**BRC1 expression regulates bud activation potential, but is not necessary or sufficient for bud growth inhibition in Arabidopsis**

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ABSTRACT

The degree of shoot branching in Arabidopsis is determined by the activation of axillary buds. Bud activity is regulated by diverse environmental and developmental signals, often mediated via plant hormones including auxin, strigolactone and cytokinin. The transcription factor, BRANCHED1 (BRC1), has been proposed to integrate these regulatory signals. This idea is based on increased branching in brc1 mutants, the effects of bud-regulating hormones on BRC1 expression, and a general correlation between BRC1 expression and bud growth inhibition. These data demonstrate the important role of BRC1 in shoot branching, but here we show that in Arabidopsis this correlation can be broken. Buds lacking BRC1 expression can remain inhibited and sensitive to inhibition by strigolactone. Furthermore, buds with high BRC1 transcript levels can be active. Based on these data, we propose that BRC1 regulates bud activation potential in concert with an auxin-transport based mechanism underpinning bud activity. In the context of strigolactone-mediated bud regulation, our data suggest a coherent feed-forward loop in which strigolactone treatment reduces the probability of bud activation by parallel effects on BRC1 transcription and the shoot auxin transport network.
INTRODUCTION

The degree of shoot branching is an excellent example of plant developmental plasticity. In flowering plants, secondary shoots are formed by the activation of axillary buds, established in the axils of leaves produced by active shoot apical meristems. Differential activation of axillary buds allows a continuum of shoot forms, from a solitary unbranched stem to a highly ramified bush. Bud activity is regulated by diverse endogenous and environmental inputs. The integration of these inputs shapes the shoot system according to genotype and environment.

The mechanisms by which buds integrate regulatory signals are poorly understood. There are two non-exclusive candidates for the integrating hub (Rameau et al., 2015). According to the auxin transport canalization-based model for bud activation, sustained bud activity requires the establishment of canalized auxin transport from the bud into the main stem (Balla et al., 2011; Bennett et al., 2006; Li and Bangerth, 1999; 2003; Prusinkiewicz et al., 2009). The auxin transport canalization hypothesis posits that an initial passive flux of auxin between an auxin source and an auxin sink results in the upregulation and polarisation of auxin transporters in the direction of the flux, canalizing auxin transport into files of cells with highly polar, high capacity auxin transport connecting the auxin source to the sink (Sachs, 1969; 1981; see also Bennett et al., 2014). Axillary buds are potential auxin sources, and the main stem acts as an auxin sink by transporting auxin rootward (Prusinkiewicz et al., 2009; Sachs 1968). Since all active apices export auxin into the stem, thereby reducing its sink strength, axillary buds effectively compete for access to a common auxin transport path down the stem to the root. The ability of a bud to activate is therefore relative, depending on its strength as an auxin source compared to the auxin sink strength of the stem, and on the degree of positive feedback between auxin flux and auxin transporter upregulation/polarization. This system can act as a hub for the integration of multiple signals, which could influence bud activity by modulating stem auxin sink strength, bud auxin source strength, or canalization feedback.

The canalization-based model of bud regulation can account for diverse shoot branching phenomena, including apical dominance. It is long-established that auxin produced by the growing primary apex is transported down the stem, inhibiting the activity of subtending axillary buds (Thimann and Skoog, 1934). Decapitation removes the auxin source, allowing sustained bud growth, which is prevented by application of auxin to the decapitated stump (Thimann and Skoog, 1934). This auxin acts indirectly since very little is transported into the bud itself (Booker et al., 2003; Brown et al., 1979; Everat-Bourbouloux and Bonnemain, 1980; Prasad et al., 2003). Restated in terms of the canalization hypothesis, apically-derived auxin in the stem inhibits sustained bud
activation by reducing stem auxin sink strength, hence preventing auxin export from buds. The canalization-based model can also explain how the plant hormone strigolactone (SL) inhibits branching. One effect of SL is to trigger removal of the PIN-FORMED1 (PIN1) auxin exporter from the plasma membrane, thereby dampening positive feedback between auxin flux and auxin transporter polarization/up-regulation, making it more difficult for buds to activate (Crawford et al., 2010; Prusinkiewicz et al., 2009; Shinohara et al., 2013). This can explain why SL treatment enhances competition between branches on a two-node stem segment, focusing growth into one branch, rather than simply inhibiting both branches (Crawford et al., 2010).

The other likely regulatory hub for bud activity is expression of TCP transcription factors of the TEOSINTE BRANCHED1 (TB1) class. TB1 was originally identified through its role in maize domestication (Doebley et al., 1997). In contrast to maize, its wild relative, teosinte, is highly branched. This is due to constitutive over-expression of TB1 in maize, which inhibits the activity of axillary buds, but also influences floral transition in branches (Hubbard et al., 2002). As a result, branches from the middle nodes of the maize primary stem develop and terminate as female inflorescences (ears), whereas in teosinte they develop as elongating branches.

Closely related genes have been characterised in several species, including FINE CULM1 (FC1) in rice and BRANCHED1 (BRC1) in Arabidopsis, pea, tomato and potato (Aguilar-Martinez et al., 2007; Arite et al., 2007; Braun et al., 2012; Dun et al., 2012; Finlayson 2007; Takeda et al., 2003; Martin-Trillo et al., 2011; Nicolas et al., 2015). In core eudicots there appear to be three co-homologues of TB1/FC1 from grasses (Citerne et al., 2013; Howarth & Donoghue, 2006; Martin-Trillo & Cubas, 2010). In Arabidopsis two of these genes, BRC1 and BRC2, have effects on branching, but it is BRC1 that has the major effect and is proposed to act as a regulatory hub (Aguilar-Martinez et al, 2007; Poza-Carrion et al., 2007). Similar to TB1, BRC1 also affects aspects of the floral transition (Aguilar-Martinez et al, 2007; Niwa et al, 2013).

Across these species, loss of function of TB1-class genes results in increased branching. Furthermore, their transcript levels correlate with bud growth inhibition and change rapidly in response to bud-regulatory treatments. It has therefore been proposed that the transcription of TB1-class genes acts as an integrated readout of bud-regulatory signals. For example, in pea the level of BRC1 transcript in buds is regulated positively by the bud-activating hormone cytokinin (CK), and negatively by SL, with both bud activity and BRC1 transcript abundance reaching intermediate levels when buds are treated with both hormones simultaneously (Braun et al., 2012; Dun et al., 2012). In this model, apical dominance is attributed to the ability of auxin to increase expression of
SL biosynthetic genes and reduce the expression and CK biosynthetic genes in the stem, thereby regulating the supply of these hormones to buds (Dun et al., 2012; 2013). Consistent with this idea, the buds of brc1 mutants of both pea and Arabidopsis have been reported to be completely insensitive to SL (Brewer et al., 2009; Braun et al., 2012).

These two models are not mutually exclusive, but they are distinct. SL signalling/synthesis mutants have both increased PIN1 accumulation in the shoot and decreased bud BRC1 transcript levels, but these effects are apparently independent (Bennett et al, 2016b). Beyond this, it has proved difficult to design experiments that distinguish between the two models. It has not been possible to separate sustained bud outgrowth from bud auxin export, although limited bud elongation can occur in the presence of auxin transport inhibitors (Brewer et al., 2015). However, interpretation of these results is problematic because, as described above, the absolute levels of auxin transport are of limited relevance to bud activation, and it is difficult to make clean local perturbations that affect the relevant relative properties of the system. Thus the observation that plants with systemically inhibited auxin transport can still respond to strigolactone is consistent with both modes of action for strigolactone (Brewer et al., 2015). This problem is exacerbated by the fact that a central tenet of auxin transport canalization is that auxin transporters polarise in a flux-correlated manner and, despite its demonstrable predictive power, this process is currently entirely mechanistically obscure, making it difficult to perturb the system in informative ways (Bennett et al., 2014).

Conversely, there are unambiguous examples in which the correlation between bud growth inhibition and BRC1 transcript levels is broken. For example, maize SL deficient mutants are highly branched but maintain constitutively high TB1 expression (Guan et al., 2012), and genetic variation in maize branching does not correlate with TB1 expression (Kebrom & Brutnell, 2015). Furthermore, in rice FC1 transcript abundance is apparently insensitive to SL (Arite et al., 2007; Minakuchi et al., 2010). In addition, in Psbrc1 and fc1 mutants, some buds remain inhibited (Arite et al., 2007; Braun et al., 2012). These data suggest that BRC1 cannot be a simple branch regulatory hub. In order to explore this issue further we have focused on the role of BRC1 in Arabidopsis bud growth inhibition, where many relevant tools are available. We show that BRC1 is neither necessary nor sufficient for bud growth inhibition in a variety of contexts. Our data suggest that BRC1 acts to modulate bud activation potential, within a wider system of bud activity control.
RESULTS

*BRC1* transcript abundance decreases as buds activate

Following floral transition, Arabidopsis buds activate in a basipetal sequence starting at the uppermost cauline (inflorescence) node, proceeding down the shoot into the rosette nodes (Alvarez et al., 1992; Hempel and Feldman, 1994). We have previously shown that buds remain inactive when a young inflorescence, bearing one or two leaves, is removed from the plant and placed in a microcentrifuge tube containing nutrient medium (Ongaro et al., 2008; Prusinkiewicz et al., 2009). The buds can be activated by decapitation of the primary shoot apex (Ongaro et al., 2008) or basal supply of CK (Müller et al., 2015). To characterise the dynamics of *BRC1* transcript abundance in this system, we quantified *BRC1* transcript levels and the initiation of bud elongation in response to bud-activating treatments.

Excised shoot apices bearing a single cauline node with its associated axillary bud were left untreated, decapitated directly above the node, or supplied basally with CK (0.1 µM benzylaminopurine (BAP)). A decrease in bud *BRC1* transcript abundance was detected 1 h after decapitation, and levels continued to drop until 3 h post-decapitation (Fig. 1A). Decreases in *BRC1* transcripts were not detected at 15 or 30 minutes post-decapitation (Fig. S1). Similarly, *BRC1* transcripts decreased in buds within 1 h of basal CK treatment (Fig. 1B). To correlate this with bud growth, time-lapse images of treated and control buds were captured every hour for 24 hours, and the length of the elongating bud stem, between the axil and the oldest bud leaf was determined. Control buds elongate slowly across this time course (Fig. 1C). However, decapitation activates rapid bud growth well within 24 hours. After log transformation of the data to allow parametric testing, significant differences in bud stem length between intact and decapitated explants were apparent from 8 h ($p < 0.05$ for independent two-sample t-tests between treatment groups from 8 h). With basal CK treatment, rapid growth begins later (Fig. 1D), with a significant CK effect not observed until 19 hours after treatment ($p < 0.05$ for independent two-sample t-tests between treatment groups from 19 h).

Accurate bud stem measurement is challenging as growing buds slightly change their orientation within the focal plane of the camera. Given this, and more generally the likely differences in sensitivity of qRT-PCR vs length measurements, it cannot be clearly established whether changes in *BRC1* transcript abundance and growth are simultaneous, or one precedes the other. Interestingly, although both bud-activating treatments trigger a significant drop in *BRC1* transcript abundance
within one hour, significant differences in bud length are detected 11 hours later for CK-activated buds compared to decapitation-activated buds.

**Bud BRC1 transcript abundance responds to an apical signal**

An obvious hypothesis is that decapitation triggers changes in BRC1 transcript levels through main stem auxin depletion. To manipulate the time at which a bud might experience changes in main stem auxin, the length of stem between the bud and the decapitation site was varied (Bennett et al., 2016a). Stem explants bearing 3 nodes were isolated and subjected to 4 different treatments: intact primary shoot apex with lowest buds collected at 0 h; intact primary shoot apex with intermediate buds removed, and lowest buds collected at 1 h; lowest buds collected 1 h after decapitation directly above the bud; and lowest buds collected 1 h after decapitation 8 mm above the bud, with intermediate buds removed (Fig. 2A).

As expected, 1 h after decapitation immediately above the bud, a substantial reduction in BRC1 transcript abundance was observed. Decapitation 8 mm above the bud also reduced BRC1 transcript levels, but they were significantly higher than those observed in buds decapitated immediately above the node (Fig. 2B). Interestingly, in intact explants from which the intermediate buds had been removed, there was also a small but significant reduction in BRC1 transcripts in the lower bud relative to the 0 h time point. This suggests that even inactive buds can influence BRC1 transcription in the other buds in the shoot system, for example by contributing a small amount of auxin to the main stem. Together, these data suggest that an apically-derived signal, such as auxin, moving basipetally in the stem regulates BRC1 transcription.

To determine whether decapitation 8 mm above the bud also delays bud activation, bud stem elongation was assessed using time-lapse imaging. There was no detectable difference in the timing of bud stem elongation between explants decapitated directly vs 8 mm above the bud (Fig. 2C). Additionally, removing the upper buds of intact explants did not significantly alter bud length. Thus the subtle differences in BRC1 transcript dynamics observed in this assay are not reflected in detectable differences in bud elongation, although this could be because of the differential sensitivity of these assays, as described above.

**Bud BRC1 transcript abundance responds to auxin in the main stem**

To assess more directly whether main stem auxin influences bud BRC1 transcript abundance, a similar experimental setup was used, but with auxin applied to the decapitated stump in lanolin paste. Explants were decapitated immediately above the node and subjected to one of three apical
auxin treatments (Fig. 3A); lanolin containing the auxin analogue 1-naphthylene acetic acid (NAA) (1 mM); lanolin with dimethyl sulphoxide (DMSO); or lanolin with DMSO for the first 3 hours after decapitation then replaced with lanolin containing 1 mM NAA. Buds were collected 6 h (Fig. 3B) or 21 h (Fig. 3C) after the start of the experiment, giving the ‘NAA from 3 h’ explants 3 h or 18 h of apical NAA respectively.

When apical auxin was supplied immediately after decapitation, $BRC1$ transcript abundance remained high (Fig. 3B,C). In contrast, the ‘NAA from 3 h’ buds showed a major reduction in $BRC1$ transcript levels at 6 h post-decapitation, to levels not significantly different from the ‘lanolin only’ buds (Fig. 3B). By 21 h post-decapitation, $BRC1$ transcript abundance in ‘NAA from 3 h’ buds returned to levels comparable to those in ‘NAA from 0 h’ buds, and significantly higher than the lanolin only control (Fig. 3C). Both apical auxin treatments prevented bud outgrowth compared to lanolin only controls (Fig. 3D), despite the transient dip in $BRC1$ transcript abundance in ‘NAA from 3 h’ buds. Together, these data indicate that bud $BRC1$ transcripts are dynamically responsive to auxin in the main stem, but only sustained down-regulation of $BRC1$ is associated with sustained bud outgrowth.

**Auxin-mediated bud inhibition is partially dependent on $BRC1$**

To investigate the relationship between auxin-mediated bud inhibition and $BRC1$ activity, we tested the sensitivity of $brc1-2$ $brc2-1$ mutant buds to apical auxin supply. Plants were grown axenically and stem segments bearing one cauline node with an associated bud less than 1.5 mm were excised and inserted between two agar slabs in a Petri dish (Chatfield et al., 2000). Auxin was applied via the apical slab at concentrations ranging from 1 to 10 µM and bud growth was monitored for 10 days (Fig. 4A,B).

In wild-type, apical auxin delayed bud activation, as shown by plotting the median number of days for buds to reach 5 mm, which increased with increasing auxin concentration (Fig. 1E). However, even with high auxin treatments, a few buds eventually activated, elongating at a rate similar to untreated buds. This can be seen by aligning the growth curves for each bud at the time-point when the bud first reached or exceeded 5 mm (defined as $t = 0$ on the graph in Fig. 4C). Thus in wild-type, auxin delays bud activation without affecting its kinetics.

The buds of $brc1-2$ $brc2-1$ mutants responded differently to apical auxin (Fig. 4B,D). They did not show the typical switching behaviour seen in wild-type, but either elongated slowly throughout the experiment, or elongated slowly and then arrested (Fig. 4B). The onset of elongation was only
slightly delayed, even by high concentrations of auxin (Fig. 4E), but the subsequent rate of elongation was inversely proportional to auxin concentration (Fig. 4D). These data suggest that \textit{BRC1} is involved in modulating bud activation dynamics in response to auxin.

\textbf{BRC1 is not necessary for bud growth inhibition}

In long-day-grown Arabidopsis, the number of primary branches formed in \textit{brc1-2 brc2-1} closely matches the number of primary axis leaves (Aguilar-Martinez et al., 2007; Bennett et al., 2016b). This could indicate that \textit{BRC1} is essential for bud growth inhibition. To test this idea, we used low nitrate availability, which suppresses branching in Arabidopsis (de Jong et al, 2014). Plants were grown with high (9 mM) or low (1.8 mM) nitrate supply. As expected, wild-type plants produced significantly more branches on high compared to low nitrate (Fig. 5C). Although \textit{brc1-2 brc2-1} mutants had significantly more branches than wild-type in both conditions, they were still capable of reducing branch number under nitrate limitation (Fig. 5C). As previously reported (Aguilar-Martinez et al 2007, Niwa et al 2013), \textit{brc1-2 brc2-1} plants had a greater proportion of floral branches in both conditions compared to wild-type (Fig. 5E). However there were no differences in the total number of primary leaf-bearing nodes either between genotypes or between treatments, and \textit{brc1-2 brc2-1} mutants had inhibited buds in both conditions (Fig. 5C,D). These data demonstrate that \textit{BRC1} is not necessary for bud growth inhibition.

To investigate this effect further in a situation with a larger number of primary leaf-bearing nodes, and therefore buds, we grew wild-type and \textit{brc1-2 brc2-1} mutants in short days and determined the number of primary leaf-bearing nodes and branches formed. The number of primary leaves was similar between the two genotypes (Figure 5A). While \textit{brc1-2 brc2-1} still produced twice as many branches as wild-type, more than half its nodes did not produce an elongated branch (Fig. 5A), confirming that \textit{BRC1} is not necessary for bud growth inhibition.

To manipulate leaf number in long-day conditions, we crossed \textit{brc1-2 brc2-1} into the \textit{flowering locus t-10 (ft-10)} mutant background, which has delayed flowering and increased vegetative node number in long days (Koornneef et al, 1991; Yoo et al., 2005). We observed that the total number of branches on wild-type and \textit{ft-10} plants was similar (Fig. 5B). Likewise, the number of branches produced by \textit{brc1-2 brc2-1} and \textit{ft-10 brc1-2 brc2-1} plants was almost identical (Fig. 5B). Thus, as in short days, \textit{brc1-2 brc2-1} double mutants have many inhibited buds in the \textit{ft-10} background. Loss of \textit{BRC1} increases the number of buds that activate, but this effect is limited, and independent of the total number of leaves, and hence buds, on the primary axis. It has previously been shown that \textit{brc1} has accelerated floral transition in its branches, and that this is suppressed by loss of \textit{FT} (Niwa et
al., 2013). Our data suggest that ft does not suppress the brc1 branch outgrowth phenotype (Fig. 5B).

**BRC1 transcript accumulation is not sufficient for bud growth inhibition**

We next investigated whether high BRC1 transcription is sufficient to prevent bud outgrowth. An oestradiol-inducible lex: BRC1 line was used to induce overexpression of BRC1 (Gonzalez-Grandio et al., 2013) by supplying 10 mM β-oestradiol basally to shoot explants bearing a single cauline node held in microcentrifuge tubes and either decapitated or left intact. After three hours treatment, oestradiol increased BRC1 transcripts in both intact and decapitated samples to levels greater than normally observed in intact explants, where buds are inhibited (Fig. 6C). Specifically, BRC1 transcript levels were significantly higher in induced vs uninduced decapitated samples, but this had no effect on branch growth (Fig. 6D). Branches grew normally despite sustained contact with the oestradiol. While it is possible that the high levels of BRC1 were not sustained, the result suggests that BRC1 transcript accumulation may not be sufficient to maintain bud growth inhibition.

To investigate situations in intact plants where growing buds might have high BRC1 transcript levels, we turned to mutants in the recently identified SMAX1-LIKE6 (SMXL6), SMXL7 and SMXL8 proteins, which are the proteolytic targets of MAX2-mediated SL signalling (Soundappan et al., 2015; Wang et al., 2015). Their over-accumulation in SL signalling mutants, such as max2, results in increased branching and constitutively low BRC1 transcript accumulation in buds. Mutation of SMXL6, 7 and 8 completely restores max2 branching to wild-type, and results in very high BRC1 transcript levels in inhibited buds (Soundappan et al., 2015; Wang et al., 2015). This supports the hypothesis that SL inhibits bud growth by up-regulating BRC1 transcription. Despite this, the smxl6-4 smxl7-3 smxl8-1 max2-1 (hereafter smxl678 max2) mutant has near wild-type levels of branching. We reasoned that BRC1 expression might be constitutively high in smxl678 max2 buds, whether active or inhibited. To test this hypothesis, buds 5-10 mm long (i.e. beginning to grow) from 4 plants of each genotype were pooled and their RNA extracted for analysis. In these apical smxl678 max2 buds, BRC1 transcript levels were extremely high compared to WT (Fig. 6A). Despite this, these apical branches on smxl678 max2 mutants grow with normal kinetics (Fig. 6B). This provides further evidence that high BRC1 expression is not sufficient to suppress bud growth, although it is possible that there are post-transcriptional effects on BRC1 activity operating specifically in these buds.
**brc1** is epistatic to **smxl6 smxl7 smxl8**

This raises interesting questions about the relationship between strigolactone, **BRC1** and the SMXL678 clade in the regulation of shoot branching. We therefore generated **smxl678 max2-1 brc1-2** quintuple mutants, and compared their branching to that of **smxl678 max2-1** quadruple mutants. Branching relationships among the parental types were as previously reported (Soundappan et al., 2015; Wang et al., 2015; Aguilar-Martinez et al., 2007). The **smxl678 max2-1** quadruple mutant had branching levels similar to wild-type, with the highly branched phenotype of **max2** being completely suppressed (Fig. 7A). Strikingly **brc1-2** is epistatic to **smxl678** (Fig. 7A), bringing the branching level of **smxl678 max2-1** up to that of **brc1**. This result is consistent with a model in which SMXL678 promote branching by down-regulating **BRC1**, but it is equally consistent with **BRC1** acting independently of SMXL678.

**BRC1** is not required for strigolactone-mediated bud inhibition

Previous comparisons of **brc1** and SL signalling mutants showed that loss of SL signalling results in higher branching levels than loss of **BRC1** (Chevalier et al, 2014; Bennett et al, 2016b), suggesting that misregulation of **BRC1** expression is insufficient to explain the full branching phenotype of SL mutants. Extending this result, we found that growth in short days exaggerates the differences between SL mutants and **brc1-2 brc2-1**, with **max4-5, d14-1** and **max2-1** having almost double the branch number of **brc1-2 brc2-1** (Fig. 7C). We also made triple mutants between **max4-5 or d14-1** and **brc1-2 brc2-1**. Consistent with previous results, the triple mutants had considerably more branches than either parent (Fig. 7B,D) (Chevalier et al, 2014). Since SL mutants typically make a branch at almost every primary node, the strong additivity suggests activation of higher order branches in the triple mutant. The triple mutants also showed additivity with respect to the parental reduced height phenotypes (Figure S2). These data provide further evidence that SL and **BRC1** act at least partially independently.

Support for the idea that SL regulates branching via **BRC1** comes from reports that **brc1** buds are SL insensitive (Brewer et al, 2009). Branching in **brc1** mutants was unaffected by growth on 5.8 µM GR24 (a strigolactone analogue). However, in the growth conditions used in that experiment very few branches were formed for any genotype regardless of treatment (Brewer et al., 2009). To explore SL response in more branch-conducive conditions, we grew plants of relevant genotypes axenically in jars on agar-solidified nutrient media supplemented with 5 µM GR24, or a solvent control (Fig. 8D). The total number of branches longer than 1cm was counted after 6 weeks (Fig. 8D). Consistent with previous reports (Crawford et al., 2010; Shinohara et al., 2013), GR24
strongly reduced branching in the SL biosynthesis mutant, \textit{max4-5}, but had no effect on the SL signalling mutant, \textit{d14-1}. Interestingly, branching in \textit{brc1-2 brc2-1} was significantly reduced by GR24 treatment, though less than in \textit{max4-5}, implying that \textit{brc1-2 brc2-1} retains partial SL sensitivity.

To test this further, we used a bud-bud competition assay. Shoot explants bearing two cauline nodes were placed in microcentrifuge tubes with media supplemented with 5 µM GR24 or a control. The explants were decapitated and branch length measured over time, mimicking the classical 2-branch pea assay developed by Snow (Snow 1925; 1929; 1931, Ongaro et al., 2008). Typically in these experiments, both buds begin to grow, and over time either both continue to grow at a similar rate, or one dominates the other, leading to highly asymmetric growth of the two branches (Ongaro et al., 2008). The relative frequency of these outcomes is assessed using a relative growth index (RGI), defined as the length of the longest branch divided by the total length of both branches. A value close to 0.5 indicates equal branch growth, whereas values closer to 1 indicate that one branch dominates the other. In this assay, SL mutants have low RGIs compared to wild-type, and GR24 treatment increases the RGI in wild-type and SL biosynthetic mutants, but not in SL signalling mutants (Fig. 8A,B,C), (Crawford et al., 2010). Untreated \textit{brc1-2 brc2-1} explants had a low RGI, similar to \textit{d14-1} and \textit{max4-5} (Fig. 8C). GR24 treatment resulted in a significant increase in RGI, of a magnitude similar to that observed in wild-type and \textit{max4-5} explants. In general, the phenotype of \textit{brc1-2 brc2-1} in this assay was both quantitatively and qualitatively similar to \textit{max4-5}. When comparing the mean length of the longer (Fig. 8A) and shorter (Fig. 8B) branches on each explant, it is clear that GR24 treatment strongly inhibits one branch in wild-type, \textit{max4-5}, and \textit{brc1-2 brc2-1} mutants, while the other branch grows relatively vigorously. In contrast, both branches of the \textit{d14-1} mutant grow strongly regardless of GR24 treatment. Therefore in this assay, \textit{brc1-2 brc2-1} mutants are fully strigolactone sensitive.
DISCUSSION

**BRC1 is not necessary or sufficient for bud growth inhibition**

The Arabidopsis *BRC1* gene, and its homologues in other species, are important regulators of shoot branching. Their loss of function results in highly branched phenotypes and their expression correlates with bud growth inhibition. Indeed, *BRC1* transcript level is often used as a marker for bud growth inhibition. This has led to the hypothesis that branch regulatory signals are integrated at the level of *BRC1* expression (Aguilar-Martinez et al., 2007; Braun et al., 2012). The data we present here suggest that this integration operates within a wider framework for bud activity control. Consistent with previous results, we show that a reduction in *BRC1* transcripts is an early event during bud activation by decapitation or by CK treatment (Fig. 1), and that bud outgrowth is associated with a sustained reduction in *BRC1* transcripts (Aguilar-Martinez et al., 2007; Martin Trillo et al., 2011; 2007; Braun et al., 2012; Nicolas et al., 2015). However, we also show that *brc1 brc2* mutants have inhibited buds (Fig. 5), and that elevated *BRC1* expression is not sufficient to prevent bud outgrowth (Fig. 6). Thus high *BRC1* transcript levels are neither necessary nor sufficient for bud inhibition.

**A bud activation threshold model for BRC1 action**

As described above, in Arabidopsis after the floral transition buds activate in a basipetal sequence. This sequence is likely generated by release of buds from auxin-mediated apical dominance, which is weakened at floral transition due to reduced auxin export from floral compared to vegetative apices (Prusinkiewicz et al., 2009). It is striking that the buds that remain inhibited in *brc1-2 brc2-1* mutants are those at the basal end of this sequence, whereas those that are active even with high *BRC1* transcription, as in the *smx678 max2* mutant, are those at the apical end. This suggests the hypothesis that *BRC1* modulates a bud activation threshold within the broader context of an auxin-mediated bud regulatory programme, such as the well-supported canalization-based mechanism described in the introduction. According to this idea, for buds with high *BRC1* expression, activation requires a highly canalization-conducive environment, such as the relatively low stem auxin concentration at apical nodes. In contrast, for *brc1-2 brc2-1* buds, which lack *BRC1* expression, bud inhibition requires a strongly canalization-inhibitory environment, such as the high main stem auxin concentration at the basal nodes (Prusinkiewicz et al., 2009). This model could explain the correlation between bud inhibition and *BRC1* expression, while at the same time being fully consistent with the evidence that *BRC1* is neither necessary nor sufficient for bud inhibition. The model also straightforwardly explains why *brc1-2 brc2-1* mutants produce similar numbers of branches, irrespective of the number of leaf-bearing nodes on the primary stem (Fig 5).
BRC1, bud activation potential and shoot system architecture

Our data support the hypothesis that BRC1 acts to modulate a bud activation threshold in concert with a systemic and relative auxin transport-mediated regulatory system for branching. BRC1 transcript levels can be locally regulated by specific environmental cues, such as light quality (Finlayson et al., 2010; Gonzalez-Grandio et al., 2013; Gonzalez-Grandio et al., 2016). Under our model, this would adjust the local activation threshold, allowing branches to activate (or be inhibited) out of the normal basipetal sequence. Thus BRC1 may be a point of integration between local, absolute regulatory inputs and systemic relative branching regulatory mechanisms, combining the advantages of both. In this way, BRC1-mediated variation in bud activation potential could also contribute to the patterns of branching observed in some species. For example, in rice the BRC1 homologue, FC1, is expressed at higher levels in the most basal bud compared to the next bud up and this correlates with their growth inhibition (Arite et al., 2007).

This threshold model is also interesting in the context of recent evidence that changes in sugar accumulation in buds correlate strongly with the down regulation of BRC1 expression and initiation of bud expansion in pea following decapitation (Mason et al., 2014). This effect can explain the initiation of bud expansion significantly ahead of main stem auxin depletion following decapitation. However, while application of apical auxin to the decapitated stump does not prevent this initial expansion, it does prevent sustained bud growth (Morris et al., 2005), despite the presumably high sugar content in the buds and the lack of a strong apical sugar sink. These observations are consistent with the idea that down-regulation of BRC1 can prime buds for activation, but this is not sufficient for sustained bud growth, which appears to require a canalization permissive environment.

Feed-forward regulation and bud activation dynamics

Although changes in BRC1 transcript levels are not required for changes in bud activity, BRC1 expression is clearly dynamically regulated during bud activation in Arabidopsis. For example, BRC1 expression is rapidly down-regulated following decapitation, likely in response to reduced auxin concentrations in the main stem. Since it is not required for bud activation, and indeed the timing of down-regulation does not correlate robustly with the timing of initiation of rapid bud elongation, what then is the function of this reduced BRC1 expression? If low stem auxin does indeed act in two distinct ways, promoting auxin transport canalization out of the bud, and down-regulating BRC1 expression, this creates what is essentially a coherent feedforward network motif. This type of regulatory circuit has been shown to increase robustness in switching mechanisms and
to affect unidirectionally the timing of switching (Mangan & Alon, 2003; Shen-Orr et al., 2002). The effects of brc1-2 brc2-1 mutation on bud auxin response are very interesting in this context. In a wild-type background, which according to our model would have both arms of the feedforward system, increased stem auxin delays bud activation, which eventually proceeds rapidly in a strong switch-like manner. In contrast, when one of the arms of the feedforward loop is removed, as in the brc1-2 brc2-1 mutant background, increased stem auxin only weakly delays initiation of growth, but the switch-like behaviour of wild-type buds is attenuated to give a more gradual response (Fig. 4).

This system differs in several important respects from the well-studied feedforward circuits, which are microbial transcriptional on-off switches (Mangan & Alon, 2003; Shen-Orr et al., 2002). In the case of bud activation, the canalization mechanism is itself driven by a positive feedback loop providing switch-like behaviour and hysteresis, making reversibility difficult. Furthermore, active buds export auxin into the stem (Balla et al., 2011; Li and Bangerth 1999; Morris, 1977; Thimann and Skoog, 1933) thereby closing a negative feedback loop by re-establishing high stem auxin. Nonetheless, a dual acting system, such as the one we propose here, is likely to confer properties to the bud activation switch not possible with a unitary pathway, and this will be important to explore further.

**BRC1 and strigolactone signalling**

Integration of *BRC1*-mediated and auxin transport canalization-mediated bud regulation also allows integration of two current models for the mechanism of action of SL in bud control. There is strong evidence that SL regulates shoot branching is by triggering depletion of PIN1 auxin exporters from the plasma membrane of cells in the shoot auxin transport network (Bennett et al., 2006; Crawford et al., 2010; Prusinkiewicz et al., 2009; Shinohara et al., 2013). This reduces the positive feedback in auxin transport canalization, making it more difficult for buds to activate.

In several dicot species, SL also up-regulates the transcription of *BRC1*, providing another mechanism by which SL regulates bud activity (Braun et al., 2012; Dun et al., 2012; 2013; Mashiguchi et al., 2009). However, it is clear that this mechanism is not sufficient to explain SL-mediated bud inhibition. Consistent with results in pea (Braun et al., 2012), our results show that SL signalling mutants are more branchy than brc1-2 brc2-1 mutants, especially in conditions that increase the number of vegetative nodes (Fig 7). Furthermore, this more discriminatory environment provides convincing evidence that triple mutants lacking *BRC1, BRC2* and SL signalling have a strongly additive phenotype (Fig. 7), resolving previously reported conflicting
results in Arabidopsis (Aguilar-Martinez et al., 2007; Chevalier et al., 2014), and extending a similar result in pea that was considered to be surprising (Braun et al., 2012). In contrast to previous reports (Brewer et al., 2009; Braun et al., 2012), using more sensitive assays we demonstrate that \textit{brc1-2} \textit{brc2-1} mutant buds are SL responsive in both whole plants and two-node bud-bud competition assays (Fig. 8). Interestingly, mutation in \textit{brc1} is epistatic to \textit{smxl678} mutation. While this could be interpreted as demonstrating a requirement for \textit{BRC1} in SL-mediated bud inhibition, this is inconsistent with our other results. Rather, this epistasis is easily explained by the bud activation threshold model. According to this model, \textit{brc1} mutation increases branching in a wild-type background by a fixed amount by reducing the bud activation threshold. In the same way \textit{brc1} mutation in the \textit{smxl678 max2} quadruple mutant background, which has wild-type levels of branching, increases branching by a fixed amount, producing \textit{brc1} branching levels.

Although SL can clearly act independently of \textit{BRC1}, the constitutively low levels of \textit{BRC1} expression in SL deficient mutants, and the constitutively high levels in \textit{smxl678}, suggest that \textit{BRC1} is indeed a transcriptional target of SL signalling. This suggests that SL affects both arms of the proposed feed-forward loop in parallel, through effects on both PIN protein trafficking and \textit{BRC1} transcription. Consistent with parallel action, \textit{brc1} mutants have wild-type PIN protein accumulation and auxin transport mutants have wild-type \textit{BRC1} expression (Bennett et al., 2016b). Thus it seems likely that, like low auxin in the main stem, SL regulates bud activity via a coherent feedforward circuit involving effects on both \textit{BRC1} transcription and auxin transport canalization.

Interestingly, in monocots, there is very little evidence that SLs act via transcriptional regulation of \textit{BRC1} family members. For example, in rice \textit{FC1} does not appear to be transcriptionally regulated by SL and SL mutants are highly branched despite wild-type levels of \textit{FC1} expression (Arite et al., 2007; Minakuchi et al., 2010). Similarly, SL-deficient maize mutants have a highly branched phenotype despite constitutively high \textit{TB1} expression (Guan et al., 2012). If this does represent a consistent difference between monocots and dicots it will be interesting to assess what the functional significance and evolutionary origin of the difference might be.
MATERIALS AND METHODS

**Plant material and growth conditions**

Arabidopsis Col-0 wild type plants were used throughout. The following lines used have been previously described: \textit{brc1-2}, \textit{brc1-2 brc2-1} (Aguilar-Martinez et al., 2007); \textit{lexa::BRC1} (Gonzalez-Grandio et al., 2013); \textit{d14-1} (Waters et al., 2012); \textit{max2-1} (Stirnberg et al., 2002); \textit{max4-5} (Bennett et al., 2006); \textit{smxl6-4 smxl7-3}, \textit{smxl6-4 smxl7-3 max2-1}, \textit{smxl6-4 smxl7-3 smxl8-1 max2-1} (Soundappan et al., 2015); \textit{ft-10} (Yoo et al., 2005). Higher order mutant combinations were generated by crossing, using of visible and PCR-based markers for genotyping.

Plants were grown on Levington’s F2 compost in glasshouses with temperature range 15-24°C. Daylight was supplemented when necessary to approximately 150 µmol photons m\(^{-2}\) s\(^{-1}\) to 16 h/8 h light/dark. For short day conditions, plants were grown in Conviron growth chambers with white fluorescent tube lighting at ~150 µmol photons m\(^{-2}\) s\(^{-1}\), 8 h/16 h light/dark, 22/18°C.

Assessment of branching responses to nitrate availability was as described in de Jong et al. (2014). Plants were grown in the glasshouse on a sand/Terra-Green mixture supplemented by \textit{Arabidopsis thaliana} salts (ATS) (Wilson et al., 1990) containing either 9 mM or 1.8 mM Nitrate.

Excised shoot explants were grown in Eppendorf tubes containing 1.95 ml ATS. The explants were transferred to Conviron growth chambers with white fluorescent tube lighting at ~150 µmol photons m\(^{-2}\) s\(^{-1}\), 16 h/8 h light/dark, 22/18°C. The tubes were kept in trays containing wet filter paper and covered with propagator lids to reduce evaporation. For experiments with time-lapse photography, Conviron growth chambers with 24 h light were used.

Plants were grown axenically as described in de Jong et al. (2014). Seven seed were sown per glass jar containing agar-solidified ATS medium supplemented with \textit{rac}-GR24 dissolved in acetone, or with solvent control.

**Bud growth assays**

Auxin dose-response assays were carried out as described in Chatfield et al. (2000). Buds were held between two agar slabs in a Petri dish, allowing apical auxin treatments, with bud growth followed over time.
Non-axenic bud growth and gene expression assays were carried essentially as described in Ongaro et al. (2008). Shoot apices of glasshouse-grown plants bearing 2-3 leaves were excised 2-3 days after bolting and placed in Eppendorf tubes as described above. Explants were allowed to acclimatise for three days, which also allowed elongation of the main stem enabling isolation of a single node by removing more basal nodes. For bud-bud competition assays, two buds were left on the explants. Explants were decapitated and/or were transferred to fresh tubes with appropriate supplements. Bud samples were collected for gene expression analysis, or bud was length measured as appropriate.

Basal hormone or gene induction treatments were carried out by supplementing the ATS solution with the appropriate hormone or inducer. Apical hormone treatments were carried out by applying lanolin paste to the decapitation site. 6-benzylaminopurine (BAP) was dissolved in DMSO and GR24 was dissolved in acetone. 1-naphthalene acetic acid (NAA) was dissolved in ethanol for application in agar and in DMSO for application in lanolin. Where NAA was mixed with lanolin, 5 µl 6X bromophenol blue-based gel loading dye (NEB) was also added to assess mixing.

**Time-lapse photography**
50 µm silica microsphere beads (Corpuscular Inc) were stained with bromophenol blue and stuck at the junction between the main stem and the bud stem, and where the bud stem meets the oldest bud leaf using lanolin. Digital cameras (Nikon) with macro lenses and 2 extender focus tubes (Meike) were used to take images every hour for 24 h. The length of the bud stem (between the two beads) was measured using the manual tracking plugin on Fiji.

**qRT-PCR**
Pools of 15-20 buds were harvested and snap frozen in liquid nitrogen. Total RNA was isolated using the RNeasy Plant Mini Kit (Qiagen) and DNase treated with Turbo DNA-free Kit (Ambion) according to manufacturers’ instructions. RNA was quantified with a NanoDrop 1000 and 1 µg RNA was used for cDNA synthesis using Superscript II (Invitrogen) primed with OligodT according to manufacturer’s instructions. Transcript levels were quantified relative to UBQ10 (UBIQUITIN 10; AT4G05320) using SYBR Green (Bioline) with 10 ng cDNA in a 10 µl reaction volume on a Light Cycler 480 II (Roche). Expression levels were calculated using the ΔΔCt method using Ct values calculated by the 2nd derivative maximum function of the Light Cycler 480 II software.
Statistics
For parametric testing, Student’s t-tests were carried out after log transformation of the data where necessary. For non-parametric tests, Wilcoxon pairwise rank-sum tests were used. For censored data (i.e. where data collection ceased before all individuals had exhibited the phenomenon being measured), Kaplan-Meier survival analysis was used with log-rank tests. Holm-Bonferroni-adjustments were made for multiple comparisons.

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AUTHOR CONTRIBUTION STATEMENT
MS TB and OL developed the concepts and approaches and designed the research. MS performed the experiments and data analysis with input from TB. MS TB and OL interpreted the results and wrote the manuscript.
REFERENCES


Fig. 1. *BRC1* transcripts decrease rapidly after bud activation.

(A,B) Transcript levels of *BRC1* in pooled samples of buds from 15-20 individual one-node explants. Each panel shows the mean of 3-5 independent biological repeats, error bars are s.e.m. (A) 6 h time course of buds with intact primary shoot apex or activated by decapitation; (B) 6 h time course of buds with DMSO control treatment or activated by 0.1 µM basal benzylaminopurine (BAP), a synthetic CK. (C,D) Mean length of the bud stem over time, n=9-12, error bars are s.e.m. (C) Buds activated by decapitation or with an intact primary shoot apex; (D) buds activated with 0.1µM basal BAP treatment or DMSO controls.
Figure 2: BRC1 transcript levels respond to an apical signal.

(A) Experimental setup: manipulation of the length of main stem above the node following decapitation. In treatment a the lowermost buds were collected before decapitation; in b the lowermost buds were collected 1h after removal of intermediate buds; in c buds were collected 1 h after decapitation directly above the lowermost bud; in d lowermost buds were collected 1 h after decapitation and removal of upper buds leaving 8 mm of stem. (B) BRC1 transcript levels 0h and 1h after decapitation with treatments according to (A), results are the mean of 4 independent biological repeats with 15-20 buds per sample, error bars are s.e.m. Asterisks indicate a statistically significant difference between groups with two-sample t-tests and Holm-Bonferroni adjustment at p < 0.1 (one asterisk) or p < 0.05 (two asterisks). (C) Length of the bud stem of buds with intact or decapitated
primary shoot apex with treatments according to (A). ‘Intact’ and ‘Decapitated < 1 mm stem’ are the same data reproduced from Fig. 1C, error bars are s.e.m. n=9-12.
Figure 3: *BRC1* transcript levels respond to apical auxin.

(A) Experimental setup: explants bearing single nodes were decapitated and 1 mM NAA (1-naphthylene acetic acid, a synthetic auxin) in lanolin or the control treatment was placed on decapitated stump at either 0 h or 3 h post decapitation. (B-C) *BRC1* transcript levels in pooled bud samples of 15-20 explants treated according to (A) collected at 6 h post decapitation (B), or 21 h post decapitation (C). Brackets indicate the results of two-sample t-tests as not significant (n.s.), or significant at p < 0.05 (two asterisks). (D) Bud length over time for each treatment, n=14-16. The data are representative of 3 independent biological repeats, error bars are s.e.m.
Figure 4: **brc1-2 brc2-1** buds have altered auxin responses

Branch lengths of isolated one-node explants of (A) wild-type Col-0 and (B) *brc1-2 brc2-1* held between 2 agar slabs with apical NAA at concentrations of 1, 2.5, 5 and 10 µM, or ethanol only. Bud length was measured for 10 days after excision of explants, n=19-20, error bars are s.e.m. Data are representative of 2 independent repeats. (C) Wild-type and (D) *brc1-2 brc2-1* growth curves of the same data aligned to the activation day (day 0) defined as the day on which branches reached 5 mm or greater. Some explants do not feature at some treatments and/or time-points since they never reached 5 mm, or did so late in the experiment, giving sample sizes between 1 and 20. (E) Median day on which buds reach 5 mm or more in length. Crosses indicate a median of at least 10 days, with insufficient buds reaching lengths greater than 5 mm for these treatments. Different letters indicate statistically significant differences at p < 0.05 with log-rank test with Holm-Bonferroni adjustment. Comparisons were carried out between genotypes of the same treatment and between treatments of the same genotype.
Figure 5: BRC1 is not necessary for bud growth inhibition

Branch numbers and vegetative node numbers scored at proliferative arrest. (A) Total number of branches and nodes for plants grown in short days, n=16-23, representative of 2 independent repeats. (B) Number of rosette and cauline branches in wild-type, ft-10, brc1-2 brc2-1 and ft-10 brc1-2 brc2-1 plants, n=39-40, representative of 2 independent repeats. (C) Number of branches, n=21-23, representative of 3 independent repeats, and (D) number of nodes, n=10, on plants grown with high (9 mM) or low (1.8 mM) nitrate supply. (E) Percentage of nodes with axillary buds that were vegetative or reproductive in plants grown with high or low nitrate supply, n=10. Letters indicate statistically significant differences at p < 0.05 with Wilcoxon rank-sum tests with Holm-Bonferroni adjustments where appropriate. Error bars are s.e.m.
Figure 6: High levels of bud *BRC1* transcripts do not prevent branch growth.

(A) *BRC1* transcript levels in the two apical-most buds (n and n-1) of wild-type and *smxl678 max2* mutants. Pooled bud samples were taken from 4 plants per genotype. (B) Branch length of apical n and n-1 branches of WT and *smxl678 max2* mutants 0 and 3 days after decapitation, n=4. Different letters indicate statistically significant differences between groups with pairwise t-tests and Holm-Bonferroni adjustment. (C-D) oestradiol-inducible *lexa::*BRC1* explants treated basally with 10 mM β-oestradiol for 90 mins before decapitation. (C) *BRC1* transcript levels in buds of induced and mock-treated explants. (D) Branch length of induced and mock-treated explants after decapitation over time, n=10-15 For (A) and (C) data are the mean of 3 biological replicates. Brackets with two asterisks indicate a statistically significant difference at p < 0.05 with a two-sample t-test. For (B) and (D) data are representative of 2 and 3 independent repeats respectively. Error bars are s.e.m.
Figure 7: Strigolactone and BRC1 affect branching additively

Total branch number at proliferative arrest. (A) Wild-type, max2-1, brc1-2, smxl6-4 smxl7-3 smxl8-1, smxl6-4 smxl7-3 smxl8-1 max2-1, and smxl6-4 smxl7-3 smxl8-1 max2-1 brc1-2 plants grown in long day growth chambers, n=9-13 (B) Wild-type, brc1-2 brc2-1, max4-5, d14-1, max4-5 brc1-2 brc2-1 and d14-1 plants grown in long day glasshouse conditions, n=23-24, representative of 3 independent repeats. (C) Wild-type, brc1-2 brc2-1, d14-1, max2-1, and max4-5 plants grown in short day growth conditions, n=16-23, data for Col-0 and brc1-2 brc2-1 is the same as for Fig 5A. (D) Photographs of representative plants of the genotypes indicated grown in long days. Different letters indicate statistically significant differences at p < 0.05 with pairwise Wilcoxon rank-sum tests with Holm-Bonferroni adjustments, error bars are s.e.m.
Figure 8: *brc1 brc2* mutants are strigolactone-sensitive.

(A–C) Bud-bud competition assay in which explants bearing two nodes are basally treated with 5 µM GR24 (a synthetic strigolactone) or acetone control and decapitated, n=19–20, error bars are s.e.m. Length of the longer branch (A) or the shorter branch (B) over time after decapitation. (C) Violin plots of the relative growth index (the length of the longest branch divided by the total length of both branches) at day 10 after decapitation, white circles indicate the median and coloured shapes indicate the probability distribution of the data. (D) Total branch number of 6-week-old plants grown axenically on media supplemented with 5 µM GR24 or acetone control. For (C–D) asterisks indicate statistically significant differences between treatments for each genotype at \( p < 0.05 \) with pairwise Wilcoxon rank-sum tests. Data for all panels are representative of 3 independent repeats.
**BRC1** expression reduces bud activation potential, but is not necessary or sufficient for bud inhibition in Arabidopsis

**Supplementary Figures**

**Figure S1.** **BRC1** transcripts do not change 30 minutes post-decapitation

Transcript levels of **BRC1** in pooled samples of buds from 15-20 individual one-node explants, 1 h time course of buds with intact primary shoot apex, or activated by decapitation. Data are the mean of 3-4 independent biological repeats, error bars are s.e.m.

**Figure S2.** Strigolactone and **BRC1** affect plant height additively

Height of the primary inflorescence shoot at proliferative arrest. Different letters indicate statistically significant differences at p < 0.05 with pairwise Wilcoxon rank-sum tests with Holm-Bonferroni adjustments, n=15-20, error bars are s.e.m.