist.

Funding

| Funder                      | Grant reference number | Author                          |
|-----------------------------|------------------------|---------------------------------|
| Danmarks Grundforskningsfond| DNRF99                 | Frederikke Gro Malinovsky       |
|                            |                        | Marie-Louise F Thomsen          |
|                            |                        | Sebastian J Nintemann           |
|                            |                        | Baptiste Bourgine               |
|                            |                        | Meike Burow                     |
|                            |                        | Daniel J Kliebenstein           |
| National Science Foundation | IOS 13391205           | Daniel J Kliebenstein           |
| National Science Foundation | MCB 1330337            | Daniel J Kliebenstein           |
The funders had no role in study design, data collection and interpretation, or the decision to submit the work for publication.

Author contributions
Frederikke Gro Malinovsky, Conceptualization, Data curation, Formal analysis, Validation, Investigation, Visualization, Methodology, Writing—original draft, Writing—review and editing; Marie-Louise F Thomsen, Formal analysis, Validation, Investigation, Writing—review and editing; Sebastian J Niinemann, Lea Møller Jagd, Investigation, Visualization, Writing—review and editing; Baptiste Bourgine, Investigation, Writing—review and editing; Meike Burow, Conceptualization, Resources, Formal analysis, Supervision, Funding acquisition, Investigation, Visualization, Methodology, Writing—original draft, Project administration, Writing—review and editing; Daniel J Kliebenstein, Conceptualization, Data curation, Formal analysis, Supervision, Funding acquisition, Visualization, Methodology, Writing—original draft, Project administration, Writing—review and editing

Author ORCIDs
Frederikke Gro Malinovsky http://orcid.org/0000-0002-9833-7968
Meike Burow http://orcid.org/0000-0002-2350-985X
Daniel J Kliebenstein http://orcid.org/0000-0001-5759-3175

Decision letter and Author response
Decision letter https://doi.org/10.7554/eLife.29353.021
Author response https://doi.org/10.7554/eLife.29353.022

Additional files
Supplementary files
- Supplementary file 1. Plant seeds used for in vitro root growth assays for the various plant species. The source, common name, order, family, genus, species, and subspecies seeds are listed for the used plant species. As well as the plate size (cm × cm), and plating distance used for the individual response assays.
  DOI: https://doi.org/10.7554/eLife.29353.018
- Transparent reporting form
  DOI: https://doi.org/10.7554/eLife.29353.019

References
Anderson GH, Veit B, Hanson MR. 2005. The Arabidopsis AtRaptor genes are essential for post-embryonic plant growth. BMC Biology 3:12. DOI: https://doi.org/10.1186/1741-7007-3-12, PMID: 15845148
Baena-González E, Rolland F, Thevelein JM, Sheen J. 2007. A central integrator of transcription networks in plant stress and energy signalling. Nature 448:938–942. DOI: https://doi.org/10.1038/nature06069, PMID: 17671505
Baena-González E. 2010. Energy signaling in the regulation of gene expression during stress. Molecular Plant 3:300–313. DOI: https://doi.org/10.1093/mp/spq043, PMID: 2080814
Barth C, Jander G. 2006. Arabidopsis myrosinases TGG1 and TGG2 have redundant function in glucosinolate breakdown and insect defense. The Plant Journal 46:549–562. DOI: https://doi.org/10.1111/j.1365-313X.2006.02716.x, PMID: 16640593
Bednarek P, Pislew ska-Bednarek M, Svatos A, Schneider B, Doubsky J, Mansurova M, Humphry M, Consonni C, Panstruga R, Sanchez-Vallet A, Molina A, Schulze-Lefert P. 2009. A glucosinolate metabolism pathway in living plant cells mediates broad-spectrum antifungal defense. Science 323:101–106. DOI: https://doi.org/10.1126/science.1163732, PMID: 19095900
Boerjan W, Cervera MT, Delarue M, Beeckman T, Dewitte W, Bellini C, Caboche M, Van Onckelen H, Van Montagu M, Inzé D. 1995. Superroot, a recessive mutation in Arabidopsis, confers auxin overproduction. The Plant Cell Online 7:1405–1419. DOI: https://doi.org/10.1105/tpc.7.9.1405, PMID: 8589625
Boller T, Felix G. 2009. A renaissance of elicitors: perception of microbe-associated molecular patterns and danger signals by pattern-recognition receptors. Annual Review of Plant Biology 60:379–406. DOI: https://doi.org/10.1146/annurev.arplant.57.032905.105346, PMID: 19400727
Bonawitz ND, Soltau WL, Blatchley MR, Powers BL, Hurlock AK, Seals LA, Weng JK, Stout J, Chapelle C. 2012. REF4 and RFR1, subunits of the transcriptional coregulatory complex mediator, are required for phenylpropanoid homeostasis in Arabidopsis. Journal of Biological Chemistry 287:5434–5445. DOI: https://doi.org/10.1074/jbc.M111.312298, PMID: 22167189

Bonawitz ND, Kim JI, Tobimatsu Y, Ciesielski PN, Anderson NA, Ximenes E, Maeda J, Ralph J, Donohoe BS, Ladisch M, Chapelle C. 2014. Disruption of Mediator rescues the stunted growth of a lignin-deficient Arabidopsis mutant. Nature 509:375–380. DOI: https://doi.org/10.1038/nature13084, PMID: 24670657

Bones AM, Rossiter JT. 2006. The enzymic and chemically induced decomposition of glucosinolates. Phytochemistry 67:1053–1067. DOI: https://doi.org/10.1016/j.phytochem.2006.02.024, PMID: 16624350

Brown PD, Tokuhisa JG, Reichelt M, Gershenson J. 2003. Variation of glucosinolate accumulation among different organs and developmental stages of Arabidopsis thaliana. Phytochemistry 62:471–481. DOI: https://doi.org/10.1016/S0031-9422(02)00549-6, PMID: 12603620

Chan EK, Rowe HC, Corwin JA, Joseph B, Kliebenstein DJ. 2011. Combining genome-wide association mapping and transcriptional networks to identify novel genes controlling glucosinolates in Arabidopsis thaliana. PLoS Biology 9:e1001125. DOI: https://doi.org/10.1371/journal.pbio.1001125, PMID: 21857804

Chresta CM, Davies BR, Hickson I, Harding T, Cosulich S, Critchlow SE, Vincent JP, Elliston R, Jones D, Sini P, James D, Howard Z, Dudley P, Hughes G, Smith L, Maguire S, Hummerson M, Malagu K, Menea K, Jenkins R, et al. 2010. AZD8055 is a potent, selective, and orally bioavailable ATP-competitive mammalian target of rapamycin kinase inhibitor with in vitro and in vivo antitumor activity288–298. Cancer Research 70:288–298. DOI: https://doi.org/10.1158/0008-5472.CAN-09-1751, PMID: 20028854

Clay NK, Adio AM, Denoux J, Jander G, Ausubel FM. 2009. Glucosinolate metabolites required for an Arabidopsis innate immune response. Science 323:95–101. DOI: https://doi.org/10.1126/science.1164627, PMID: 19095898

Cni KDJ. 2016. Chapter Three – Nonlinear selection and a blend of convergent, divergent and parallel evolution shapes natural variation in glucosinolates. Advances in Botanical Research 80:31–55. DOI: https://doi.org/10.1016/bs.abr.2016.06.002

Crocoll C, Halkier BA, Burow M. 2016. Analysis and Quantification of Glucosinolates. Current Protocols in Plant Biology: John Wiley & Sons, Inc.

Crozet P, Margalha L, Confraria A, Rodrigues A, Martinho C, Adamo M, Elias CA, Baena-González E. 2014. Mechanisms of regulation of SNF1/AMPK/SnRK1 protein kinases. Frontiers in Plant Science 5:190. DOI: https://doi.org/10.3389/fpls.2014.00190, PMID: 24904600

Daxenbichler ME, Spencer GF, Schroeder WP. 1980. 3-Hydroxypropylglucosinolate, a new glucosinolate in seeds of Erysimum hieracifolium and Malcolmia maritima. Phytochemistry 19:813–815. DOI: https://doi.org/10.1016/0031-9422(80)85117-X

Daxenbichler ME, Spencer GF, Carlson DG, Rose GB, Brinker AM, Powell RG. 1991. Glucosinolate composition of seeds from 297 species of wild plants. Phytochemistry 30:2623–2638. DOI: https://doi.org/10.1016/0031-9422(91)85112-D

Deprost D, Truong H-N, Robaglia C, Meyer C. 2005. An Arabidopsis homolog of RAPTOR/KOG1 is essential for early embryo development. Biochemical and Biophysical Research Communications 326:844–850. DOI: https://doi.org/10.1016/j.bbrc.2004.11.117

Deprost D, Yao L, Sormani R, Moreau M, Letterreux G, Nicolai M, Bedu M, Robaglia C, Meyer C. 2007. The Arabidopsis TOR kinase links plant growth, yield, stress resistance and mRNA translation. EMBO Reports 8:864–870. DOI: https://doi.org/10.1038/sj.embor.7400437, PMID: 17721444

Dolan L, Davies J. 2004. Cell expansion in roots. Current Opinion in Plant Biology 7:33–39. DOI: https://doi.org/10.1016/j.pbi.2003.11.006, PMID: 14732439

Edger PP, Heidel-Fischer HM, Bekbaert M, Rota J, Glöckner G, Platts AE, Heckel DG, Der JP, Wafula EK, Tang M, Hofberger JA, Smithson A, Hall JC, Blanchette M, Bureau TE, Wright SI, dePamphilis CW, Erick Schranz M, Barker MS, Conant GC, et al. 2015. The butterfly plant arms-race escalated by gene and genome duplications. PNAS 112:8362–8366. DOI: https://doi.org/10.1073/pnas.1503926112, PMID: 26108883

Fang J, Reichelt M, Hidalgo W, Agnolet S, Schneider B. 2012. Tissue-specific distribution of secondary metabolites in rapeseed (Brassica napus L.). PLoS One 7:e48006. DOI: https://doi.org/10.1371/journal.pone.0048006, PMID: 23135359

de Mendiburu F. 2010. Agricolae: statistical procedures for agricultural research. https://CRAN-R-project.org/package=agricolae

Feng Y, He D, Yao Z, Kliosky DJ. 2014. The machinery of macroautophagy. Cell Research 24:24–41. DOI: https://doi.org/10.1038/cr.2013.168, PMID: 24366339

Francisco M, Joseph B, Caligasan H, Li B, Corwin JA, Lin C, Kerwin RE, Burow M, Kliebenstein DJ. 2016a. Genome wide association mapping in arabidopsis thaliana identifies novel genes involved in linking allyl glucosinolate to altered biomass and defense. Frontiers in Plant Science 7:1010. DOI: https://doi.org/10.3389/fpls.2016.01010, PMID: 27462337

Francisco M, Joseph B, Caligasan H, Li B, Corwin JA, Lin C, Kerwin R, Burow M, Kliebenstein DJ. 2016b. The defense metabolite, allyl glucosinolate, modulates arabidopsis thaliana biomass dependent upon the endogenous glucosinolate pathway. Frontiers in Plant Science 7:774. DOI: https://doi.org/10.3389/fpls.2016.00774, PMID: 27313596

Fujika Y, Noda NN, Fujii K, Yoshimoto K, Ohsumi Y, Inagaki F. 2008. In vitro reconstitution of plant Atg8 and Atg12 conjugation systems essential for autophagy. Journal of Biological Chemistry 283:1921–1928. DOI: https://doi.org/10.1074/jbc.M706214200, PMID: 18039664
Gallii G, Avin-Wittenberg T, Angelovic R, Fernie AR. 2014. The role of photosynthesis and amino acid metabolism in the energy status during seed development. *Frontiers in Plant Science* 5:447. DOI: https://doi.org/10.3389/fpls.2014.00447, PMID: 25232362

García-Martínez JM, Moran J, Clarke RG, Gray A, Cosulich SC, Chresta CM, Alessi DR. 2009. Ku-0063794 is a specific inhibitor of the mammalian target of rapamycin (mTOR). *Biochemical Journal* **421**:29–42. DOI: https://doi.org/10.1042/BJ20090489, PMID: 19402821

Hao Y, Cui H. 2012. SHORT-ROOT regulates vascular patterning, but not apical meristematic activity in the Arabidopsis root through cytokinin homeostasis. *Plant Signaling & Behavior* **7**:314–317. DOI: https://doi.org/10.4161/psb.19118, PMID: 22476466

Henriques R, Bögre L, Horváth B, Magyar Z. 2014. Balancing act: matching growth with environment by the TOR signalling pathway. *Journal of Experimental Botany* **65**:2691–2701. DOI: https://doi.org/10.1093/jxb/eru049, PMID: 24567496

Inoue Y, Suzuki T, Hattori M, Yoshimoto K, Ohsumi Y, Moriyasu Y. 2006. AtATG genes, homologs of yeast autophagy genes, are involved in constitutive autophagy in Arabidopsis root tip cells. *Plant and Cell Physiology* **47**:1641–1652. DOI: https://doi.org/10.1093/pcp/pcl031, PMID: 17085765

Jensen LM, Jepsen HS, Halliker BA, Kliebenstein DJ, Burrow M. 2015. Natural variation in cross-talk between glucosinolates and onset of flowering in Arabidopsis. *Frontiers in Plant Science* **6**:697. DOI: https://doi.org/10.3389/fpls.2015.00697, PMID: 26442014

Jia J, Zhu F, Ma X, Cao Z, Cao ZW, Li Y, Li YX, Chen YZ. 2009. Mechanisms of drug combinations: interaction and network perspectives. *Nature Reviews Drug Discovery* **8**:111–128. DOI: https://doi.org/10.1038/nrd2683, PMID: 19180105

Katz E, Nisani S, Yadav BS, Woldemariam MG, Shai B, Obolski U, Ehrlich M, Shani E, Jander G, Chamovitz DA. 2015. The glucosinolate breakdown product indole-3-carbinol acts as an auxin antagonist in roots of Arabidopsis thaliana. *The Plant Journal* **82**:547–555. DOI: https://doi.org/10.1111/tpj.12824, PMID: 25758811

Kaufmann A, Beier V, Franquelin HG, Wollert T. 2014. Molecular mechanism of autophagic membrane-scaffold assembly and disassembly. *Cell* **156**:469–481. DOI: https://doi.org/10.1016/j.cell.2013.12.022, PMID: 24485455

Kerwin RE, Jimenez-Gomez JM, Fulop D, Harmer SL, Maloof JN, Kliebenstein DJ. 2011. Network quantitative trait loci mapping of circadian clock outputs identifies metabolic pathway-to-clock linkages in Arabidopsis. *The Plant Journal* **64**:471–485. DOI: https://doi.org/10.1007/s00118-010-1806-8, PMID: 21343415

Khokon MA, Jahan MS, Rahman T, Hossain MA, Muroyama D, Minami I, Munemasa S, Mori IC, Nakamura Y, Murata Y. 2011. Allyl isothiocyanate (AITC) induces stomatal closure in Arabidopsis. *Plant, Cell & Environment* **34**:1900–1906. DOI: https://doi.org/10.1111/j.1365-3040.2011.02385.x, PMID: 21711355

Kim JI, Ciesielski PN, Donohoe BS, Chapple C, Li X. 2014. Chemically induced conditional rescue of the reduced epidermal fluorescence8 mutant of Arabidopsis thaliana reveals rapid restoration of growth and selective turnover of secondary metabolite pools. *Plant Physiology* **164**:584–595. DOI: https://doi.org/10.1104/pp.131.229393, PMID: 24381065

Kliebenstein DJ, Lambrix VM, Reichelt M, Gershenzon J, Mitchell-Olds T. 2001a. Gene duplication in the diversification of secondary metabolism: tandem 2-oxoglutarate-dependent dioxygenases control glucosinolate biosynthesis in Arabidopsis. *The Plant Cell Online* **13**:681–693. DOI: https://doi.org/10.1101/tpc.13.3.681, PMID: 11251105

Kliebenstein DJ, Gershenzon J, Mitchell-Olds T. 2001b. Comparative quantitative trait loci mapping of aliphatic, indolic and benzyl glucosinolate production in Arabidopsis thaliana leaves and seeds. *Genetics* **159**:359–370. PMID: 11560911

Kliebenstein DJ, Kroymann J, Brown P, Figuth A, Pedersen D, Gershenzon J, Mitchell-Olds T. 2001c. Genetic control of natural variation in Arabidopsis glucosinolate accumulation. *Plant Physiology* **126**:811–825. DOI: https://doi.org/10.1104/pp.126.2.811, PMID: 11402209

Kliebenstein DJ, Pedersen D, Barker B, Mitchell-Olds T. 2002. Comparative analysis of quantitative trait loci controlling glucosinolates, myrosinase and insect resistance in Arabidopsis thaliana. *Genetics* **161**:325–332. PMID: 12019246

Kliebenstein DJ, D’Auria JC, Behera AS, Kim JH, Gunderson KL, Breen JN, Lee G, Gershenzon J, Last RL, Jander G. 2007. Characterization of seed-specific benzoyloxyglucosinolate mutations in Arabidopsis thaliana. *The Plant Journal* **51**:1062–1076. DOI: https://doi.org/10.1111/j.1365-313X.2007.03205.x, PMID: 17651367

Lambrix V, Reichelt M, Mitchell-Olds T, Kliebenstein DJ, Gershenzon J. 2001. The Arabidopsis epiphiophor protein promotes the hydrolysis of glucosinolates to nitriles and influences Trichoplusia ni herbivory. *The Plant Cell Online* **13**:2793–2807. DOI: https://doi.org/10.1105/tpc.13.12.2793, PMID: 11752388

Lastdrager J, Hanson J, Smeekens S. 2014. Sugar signals and the control of plant growth and development. *Journal of Experimental Botany* **65**:799–807. DOI: https://doi.org/10.1093/jxb/eru474, PMID: 24453229

Le Bars R, Marion J, Le Borgne R, Satiat-Jeunemaitre B, Bianchi MW. 2014. ATGS defines a phagophore domain connected to the endoplasmic reticulum during autophagosome formation in plants. *Nature Communications* **5**:4121. DOI: https://doi.org/10.1038/ncomms5121, PMID: 24947672

Lenth RV. 2016. Least-Squares Means: The R Package lsmeans. *Journal of Statistical Software* **69**:1–33. DOI: https://doi.org/10.18637/jss.v069.i01

Li WW, Li J, Bao JK. 2012. Microautophagy: lesser-known self-eating. *Cellular and Molecular Life Sciences* **69**:1125–1136. DOI: https://doi.org/10.1007/s00018-011-0865-5, PMID: 22080117

Liu Y, Bassham DC. 2010. TOR is a negative regulator of autophagy in Arabidopsis thaliana. *PLoS One* **5**:e11883. DOI: https://doi.org/10.1371/journal.pone.0011883, PMID: 20686696
Liu Q, Wang J, Kang SA, Thoeran CC, Hur W, Ahmed T, Sabatini DM, Gray NS. 2011. Discovery of 9-(6-
aminopyridin-3-yl)-1-[(3-trifluoroethyl)methylpheno]benzo[hl]1,2-naphthyridine-2(1H)-one (Torin2) as a potent,
selective, and orally available mammalian target of rapamycin (mTOR) inhibitor for treatment of cancer. Journal of Medicinal Chemistry 54:1473–1480. DOI: https://doi.org/10.1021/jm101520v, PMID: 21322566

Menand B, Desnos T, Nussaume L, Berger F, Bouchez D, Meyer C, Robaglia C. 2002. Expression and disruption of the Arabidopsis TOR (target of rapamycin) gene. Plant Cell 14:6422–6427. DOI: https://doi.org/10.1016/j.plants.2002.06.005

Mikkelsen MD, Naur P, Halkier BA. 2004. Arabidopsis mutants in the C-S lyase of glucosinolate biosynthesis establish a critical role for indole-3-acetaldoxime in auzin homeostasis. The Plant Journal 37:770–777. DOI: https://doi.org/10.1111/j.1365-313X.2004.02002.x, PMID: 14871316

Montané MH, Menand B. 2013. ATP-competitive mTOR kinase inhibitors delay plant growth by triggering early differentials of meristematic cells but no developmental patterning change. Current Experimental Botany 64:4361–4374. DOI: https://doi.org/10.1093/jxb/ert242, PMID: 23963679

Moreau M, Azzopardi M, Clément G, Dobrenel T, Marchive C, Renne C, Martin-Magniette ML, Taconnat L,
Renou JP, Robaglia C, Meyer C. 2012. Mutations in the Arabidopsis homolog of LST8/G kinase, a partner of the target of Rapamycin kinase, impair plant growth, flowering, and metabolic adaptation to long days. The Plant Cell Online 24:463–481. DOI: https://doi.org/10.1105/tpc.111.091306, PMID: 22037851

Möckli N, Deplazes A, Hassa PO, Zhang Z, Peter M, Hottiger MO, Stagljar I, Auerbach D. 2007. Yeast split-
ubiquitin-based cytosolic screening system to detect interactions between transcriptionally active proteins. BioTechniques 42:725–730. DOI: https://doi.org/10.2144/000112455, PMID: 17612995

Nanono RT, Piślewski-Bednarek M, Yamada K, Edger PP, Miyahara M, Kondo M, Böttcher C, Mori M, Nishimura M, Schulze-Lefert P, Hara-Nishimura I, Bednarek P. 2017. PYK10 myrosinase reveals a functional coordination between endoplastic reticulum bodies and glucosinolates in Arabidopsis thaliana. The Plant Journal 90:204–220. DOI: https://doi.org/10.1111/tpj.13377, PMID: 27612205

Park D, Jeong H, Lee MN, Koh A, Kwon O, Yang YR, Noh J, Suh PG, Park H, Ryu SH. 2016. Resveratrol induces autophagy by directly inhibiting mTOR through ATP competition. Scientific Reports 6:21772. DOI: https://doi.org/10.1038/srep21772, PMID: 26902888

Pearce LR, Alton GR, Richter DT, Kath JC, Lingardo L, Chapman J, Hwang C, Alessi DR. 2010. Characterization of PF-4708671, a novel and highly specific inhibitor of p70 ribosomal S6 kinase (S6K1). Biochemical Journal 431:245–255. DOI: https://doi.org/10.1042/Bj20100124, PMID: 20704653

Perilli S, Sabatini S. 2010. Analysis of root meristem size development. Methods in Molecular Biology 655:177–187. DOI: https://doi.org/10.1007/978-1-60761-765-5_12, PMID: 20734261

Petersen BL, Chen S, Hansen CH, Olsen CE, Halkier BA. 2002. Composition and content of glucosinolates in developing Arabidopsis thaliana. Planta 214:562–571. DOI: https://doi.org/10.1007/s004250100659, PMID: 11925040

Rexin D, Meyer C, Robaglia C, Veit B. 2015. TOR signalling in plants. Biochemical Journal 470:1–14. DOI: https://doi.org/10.1042/Bj20150505, PMID: 26251442

Ryabovol VV, Minibayeva VF. 2016. Molecular mechanisms of autophagy in plants: role of ATG8 proteins in formation and functioning of autophagosomes. Biochemistry 81:348–363. DOI: https://doi.org/10.1134/S0006297916040052, PMID: 27293092

Sabatini S, Heidstra R, Wildwater M, Scheres B. 2003. SCARECROW is involved in positioning the stem cell niche in the Arabidopsis root meristem. Genes & Development 17:354–358. DOI: https://doi.org/10.1101/gad.252503, PMID: 12569126

Sablowski R, Carnier Dornelas M, Dornelas MC. 2014. Interplay between cell growth and cell cycle in plants. Journal of Experimental Botany 65:2703–2714. DOI: https://doi.org/10.1093/jxb/ert354, PMID: 24218325

Schneider CA, Rasband WS, Eliceiri KW. 2012. NIH Image to ImageJ: 25 years of image analysis. Nature Methods 9:671–675. DOI: https://doi.org/10.1038/nmeth.2089, PMID: 22930834

Sheen J. 2014. Master regulators in plant glucose signaling networks. Journal of Plant Biology 57:67–79. DOI: https://doi.org/10.1007/s12374-014-0902-7, PMID: 25530701

Shibutani ST, Yoshimori T. 2014. A current perspective of autophagosome biogenesis. Cell Research 24:58–68. DOI: https://doi.org/10.1038/cr.2013.159, PMID: 24296784

Smith AM, Stitt M. 2007. Coordination of carbon supply and plant growth. Plant, Cell & Environment 30:1126–1149. DOI: https://doi.org/10.1111/j.1365-3040.2007.01708.x, PMID: 17661751

Stagljar I, Korostensky C, Johnsson N, te Heesen S. 1998. A genetic system based on split-ubiquitin for the analysis of interactions between membrane proteins in vivo. PNAS 95:5187–5192. DOI: https://doi.org/10.1073/pnas.95.9.5187, PMID: 9560251

Senderby IE, Hansen BG, Bjarnholt N, Ticconi C, Halkier BA, Kliebenstein DJ. 2007. A systems biology approach identifies a R2R3 MYB gene subfamily with distinct and overlapping functions in regulation of aliphatic glucosinolates. PLoS One 2:e1322. DOI: https://doi.org/10.1371/journal.pone.0001322, PMID: 18094747

Senderby IE, Geu-Flores F, Halkier BA. 2010. Biosynthesis of glucosinolates—gene discovery and beyond. Trends in Plant Science 15:283–290. DOI: https://doi.org/10.1016/j.tips.2010.02.005, PMID: 20303821

RStudio Team. 2015. RStudio: Integrated Development for R. https://www.rstudio.com/

R Core Team. 2017. R: A language and environment for statistical computing. http://www.R-project.org/

Thompson AR, Doelling JH, Suttangkakul A, Vierstra RD. 2005. Autophagic nutrient recycling in Arabidopsis directed by the ATGB and ATG12 conjugation pathways. Plant Physiology 138:2097–2110. DOI: https://doi.org/10.1104/pp.105.060673, PMID: 16040659
Velikkakath AK, Nishimura T, Oita E, Ishihara N, Mizushima N. 2012. Mammalian Atg2 proteins are essential for autophagosome formation and important for regulation of size and distribution of lipid droplets. *Molecular Biology of the Cell* **23**:896–909. DOI: https://doi.org/10.1091/mbc.E11-09-0785, PMID: 22219374

Walczak M, Martens S. 2013. Dissecting the role of the Atg12-Atg5-Atg16 complex during autophagosome formation. *Autophagy* **9**:424–425. DOI: https://doi.org/10.4161/auto.22931, PMID: 23321721

Windsor AJ, Reichelt M, Figuth A, Svatos A, Kroymann J, Kliebenstein DJ, Gershenzon J, Mitchell-Olds T. 2005. Geographic and evolutionary diversification of glucosinolates among near relatives of Arabidopsis thaliana (Brassicaceae). *Phytochemistry* **66**:1321–1333. DOI: https://doi.org/10.1016/j.phytochem.2005.04.016, PMID: 15913672

Xie DX, Feys BF, James S, Nieto-Rostro M, Turner JG. 1998. COI1: an Arabidopsis gene required for jasmonate-regulated defense and fertility. *Science* **280**:1091–1094. DOI: https://doi.org/10.1126/science.280.5366.1091, PMID: 9582125

Xiong Y, McCormack M, Li L, Hall Q, Xiang C, Sheen J. 2013. Glucose-TOR signalling reprograms the transcriptome and activates meristems. *Nature* **496**:181–186. DOI: https://doi.org/10.1038/nature12030, PMID: 23542588

Yu K, Shi C, Toral-Barza L, Lucas J, Shor B, Kim JE, Zhang WG, Mahoney R, Gaydos C, Tardio L, Kim SK, Conant R, Curran K, Kaplan J, Verheijen J, Ayral-Kaloustian S, Mansour TS, Abraham RT, Zask A, Gibbons JJ. 2010. Beyond rapalog therapy: preclinical pharmacology and antitumor activity of WYE-125132, an ATP-competitive and specific inhibitor of mTORC1 and mTORC2. *Cancer Research* **70**:621–631. DOI: https://doi.org/10.1158/0008-5472.CAN-09-2340, PMID: 20068177

Zhuang X, Chung KP, Jiang L. 2016. Origin of the Autophagosomal Membrane in Plants. *Frontiers in Plant Science* **7**:1655. DOI: https://doi.org/10.3389/fpls.2016.01655, PMID: 27867391

Züst T, Heichinger C, Grossniklaus U, Harrington R, Kliebenstein DJ, Turnbull LA. 2012. Natural enemies drive geographic variation in plant defenses. *Science* **338**:116–119. DOI: https://doi.org/10.1126/science.1226397, PMID: 23042895