Polarized localization and borate-dependent degradation of the *Arabidopsis* borate transporter BOR1 in tobacco BY-2 cells

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Abstract

In *Arabidopsis* the borate transporter BOR1, which is located in the plasma membrane, is degraded in the presence of excess boron by an endocytosis-mediated mechanism. A similar mechanism was suggested in rice as excess boron decreased rice borate transporter levels, although in this case whether the decrease was dependent on an increase in degradation or a decrease in protein synthesis was not elucidated. To address whether the borate-dependent degradation mechanism is conserved among plant cells, we analyzed the fate of GFP-tagged BOR1 (BOR1-GFP) in transformed tobacco BY-2 cells. Cells expressing BOR1-GFP displayed GFP fluorescence at the plasma membrane, especially at the membrane between two attached cells. The plasma membrane signal was abolished when cells were incubated in medium with a high concentration of borate (3 to 5 mM). This decrease in BOR1-GFP signal was mediated by a specific degradation of the protein after internalization by endocytosis from the plasma membrane. Pharmacological analysis indicated that the decrease in BOR1-GFP largely depends on the increase in degradation rate and that the degradation was mediated by a tyrosine-motif and the actin cytoskeleton. Tyr mutants of BOR1-GFP, which has been shown to inhibit borate-dependent degradation in *Arabidopsis* root cells, did not show borate-dependent endocytosis in tobacco BY-2 cells. These findings indicate that the borate-dependent degradation machinery of the borate transporter is conserved among plant species.
Introduction
Boron is one of the essential nutrients for plants, and boron deficiency is a major cause of reduced crop production. Large quantities of boron are toxic to plants, and boron toxicity is a worldwide problem in food production. Two different classes of borate transporters were discovered in Arabidopsis thaliana. One of them, a transporter named BOR1, is a plasma membrane borate exporter in Arabidopsis root cells, and is essential for efficient xylem loading of boron. BOR1 and its paralogs are also involved in boron toxicity tolerance in plants.

It was reported that the level of BOR1 is tightly regulated by the concentration of borate in the growth environment. At low concentrations of borate, BOR1 is stably localized to the proximal side of plasma membrane in root cells but is degraded upon application of high concentrations of borate. This degradation occurred after endocytosis of the transporter from the plasma membrane and the endocytosed transporter was transported from early endosome to multivesicular body, ubiquitinated and finally targeted to vacuoles for degradation. A similar boron-dependent decrease in borate transporter levels was also observed in rice, although in this case the mechanism of the reduction was not elucidated. Thus it was not clarified whether the borate-induced endocytotic degradation of BOR1 that is found in Arabidopsis root cells is conserved among different plant species and different types of plant cells.

The tobacco BY-2 cell line is widely used as a model for the analysis of the cell cycle and protein trafficking in plant cells. This cell line is advantageous for conducting pharmacological studies because of the small size of its cell clumps as well as its ability to grow in liquid suspension. To obtain an insight into the regulation of borate transporter levels and borate sensing machinery, we investigated the localization and degradation of BOR1-green fluorescent protein (GFP) fusion in tobacco BY-2 cells in the presence of high concentrations of borate and analyzed the effect of inhibitors for protein synthesis, protein degradation and intracellular trafficking on its degradation.

Results
Expression of the GFP fusion and Arabidopsis borate transporter BOR1 in tobacco BY-2 cells
We expressed the Arabidopsis BOR1-GFP fusion construct in tobacco BY-2 cells under the control of the cauliflower mosaic virus 35S RNA promoter. Examination of transformed cells at growing phase using an epifluorescence microscope indicated that the fluorescence localized at the most peripheral part of the cells as well as intracellular dots (Figure 1A). In contrast, cells grown to the stationary-phase showed faint punctate distribution of the fluorescence in the cell (Figure 1B). To examine whether the peripherally-localized BOR1-GFP in rapidly-growing cells actually localized to the plasma membrane we stained the cells with FM4-64 for 15 min on ice and compared the pattern of FM4-64 and the GFP fluorescence using a spinning disk confocal microscope (Figure 1C, 0 min). The peripheral staining pattern of FM4-64 is almost completely identical to that of the fluorescence of GFP. When the FM4-64 incubated cells were further incubated at room temperature for 30 min, we observed internalization of the FM4-64 signal. Under this condition we did not observe the internalization of BOR1-GFP signal (Figure 1C, 30 min). These observations suggest that BOR1-GFP is localized to the plasma membrane in transformed tobacco BY-2 cells.

We also analyzed the distribution of the fluorescence protein after fractionation of the cell lysate into membranous organelle and soluble fractions. As shown in Figure 1D, BOR1-GFP was enriched in the precipitated fractions. This confirmed that BOR1-GFP was
targeted to the membranes in tobacco BY-2 cells. Taken together, we concluded that a significant portion of BOR1-GFP expressed in tobacco BY-2 cells is targeted to the plasma membrane as in the case of *Arabidopsis* root cells.

**Borate-dependent decrease of BOR1-GFP in transformed tobacco cells**

In order to address whether BOR1-GFP is degraded upon supply of high concentrations of borate in tobacco cells as observed in *Arabidopsis*, we first analyzed the effect of borate on the growth of tobacco cells that expressed the BOR1-GFP. Tobacco cells were grown for a week in medium containing varied concentrations of borate (0.1 to 20 mM) and the volume of cells was measured. No difference in the response to cell growth was observed between non-transformed and BOR1-GFP expressing cells. Medium containing up to 5 mM borate caused little defect in the growth of cells (Figure 2).

We then treated the transformed cells with 5 mM borate for up to 90 min and the fluorescence of BOR1-GFP was monitored under an epifluorescence microscope (Figure 3A). We found that the BOR1-GFP signal at the peripheral part of the cells decreased significantly during incubation. The same decrease in signal was also observed in the presence of 3 mM borate. Interestingly, lower concentrations of borate (0.1 mM) did not cause a loss of the BOR1-GFP signal. This decrease is specific for BOR1-GFP as the same treatment did not decrease the fluorescence of plasma membrane intrinsic protein (PIP)-GFP, which is a plasma membrane aquaporin fused with GFP.

We also analyzed the fate of BOR1-GFP after separation of cellular proteins by SDS/PAGE and recorded the fluorescence (Figure 3B). We found that BOR1-GFP signal was decreased in the presence of higher concentrations of borate. This decrease was specific for BOR1-GFP as this protein was not decreased without exposure to borate and the level of PIP-GFP was not changed even with high concentrations of borate. Quantification of the intensities of the BOR1-GFP band allowed us to estimate the kinetic parameters of the decline in the signal intensity. The decrease ratio fitted well into an index recurrence curve with a lag time of 11.7 min and half life of 62 min (Figure 3C), suggesting that the degradation kinetics is first order. The presence of a lag time suggested that several steps of cellular events must have taken place before degradation of the BOR1-GFP.

**Figure 2.** Effect of borate on the growth of BY-2 cells. Seven-day old tobacco BY-2 cells were subcultured into medium containing the indicated concentration of borate and shaken for a week. The volume of cells in the culture was measured. Gray bar; standard borate concentration medium. Error bars, SD.

**Figure 3.** Higher concentrations of borate in the medium induced the degradation of BOR1-GFP. A. Fluorescence images of transformed cells expressing BOR1-GFP in medium containing a high concentration of borate (5 mM, +Borate) and a normal concentration of borate (0.1 mM, -Borate). Images were collected with identical exposure conditions using a confocal microscope. A representative gel image is shown. B. Time course of the decrease in BOR1-GFP signal in the presence of borate. Upper gel; cells expressing BOR1-GFP were incubated in medium with normal or high borate concentrations for the indicated times. Lower gel; cells expressing PIP-GFP were incubated in high borate medium for the indicated times. Fluorescence of GFP fusion proteins after separation by SDS-PAGE was detected by scanning the gel with an image analyzer. A representative gel image is shown. C. Quantification of the degradation of BOR1-GFP. Means of three independent experiments are shown. The gray line is a best-fit curve of the exponential decrease of BOR1-GFP.

**Polarized localization of BOR1-GFP**

As described above, BOR1-GFP but not PIP-GFP was degraded upon addition of borate. In *Arabidopsis* roots BOR1-GFP behavior differs from some other plasma membrane proteins in both
Figure 4C

**A** Polarized localization of BOR1-GFP in tobacco BY-2 cells. A. Confocal images of PIP-GFP and BOR1-GFP at the four-cell stage. About half to two-thirds of cell clumps are shown. GFP fluorescence of PIP-GFP at the plasma membrane that faces to the medium showed much higher intensities than that of BOR1-GFP. B. Schematic representation of each cell membrane face in the cell clump. Only half of the 4-cell stage cell clump was indicated. A1, A2, E1 and E2 indicate the face of cells defined for the analysis of polarized localization. C. Comparison of the relative density of fluorescence at each face of PIP-GFP and BOR1-GFP expressing cells. Means (with SD) density relative to the E1 face (n=8) are shown.

To address whether localization of BOR1-GFP and PIP-GFP differs in tobacco BY-2 cells, we compared the localization of these proteins in tobacco BY-2 cells at the early log-phase stage (Figure 4).

With our culture conditions, BY-2 cells at the stationary phase are largely unicellular. Cells at this stage are approximately three-times longer than cells at mid-log phase. When such cells are transferred to fresh medium, they start to divide at the center resulting in two daughter cells which elongate slightly and further divide at the center of each daughter cell without separation. At this stage, a linearly attached clump of cells contain four cells. The clump has four cell walls with a distinct location and age; the one between the two cells that generated at the first cell division, the one between the two cells that generated at the second cell division, the one exposed to the culture medium in the center of two cells, and the one exposed to medium of the two extreme cells containing tips of the cell wall clump. We name these four faces of cells as A1, A2, E1 and E2, respectively (Figure 4B).

We measured the density of the green fluorescence of these four faces from cells expressing BOR1-GFP and PIP-GFP and the distribution of their relative density (relative to the E1 face). In both cases, the fluorescence density at E1 and E2 was almost the same but in the case of PIP-GFP, the densities at A1 and A2 were approximately 1.5- and 2-fold higher than that of E1, respectively (Figure 4C). These fluorescence intensities between cells suggest that PIP-GFP is distributed nearly evenly in the plasma membrane as the A1 and A2 faces have two sheets of plasma membrane. In contrast, the density of BOR1-GFP at both A1 and A2 faces were approximately 5-fold higher than that of E1 (Figure 4C). This observation indicated the polarized localization of BOR1-GFP and clearly showed that this fusion protein accumulates to the plasma membrane that is facing neighboring cells in growing tobacco BY-2 cells.

**Inhibition of protein synthesis partially suppresses borate-induced degradation**

The observation that BOR1-GFP is degraded upon addition of borate in the medium (Figure 3) suggest two possible scenarios for the borate-dependent decrease in BOR1-GFP. The first possible case is that the decrease of BOR1-GFP in the presence of high concentrations of borate is the result of inducible degradation. Another possibility is that the rate of the degradation of BOR1-GFP is rapid in BY-2 cells even in the absence of borate but a high rate of biosynthesis of BOR1-GFP in the normal medium maintained the level of the transporter.

To address the likely scenario causing the BOR1-GFP decrease in tobacco cells under high borate conditions, we treated the transformed cells with a protein biosynthesis inhibitor cycloheximide (CHX) in either normal or high-boron medium and monitored the level of BOR1-GFP (Figure 5). In the normal borate medium, addition of CHX did not change the level of BOR1-GFP significantly over 2 hours. This observation indicated that the turnover of BOR1-GFP in normal medium is slow and most of the BOR1-GFP showing the green fluorescence is the result of the accumulation of this protein. This also indicated that the decrease in BOR1-GFP in high borate medium is a result of inducible degradation.

We observed partial inhibition of BOR1-GFP degradation in the presence of CHX (Figure 5). This observation indicated that continuous synthesis of degradation machinery is necessary for the continuation of the degradation. The partial suppression of the degradation also indicated that the machinery for the degradation is already present in cells that were cultured in the normal medium.

**Endocytosis of BOR1-GFP under high boron condition**

It was shown previously that endocytosis is the first step of BOR1-GFP degradation in *Arabidopsis* [1]. To assess whether similar machinery operates in tobacco BY-2 cells in the presence of high concentrations of borate, we first analyzed the BOR1-GFP signal...
in cells cultured for 1 to 2 hours under high borate conditions using a confocal laser scanning microscope (CLSM). We found intracellular puncta in cells treated with high borate (Figure 6A). We then analyzed the colocalization of the BOR1-GFP signal with an early endosome that can be marked with FM4-64. Tobacco cells expressing BOR1-GFP were incubated for 2 hours in a medium containing 3 mM borate and then the cells were stained with FM4-64 on ice, washed with cold medium containing 3 mM borate, after which they were returned to room temperature to resume intracellular trafficking as described\(^1\). Five to 10 min after FM4-64 incubation, dotted structures emitting red fluorescence, which represent an early endosome distinct from the trans-Golgi network (TGN) in tobacco cells\(^1\), was observed (Figure 6B, left). Most of the early endosome signal colocalized with intracellular dots of BOR1-GFP, suggesting that BOR1-GFP passes through early endosome during degradation as in the case of Arabidopsis cells.

It is also known that FM4-64 is transported from the early endosome to the TGN, and thereafter targeted to the vacuolar membrane. To evaluate whether BOR1-GFP passes through the TGN, we expressed SCAMP2-mRFP\(^{14}\) in cells expressing BOR1-GFP. SCAMP2-mRFP localizes in the plasma membrane, the TGN and the secretory vesicle cluster, which is an exocytic compartment derived from the TGN\(^{14}\). No colocalization of green and red intracellular dots was observed when cells were treated with 3 mM borate for 2 hours (Figure 6C). These findings suggested that the BOR1-GFP is internalized to endosomes and then sorted to a degradation pathway from the endocytic route, which is directed to the TGN.

### Traffic and other inhibitors that affect the degradation

It was previously reported that the treatment of Arabidopsis roots with brefeldin A (BFA), which is an inhibitor for both endocytosis and exocytosis, prevents the borate-dependent degradation of BOR1-GFP\(^2\). To address whether this compound is also effective in tobacco BY-2 cells and to assess the mechanism of the degradation, we analyzed borate-induced degradation in the presence of various inhibitors (Figure 6). We used BFA, 2,3,5-triiodobenzoic acid (TIBA), a compound that stabilizes the actin bundle\(^3\); tyrphostin A23, which is a tyrosine kinase inhibitor, which can inhibit endocytosis by competing with the interaction of the AP-2 adaptor complex and Tyr-containing endocytosis motifs in mammalian cells and also can inhibit the endocytosis of rice SCAMP1-YFP and FM4-64 in tobacco BY-2 cells\(^{4,17}\); ikarugamycin, which is an inhibitor of clathrin-mediated endocytosis in tobacco\(^4\); colchicine, which is an inhibitor of microtubule depolymerization; latrunculin B, which inhibits actin polymerization; 2,3-butanedione monoxime (BDM), which is a general myosin ATPase inhibitor and inhibits the endocytosis of SCAMP2-mRFP in tobacco BY-2 cells\(^{17}\); MG-132, which is an inhibitor of proteasomes at low concentrations and also inhibits other proteases; and K-252a, which is a general Ser/Thr type protein kinase inhibitor.

Three hours after application of inhibitor solution to the tobacco cell culture, borate was added at a final concentration of 5 mM and further incubated for 2 hours. Thereafter, the amount of BOR1-GFP was quantified (Figure 7A). We found that BFA and TIBA inhibited degradation almost completely (>80% remained) at a concentration that inhibits the endocytosis of SCAMP2-YFP\(^{14}\). A slightly weaker effect of tyrphostin A23 was observed at 400 μM, which is a concentration similar to that used in the works of Leborgne-Castel et al.\(^{18}\) and Lam et al.\(^{17}\). In this case, approximately 70% of BOR1-GFP remained in the cell. We also found partial inhibition of degradation by BDM at 5 mM, the concentration that inhibits the endocytosis of SCAMP2-YFP\(^{14}\). We also observed partial inhibition by MG-132 at 400 μM. In these two cases, approximately 40% of BOR1-GFP remained in the cell. Other inhibitors tested did not cause any detectable effect on the extent of degradation.

Time-course analysis of the decrease of BOR1-GFP in the presence of BFA and tyrphostin A23 (Figure 7B) clearly showed that most of the degradation was inhibited up to 3 hours. We next addressed whether the inhibition occurred at the point of endocytosis from the plasma membrane or at intermediate compartments. Fluorescence images were collected in transformed tobacco cells in the presence of both inhibitors and borate. We found that the images were not significantly different up to 3 hours in the presence of either BFA or TIBA (Figure 7C). These observations suggested that both BFA and TIBA inhibited the early event of endocytosis. We could not...
Figure 6. Endocytotic transport of BOR1-GFP during borate-induced degradation. A. Two hours after incubating the cells in medium containing 3 mM borate, cells were incubated with FM4-64 for 30 min and analyzed with a confocal fluorescent microscope. B. Many, but not all intracellular BOR1-GFP dots colocalized with puncta of FM4-64. C. BOR1-GFP did not pass through the trans-Golgi network (TGN) during degradation. BOR1-GFP was transiently expressed in cells expressing SCAMP2-mRFP, which is a marker for TGN and the secretory vesicle cluster \(^{14}\). Co-expressed cells were incubated in medium containing 3 mM borate for 2 hours and the fluorescence was analyzed by confocal laser scanning microscopy. No clear overlap of the intracellular puncta of SCAMP2-mRFP and BOR1-GFP was seen.
completely. These observations indicated that these inhibitors have a direct effect against the endocytosis of BOR1-GFP and were not a side effect of these chemicals. These observations also suggested that actin dynamics, GDP-GTP exchange of Arf small GTPase, and either tyrosine phosphorylation or AP-2 adaptor interaction are necessary for the degradation of BOP1-GFP in tobacco BY-2 cells.

Tyrosine mutations prevented borate-induced degradation

It was shown that both polarized localization and borate-dependent degradation are mediated by specific Tyr residues in BOR1-GFP in Arabidopsis root cells. To address whether these Tyr residues also contribute to the localization and borate-induced degradation, we investigated fluorescence images in the presence of tyrphostin A23 because treatment of BY-2 cells with this compound increased cellular fluorescence significantly and such fluorescence was independent of the expression of BOR1-GFP.

To address whether the inhibition of endocytosis required BY-2 cells to be pretreated with BFA, tyrphostin A23 or TIBA, we investigated the timing of adding the inhibitors (Figure 7D). We compared the level of BOR1-GFP without inhibitors with BOR1-GFP levels when these inhibitors were added into the culture. We found that addition of inhibitors at the same time, in the lag time as well as during the degradation phase, prevented the decrease in BOR1-GFP almost completely. These observations indicated that these inhibitors have a direct effect against the endocytosis of BOR1-GFP and were not a side effect of these chemicals. These observations also suggested that actin dynamics, GDP-GTP exchange of Arf small GTPase, and either tyrosine phosphorylation or AP-2 adaptor interaction are necessary for the degradation of BOP1-GFP in tobacco BY-2 cells.

Figure 7. Inhibitors that prevent borate-dependent degradation. A. Cells incubated with or without inhibitors for 30 min were further incubated in medium containing 5 mM of borate for two hours in the presence of the same inhibitors. The relative intensity of BOR1-GFP fluorescence was quantified as described in the Figure 3 legend. The final concentration of inhibitors in the culture and the number of replicates were; brefeldin A (BFA), 25 μM, n=8; 2,3,5-triodobenzoic acid (TIBA), 50 μM, n=5; tyrphostin A23, 100 μM, n=5; MG-132, 400 μM, n=5; 2,3-butanedione monoxime (BDM), 40 mM, n=3; Ikarugamycin, 20 μM, n=3; Latrunculin A, 100 μM, n=3; colchicine, 200 μM, n=3. Error bars, SD. ** and * represent the significant difference relative to no inhibitor (p<0.01 and <0.05 respectively). B. Time-course analysis of the degradation of BOR1-GFP in the presence of either TIBA, tyrphostin A23 or in the absence of an inhibitor. C. Changes in the fluorescence pattern of BOR1-GFP in the absence and presence of BFA or TIBA in the presence of 5 mM borate. D. Timing of the addition of inhibitors did not affect the subsequent degradation. Cells were incubated in medium containing 5 mM of borate. At the same time (0 min) or the other indicated times, either BFA, tyrphostin A23 or TIBA was included in the medium and further incubated for up to 120 min. The intensity of the BOR1-GFP band relative to that at the time when the inhibitor was included is shown. Error bars, SD.
This observation indicated that the polarized localization of BOR1-GFP in tobacco cells is largely independent of the tyrosine residues in the BOR1 protein.

When tobacco cells expressing the mutant protein were incubated in medium containing 5 mM borate, we still observed strong fluorescence of GFP signals. We did not detect any difference in the pattern and intensity of the fluorescence signals between cells incubated in normal medium and high borate medium (Figure 8C). Under the same condition, most of the wild-type BOR1-GFP was degraded in the cell (Figure 3, Figure 8C). This observation indicated that Y373, Y398, and/or Y405 residues of BOR1-GFP are involved in borate-induced degradation in tobacco BY-2 cells as in the case in Arabidopsis root cells.

Discussion

We found that the Arabidopsis borate transporter BOR1-GFP fusion protein was targeted to the plasma membrane of rapidly growing tobacco BY-2 cells. The plasma membrane localization of BOR1-GFP was different from that of PIP-GFP. BOR1-GFP was more abundant in the plasma membranes that face neighboring cells (Figure 4A, B). This observation indicates that BOR1-GFP showed a polarized localization in tobacco BY-2 cells. This observation further indicates that plasma membranes of BY-2 cells facing the medium and the neighboring cells are distinct in nature. It was shown in Arabidopsis root cells that BOR1-GFP is localized in the proximal region of the plasma membrane. However, polarized localization in BY-2 cells and Arabidopsis root cells seems to be mediated by distinct mechanisms because the latter, but not the former, is sensitive to the mutation of Tyr residues in the cytoplasmic loop of BOR1 protein (Figure 8A, B).

The concentrations of borate that caused the degradation of BOR1-GFP in BY-2 cells was >3 mM, which is higher than that in Arabidopsis plants (100 μM). This difference might correlate with the difference of high-boron sensitivity of the growth between the two species. BY-2 cells can grow normally up to 5 mM borate (Figure 2D) whereas 300 μM borate caused about 50% reduction of the growth of Arabidopsis plants. Thus it will be interesting to investigate in the future what attributes of plants including tobacco cells and Arabidopsis plants determine sensitivity to high borate levels.

The borate-induced degradation of BOR1-GFP was inhibited by the protein synthesis inhibitor CHX (Figure 5). We observed previously that CHX treatment did not inhibit endocytosis of SCAMP2-YFP in tobacco BY-2 cells. Therefore, the effect of CHX on the inhibition of BOR1-GFP is not a general response in tobacco cells. The inhibition by CHX on BOR1-GFP degradation was not complete and approximately 50% of the protein was degraded under CHX treatment (Figure 5). This observation suggested that BY-2 cells used...
pre-existing machinery for the degradation of BOR1-GFP. However, the degradation the machinery was limited in the cell and de novo synthesis of the machinery is required for the continuation of the degradation.

The plasma membrane accumulation of BOR1-GFP was abolished when cells were incubated in medium containing high concentrations of borate. The decline in the BOR1-GFP was mediated by endocytosis and further degradation (Figure 3 and Figure 6) and was dependent on the tyrosine residues (Figure 8C) as observed in Arabidopsis root cells. These observations suggest that boron sensing mechanisms as well as the borate-dependent induction of the endocytosis of BOR1-GFP is conserved in different plant species. However, there are some differences in the endocytic trafficking between tobacco cells and Arabidopsis root cells. The most pronounced difference is the organization of organelles in these cells. In tobacco BY-2 cells, early endosomes and the TGN are separated and BOR1-GFP passed through the early endosome but not the TGN during degradation (Figure 6B, C). In contrast, early endosomes cannot be distinguished from the TGN in Arabidopsis root cells. Therefore, differentiation of secretory organelles is not completely identical in these two species. Further analysis of the degradation of BOR1-GFP in either cultured Arabidopsis cells or tobacco root cells will give a clearer picture as to whether the difference in trafficking paths originate from differences between species or the differentiation state of the cells.

The endocytosis-dependent degradation of BOR1-GFP was inhibited almost completely upon addition of the inhibitors BFA, TIBA and tyrphostin A23. BFA is known as an inhibitor of the GDP-GTP exchange factor for ADP-ribosylation factor and has many effects on different plant cells. On tobacco BY-2 cells, we have shown that BFA inhibits both endocytosis of SCAMP2-YFP, Golgi protein localization and export from the endoplasmic reticulum. It was also shown that BFA inhibits the endocytosis of BOR1-GFP in Arabidopsis root cells. The fact that BFA inhibited the endocytosis of BOR1-GFP in tobacco BY-2 cells in the presence of borate (Figure 7) indicates that some machinery for the borate-dependent endocytotic degradation is conserved between Arabidopsis root cells and tobacco BY-2 cells.

TIBA is known to inhibit the endocytosis of the auxin transporter and its function is to stabilize the actin cytoskeleton at 25 μM in tobacco BY-2 cells. The fact that TIBA at a similar concentration range inhibited the degradation of BOR1-GFP almost completely (Figure 7) suggested that the actin cytoskeleton is involved in the endocytotic degradation of BOR1-GFP. The partial inhibition of degradation by BDM (Figure 7), which is an inhibitor of myosin motors also supports the idea that an actin-dependent mechanism is a key part of the machinery for the endocytotic degradation of BOR1-GFP.

Another inhibitor that prevented borate-dependent degradation was tyrphostin A23. This chemical is known to inhibit Tyr-signal mediated endocytosis by inhibiting the recognition of the AP-2 adaptor complex signal in mammalian cells. The requirement of Tyr residues in BOR1 for endocytosis (Figure 8C) agrees well with the action of tyrphostin A23 in mammalian cells. This compound is also known to inhibit endocytosis in several plant species. It was reported recently that this compound inhibited elictor-induced endocytosis in tobacco BY-2 cells and this event seems related to the induction of clathrin-mediated endocytosis. However, ikarugamycin, which is known to inhibit clathrin-mediated endocytosis in both animal cells and tobacco protoplasts, did not inhibit the degradation of BOR1-GFP. It was reported recently that AP-2 is involved not only in clathrin-mediated endocytosis but also in clathrin-independent endocytosis in mammalian cells. Therefore further investigation is necessary to elucidate whether clathrin is involved in the endocytosis of BOR1-GFP.

In addition to these inhibitors, we observed that MG-132 also inhibited the degradation of BOR1-GFP. It was reported that this compound also inhibits the degradation of interleukin-2 through the inhibition of endocytosis of its receptor in mammalian cells. As MG-132 is a potent inhibitor of proteasomes, our finding raises the possibility that a proteasome system is somehow involved in BOR1-GFP degradation. However, our data does not rule out the possibility that the target of MG-132 on BOR1-GFP degradation is different from proteasomes as we needed a very high concentration (400 μM) of this compound to prevent BOR1 degradation and MG-132 can inhibit other proteases, such as calpain in animals, at a high concentration. Thus, future experiments will be necessary to ascertain whether proteasomes are involved in the degradation of BOR1-GFP and to show which molecule is the real target of MG-132 on BOR1-GFP degradation in tobacco cells.

Materials and methods
Plasmids
Plasmids for the expression of wild-type and mutant BOR1-GFP, and SCAMP2-mRFP have been described previously. The expression plasmid for PIP-GFP was a kind gift from Dr. M. Maeshima (Nagoya University). This plasmid contains radish plasma membrane type aquaporin PAQ1 and GFP fusion protein under an enhancer-duplicated cauliflower mosaic virus 35S promoter.

Chemicals
The following chemicals were used to test the inhibition of the degradation: BFA (Wako Pure Chemicals, Osaka, Japan), CHX (Wako Pure Chemicals, Osaka, Japan), TIBA (Aldrich Chem. Co. USA), Tyrphostin A23 (Sigma-Aldrich, Inc, Missouri, USA), colchicine (Tokyo Kasei Co., Tokyo, Japan), K-252a (Sigma-Aldrich, Inc, Missouri, USA), BDM (Sigma-Aldrich, Inc, Missouri, USA), Latruncurin A (Calbiochem, San Diego, USA), ikarugamycin (BIOMOL Research Labs Inc. USA), MG-132 (Calbiochem Inc, San Diego USA), wortmannin (Calbiochem Inc. San Diego USA), 3MA (Sigma-Aldrich, Inc., Missouri, USA) and E-64 (Peptide Institute, Osaka, JAPAN).

Cell culture and transformation
The culture of tobacco BY-2 cells (a kind gift from Drs. K. Nakamura (Nagoya University) and T. Nagata (University of Tokyo) and the transformation of this cell line were as described. Cells were cultured weekly and 3 or 4 day-old cultures were used unless otherwise stated. For transient transformation of cells expressing BOR1-GFP to express SCAMP2-mRFP in the same cell, Agrobacterium tumefaciens EHA105 harboring a binary plasmid was co-cultured with BOR1-GFP expressing tobacco cells for 2 days, washed with fresh
medium and further incubated in medium containing 200 mg/L ceftaxime for a day. Thereafter fluorescence of cells was visualized using CLSM.

**Quantification of GFP fusion proteins**

Cells expressing GFP fusion proteins were lysed with equal volumes of buffer consisting of 0.4 M Sorbitol, 0.1 M KOAc, 6 mM EGTA, 4 mM EDTA, 40 mM HEPES-KOH, 2 mM DTT, 1 drop/100 ml protease inhibitor tablet (Complete EDTA-free, Roche), pH 7.3 at room temperature. Cells were lysed by sonication and the resulted cell lysate was centrifuged at 300 x g for 10 min. The resulting supernatant was used as the cell lysate. Proteins in the cell lysate was mixed with 1/5 volume of a buffer consisting 0.25 M Tris-HCl, 0.5 M dithiothreitol, 50% glycerol, 10% SDS, 0.025% bromophenol blue, pH 6.8 and loaded on SDS-polyacrylamide gels without heating. Fluorescence images of the gels were recorded using Typhoon 9600 scanner (GE Healthcare, Tokyo, Japan) at the following setting; filter 520SPCy2, excitation 488 nm blue laser at PMT 650, medium sensitivity). The intensities of bands were quantified using Image Quant software (GE Healthcare).

**Microscopic analysis**

For epifluorescence detection, cells were mounted on glass slides and observed using IX40 inverted microscope with an LCPlanFL 20X lens using NBY excitation/emission filter set and a DP-70 camera (Olympus Co. Tokyo Japan). For time-lapse image collection, cell suspension was placed into in a well of glass-bottom 96 well micro-plates (EZVIEW® Glass Bottom Culture Plates LB, Asahi Techno Glass Co. Tokyo, Japan). Cells in the plate wells were observed using an IX80 inverted fluorescence microscope equipped with DSU spinning disk confocal unit and an Uplan SApo 40×/0.90 lens using NBY or RFP excitation/emission filter sets (Olympus). Images were collected using an iXON+-DU888E-CS0-BV camera (Andor) under the control of Meta Imaging Series 7.6.3 software (Molecular Devices Inc. Sunnyvale, CA, USA).

For the colocalization analysis, cells were mounted on slides with culture medium. These cells were observed using a CLSM system (LSM510 META, Axioplan2 Imaging; Carl Zeiss) with a Plan-Apochromat lens (633 1.4 oil differential interference contrast; optical slices of 1 mm). We used a 25-mW argon laser (power, 5%) with 488-nm excitation and a 505–530-nm band-pass filter for GFP8. For FM4-64 (Molecular Probes, Eugen, USA) we used a 650-nm long-pass filter and 488-nm excitation. A 560-nm long-pass filter and 1-mW He-Ne laser at 80% power with 543-nm excitation was used for Alexa Fluor 568 and mRFP. Crosstalk was prevented using a multitrack configuration with line sequential scanning. Composite figures were prepared using Zeiss LSM Image Browser software.

For the measurement of intensity, confocal images at 256 bit tiff file was quantified using ImageJ software. If required, relative density to the E1 face was calculated in each cell clump.

**Borate treatment to monitor endocytosis**

An aliquot of 0.5 M boric acid in water was added to 96.5 ml of suspension culture of three-day old tobacco cells expressing BOR1-GFP in a 300 ml Erlenmeyer flask at a concentration of 3 or 5 mM (as indicated in the legends to figures) and further cultured with the same condition of regular culture. For the control, cells grown in a normal culture medium, which contains 0.1 mM borate, was used. At the indicated time, a 0.5 or 1 ml aliquot of the culture was removed from the flask and used for the analysis.

**Statistical analyses**

For the comparison of the two values student’s t-test was employed using Microsoft Excel software. For the comparison of difference with multi values, Tukey’s test was used.

**Author contributions**

TF and KM conceived the study and designed the experiments. NY, TG, SA, KT and KM carried out the research. TG, NY and KM prepared the first draft of the manuscript and TF contributed the preparation of the manuscript. All authors were involved in the revision of the draft manuscript and have agreed to the final content.

NY and TG contributed equally to the work.

**Competing interests**

No competing interests were disclosed.

**Grant information**

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References


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Agustín González-Fontes
Department of Physiology, Anatomy and Cellular Biology, Pablo de Olavide University, Seville, Spain

This article is an extension and confirmation of findings on the degradation of the borate transporter BOR1 in Arabidopsis thaliana tobacco BY-2 cells. From elegant and conclusive techniques used in previous works by the Fujiwara and Matsuoka groups, the authors demonstrated that, under high B supply (3-5 mM), the BOR1 transporter in transformed tobacco BY-2 (BOR1-GFP) cells was also transported from the plasma membrane to the vacuole via endocytosis for degradation, this borate-dependent process being specific for BOR1-GFP. However, unlike in Arabidopsis, BOR1-GFP did not pass through the trans-Golgi network during degradation.

Interestingly, the authors showed that de novo synthesis of the borate-dependent degradation machinery of the borate transporter BOR1 is required for maintaining the degradation. However, it is noteworthy that the threshold B concentration for the onset of borate-dependent degradation in tobacco BY-2 cells was 3 mM, but in Arabidopsis root cells is only 0.1 mM B. The authors should give some additional explanation for this significant discrepancy, other than that the plant species have different high-boron sensitivity.

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 28 October 2013
doi:10.5256/f1000research.751.r2177

Liwen Jiang
School of Life Sciences, Chinese University of Hong Kong, Hong Kong, China

This is an interesting study on the degradative pathway and dynamics of BOR1-GFP in transgenic tobacco BY-2 cells. The conclusions are largely supported by the provided data. I welcome its publication as it is informative and useful for the study of endocytosis in plant cells.

Some minor suggestions:

1. Add scale bars to some of the Figures (e.g. Figures 1C, 3A, 4A, 7C, 8A).
2. Please indicate the molecular weight (kDa) of the detected proteins in Figures 1D, 3B and 5A.
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.

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Referee Report 23 October 2013

**Tomomichi Fujita**
Faculty of Science, Hokkaido University, Sapporo, Japan

This paper demonstrates that in a heterologous plant system the Arabidopsis borate transporter BOR1 was polarly localized in tobacco BY-2 cells and was degraded in a borate-dependent manner as has been reported in Arabidopsis roots. Pharmacological analysis suggested both common and distinct routes for the BOR1 degradation between Arabidopsis roots and tobacco BY-2 cells. The authors also explored Tyr mutants of the BOR1-GFP in BY-2 cells and found an interesting, distinct response from Arabidopsis. Thus I think this article has enough quality of science to be approved in F1000Research. I have some comments and questions that should be considered and explained for the improvement of the article.

1. Fig. 1 A: I can see some green dots from BOR1-GFP in the cytosol. According to the rest of the results (e.g. Fig. 1C, 3A (-Borate), 4A (Bor1-GFP)) the fluorescent signal in the low concentration of borate media and in the log phase of the BY-2 line is mainly detected in plasma membranes but not in the cytosol; could you explain why you think this difference occurred?
2. Fig. 1 D: The authors fractionated the cell lysate to show that BOR1-GFP is targeted to the membrane fractions; it would be interesting to know whether either fraction between P10 and P100 contains more plasma membranes or more microsomes. It would also be helpful if the authors could add control experiments for each fraction using protein markers that are known to localize in plasma membranes, microsomes, or the soluble fraction.
3. p.4 left column: The authors used PIP-GFP, and then later SCAMP2-mRFP. It would be better to describe the origins (Arabidopsis or tobacco) of these in the Materials and Methods section.
4. Fig.2: The authors measured the cell volume of BY-2 cells. Please add the methods used and how many times the measurement was replicated (n=?).
5. Fig. 6A: In the picture, please add the note ‘Bor1-GFP’ as indicated in Fig. 6B and C.
6. Fig. 6C legend: Please check if the sentence ‘BOR1-GFP was transiently expressed in cells expressing SCAMP2-mRFP...’ is not instead supposed to read ‘SCAMP2-mRFP was transiently expressed in cells expressing BOR1-GFP.’
7. Fig. 6C: The authors examined the endocytic route for BOR1 degradation. Based on the observation in the merged picture, they concluded that since no overlapping of green and red dots was observed, co-localization of SCAMP2 and BOR1 was unlikely. However, I see only a few dots in the original picture of BOR1-GFP that they presented, and I wonder if this is why in the merged picture there were few yellow signals indicating an overlap of SCAMP2 and BOR1. The authors should show an appropriate BOR1-GFP image with more dots of BOR1-GFP, and the SCAMP2 image should be merged so that the colocalization can be observed. Additionally, even in the current merged picture, I can see at least one yellow dot at the periphery of the bottom-side plasma membrane; how is this dot explained?
8. p.6 right column, line-16: (Figure 6) should be (Figure 7).
9. Fig. 7C: The authors examined whether inhibition occurred at the point of endocytosis from the plasma membrane or at intermediate compartments. However, it is hard to distinguish if there are any intermediate compartments or not at the presented magnification/resolution of the images. Especially in the images for the TIBA treatment, as the relatively strong fluorescent signal was probably recognized in a surrounding region of the nucleus. Please show better images and explain if this is related to the intermediate compartments or something else.

10. p.11 left column, line 7 from below: ‘GFP8’ could be ‘GFP’?

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.