RESEARCH ARTICLE

Strong morphological defects in conditional Arabidopsis abp1 knock-down mutants generated in absence of functional ABP1 protein [version 1; referees: 1 approved, 2 approved with reservations]

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Abstract
The Auxin Binding Protein 1 (ABP1) is one of the most studied proteins in plants. Since decades ago, it has been the prime receptor candidate for the plant hormone auxin with a plethora of described functions in auxin signaling and development. The developmental importance of ABP1 has recently been questioned by identification of Arabidopsis thaliana abp1 knock-out alleles that show no obvious phenotypes under normal growth conditions. In this study, we examined the contradiction between the normal growth and development of the abp1 knock-outs and the strong morphological defects observed in three different ethanol-inducible abp1 knock-down mutants (abp1-AS, SS12K, SS12S). By analyzing segregating populations of abp1 knock-out vs. abp1 knock-down crosses we show that the strong morphological defects that were believed to be the result of conditional down-regulation of ABP1 can be reproduced also in the absence of the functional ABP1 protein. This data suggests that the phenotypes in abp1 knock-down lines are due to the off-target effects and asks for further reflections on the biological function of ABP1 or alternative explanations for the missing phenotypic defects in the abp1 loss-of-function alleles.
Introduction
The naturally occurring auxin, indole-3-acetic acid, plays a central role in plant growth and development alone or in orchestration with other plant hormones. Proper sensing and interpretation of fluctuating cellular auxin signals is necessary for mediating a diverse range of developmental and cell biology responses (Enders & Strader, 2015; Grunewald & Friml, 2010; Paciorek et al., 2005; Petrasek et al., 2006). In the early screens for auxin receptors, Auxin Binding Protein 1 (ABP1) has been identified based on its ability to bind auxin with high affinity (Hertel et al., 1972; Löbler & Klämbt, 1985) and soon became a prime candidate for an extracellular auxin receptor based mainly on electrophysiological studies utilizing antibodies against ABP1 that showed rapid ABP1-mediated modulation of plasma membrane ion transport in an early step of auxin action (Barbier-Brygoo et al., 1989; Leblanc et al., 1999). Over the next decades, the auxin-binding activity of ABP1 has been characterized in detail by biochemical studies (Batt et al., 1976; Napier et al., 2002; Napier & Venis, 1995; Ray et al., 1977) and its protein structure including the auxin-binding pocket has been revealed (Woo et al., 2002). Phylogenetic studies have shown that ABP1 homologues are present in the genomes of all plant species from bryophytes to flowering plants (Tromas et al., 2010) with more than one copy present e.g. in the genome of maize, rice, poplar or the moss Physcomitrella patens (http://phytozome.jgi.doe.gov/pz/portal.html).

Since its discovery, however, the biological importance of the ABP1 protein as a plasma membrane auxin receptor has been a matter of debate, in part because of its predominant subcellular localization in the endoplasmatic reticulum (ER) in maize where the conditions for auxin binding are unfavorable (Habets & Offringa, 2015; Napier et al., 2002). Recently, these discussions were revived by the isolation of two new Arabidopsis abp1 knock-out alleles, abp1-c1 and abp1-TD1 (Gao et al., 2015) that show no obvious phenotypes under standard growth conditions. The contradiction between this observation and the previously published embryo-lethal phenotypes of abp1 mutants (Chen et al., 2001; Tzafir et al., 2004) has recently been clarified by proving that the embryo-lethality of the originally reported alleles abp1-1 and abp1-1s was caused by disruption of the tightly-linked neighboring gene BELAYA SMERT (BSM) rather than knock-out of ABP1 (Dai et al., 2015; Michalko et al., 2015). This correction and the demonstration of normal embryo development in the abp1 knock-outs (Michalko et al., 2015) suggest that ABP1 plays no essential role in early Arabidopsis embryogenesis.

The ongoing discussion focuses on the relevance of ABP1 in auxin signaling and other post-embryonic auxin-related biological processes that have been demonstrated using different genetic tools, namely the conditional knock-down (KD) lines, the abp1-5 weak allele harboring a point mutation in the ABP1 auxin-binding pocket and gain-of-function alleles, all of which often provided internally consistent results (Braun et al., 2008; Covanova et al., 2013; David et al., 2007; Grones et al., 2015; Robert et al., 2010; Sassi et al., 2014; Tommas et al., 2013; Xu et al., 2010; Xu et al., 2014).

Conditional ABP1 KD lines SS12S6, SS12K9 and abp1-AS have been generated using two fundamentally different approaches of gene or protein down-regulation. In the SS12S6 and SS12K9 lines, ABP1 was inactivated by inducible over-expression of a recombinant immunoglobulin fragment termed single-chain fragment variable (scFv) (Conrad & Fiedler, 1998). This construct, consisting of the heavy- and light-chain variable domains of a well-characterized anti-ABP1 monoclonal antibody mAb12 (David & Perrot-Rechenmann, 2001; David et al., 2007; Leblanc et al., 1999) linked by a flexible peptide was additionally fused to the sequence encoding the 3’KDEL motif to mediate scFv ER-retention in the SS12K9 line, while the SS12S-encoded scFv12 was meant to be secreted to the apoplast. In planta-produced scFv12 was able to pull down ABP1, and reciprocally immunoprecipitation of ABP1 using another antibody was shown to pull down scFv12 (Trommas et al., 2009). An antisense approach was utilized in the abp1-AS line, where inducible over-expression of full-length ABP1 antisense cDNA led to the formation of duplexes with its sense mRNA, thus preventing ABP1 translation, and potentially also transcription by RNA interference mechanism (Meister & Tuschl, 2004; Tufarelli et al., 2003). Both antibody- and antisense-based lines use the ethanol-inducible system, which is well established and widely used for the conditional expression of plant genes (Deveaux et al., 2003; Roslan et al., 2001).

These three abp1 knock-down lines have been instrumental to connect ABP1 function to multiple cellular and developmental processes. For example, they showed defects in shoot and root growth (Braun et al., 2008; Tommas et al., 2009), cell wall re-modeling (Paque et al., 2014) or clathrin-mediated endocytosis of PIN auxin efflux carriers (Dhomoukhe et al., 2007; Robert et al., 2010). In contrast, the abf1 gain-of-function transformants promote PIN internalization both in tobacco and Arabidopsis (Grones et al., 2015; Robert et al., 2010). Contrasting effects of ABP1 KD and gain-of-function lines were shown also in the case of auxin effect on the control of leaf epidermal pavement cells morphogenesis (Braun et al., 2008; Nagawa et al., 2012) on ROP GTPase activation (Xu et al., 2010) and on microtubule rearrangement (Chen et al., 2014; Xu et al., 2014). Furthermore, analysis of ABP1 variants with mutations in the auxin-binding pocket demonstrated the importance of axuin-binding to ABP1 for its gain-of-function phenotypes (Grones et al., 2015). Altogether, these studies provided an internally consistent picture showing involvement of ABP1 signaling in multiple physiological and cellular processes. These observations were further supported by the finding that loss-of-function mutants in TMK receptor-like protein kinases, that were recently shown to interact with ABP1 in an auxin-inducible manner, show similar phenotypes with abp1 KD mutants (Xu et al., 2014) which was consistent with the importance of the ABP1/TMK complex-mediated auxin perception in plant development. Recent identification of wild-type looking Arabidopsis abp1 loss-of-function alleles by Gao et al. (2015) thus questions the interpretation of data obtained in the aforementioned studies.

Here, we address the missing phenotypes in the true abp1 null alleles in relation to the strong and consistent morphological defects observed in the conditional abp1 knock-down lines. We show that the morphological phenotypes in SS12S6, SS12K9 and abp1-AS can be generated in the absence of functional ABP1 protein and we discuss possible underlying causes of this.
Material and methods

Plant material and growth conditions

Arabidopsis thaliana mutants used in this study were: abp1-c1, abp1-TD1 (Gao et al., 2015), abp1-AS, SS12K9, SS12S6 (Braun et al., 2008; David et al., 2007). A. thaliana Col-0 wild type seeds were obtained from The Nottingham Arabidopsis Stock Centre (NASC, http://www.arabidopsis.info). For in vitro experiments, seeds were surface-sterilized with chlorine vapor, vernalized for 2 days in the dark at 4°C and grown on 1/2 MS 0.8% agar medium with or without 1% w/v sucrose (pH 5.9) on vertical Petri dishes under long day conditions (16 h light/8 h dark) or in complete darkness at 21°C. A sterilized microtube with 500 µl 5% ethanol was placed at the bottom of the plate to induce expression of abp1-AS, SS12K9 and SS12S6 constructs before germination. Plates with 5-day old etiolated or 7-day old light-grown seedlings were scanned on a flatbed scanner, phenotyped by visual examination and used for DNA/RNA extraction.

Genotyping mutants

Ethanol-inducible ABP1 down-regulating lines (abp1-AS, SS12K9, SS12S6) were genotyped for the presence of the alcR gene encoding the transcriptional regulator of the ethanol-inducible system using primers alcR_for and alcR_rev (Table 1). Fragments amplified from abp1-c1 with primer pairs ABP1-U409F + ABP1-586R or ABP1-5P + ABP1-586R were digested with BstI, which cuts the WT fragment once and does not cut the mutant fragment; abp1-TD1 was genotyped as described previously (Gao et al., 2015). Genomic DNA was isolated using the CTAB extraction method. GoTaq G2 polymerase (Promega) and Bio-Rad T100 Thermal Cycler were used for PCR under following conditions: initial denaturation 5 min 98°C, 35–45 cycles (denaturation 30 s at 98°C; annealing 30 s at 55°C, elongation 1 min at 72°C); final elongation 5 min at 72°C. Restriction analysis was performed by adding the restriction enzyme directly to unpurified PCR reaction. Alternatively, Phire Plant Direct PCR Kit (Thermo Scientific by Finnzymes) and QIAquick Gel Extraction Kit (QIAGEN) were used following manufacturer’s instructions to genotype the SS12K9 × abp1-c1 line.

Quantitative RT-PCR

Total RNA from approximately twenty 8-day old seedlings frozen in liquid nitrogen was extracted using the TRIzol reagent (Invitrogen, Carlsbad, CA, USA) and purified using RNeasy Mini Kit (Qiagen) according to manufacturer’s instructions. 2 µg of purified total RNA were used for a reverse transcription reaction using the iScript cDNA Synthesis Kit (BioRad). qRT-PCR was performed using the LightCycler 480 SYBR Green I Master chemistry (Roche) in a LightCycler480 II thermal cycler (Ser. no. 5659, Roche) according to manufacturer’s instructions. cDNA diluted 1:10 in water was used as a template to prepare 5 µL reaction mixture (final volume). Primers used for the quantitative RT-PCR were designed using QuantPrime (http://www.quantprime.de). The ABP1 cDNA fragment (84 bp in length) was amplified with ABP1-1E and ABP1-586R primers. Arabidopsis Tubulin beta chain 2 (TUB2, At5g62690) amplified with TUB2-F and TUB2-R primers was used as a reference gene (Dataset 1). Gene expression was calculated with the 2^(-ΔΔCT) method (Livak & Schmittgen, 2001). Results are expressed as the average +/- standard deviation of 2 biological and three technical replicates. Sequences of primers used for genotyping and qRT-PCR analysis are listed in Table 1.

| Table 1. Primer sequences used in this study. |
|---------------------------------------------|
| ABP1-U409F: CCTCATCACACAACAAAGTCACTC |
| ABP1-586R: GGAGCCAGCAACAGTGATGTG |
| ABP1-5P: ATGATCGTACTTTCTGTTGGTTCC |
| ABP1-2E: TGGCCAACTGTAAGGAATATTAG |
| pSKTAIL-L3: ATACGCGAGCTGAATTGGTCG |
| AlcR F: AGAACAAGAAGAAGCCAGGA |
| AlcR R: GCGTGAGAGAAAGATGA |
| TUB2 F: TAAACACTGGCCAAAGGGGACAC |
| TUB2 R: ACAACCTGGAAACCTGGAGAC |

Results

Dataset 1. Scans of ethanol-induced F2 seedlings of crosses (A) SS12S6 × abp1-c1, (B) SS12S6 × abp1-TD1, (C) abp1-AS × abp1-c1, (D) abp1-AS × abp1-TD1, (E) SS12K9 × abp1-c1 and (F) SS12K9 × abp1-TD1 that were used for phenotyping and genotyping (Figure 1 and Figure 2).

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Dataset 2. Agarose gel images from the PCR genotyping of the F2 crosses (A) SS12S6 × abp1-c1, (B) SS12S6 × abp1-TD1, (C) abp1-AS × abp1-c1, (D) abp1-AS × abp1-TD1, (E) SS12K9 × abp1-c1 and (F) SS12K9 × abp1-TD1 (Figure 3).

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All crosses were genotyped for the presence of the alcR transcriptional regulator (first row of the gel images) which is an integral part of the ethanol-inducible cassette in abp1 knockdown lines. The presence of point mutation in abp1-c1 crosses was genotyped by restriction analysis of ABP1 PCR product as described in Gao et al. (2015) (second row of the gel images of abp1-c1 crosses). The presence of the T-DNA insertion in abp1-TD1 crosses was genotyped according to Gao et al. (2015) (second and third row of gel images of abp1-TD1 crosses). GeneRuler DNA ladder mix #0331 (Thermo Scientific) was used as a fragment size standard to determine the approximate size of DNA fragments. Fragment sizes of 1000 bp and 500 bp are indicated.

Dataset 3. Source qPCR data (Figure 3c).

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Individual samples are annotated with their position on a 384-well plate (column A), the cDNA (column B) and primer pair (column C); the Cp value of each sample is shown in column D. The experiment was performed in two biological (1 or 2 at the last position in column B) and three technical replicates. Figure 3c shows gene expression calculated with the 2ΔΔCt method (Livak & Schmittgen, 2001) from values of ABP1-2E and TUB as a reference gene; using ABP1-5P and/or EF as a reference gene instead gave similar results.
Figure 1. Strong morphological defects in conditional abp1 knock-down lines correlate with the presence of the ethanol-inducible cassette and segregate normally when crossed with abp1-c1 knock-out allele. (A) abp1-AS × abp1-c1 F2 plants grown for 7 days in the presence of 5% ethanol segregate strong morphological defects characteristic of the abp1 conditional knock-down (KD) alleles approximately in a 3:1 ratio. (B) alcR-specific PCR bands amplified from the genomic DNA of abp1-AS × abp1-c1 F2 plants shown in (A) demonstrate that the KD phenotype is caused by the presence of the ethanol-inducible insertion. (C) Phenotypes of the scFv12-based KD lines segregate similarly in F2 crosses with abp1-c1, while altered segregation ratios can be observed in F2 of all three KD alleles crossed to abp1-TD1, which is most apparent in seedlings grown for 5 days in the dark (grey background).

Figure 2. Mendelian segregation of strong ethanol-inducible phenotypes in the F2 generation of abp1 knock-out x knock-down crosses is independent of abp1 mutant background. (A) Representative abp1-AS × abp1-c1 F2 plants, (B) PCR products amplified from their genomic DNA and (C) segregation ratios from all crosses show that the ethanol-inducible phenotypes segregate independently of the presence of abp1 knock-out alleles following approximately Mendelian rules for di-hybrid crosses. Homozygous abp1 knock-out mutants with the inducible KD phenotype could be found in all crosses (plants 2,5,8 in (A) and (B), red numbers in (C)), suggesting that the phenotype does not require a functional ABP1 gene. Strong deviations from the expected Mendelian segregation were detected in the SS12K9 x abp1-c1 cross, indicating genetic linkage between ABP1 locus and the inserted ethanol-inducible scFv construct.
Segregation of strong morphological defects in conditional abp1 knock-down alleles crossed with abp1-TD1 and abp1-c1 knock-out alleles

To investigate the contradiction between missing phenotypic defects in the loss-of-function abp1 alleles and strong morphological defects of conditional ABP1 down-regulating lines (knock-down; KD), we decided to cross both types of lines to test three possible scenarios: 1) The absence of the strong morphological defects in the abp1-c1 or abp1-TD1 alleles is caused by an adaptation of the plants to the permanent loss of the ABP1 function, which compensates for this deletion; 2) the strong morphological phenotypes induced in the KD lines do not require functional ABP1 and are caused by off-target effects; or 3) both abp1-TD-1 and abp1-c1 lines contain background mutation(s) that suppress the phenotypes caused by the absence of ABP1.

We crossed each of the conditional lines with abp1-TD1 and abp1-c1 null mutants and with an ABP1-WTc1 line as a control and analyzed seedling phenotypes of ethanol induced F2 segregating plants (Figure 1a). We hypothesized that in case of an adaptive process, the conditional abp1 KD phenotypes (short wavy roots and epinastic cotyledons) would not be manifested in homozygous abp1 null background, resulting in a 9/16 KD and 7/16 WT phenotype segregation ratio. If the inducible phenotypes in the KD lines are caused by off-target effects; or 3) both abp1-TD-1 and abp1-c1 lines contain background mutation(s) that suppress the phenotypes caused by the absence of ABP1.

Segregation of the morphological phenotypes in the F2 plants from different crosses is summarized in Figure 1b. These observations show that strong phenotypes in both the abp1 antisense-based and the scFv12-based conditional knock-down lines segregate approximately 75% in the F2 crosses with abp1-c1. This observation favors the scenario that the strong morphological defects in the KD lines are not influenced by the presence or absence of the functional ABP1 gene copy. The F2 phenotypic segregation is however shifted in favor of WT-looking plants in all three KD lines crossed to abp1-TD1. This segregation shift may be ascribed to partial transcriptional silencing of the ethanol-inducible constructs due to the presence of multiple 35S promoters/enhancers in the constructs themselves as well as the tandem T-DNA insertion in abp1-TD1.

We genotyped all analyzed F2 plants for the presence of the alcR transcriptional regulator, which is an integral part of the ethanol-inducible system and verified that the observed morphological defects were indeed correlating with the presence of the ABP1 KD constructs (Figure 1c). About 5% of seedlings from all lines showed WT phenotype despite being positive for the presence of alcR or vice versa. As this phenomenon was independent of ABP1 genetic background and could not be reproduced in F3 progeny (Supplementary Figure 2), we put it down to biological variability and/or occasional silencing of the ethanol-inducible constructs.

Strong morphological defects in conditional abp1 knock-down alleles can be manifested in homozygous abp1 knock-out alleles

To investigate whether the abp1 KD phenotypes can be observed in the absence of a functional copy of the At4g02980 ABP1 gene we further genotyped the respective abp1 mutations in F2 seedlings of all crosses (Figure 2). As summarized in Figure 2c, in all crosses

Figure 3. Strong inducible knock-down phenotypes in the absence of functional ABP1 gene confirmed in the SS12K9 × abp1-c1 F3 progeny. (A) Representative seedlings of the ethanol-induced F3 progeny of one of the SS12K9 × abp1-c1 F2 plants (plant A) that showed KD phenotype in the absence of the functional ABP1. All F3 seedlings manifesting KD phenotype were homozygous for abp1-c1 mutation. (B) Genotyping of the plants shown in A. The image is assembled from different regions of two gels that were copy-pasted next to each other in order to save space. (C) qRT-PCR analysis of KD-phenotype positive F3 seedlings of both lines revealed that ABP1 transcript levels are reduced by about 80% like in the original abp1-c1 mutant. Altogether these data confirm that in the abp1 down-regulating lines the KD phenotype can be manifested without the ABP1 function. In (C) average of two biological and three technical replicates +/- SD is shown. PC- positive control.
we were able to identify multiple homozygous *abp1* mutants that showed the strong KD phenotype following ethanol induction. This analysis demonstrates that strong morphological phenotypes in *abp1* antisense-based (*abp1-AS*) and scFv12 antibody-based (SS12S6, SS12K9) conditional KD lines can be generated also in a null *abp1* background.

In case of the crosses SS12K9 × *abp1-c1* and SS12K9 × *abp1-TD1* we observed a lower level of allelic segregation between the *abp1* mutations and the KD construct in their F2 progeny (Figure 2c). Out of 28 genotyped plants with WT phenotype, 24 (85.7%) were homozygous for *abp1* mutation and did not contain the ethanol-inducible KD cassette. These results point towards genetic linkage between these two loci, most likely caused by the positional effect of the KD cassette located close to the *ABP1* locus on the chromosome 4. Nevertheless, some level of genetic recombination was happening between the two loci in the crosses as demonstrated by the identification of three F2 SS12K9 × *abp1-c1* plants showing KD phenotype that were homozygous for *abp1-c1* mutation (Figure 2c). This analysis confirms that also SS12K9 conditional KD construct can generate strong morphological phenotypes in the homozygous *abp1-c1* knock-out alleles despite the insertion position being linked to the *ABP1* locus. Altogether these data are consistent with results obtained by the other crosses and further support that morphological phenotypes in the *abp1* knock-down lines can be generated in the absence of the functional ABP1.

**Analysis of F3 generation confirms SS12K9-induced strong morphological defects in absence of ABP1 function**

Next we tested the occurrence of the strong KD-induced morphological phenotypes in the absence of the functional ABP1 in the next generation by analyzing the F3 progeny following the SS12K9 × *abp1-c1* plants showing strong KD phenotype. We confirmed that the F3 progeny was homozygous for the *abp1-c1* mutation and segregated the ethanol-inducible construct approximately in a 3:1 ratio (Figure 3b). After induction with ethanol, the analyzed F3 population of SS12K9 × *abp1-c1* plant A segregated into 27 plants (67.5%) with KD phenotype and 13 WT looking plants (32.5%) (Figure 3). The F3 population of plant B segregated into 18 plants with KD phenotype (81.2%) and 4 WT looking plants (18.2%) (data not shown). Genotyping of all F3 plants with ethanol-inducible phenotypes revealed that they contain KD construct in the homozygous *abp1-c1* background (Figure 3b). Notably, among the 17 analyzed WT looking F3 seedlings we also identified two plants that contain the ethanol-inducible construct in homozygous *abp1-c1* background (Figure 3b) suggesting that in these plants the functionality of the construct was affected, most probably by its silencing. Nonetheless, the majority of the plants containing the ethanol-inducible construct generated the strong morphological phenotypes even in the *abp1-c1* homozygous background.

We also analyzed the ABP1 expression in WT, *abp1-c1* and SS12K9 × *abp1-c1* F3 seedlings by quantitative RT-PCR just to verify that introducing KD alleles does not influence, in any way, the *ABP1* expression (Figure 3c). We observed ca. 80% decrease in *ABP1* transcript levels in *abp1-c1*. We assume that this difference - somewhat surprising, since the CRISPR-induced small deletion does not necessarily decrease transcript levels - is probably caused by the decreased stability of the mutant mRNA. SS12K9 × *abp1-c1* F3 plants positive for the KD phenotype and homozygous for *abp1-c1* showed the same 80% decrease in *ABP1* transcription.

In summary, the phenotypic, genotypic and expression analyses consistently showed that all three conditional *abp1* knock-down alleles can generate strong morphological defects also in the absence of the functional ABP1 protein.

**Discussion**

**Strong morphological phenotypes in *abp1* conditional knock-down alleles are not caused by ABP1 down-regulation**

All three available conditional *abp1* knock-down alleles have been extensively characterized and used to link number of developmental and cellular processes to the ABP1-mediated signaling (for overview, see Grones & Friml, 2015). They are based on two unrelated strategies for down-regulation of the protein’s functionality: the antisense (*abp1-AS*) and the scFv12 monoclonal antibody expression (SS12S6, SS12K9), which suppress the protein functionality by entirely different mechanisms and at different levels (Tomaras et al., 2009). All three lines showed consistent and reproducible results in a number of different laboratories and a number of developmental, physiological and cellular processes.

Nonetheless, our analysis, made possible by the newly available *abp1* knock-out lines (Gao et al., 2015), strongly suggests that these observed and described effects are not caused by conditional down-regulation of the ABP1. This is supported by the fact that all three constructs show the same strong conditional phenotypes in two different homozygous *abp1* null alleles. This means that even in the absence of the functional ABP1 protein, the ethanol-inducible constructs are inducing phenotypic defects that were originally ascribed to the down-regulation of ABP1. Therefore, results generated using these lines need to be critically re-interpreted.

**Possible modes of action of *abp1* conditional knock-down lines**

All three types of *abp1* KD Arabidopsis lines generate indistinguishable morphological phenotypes. How it is possible that independent lines using fundamentally different approaches for functional down-regulation of a unique target would have in fact the same off-target effects; we do not know. One possible explanation is that the morphological defects are an artifact of the ethanol-inducible expression system. However, control lines generated in parallel using the same vector and expressing the UIDA reporter gene did not exhibit any significant growth and developmental alterations (Braun et al., 2008). Furthermore, a number of authors have used the same ethanol-inducible system to control the expression of distinct genes and to the best of our knowledge, there are no reports describing similar phenotypes by using the ethanol-inducible system for other genes in other studies (Battaglia et al., 2006; Deveaux et al., 2003; Laufs et al., 2003; Peaucelle et al., 2008; Roslan et al., 2001). This system was also used to successfully rescue mutant defects after ethanol induction of gene expression e.g. for LEAFY (Maizel & Weigel, 2004) or for N-myristoyltransferase (Pierre et al., 2007) indicating that it is not responsible per se of the phenotypes observed with the ethanol inducible ABP1 AS
and scFv12 constructs. In tobacco plants and BY-2 cells, tetracycline de-repressible promoter-driven expression of the SS12S and SS12K constructs resulted in similar growth defects as their ethanol-inducible expression in Arabidopsis (Braun et al. 2008; David et al., 2007), suggesting that the observed phenotypes are tightly correlated to the scFv12 action. The expression of the scFv12 in the cytosol had however no effect on cell proliferation in BY2 cells indicating that expression of scFv12 per se is not sufficient to generate severe phenotypes whatever its cellular localisation and that scFv12 effects are correlated to its secretion and/or retention in the ER that are known location of ABP1 (David et al., 2007).

Another possibility is that both the antisense- and antibody-based lines have off-target(s) either on the very same gene(s) or elements of a common genetic pathway. Such a hypothesis would be supported by strict similarities in the phenotypes resulting from ABP1 antisense and scFv12 expression and by the fact that opposite and auxin-related defects were observed in both constitutive and conditional gain-of-function Arabidopsis transgenic plants as well as transitionally expressing tobacco cells (Grones et al., 2015; Robert et al., 2010). ABP1 is placed within the superfamily of cupins based on the presence of cupin-like motifs HXH(X)_nG and P(X)_mH(X)_nN (where X is any amino-acid residue) and a β-barrel jellyroll fold subunit structure (Dunwell et al., 2004; Woo et al., 2002). The epitope recognized by the scFv12 might be present in proteins belonging to this functionally highly diverse protein superfamily. On the other hand, the sequence similarity of even the closest ABP1 homologues in Arabidopsis does not seem to be sufficiently high to be targeted by the abp1-AS constructs, thus this explanation is unlikely as well.

We also cannot completely rule out that the WT phenotype of the abp1 knock-out mutants is caused by suppressor mutation(s). However, we do not consider it very likely, as this would imply that the similar mutation(s) or mutations with similar effects are present in the genetic background of both abp1-c1 and abp1-TDI, which are independent alleles from independent mutant collections.

In summary, we do not understand how it is possible that the used abp1 knock-down alleles generate the similar strong morphological phenotypes also in absence of the functional ABP1 protein. All possible explanations we could come up with are unlikely, including common off-targets in abp1 antisense and antibody KD lines or common suppressor mutations in two different abp1 knock-out alleles. Thus, more experimentation is needed to figure out what really happens in the different abp1 KD lines and how it is possible that they independently generate phenotypes that are so consistent.

Whatever the explanation at the end will be, in light of the presented data it seems obvious that these lines do not act solely by down-regulating the ABP1 function, despite the accumulation of well-fitting data from independent and complementary approaches. It is a sobering realization that even when you use independent approaches with all standard controls performed, there is no real guarantee that the observations will not lead you amiss.

Data availability
F1000Research: Dataset 1. Scans of ethanol-induced F2 seedlings of crosses (A) SS12S6 × abp1-c1, (B) SS12S6 × abp1-TDI, (C) abp1-AS × abp1-c1, (D) abp1-AS × abp1-TDI, (E) SS12K9 × abp1-c1 and (F) SS12K9 × abp1-TDI that were used for phenotyping and genotyping (Figure 1 and Figure 2), 10.5256/f1000research.7654.d110722 (Michalko et al., 2016a).

F1000Research: Dataset 2. Agarose gel images from the PCR genotyping of the F2 crosses (A) SS12S6 × abp1-c1, (B) SS12S6 × abp1-TDI, (C) abp1-AS × abp1-c1, (D) abp1-AS × abp1-TDI, (E) SS12K9 × abp1-c1 and (F) SS12K9 × abp1-TDI (Figure 3), 10.5256/f1000research.7654.d110723 (Michalko et al., 2016b).

F1000Research: Dataset 3. Source qPCR data (Figure 3c), 10.5256/f1000research.7654.d110724 (Michalko et al., 2016c).

Author contributions
JF, JM, CP and MG designed the experiments and wrote the manuscript, JM and MG performed most experiments and analyzed the data. All authors have seen and agreed to the final content of the manuscript.

Competing interests
No competing interests were disclosed.

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Supplementary material

**Supplementary Figure 1.** Theoretical genotype and phenotype segregation in F2 progeny of the abp1 knock-out × knock-down cross. Expected genotype and phenotype segregation ratios for three possible scenarios are shown. S/s = ethanol-inducible cassette positive/negative, A/a = wild-type ABP1/abp1 knock-out. Genotypes manifesting wild-type phenotype are shown on white background, KD phenotype on green background, genotypes that might exhibit both WT and KD phenotypes are on pale green background.

| 1st scenario (adaptation) | 2nd scenario (off-targets) | 3rd scenario (suppressor mutations) |
|---------------------------|---------------------------|-----------------------------------|
| SSAA SSAA SSAA SSAA      | SSAA SSAA SSAA SSAA      | SSAA SSAA SSAA SSAA              |
| SSAA SSAA SSAA SSaa      | SSAA SSAA SSaa ssAA      | SSAA SSaa ssAA ssAA              |
| SSAA SSAA ssAA ssAA      | SSAA ssAA ssAA ssaa      | SSAA ssaa ssaa ssaa              |
| ssAA sscc ssaa ssaa      | ssAA sscc ssaa ssaa      | ssAA sscc ssaa ssaa              |

**Supplementary Figure 2.** Restoration of the ethanol-inducible phenotype in the progeny of the F2 WT-looking plant from the cross SS12K9 × abp1-c1 containing knock-down cassette and wild-type ABP1 version. In the presence of 5% ethanol, F3 progeny show homogenous KD phenotype indicating that silencing of the construct might be responsible for wild-type phenotype of this plant in the F2 generation.

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Version 1

Referee Report 09 February 2016
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In their m/s, the authors present an analysis of genetic interactions between published ABP1 knockdown lines (based on antisense and immune-suppression approaches) and the Arabidopsis ABP1 locus. After crossing three different knockdown lines into recently described abp1 knockout alleles, followed by segregation analysis of resulting F2 and F3 progeny, the authors came to the conclusion that growth defects of their knockdown lines do not depend on a functional ABP1 locus.

These findings represent a valuable contribution to ongoing attempts, trying to clarify ABP1 issues. Nevertheless, off-target loci recognized in the knockdown lines remain mysterious. Are there any ABP1-related loci (cupins?), expression of which could be affected by these knockdown lines? Perhaps this could be tested in the abp1-AS line.

Phenotypes of the knockdown lines are reminiscent of mutants with altered auxin responses. Did the authors look into expression of some of the characterized auxin-related loci? This could give us a better idea about the genetic determinants, causing the phenotypes in these knockdown lines.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Competing Interests: No competing interests were disclosed.

Referee Report 05 February 2016
doi:10.5256/f1000research.8243.r12004

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The authors have produced a very thorough analysis of a set of well-known conditional ABP1 knock-down Arabidopsis lines. Using crosses with the recently-described abp1 knock-out lines from the work of Gao et al. (2015), the manuscript reveals that the phenotypes previously associated with loss of ABP1 can be induced even in genetic backgrounds that lack ABP1. These findings cover results from the two
independent strategies used to create the conditional knock-down phenotypes, conditional immunosuppression and conditional antisense expression.

The assay used for this work was Arabidopsis root growth, not the very detailed phenotypes described recently from the conditional lines such as PIN endocytosis, ROP GTPase activation or microtubule rearrangement. However, impaired root growth was part of the initial suite of phenotypes associated with induced knock-down. The assays show very clearly that growth impairment correlates with the inheritance of the inducible cassette even in the absence of a functional ABP1 gene. We can surmise that the detailed subcellular phenotypes listed above are associated with the gross morphological changes recorded here. There is clearly some intriguing physiology associated with the switch induced by ethanol treatment, but ABP1 is not part of that story.

The title and abstract are appropriate (but see note below*), the work is done well, the data is presented clearly and fully, the text is very well structured and is easy to follow. There is a thorough introduction which explains how and why experimentation developed to tackle tangible problems linked to existing knowledge and understanding of ABP1 genetic resources. It is shown that the science was not misled by carelessness or device, that a spectrum of consistencies coloured the hypotheses covering ABP1 activity, and that these hypotheses have now been proven incorrect thanks to improved technologies, vigilance and critical reappraisal.

“The abstract ends with a sentence “…asks for further reflections on the biological function of ABP1 or alternative explanations for the missing phenotypic defects in the abp1 loss-of-function alleles.” I think that this would be better if it read “… AND alternative explanations for the intriguing phenotypes previously associated with loss of ABP1 activity”. Or similar. I understand that there is a requirement on all sides of the debate to be objective and critical about their data and no harm is done by registering this. However, the balance of probabilities needs to be recognised and, in my view, much more has to be gained from encouraging discovery of the root cause of the many fascinating phenotypes thrown up by the ABP1 KD lines than by searching absent defects. Therefore a small rewording at the end of the abstract is recommended. Otherwise I have no changes to suggest.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Competing Interests: No competing interests were disclosed.

Referee Report 26 January 2016

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In this manuscript, Michalko et al. investigates the basis of the developmental phenotypes observed in knock-down (KD) lines of the Arabidopsis ABP1 gene. By analysing segregation of defects in root elongation in the F2 and F3 generation of crosses between three KD abp1 alleles and two recently reported abp1 knock-out (KO) alleles, they conclude that the previously reported phenotypes of the KD lines are not due to loss of ABP1 function. The manuscript is clearly written and the experiments are thoroughly carried out providing an important contribution to the ABP1 saga.
Although I sympathise with the authors difficulty in identifying an explanation for how the KD lines can lead to the observed abnormalities, one experiment that should be done would in my opinion be a test of the expression of the closest $ABP1$-like genes that can be identified.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

**Competing Interests:** No competing interests were disclosed.