Transport-coupled ubiquitination of the borate transporter BOR1 for its boron-dependent degradation

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Abstract
Plants take up and translocate nutrients through transporters. In Arabidopsis thaliana, the borate exporter BOR1 acts as a key transporter under boron (B) limitation in the soil. Upon sufficient-B supply, BOR1 undergoes ubiquitination and is transported to the vacuole for degradation, to avoid overaccumulation of B. However, the mechanisms underlying B-sensing and ubiquitination of BOR1 are unknown. In this study, we confirmed the lysine-590 residue in the C-terminal cytosolic region of BOR1 as the direct ubiquitination site and showed that BOR1 undergoes K63-linked polyubiquitination. A forward genetic screen identified that amino acid residues located in vicinity of the substrate-binding pocket of BOR1 are essential for the vacuolar sorting. BOR1 variants that lack B-transport activity showed a significant reduction of polyubiquitination and subsequent vacuolar sorting. Coexpression of wild-type (WT) and a transport-defective variant of BOR1 in the same cells showed degradation of the WT but not the variant upon sufficient-B supply. These findings suggest that polyubiquitination of BOR1 relies on its conformational transition during the transport cycle. We propose a model in which BOR1, as a B transceptor, directly senses the B concentration and promotes its own polyubiquitination and vacuolar sorting for quick and precise maintenance of B homeostasis.

Introduction
Plants take up and translocate mineral nutrients via roots using specific transporters. To deal with the fluctuating nutrient availability in the soil environment, plants tightly regulate the abundance of transporters by sensing nutrient concentrations. When a nutrient is limited, plant roots
express a set of transporters for efficient transport. In this state, resupply of the nutrient could potentially cause rapid accumulation to a toxic level in plant tissues. To avoid this problem, several nutrient transporters in plants are downregulated by degradation through ubiquitination in response to elevated nutrient concentrations (Barberon et al., 2011; Bayle et al., 2011; Kasai et al., 2011; Lin et al., 2013; Rodriguez-Furlan et al., 2019).

Boron (B) is essential for plant growth and survival due to its function in cross-linking pectin at the rhamnogalacturonan II (RG-II; Yoshinari and Takano, 2017). Boron is taken up mainly as boric acid ([B(OH)₃]) by plant roots from the soil solution by passive diffusion and channels (Funakawa and Miwa, 2015; Yoshinari and Takano, 2017). Under low-B conditions, a boric acid channel NIP5;1 and the borate anion ([B(OH)₄]⁻) exporter BOR1 play key roles in B uptake and translocation in roots of Arabidopsis thaliana (Takano et al., 2002; Takano, 2006; Takano et al., 2010; Miwa et al., 2013; Yoshinari and Takano, 2017). In various root cells, including epidermal and endodermal cells, NIP5;1 and BOR1 are localized to the plasma membrane in a polar fashion toward the soil- and stele-side, respectively (Takano et al., 2010; Yoshinari and Takano, 2017). The polar localization of NIP5;1 and BOR1 supports directional transport of B toward the root stele (Wang et al., 2017; Yoshinari et al., 2019). BOR2, one of the BOR1 homologs in Arabidopsis thaliana, is similarly a B exporter for B translocation in the roots, and it is also involved in cross-linking of the RG-II to support root cell elongation under low-B conditions (Miwa et al., 2013).

Similar to most essential nutrients, excessive supply of B inhibits plant growth (Aibara et al., 2018). B toxicity affects various cellular activities, and B accumulation often causes necrosis of tissues (Landi et al., 2019). Since B accumulates at the end of the transpiration stream, high-B tolerance is related to the reduced accumulation of B in the shoot tissue, at least in wheat (Triticum aestivum), barley (Hordeum vulgare), and Arabidopsis (Nable, 1988; Nable et al., 1990; Reid et al., 2004; Reid, 2007). In Arabidopsis and barley, B exclusion from roots by borate exporters BOR4 and Bot1, respectively, protects plants from B accumulation and subsequent toxicity (Miwa et al., 2007; Sutton et al., 2007). In Arabidopsis, transport activity of B toward shoots is upregulated under low-B conditions and is rapidly downregulated after high-B resupply (Takano et al., 2005; Takano, 2006). This regulation is thought to be dependent on the abundance of NIP5;1 and BOR1. The abundance of the NIP5;1 protein is controlled through B-induced mRNA degradation dependent on the 5′ untranslated region (5′UTR; Tanaka et al., 2011, 2016). By contrast, abundance of the BOR1 protein is controlled by two different mechanisms. High concentrations of B repress the translation of BOR1 dependent on the 5′UTR (Aibara et al., 2018). In addition, high concentrations of B rapidly induce the ubiquitination of BOR1 followed by vacuolar transport and degradation (Takano et al., 2005; Kasai et al., 2011). Importantly, transgenic Arabidopsis plants expressing a ubiquitination-defective BOR1 variant (K590A) accumulated higher concentrations of B in shoots and exhibited a significant reduction of shoot growth under high-B conditions (Aibara et al., 2018). Plants expressing BOR1 lacking both endocytic degradation and translational repression exhibited further accumulation of B and a reduction of shoot growth. These results demonstrated the physiological importance of both mechanisms of BOR1 downregulation under toxic-B
conditions. It should be noted that the translational repression occurs at higher B concentrations than the degradation of BOR1 protein (Aibara et al., 2018). Under continuous toxic-B conditions, stopping the de novo protein synthesis could be more cost-effective to control the BOR1 abundance. However, when the B concentration in soil fluctuates between low and sufficient ranges, the turnover of existing BOR1 protein should be particularly important because BOR1 is a key factor that determines the concentration of B in the soil for efficient root-to-shoot translocation under low-B conditions (Takano et al., 2002).

Subcellular localization and abundance of BOR1 are controlled by membrane trafficking. BOR1 in the plasma membrane is constantly endocytosed toward the trans-Golgi network/early endosome (TGN/EE; Viotti et al., 2010; Yoshinari et al., 2018). Under low-B conditions, the constitutive endocytosis is dependent on the AP2 clathrin-adaptor protein complex and DYNAMIN-RELATED PROTEIN 1A (DRP1A), and subsequent recycling maintains polar localization in the plasma membrane (Yoshinari et al., 2016, 2019). Upon high-B supply, however, BOR1 is further transported from the TGN/EE to luminal vesicles of the multivesicular body/late endosome (MVB/LE) and then to the vacuole for degradation (Takano et al., 2005; Viotti et al., 2010; Yoshinari et al., 2018). Interestingly, the B-induced degradation is dependent on DRP1A but not on the AP2 complex (Yoshinari et al., 2016, 2019), indicating differential routes of BOR1 endocytosis from the plasma membrane. Upon high-B supply, BOR1 is ubiquitinated presumably at a lysine residue (K590) in the C-terminal cytosolic tail (Kasai et al., 2011). It appears that ubiquitination triggers AP2-independent endocytosis specialized for vacuolar transport. However, the machinery of B sensing that promotes ubiquitination of BOR1 has not been elucidated.

In the present study, we analyzed the ubiquitination status of BOR1 upon high-B supply in depth and demonstrated that BOR1 undergoes K63-linked polyubiquitination at the K590 residue. We performed a forward genetic screen to identify a putative B-sensor/receptor and unexpectedly found that amino acid residues located in the substrate-binding pocket of BOR1 are essential for ubiquitination of BOR1. We propose a model in which BOR1 acts as a B-transceptor regulating vacuolar transport of itself.

Results

Polyubiquitination at K590 promotes vacuolar transport of BOR1

Our previous report suggested that endocytic degradation of BOR1 is mediated by mono- or diubiquitination under high-B conditions (Kasai et al., 2011). A BOR1 variant with a lysine-to-alanine substitution at the 590th amino acid (K590A) is not ubiquitinated under high-B conditions, indicating that the K590 residue is essential for high-B-induced ubiquitination (Kasai et al., 2011). However, no evidence of direct ubiquitination of the K590 has been shown. Hence, we initially attempted to investigate whether the K590 is ubiquitinated in response to a high-B treatment. To rule out the possibility that the electric charge alteration by the K590A mutation caused an abnormal conformation change of BOR1, we tested the vacuolar transport and ubiquitination of a BOR1 variant (K590R), which possesses the same positive charge as lysine. Similar to the previous results with K590A, time-lapse imaging of root epidermal cells after high-B supply (100 μM boric acid) showed that BOR1(K590R)-GFP remained in the plasma membrane for 120 min, while wild-type (WT) BOR1-GFP was rapidly internalized and degraded (Figure 1A and B). This result eliminates the possibility that the observed change of trafficking in the K590A mutant is due to the uncharged alanine.

Next, to examine whether the K590 residue is directly ubiquitinated, we performed liquid chromatography-tandem mass spectrometry (LC-MS/MS) of chymotrypsin-digested peptides of BOR1-GFP. The total protein was extracted from transgenic plants harboring proSS5:BOR1-GFP and the BOR1-GFP protein was enriched by immunoprecipitation (IP) using an anti-GFP antibody (Supplemental Figure 1). The footprint of ubiquitin (RGG, Arg-Gly-Gly) was detected at the K590 residue of the peptide fragment [590K(RGG)GAHLQDLDAAEY] (Figure 1C; Supplemental Figure 1B).

To analyze the ubiquitination status of WT and K590R-substituted BOR1-GFP, we conducted IP of BOR1-GFP and detected ubiquitin-conjugated proteins using anti-ubiquitin antibodies (Figure 1D; Supplemental Figure 2 and 3). Immunoblotting with an antibody against ubiquitin, P4D1, showed ladder-like signals from ~100 kDa to ~200 kDa above the position of WT BOR1-GFP (98 kDa) detected by an anti-GFP antibody. The signals detected by the P4D1 antibody in lanes with WT BOR1-GFP increased in response to high-B treatment (Figure 1D, Lane 4; Supplemental Figures 2 and 3) and did not appear in the corresponding lanes with Col-0 control and BOR1(K590R)-GFP (Figure 1D, Lane 1, 2, 5, and 6, respectively; Supplemental Figures 2 and 3). Importantly, immunoblotting with a chain type-specific antibody against K63-linked ubiquitination, Apu3, also showed the ladder-like signals above the position of WT BOR1-GFP in the sample treated with high-B (Figure 1D, Lane 4). We also tested whether BOR1 undergoes K48-linked ubiquitination, which is known to act in proteasome-mediated proteolysis, using a chain type-specific antibody Apu2. However, we did not detect any positive signals, which would have been indicative of K48-linked ubiquitination of BOR1 (Supplemental Figure 3). These data suggest that BOR1 undergoes K63-linked polyubiquitination in response to high-B treatments. We conclude that BOR1 undergoes ubiquitination at the K590 residue and that high-B concentrations induce K63-linked polyubiquitination.

Genetic screening identifies amino acid residues within BOR1 required for vacuolar transport

To identify the “boron sensor/receptor” that controls BOR1 ubiquitination in response to the elevation of boric acid
concentrations, we performed a forward genetic screen using an Arabidopsis thaliana transgenic line harboring pro35S:BOR1-GFP(-50UTR), which constitutively expresses BOR1-GFP without translational repression under high-B conditions (Aibara et al., 2018). When plants were grown on high-B medium (500 μM boric acid), BOR1-GFP was degraded, and thus showed weaker fluorescence than on low-B (0.5 μM) medium (Figure 2A). We screened for mutants in an ethyl-methane sulfonate (EMS) treated M2 population showing strong fluorescence of BOR1-GFP under the high-B condition (Figure 2A). Through the screen, we isolated 44 candidates showing a bright fluorescence of BOR1-GFP under high-B conditions from the pool of 70,000 M2 seedlings (Figure 2B).

Among the 44 candidates, 7 showed severe developmental defects and did not produce progenies. We then sequenced the BOR1-GFP transgene in the 37 candidates and identified 7 missense mutations within the protein-coding sequence. These BOR1 variants can be separated into two groups in terms of their subcellular localization, a plasma membrane localization-type (PM-type; A315V, G356S, P359S, and P362S) and an intracellular-localization type (G201R, V250F, and S251F). V250F and S251F were isolated repeatedly from distinct M2 pools, indicating that the screen reached near saturation. G201 is located in the transmembrane domain (TMD) 6, V250, and S251 are located in the TMD 7, A315 is located in the TMD 8, and G356, P358, and P362 are located in the TMD 10 or its vicinity (Figure 2C). It should be noted that the K590 residue, which is encoded as AAA, is theoretically not affected by EMS, which frequently produces G to A (C to T) base pair changes through N-7 alkylation of guanine rather than other nucleotides (Sega, 1984).
In contrast to WT, the intracellular-type BOR1-GFP variants colocalized with intracellular compartments stained by ER-Tracker Red in the lateral root cap cells (Supplemental Figure 4A) and showed a network pattern in cotyledon epidermal cells (Supplemental Figure 4B). These results suggest that G201R, V250F, and S251F substitutions abolished proper folding of BOR1-GFP and consequently disturbed the exit of BOR1-GFP from the endoplasmic reticulum (ER; Ellgaard and Helenius, 2003). Furthermore, these variants were not degraded under a high-B condition (Supplemental Figure 4C), indicating that initiation of B-induced vacuolar transport of BOR1 does not take place at the ER membrane or requires properly folded BOR1 protein.

Next, to characterize the PM-type BOR1-GFP variants, we grew the mutant seedlings under two different B concentrations and observed their subcellular localization and fluorescence intensity in primary root tips (Figure 2D). WT BOR1-GFP showed a decrease of the fluorescence intensity under a high-B condition (500 μM boric acid; Figure 2D). WT BOR1-GFP was faintly observed under a high-B condition, while the A315V, G356S, P359S, and P362S mutants still showed a clear localization in the plasma membrane (Figure 2D).
Intriguingly, all of the substitutions are not in the proximity of the K590 ubiquitination site but located in the TMDs.

**Substitutions of amino acids located within the substrate-binding pocket of BOR1-affected degradation and B-transport activity**

Next, we investigated the roles of the amino acid residues A315, G356, P359, and P362 in the B-transport function. These four amino acid residues are located in the putative substrate-binding pocket of BOR1, as indicated in a three-dimensional model reported in previous studies (Figures 3A and 3B; Coudray et al., 2016; Thurtle-Schmidt and Stroud, 2016). A multiple alignment indicated that G356, P359, and P362 are highly conserved in BOR1 homologs in plants, protists, and fungi (Figure 3C; Supplemental Figure 5). By contrast, A315 is conserved in clade I-BORs including BOR1 orthologs, but is less conserved in clade II-BORs including AtBOR4, which functions in B exclusion at high-B conditions (Miwa et al., 2007; Wakuta et al., 2015), and it is not conserved in the homologs in mosses (clade III-BORs), protists, and fungi (Figure 3C; Supplemental Figure 5; Supplemental Data set 1; Supplemental File 1). These substitutions in the vicinity of the substrate-binding pocket raise the question whether the degradation defect of PM-type variants is linked to the transport activity of BOR1.

To examine the involvement of these residues in borate transport activity, we introduced a series of BOR1 variants (A315V, G356A, P359A, and P362A) fused with Xpress-tag into the yeast (Saccharomyces cerevisiae) strain, that has a significantly reduced B-export capacity (Takano et al., 2007). In this and following experiments, we substituted G356, P359, and P362 to A to keep the nonpolar nature of the amino acids, although the substitutions we identified in the screening were S substitutions. The expression of functional BOR1 confers viability to yeast cells under toxic-B conditions by reducing B-concentrations in the cells (Takano et al., 2007). To test this hypothesis, we sought additional amino acid residues that would directly play a role in B-transport. A previous report identified that the D347, N391, and Q396 amino acid residues, which surround the substrate-binding pocket of S. cerevisiae Bor1 (ScBor1), are important for its B-transport activity (Thurtle-Schmidt and Stroud, 2016). The D347 residue corresponds to E241 of the bacterial uracil transporter UraA and the E681 residue of human Band3/anion exchanger 1 (AE1; Supplemental Figure 7). The E681 residue of Band3/AE1 is considered to be the proton-binding site, which is essential for its transport activity (Reithmeier et al., 2016). The B-tolerance assay showed that the D347A substitution completely inhibited the B-transport activity of ScBor1, while N391A and Q396A partially inhibited it (Thurtle-Schmidt and Stroud, 2016). The D347 residue of Band3/AE1 is considered to be the proton-binding site, which is essential for its transport activity (Reithmeier et al., 2016). The B-tolerance assay showed that the D347A substitution completely inhibited the B-transport activity of ScBor1, while N391A and Q396A partially inhibited it (Thurtle-Schmidt and Stroud, 2016). Because D347, N391, and Q396 of ScBor1 correspond to D311, N355, and Q360 of AtBOR1, respectively, their substitutions to alanine residues Mup1, Can1, Gap1, and uracil transporter Fur4 indicated that substrate-binding to the transport site induces a conformational transition and promotes their ubiquitination (Guiney et al., 2016; Gournas et al., 2017). Based on this regulatory model of yeast transporters, we hypothesized that substrate binding and conformational transition of BOR1 are involved in B-induced ubiquitination of K590.

To test this hypothesis, we sought additional amino acid residues that would directly play a role in B-transport. A previous report identified that the D347, N391, and Q396 amino acid residues, which surround the substrate-binding pocket of S. cerevisiae Bor1 (ScBor1), are important for its B-transport activity (Thurtle-Schmidt and Stroud, 2016). The D347 residue corresponds to E241 of the bacterial uracil transporter UraA and the E681 residue of human Band3/anion exchanger 1 (AE1; Supplemental Figure 7). The E681 residue of Band3/AE1 is considered to be the proton-binding site, which is essential for its transport activity (Reithmeier et al., 2016). The B-tolerance assay showed that the D347A substitution completely inhibited the B-transport activity of ScBor1, while N391A and Q396A partially inhibited it (Thurtle-Schmidt and Stroud, 2016). Because D347, N391, and Q396 of ScBor1 correspond to D311, N355, and Q360 of AtBOR1, respectively, their substitutions to alanine were expected to disturb the B-transport activity of BOR1 (Figure 4A; Supplemental Figure 7).

To investigate the relationship between the transport activity and ubiquitination of BOR1, we generated AtBOR1 variants carrying D311A, N355A, and Q360A substitutions and assessed their B-transport activity in yeast (Figure 4B and C). Immunoblotting and immunostaining against the Xpress-tag showed comparable expression levels and PM localization of the BOR1 variants in yeast cells (Supplemental Figures 6E–G). The B-tolerance assay showed that D311A completely inhibited the transport activity of BOR1, whereas N355A and Q360A did not or only slightly inhibited it (Figure 4B). Consistent with these results, the B-export assay showed that the D311A mutation completely inhibited the B-transport activity, whereas N355A and Q360A slightly...
inhibited it (Figure 4B and C, \( P < 0.05 \) by one-way ANOVA followed by Tukey–Kramer’s post hoc test).

**Transport activity is necessary for high B-induced ubiquitination of BOR1**

To investigate the contribution of amino acid residues located in the substrate-binding pocket on B transport and degradation of BOR1 in *Arabidopsis thaliana*, we introduced *proBOR1:BOR1-GFP* with D311A, N355A, and Q360A substitutions into *Arabidopsis thaliana*. We used a *bor1-3 bor2-1* double knockout mutant as background to avoid possible interaction between BOR1-GFP variants and endogenous BOR2 (Miwa et al., 2013). We also introduced *proBOR1:BOR1-GFP* with A315V, G356A, P359A, and P362A substitutions to validate the importance of the amino acid residues identified by the genetic screening using *pro35S:BOR1-GFP* constructs. All of the BOR1-GFP variants localized in the plasma membrane in a polar fashion toward
with liquid medium containing 1 mM of boric acid for 50 min at 30°C. Plant growth of the bor1-3 bor2-1 double mutants under a low-B condition was fully restored by introduction of WT BOR1-GFP (Figure 5D and E). It is likely that the BOR1 variants partially defective in B transport complemented the shoot growth due to a relatively higher expression in transgenic plants. The G356A variants, which showed no or slightly reduced B-transport activity to WT in yeast cells (Figure 3D and E), fully restored the shoot and root growth (Figure 5D and E). The P362A variants, which showed no apparent reduction in the tolerance of yeast cells at a high B condition (Figure 3D and E), restored the shoot growth fully, but root growth partially (Figure 5D and E). The A315V variants, which showed similar B-transport activity to WT in yeast cells, fully restored the shoot and the root growth in a line (#8) but not fully in the other line (#7; Figure 5D and E). Taken together, our analyses in yeast and plants suggest that D311 and P359 are essential, N355 and Q360 are involved, and A315, G356, and P362 are less or not important for B-transport activity. The defective complementation of the root growth by the transport-competent A315V and P362A variants might be related to their reduced polar localization (Figure 5B).

To address the relationship between B-transport activity and B-induced ubiquitination of BOR1, we examined whether the BOR1-GFP variants undergo polyubiquitination after high-B supply in planta using antibodies against general ubiquitin (P4D1) and K63-linked polyubiquitination (Apu3; Figure 6A; Supplemental Figures 9 and 10). The variants A315V, N355A, G356A, Q360A, and P362A showed reduced rates of polyubiquitination upon high-B supply compared to WT; 22 ± 10%, 44 ± 1.7%, 18 ± 4.2%, 61 ± 31%, and 18 ± 7.4% of WT, respectively (P < 0.05 vs. WT by one-way ANOVA).
Figure 5 Boron-transport activity of BOR1 variants in plants. (A) Representative confocal images of BOR1-GFP variants in the primary root tip of Arabidopsis seedlings grown on 0.5 μM B MGRL medium for 5 days. Scale bars represent 50 μm. (B) Box plot representation of polarity indexes of BOR1-GFP variants. Letters above box plots indicate significant differences determined by the nonparametric Kruskal–Wallis test (*P < 0.05, ***P < 0.001, and ****P < 0.0001, n.s. means no significant difference). n = 49–87 cells from 3 to 5 different roots. Box plot center lines show the medians. Box limits indicate the 25th and 75th percentiles. Whiskers are extended to the highest and the lowest values. (C) Fluorescence intensity of BOR1-GFP. The total fluorescence of BOR1-GFP in the root tip was obtained by taking a Z-stack in 2 μm intervals over a total distance of 80 μm for each individual root by a 20× dry objective lens equipped with LSM800 (Zeiss). Error bars represent mean ± SD. n = 4–8 plants. (D and E) Leaf fresh weight (D) and root length (E) values relative to WT Col-0 plants grown in the same plate. Error bars represent mean ± SD. Significant differences were determined by Student’s t-test (*P < 0.05, **P < 0.01, ***P < 0.001). n = 3 sets of plates.
with Tukey–Kramer’s post hoc test), while D311A and P359A did not show such ladder-like signals; 2.8 ± 2.9% and 3.2 ± 2.7% of WT, respectively (Figure 6B).

We also tested whether the BOR1-GFP variants are degraded in response to high-B supply (Figure 6C–E; Supplemental Figure 11). The vacuolar sorting of BOR1-GFP...
was examined using the fact that GFP is relatively stable in the vacuole in the absence of light (Tamura et al., 2003; Takano et al., 2010). When the plants were treated with 100 μM B for 2 h in the dark, the WT and Q360A variant showed strong GFP fluorescence in the vacuole, while the N355A and P362A variants showed GFP fluorescence in punctate structures, presumably endosomes, and faint GFP fluorescence in the vacuole (Figure 6C). Additionally, we performed time course analysis under a normal light condition and quantified total GFP fluorescence in primary root tips (Figure 6D and E; Supplemental Figure 11). Within 60 min after 100 μM B supply, WT BOR1-GFP and N355A and Q360A variants showed a clear serial decrease of fluorescence, while D311A, A315V, G356A, P359A, and P362A showed relatively stable fluorescence in the plasma membrane (Figure 6D). A quantitation revealed that 19.3 ± 1.1% of WT BOR1-GFP fluorescence remained after 3 h (Figure 6E and Supplemental Figure 11). Within 60 min after 100 μM B supply, WT BOR1-GFP and N355A and Q360A variants showed a clear serial decrease of fluorescence, while D311A, A315V, G356A, P359A, and P362A showed relatively stable fluorescence in the plasma membrane (Figure 6D). A quantitation revealed that 19.3 ± 1.1% of WT BOR1-GFP fluorescence remained after 3 h (Figure 6E and Supplemental Figure 11). Q360A showed a fluorescence decrease rate comparable to WT; 25.3 ± 2.0% at 3 h (P = 0.97 vs. WT by two-way ANOVA with Tukey–Kramer’s post hoc test). G356A, P359A, P362A, and N355A showed significantly slower rates of decrease than WT BOR1; 71.7 ± 7.8%, 76.3 ± 35%, 74.5 ± 11%, and 53.4 ± 10% at 3 h, respectively (P < 0.0001 vs. WT by two-way ANOVA with Tukey–Kramer’s post hoc test). Strikingly, D311A and A315V variants did not show a decrease within 3 h (P > 0.75 between time points by two-way ANOVA with Tukey–Kramer’s post hoc test).

The relationship between the B-dependent polyubiquitination, degradation, and the B-transport activity in yeast cells was analyzed based on their quantified values among the BOR1 variants (Figure 6F and G; Supplemental Table 1). As expected, the rates of B-dependent polyubiquitination and degradation showed a positive correlation, confirming that the polyubiquitination leads to vacuolar transport and degradation of BOR1 (Figure 6F). As an exception, the P359A variants showed degradation at a slightly smaller scale than WT, but no apparent ubiquitination could be observed. It is possible that ubiquitinated BOR1-GFP proteins are rapidly subjected to vacuolar transport and degradation, and the minor amount of the ubiquitinated P359A variant might be below the detection limit.

We then compared the B-transport activity in yeast cells and the rate of degradation (Figure 6G). The D311A variant was completely inactive in B-transport and not transported to the vacuole. The D311A, P359A, N355A, and Q360A variants and WT showed a positive correlation between the B-transport activity and the rate of degradation. These results support the hypothesis that substrate binding and the B-transport cycle of BOR1 are linked to B-induced ubiquitination. It should be noted that the A315V, G356A, and P362A variants were transport competent, but fully or partially defective in ubiquitination and degradation. These variants might have defects in specific conformations (see “Discussion” section).

To rule out the possibility that the transport-defective variants do not show degradation because of the aberrant B concentrations inside or outside the cells, BOR1(D311A)-GFP and BOR1(WT)-mCherry were expressed in the same plants. In a control experiment, BOR1(WT)-GFP and BOR1(WT)-mCherry similarly showed endocytosis and degradation in the same epidermal cells within 90 min after 100 μM B supply (Figure 7A–C). By contrast, BOR1(D311A)-GFP was retained in the plasma membrane in cells where BOR1(WT)-mCherry was mostly transferred into endosomes (Figure 7A and B). BOR1(WT)-mCherry remained to a slightly higher extent in the cells with BOR1(D311A)-GFP than in those with BOR1(WT)-GFP (Figure 7B). This could be because of dimerization with each other. A quantification of the colocalization by Pearson’s correlation coefficient confirmed that BOR1(WT)-mCherry behaved similar to BOR1(WT)-GFP but differently compared to BOR1(D311A)-GFP (Figure 7C). These results suggest that B-transport activity rather than the B concentration inside or outside the cells controls B-induced endocytosis and degradation of BOR1.

The K590 residue is located closely to the last TM domain (Figure 2C) and its accessibility by an unknown ubiquitin ligase is expected to be influenced by conformational transition of BOR1 during B-transport. To test the importance of the position of the K590 residue, we generated BOR1-GFP constructs whose C-terminal regions including K590 were tandemly repeated (Supplemental Figure 12). If the ubiquitination of the K590 residue is independent of its position, the distal lysine residue in the extended C-terminal tail would be ubiquitinated in response to high-B supply and lead BOR1 to the vacuole. For convenience, the proximal (original) K590 residue is termed proximal K and the K590 residue in the repeated sequence is termed distal K. The C-terminal extended BOR1-GFP with proximal K (WT) was localized in the plasma membrane under the low-B condition and degraded in response to high-B supply regardless of the distal K presence (Supplemental Figure 12B and C; WT/WT, WT/K590R). The C-terminal extended BOR1-GFP with proximal K590R was localized in the plasma membrane under the low-B condition and it was not degraded in response to high-B supply regardless of the distal K presence (Supplemental Figure 12B and C; K590R/WT, K590R/K590R). Although the structure around the distal K is unclear, these data support the importance of the position of the ubiquitination site for B-induced degradation of BOR1. It is likely that conformational transition of BOR1 during B transport affects the local structure around the K590 residue that is close to the TM domain and provides a suitable environment for recognition by a ubiquitin ligease.

Discussion

Several plant transporters have been shown, or proposed, to have receptor functions and thus are called transceptors. For instance, the Arabidopsis dual-affinity nitrate transporter CHL1/NRT1.1 functions in nitrate sensing, which controls
the induction of nitrate-responsive genes including a high-affinity nitrate transporter NRT2.1 (Ho et al., 2009); the nitrate sensing and transport functions of CHL1/NRT1.1 can be decoupled by a mutation. In addition, the Arabidopsis iron transporter IRT1, which transports various metals and is degraded upon noniron metal stress, contains a histidine-rich stretch in the cytosolic loop region that directly binds noniron metals, is phosphorylated by the CIPK23 kinase, and subsequently is ubiquitinated by IDF1, a RING-type E3 ubiquitin-ligase (Dubeaux et al., 2018). Substitution of four histidines to alanines (4HA) in the histidine-rich stretch does not affect IRT1 transport activity but diminishes its phosphorylation and ubiquitination. Importantly, the expression of IRT1(4HA) made plants hypersensitive to noniron metal excess. These results indicate that the metal-binding site of IRT1 regulates its own degradation (Figure 8). We, therefore, propose that BOR1 is a transceptor for the maintenance of B homeostasis with a different sensing mechanism from known plant transceptors.

In fungi, many nutrient transporters are downregulated via ubiquitination, subsequent endocytosis, and degradation in the vacuole under substrate supply at high concentrations (Babst, 2020). In budding yeast S. cerevisiae, arrestin-related adaptors, Arts, recruit a ubiquitin ligase Rsp5 to specific transporters in the plasma membrane for ubiquitination (Lin et al., 2008). Amino acid permeases such as Gap1, Mup1, Fur4, and Can1 harbor degrons in their cytosolic tail regions (Keener and Babst, 2013; Ghaddar et al., 2014; Guiney et al., 2016). It was proposed that the specific degron, where Arts and Rsp5 are recruited to, is masked at a ground state (state under a low substrate concentration), while it gets exposed at an activated state (state under high substrate concentration) due to a conformational transition accompanied with substrate transport (Guiney et al., 2016). BOR1 is distantly related to the amino acid permeases in the same amino acid-polyamine organocation (APC) superfamily, and they share an overall architecture (Vastermark et al., 2014). Therefore, it is reasonable that similar
mechanisms of transport-coupled ubiquitination operate in the downregulation of yeast amino acid permeases and plant BOR1.

In this study, we first addressed the characteristics of BOR1 ubiquitination and observed that BOR1 undergoes K63-linked polyubiquitination in a B-dependent manner. Lysine 63-linked polyubiquitination is the second most abundant type of polyubiquitination and functions in a variety of cellular processes in eukaryotes (Liu et al., 2018). In plant cells, abundance of membrane proteins in the plasma membrane such as IRT1, the auxin transporter PIN2, and the brassinosteroid receptor BRI1 are controlled by accelerated endocytosis and/or vacuolar transport dependent on K63-linked polyubiquitination (Leitner et al., 2012; Martins et al., 2015; Dubeaux et al., 2018). In BOR1 protein, the K590 residue located in the cytosolic C-tail region is essential for B-dependent ubiquitination and vacuolar transport, while there was no evidence of direct ubiquitination of the residue (Kasai et al., 2011). Although BOR1 was described to undergo mono- or diubiquitination in the previous study (Kasai et al., 2011), we detected ladder-like signals recognized by a specific antibody against K63-linked ubiquitination (Figure 1D). Furthermore, we demonstrated that the K590 residue is directly modified with ubiquitin by MS analysis (Figure 1C). These results suggest that BOR1 undergoes K63-linked polyubiquitination at the K590 residue. Recently, we demonstrated that the rapid endocytosis of BOR1 for high B-induced vacuolar transport is independent of the AP-2 clathrin adaptor complex (Yoshinari et al., 2019). These findings imply that an unknown ubiquitin-binding endocytic adaptor binds to K63-linked polyubiquitin attached to BOR1. A recent study proposed that TOM1-LIKE6 (TOL6), a plasma membrane-associated ubiquitin receptor, and its paralogs act as K63-linked ubiquitin adaptors facilitating vacuolar transport of plasma membrane proteins (Moulinier-Anzola et al., 2020). Indeed, vacuolar transport of BOR1 is compromised in a higher order mutant of TOLs (tol23569; Yoshinari et al., 2018). TOL proteins associated with the plasma membrane might recognize BOR1 with K63-linked polyubiquitin for vacuolar transport.

Recently, crystal structural analyses revealed the three-dimensional structure of BOR1 in an occluded conformation at a 4.1 Å resolution and ScBor1p in an inward-open conformation at a 5.9 Å resolution (Coudray et al., 2016; Thurtle-Schmidt and Stroud, 2016). The structure of BOR1 can be distinguished into two domains, the “gate domain” and “core domain”. The gate domain is regarded as a rigid scaffold and is required for oligomerization, while the core domain moves to achieve the rocking bundle (Coudray et al., 2016) or the elevator transport mechanism (Thurtle-Schmidt and Stroud, 2016). In the core domain, the borate anion is thought to be stabilized in the space between the positive dipoles, TMD 3 and 10 (Figure 4A; Supplemental Figure 7). Our genetic screen using BOR1-GFP identified G201, V250, and S251 residues located in the gate domain as important residues for ER exit (Supplemental Figure 4). These amino acid residues are probably essential for proper protein folding or structural stability of BOR1. In the same genetic screen and targeted mutagenesis, we determined that mutations at A315, D311, N355, G356, P359, Q360, and P362 disturbed ubiquitination and vacuolar transport of BOR1 to different degrees (Figures 2–6). These amino acid residues are located around the N-terminus of TMD 10 which composes the putative borate-binding pocket in the core domain (Figure 4A; Supplemental Figure 7). Intriguingly, a substitution of the putative proton-binding amino acid residue D311 in the TMD 8 completely abolished transport activity, ubiquitination, and vacuolar transport of BOR1 (Figures 4–6). The transport-deficient D311A variant was not transported to the vacuole when B transport was secured by the WT BOR1 expressed in the same cells (Figure 7). Furthermore, a positive correlation was observed between B-transport activity, ubiquitination, and degradation among the transport-defective D311A, P359A, N355A, and Q360A variants and WT (Figure 6F and G; Supplemental Table 1). By analogy to yeast amino acid permeases, these results suggest that the degradation of BOR1 is via transport-coupled ubiquitination (Figure 8). When the extracellular concentration of borate/boric acid is relatively low, the transport cycle would proceed as follows: (1) The
borate anion accesses the substrate-binding site during the inward-open state, (2) the conformation of BOR1 changes to the occluded state and then to the outward-open state, (3) the borate anion is released from the substrate-binding pocket and the conformation immediately returns to the inward-open state. However, when the extracellular concentration of borate/boric acid is relatively high, the borate anion would remain longer in the substrate-binding pocket to keep the outward-open state. We speculate that an unknown E3 ubiquitin ligase can access and conjugate the polyubiquitin chain to the K590 residue, which is exposed to the cytosol only in the outward-open state (Figure 8).

In our mutant analysis, the A315V, G356A, and P362A mutations did not strongly affect the B-transport activity in yeast, but inhibited ubiquitination and vacuolar transport of BOR1 (Figures 3 and 6). Presumably, these mutations affected specific structures of BOR1 that are involved in the ubiquitination but play only a minor role in B-transport. According to an in silico prediction, the A315V substitution leads to the formation of an aberrant hydrogen bond between the valine residue and S361 (Supplemental Figure 13). This abnormal hydrogen bond may affect the conformation of BOR1 in the outward-open state without inhibiting the B-transport activity. We speculate that valine at this position might add a specific physiological function to certain BOR1 homologs, because the sequence is observed in AtBOR5 and plant orthologs (Figure 3C; Supplemental Figure 5). Interestingly, D311A, A315V, N355A P359A, and P362A mutations affected polar localization of BOR1 toward the stele in addition to ubiquitination (Figures 5B). These mutations might affect the structure of the later C-tail region (637–704 aa), which is recognized by the AP-2 clathrin adaptor complex and is required for polar localization of BOR1 (Kasai et al., 2011; Yoshinari et al., 2019), in addition to the region proximal to the last TMD. Further analysis is required to understand the structural impact of the mutations on the cytosolic C-tail movement during the transport cycle.

In conclusion, our study revealed that the ubiquitination and degradation of the BOR1 transporter are coupled to the borate transport cycle. We propose that BOR1 is a B receptor and is capable of rapid self-regulation. This transport-coupled ubiquitination model for nutrient receptors appears to be conserved in organisms including plants and fungi to cope with a sudden increase of nutrient availability.

Materials and methods

Plant materials

To construct 35S promoter:BOR1-GFP, the BOR1-GFP sequence in a pAT30 entry clone (Takano et al., 2010) was subcloned into pMDC32 (Curtis and Grossniklaus, 2003), a gateway destination vector containing a dual 35S promoter, by LR recombination (Invitrogen). The resulting construct was introduced into Arabidopsis thaliana Col-0 using the Agrobacterium-mediated floral dip method (Clough and Bent, 1998). Mutant constructs carrying A315V, G356A, P359A, P362A, D311A, N355A, Q360A, or K590R were generated by site-directed mutagenesis into pAT30 (BOR1-GFP in pENTR D-TOPO) using primers listed in Supplemental Table 2 and subsequent LR recombination with pAT100, a gateway destination vector containing proBOR1 (Takano et al., 2010). The proBOR1:BOR1-GFP construct (Takano et al., 2010) and its mutant constructs were introduced into the Arabidopsis thaliana bor1-3 bor2-1 double knockout mutant (Kasai et al., 2011). For construction of the C-terminally extended BOR1-GFP, DNA fragments corresponding to the BOR1 C-terminal region were amplified from entry vectors containing WT and K590-mutated BOR1-GFP (BOR1[WT or K590]::GFP in pENTR D-TOPO), and the entry vectors were linearized by inverse PCR using primers listed in Supplemental Table 2. The DNA fragments for the C-terminal region were integrated into the linearized BOR1-GFP entry vectors by the In-Fusion technique (Clontech). For observation of BOR1(WT/D311A)-GFP and BOR1(WT)-mCherry in the same cells, the F1 generation plants of a cross between BOR1(WT/D311A)-GFP/bor1-3 bor2-1 and BOR1-mcherry/bor1-1 (Miwa et al., 2013) were used. Plant materials used in this study were also listed in Supplemental Data Set 2.

Plant growth conditions

Seeds were surface-sterilized with 70% (v/v) ethanol, washed with sterile water four times, and sown on modified MGRL medium (Takano et al., 2005) solidified with gellan gum (Wako Pure Chemicals, Osaka, Japan) in plastic plates. The plates were incubated at 4°C for 3 days and placed vertically in the plant growth chamber at 22°C with 16 h at a light intensity of ~120 μmol m–2 s–1 (emitted by white fluorescent lamps) and 8 h dark. The concentration of boric acid in the media is described in each figure legend.

Confocal microscopy and quantification of fluorescence of plants

Confocal images of BOR1-GFP expressed in roots were with confocal laser scanning microscopes, Leica TCS-SP8, Zeiss LSM510, or Zeiss LSM800, equipped with a 40× water-immersion lens or a 63× oil-immersion lens. GFP was excited with a 488 nm wavelength by a diode or argon laser and detected with a 500–530 nm (SP8) or 505–550 nm (LSM800) window of wavelength. For visualization of the ER, seedlings were incubated with 4 μM ER Tracker Red (Invitrogen, CA) for 5 min. ER Tracker Red was excited with a 561 nm wavelength of a diode laser and detected with a 560–600 nm window of wavelength (SP8).

For the time course analysis of BOR1-GFP WT and K590R (Figure 1), seedlings were placed in a cover glass bottom chamber (D11131H, Matsunami, Osaka, Japan), and the roots were covered with MGRL medium containing 1% (w/v) gellan gum and 100 μM boric acid. Images were taken at the indicated time points using a TCS SP-8 system (Leica). The fluorescence of BOR1-GFP in the plasma membrane was quantified by Fiji/ImageJ software as described previously (Yoshinari and Takano, 2020).
For the time course analysis of BOR1-GFP variants harboring amino acid substitutions in the vicinity of the substrate-binding pocket (Figure 6) and C-terminally extended BOR1-GFP variants (Supplemental Figure 12), seedlings were grown for 5 days on modified MGRL medium with 0.5 μM B and then floated on liquid MGRL medium containing 100 μM boric acid for 0, 60, and 180 min. The total fluorescence of BOR1-GFP in the root tip was obtained by taking a Z-stack in 2 μm intervals over a total distance of 80 or 100 μm for each individual root with an LSM800 (Zeiss) using a 20× dry objective lens or a TCS-SP8 (Leica) using a 40× water-immersion objective lens. The fluorescence of BOR1-GFP in the sum image of the root tip region (see Supplemental Figure 11) was quantified by Fiji/ImageJ software. Comparison of expression levels of BOR1-GFP variants under a low-B condition (Figure 5A) was performed in a similar way.

The polarity index was calculated as described previously (Yoshinari and Takano, 2020). Briefly, confocal images of BOR1-GFP in the epidermal cells of the meristematic to transition zones in primary root tips were used for the calculation. Plasma membrane regions at the apical and basal domains were selected with 3-pixel-wide straight lines by Fiji/ImageJ software. The integrated density (total fluorescence) in the inner half of the plasma membrane was divided by that in the outer half of each cell.

For the analysis of high B-induced degradation of BOR1-mCherry and BOR1 (WT/D311A)-GFP (Figure 7), seedlings were grown for 5 days on modified MGRL medium with 0.5 μM B and then floated on liquid MGRL medium containing 100 μM boric acid for 2 or 90 min. The fluorescence of mCherry and GFP in the plasma membrane was observed with a TCS-SP8 (Leica) and quantified as described previously (Yoshinari and Takano, 2020). Pearson's correlation coefficient was calculated using Fiji/ImageJ with the PSC Colocalization plugin (French et al., 2008).

**Mutant screening**

For the mutant screening, approximately 12,500 T4 seeds of pro35S:bor1-gfp were mutagenized with 0.3% ethylmethanesulfonate (EMS) overnight and two pools of M2 seeds were obtained. M2 seedlings were grown on solid medium containing 500 μM B for 7 days. GFP images were visualized by the macro zoom microscope MVX-10 (Olympus) with a CCD camera CoolSNAP EZ (Photometrics).

**LC-MS/MS analysis of BOR1-GFP protein**

Total proteins were extracted from 22-day-old transgenic plants harboring pro35S:bor1-gfp (5′UTR) grown with 0.5 μM B. The homogenization was carried out in buffer (250 mM Tris- HCl [pH 8.5], 290 mM sucrose, 1 mM Pefabloc<sup>®</sup> SC [Roche], 1.25% [v/v] PSC-protector solution [Roche], complete ULTRA Mini EDTA-free [Roche], 5 mM DTT) with a multibeads shocker (Yasui Kikai, Osaka, Japan). Cell debris was removed by centrifugation at 10,000g at 4°C for 15 min three times. The microsome fraction was isolated from the supernatant by ultracentrifugation (100,000 g, 10 min, 4°C) and resuspended in lysis buffer (150 mM NaCl, 1% [v/v] Triton X-100, 50 mM Tris-HCl [pH 8.0], 1% [v/v] 3-[(3-cholamidopropyl)dimethylammonio]-1-propanesulfonate [CHAPS]) using a Potter-Elekhjem tissue grinder. BOR1-GFP was isolated by μMACSTM anti-GFP microbeads (Miltenyi Biotec) according to the manufacturer’s instructions. BOR1-GFP protein was collected together with the microbeads in the lysis buffer by dissociating the column from the μMACSTM Separator magnet. The protein–microbead suspension was incubated with the lysis buffer containing 230 μM B on ice for 30 min right before SDS-PAGE. Immunoprecipitated proteins were separated by NuPAGE 4%–12% Bis-Tris Gel (Invitrogen) and stained with Flamingo fluorescent gel stain solution (Bio-Rad). The band of the BOR1-GFP monomer was cut and placed into acetonitrile. The gel was dried in a vacuum concentrator and treated with a reducing solution (10 mM DTT, 50 mM ammonium bicarbonate) at 56°C for 15 min. After supernatant removal, the gel was treated with alklylation solution (55 mM 2-iodoacetamide, 50 mM ammonium bicarbonate) for 30 min. The supernatant was removed again, and the gel was washed with 50 mM ammonium bicarbonate three times and dried up in a vacuum concentrator. After drying, the gel was treated with an enzyme solution (10 μg/mL chymotrypsin sequencing grade [Roche], 100 mM Tris-HCl [pH 7.9], 10 mM CaCl<sub>2</sub>) overnight at 25°C. The supernatant was collected and concentrated by vacuum centrifugation. To remove debris, the protein solution was filtered with an Ultra free-MC 0.45 μm Filter unit (Merck Millipore). Modification of BOR1-GFP was detected by Thermo Scientific Q Exactive Plus Orbitrap LC-MS/MS (Thermo Scientific). The MS/MS spectra were analyzed by Thermo Scientific Proteome Discoverer with MASCOT (Matrix Science, MA) and Sequent HT (Thermo Scientific) algorithms.

**Immunoblot analysis of ubiquitination**

For immunoprecipitation of BOR1-GFP, transgenic plants harboring probor1:bor1-gfp were used. Shoot tissues were removed, and root tissues were harvested. Root tissues (0.2–0.6 g) were lysed in 0.5–2.0 mL of extraction buffer (50 mM Tris-HCl pH7.5, 5 mM DTT, 100 mM NaCl, 10% [v/v] glycerol, 2% [v/v] IGEPIAL, 20 mM N-ethylmaleimide, complete mini EDTA-free [Roche], 0.25 mg/mL Pefabloc<sup>®</sup> SC) by using a mortar and a pestle or a multibeads shocker. Cell debris was removed by centrifugation at 10,000g at 4°C for 15 min three times. Anti-GFP IP was carried out following the manufacturer's instructions (Miltenyi Biotec) with slight changes. Immunoprecipitated proteins were eluted with 70 μL of preheated elution buffer (50 mM Tris-HCl pH 6.8, 1% [w/v] SDS, 0.005% [w/v] Bromophenol Blue, 10% [v/v] glycerol, and 100 mM DTT, 95°C). Immunoprecipitated proteins were separated by Bolt<sup>TM</sup> 4%–12% Bis-Tris SDS-PAGE gel (Invitrogen) or NuPAGE<sup>®</sup> 4%–12% Bis-Tris SDS-PAGE gel (Invitrogen) with MOPS buffer and then transferred to an Immobilon PVDF membrane. Anti-ubiquitin mouse monoclonal antibody, P4D1
(Cat#: sc-8017, Lot#: F2320 or sc-8017 HRP, Lot#: A0919 [Santa Cruz Biotechnology], 1:2,000 dilution), anti-GFP monoclonal antibody (Cat#: 4363-24, Lot#: M9R8752 [Nacalai Tesque, Kyoto], 1:2,000 dilution), anti-ubiquitin, K63 specific, Apu3 (Cat#: 05-1308, Lot#: 3137755 [Merck], 1:2,000 dilution), and anti-ubiquitin K48 specific, Apu2 (Cat#: 05-1307 [Merck], 1:1,000 dilution) were diluted in Can Get Signal Solution 1 (Toyobo, Osaka, Japan). Anti-mouse IgG HRP-conjugated antibody (Cat#:115-035-003, Lot#: 138339 [Jackson ImmunoResearch, PA], 1:200,000 dilution) and anti-rabbit IgG HRP-conjugated antibody (Cat#: 111-035-144, Lot#: M9M6770 [Jackson ImmunoResearch]) were diluted in Can Get Signal Solution 2 (Toyobo). For detection of luminescence, Luminata Forte Western HRP substrate (Millipore) was used.

For quantification of polyubiquitination rates, ladders in the range of 100–250 kDa for polyubiquitination; 98 kDa for nonubiquitinated BOR1-GFP signals, respectively. Mean intensities of the corresponding nonubiquitinated BOR1-GFP signals were defined as polyubiquitination and nonubiquitinated BOR1-GFP signals, respectively. Mean intensities of the area (100–250 kDa for polyubiquitination; 98 kDa for nonubiquitinated BOR1-GFP) were quantified by Fiji/ImageJ software (Schindelin et al., 2012). Background signals (from a nonloaded region) were subtracted before calculation. The mean intensity of the polyubiquitination was divided by the mean intensity of the corresponding nonubiquitinated BOR1-GFP.

**Boron transport assay in yeast**

The S. cerevisiae bor1Δ strain Y01169 (MATa his3 leu2 met15 ura3 YNL275W::kanMX4; Takano et al., 2007), which lacks endogenous B export activity was used. To construct the vector to express BOR1 in yeast, BOR1 CDS was amplified using a BOR1 cDNA as a template with specific primers (Supplemental Table 2). The cDNA was subcloned into the pYES2 NT/A vector (Invitrogen) digested by KpnI and XhoI using the In-Fusion technique (Clontech). Constructs carrying A315V, G356A, P359A, P362A, D311A, N355A, or Q360A mutations were generated by site-directed mutagenesis of pTS32 (BOR1 in pYES2 NT/A) using primers listed in Supplemental Table 2. To determine the B export activity, an empty vector control (pYES2 NT/A) or constructs with respective AtBOR1 variants (WT, D311A, A315V, N355A, G356A, P359A, and P362A) were introduced into the strain and selected on SD-Ura + Raffinose media at 30°C overnight with shaking. The 3 mL of fresh SD-Ura + Galactose liquid medium containing 100 μL of the culture media and incubated at 28°C overnight with shaking. For the high-B shift assay, the same amount of yeast cells expressing WT-BOR1 was incubated in SD-Ura + Galactose liquid medium containing 0, 10, or 20 mM boron acid for 1 h before the following procedure.

The 1 mL of yeast cells (OD<sub>600</sub> = 1.0) was transferred to 1.5 mL tubes and collected by centrifugation at 15,000 g for 1 min. After removal of the supernatant, cell pellets were suspended in 100 μL of ice-cold protease inhibitor mix (2 mM phenylmethylsulfonyl fluoride [PMSF], 2 μM leupeptin, 0.1 μM aprotinin, 1 μM pepstatin A, 1 μM benzamidine, 1 μM 1,10-phenanthroline, 1 μM 2-mercaptoethanol) was added to the cell suspension. After incubation for 10 min on ice, 50 μL of 50% trichloroacetic acid (TCA) solution was added, and cells were incubated for 10 min on ice. Samples were centrifuged at 15,000 g for 5 min at 4°C. After removal of supernatant, cell pellets were resuspended in 50 μL of sample buffer 1 (100 mM Tris-HCl [pH 6.8], 4 mM EDTA, 4% SDS, 20% glycerol, 0.02% bromophenol blue). Finally, 50 μL of sample buffer 2 (1 M Tris base, 2% β-mercaptoethanol) was added and heated at 98°C for 3 min.

Equivalent volumes of the samples were separated by SDS-PAGE using a 10% acryliclamide gel with Tris-glycine running buffer. Proteins were transferred to an Immobilon PVDF membrane (Merck Millipore) with Tris-glycine/methanol transfer buffer by a semi-dry blotting method. Anti-Xpress tag mouse monoclonal antibody (Invitrogen, 1:5,000 dilution) and anti-mouse IgG HRP-conjugated antibody (Jackson Immuno Research, 1:100,000 dilution) were used as primary and secondary antibodies, respectively. For
the dilution of antibodies, Can Get Signal solutions (Toyobo, Osaka, Japan) were used. Total proteins were stained with Coomassie Brilliant Blue (CBB).

**Yeast immunofluorescence analysis**

Yeast in the logarithmic growth phase in SD + Galactose liquid medium were fixed with 3.75% (w/v) paraformaldehyde for 30 min at 30°C with shaking. Cells were washed with 0.1 M potassium phosphate buffer (pH 7.5) and then with 1.2 M sorbitol/0.1 M potassium phosphate buffer (pH 7.5). To lyse the cell wall, cells were incubated with 2.5 mM DTT for 10 min at 30°C and then with 150 μg/mL zymolase (Nacalai Tesque) and 0.5% 2-mercaptoethanol for 10 min at 30°C. Cells were collected by centrifugation and washed with 0.1 M sorbitol/0.1 M potassium phosphate buffer (pH 7.5). The 50 μL of cell suspension was put onto a MAS-coated slide glass (Matsunami Glass, Osaka, Japan) and treated with ice-cold methanol for 6 min and then by ice-cold acetone for 30 min. The sample on the sliding glass was dried completely and stored at room temperature. Before immunostaining, cells were rehydrated with wash buffer (5 mg/mL bovine serum albumin, 1X PBS). As a primary antibody, mouse anti-Xpress tag antibody (Invitrogen, 1:1,000) was added and incubated overnight at 4°C. The primary antibody was removed and the sample was washed with 100 μL of wash buffer four times. As a secondary antibody, anti-mouse IgG antibody conjugated with CF568 (Biotium, 1:1,000) was added and incubated for 2 h at 37°C. Cells were washed with 100 μL of wash buffer four times and with 1X Phosphate-buffered saline (PBS) three times. Confocal imaging of yeast cells was performed with a Leica and with 1X Phosphate-buffered saline (PBS) three times. Cells were washed with 100 μL of wash buffer four times. As a secondary antibody, anti-mouse IgG antibody conjugated with CF568 (Biotium, 1:1,000) was added and incubated for 2 h at 37°C. Cells were washed with 100 μL of wash buffer four times and with 1X Phosphate-buffered saline (PBS) three times. Confocal imaging of yeast cells was performed with a Leica TCS-SP8 equipped with a 63× glycerol immersion lens. For excitation and detection of CF568 signals, 552 nm and 560–650 nm wavelengths were used, respectively.

**Multiple alignment and phylogenetic analysis**

A neighbor-joining phylogenetic tree of the BOR family based on protein sequences was constructed using a multiple sequence alignment of the MUSCLE algorithm in the MEGA X software (version 10.1.8; Molecular Evolutionary Genetics Analysis). Probabilities (%) of 1,000 bootstrap trials for each node were labeled in the tree. *Homo sapiens* AE 1 (HsAE1) was used as the outgroup on the basis of the structural similarity to BORs.

**Statistics**

For statistical analysis, we used Prism 8 software (version 8.4.2; GraphPad Software, CA). Detailed ANOVA and t-test results are described in Supplemental Data Set 3.

**Accession numbers**

Gene models used in this article can be found in the Arabidopsis Genome Initiative database under the following accession numbers: BOR1 (AT2G47160), BOR2 (AT3G62270), and NIP5;1 (AT4G10380).

**Supplemental data**

Supplemental Figure 1 LC-MS/MS analysis. 
Supplemental Figure 2 Additional data of BOR1-GFP ubiquitination. 
Supplemental Figure 3 K48-linked ubiquitination of BOR1-GFP was not detected. 
Supplemental Figure 4 Ectopic localization of G201R, V250F, and S251F mutants of BOR1-GFP. 
Supplemental Figure 5 Phylogenetic analysis of BOR family proteins. 
Supplemental Figure 6 Expression of Xpress-His6-BOR1 variants in yeast. 
Supplemental Figure 7 Comparison of substrate-binding pockets of AtBOR1, UraA, and Band3/AE1. 
Supplemental Figure 8 Growth phenotype of transgenic plants expressing BOR1-GFP variants. 
Supplemental Figure 9 Replication data of immunoblotting used for quantification of ubiquitination. 
Supplemental Figure 10 K63-linked polyubiquitination is affected in BOR1-GFP variants. 
Supplemental Figure 11 Total fluorescence of BOR1-GFP variants in the primary root tips. 
Supplemental Figure 12 C-terminal tail including K590 is not sufficient for B-induced degradation of BOR1. 
Supplemental Figure 13 In silico prediction of the A315V substitution impact on BOR1 structure. 
Supplemental Table 1 Relative B-transport activity, polyubiquitination, and B-induced degradation rate of the BOR1-GFP variants. 
Supplemental Table 2 Primer list. 
Supplemental File 1 Text file of the alignment used for the phylogenetic analysis shown in Supplemental Figure 5. 
Supplemental Data Set 1 Coordinates for phylogenetic analysis shown in Supplemental Figure 5. 
Supplemental Data Set 2 Plant materials. 
Supplemental Data Set 3 ANOVA and t-test results.

**Acknowledgments**

We thank Atsushi Toyoda (The University of Tokyo) for vector construction; Kayo Konishi and Tomoko Shimizu (Hokkaido University) for preparation of transgenic lines; Takehiro Kamiya and Maki Saiki (The University of Tokyo) for ICP-MS analysis; Toshiaki Ito (Hokkaido University), Martina Beck, Ji Zhou, Silke Robatzek (The Sainsbury Laboratory, TSL), Tetsuya Kurata, Tomoaki Sato (Nagoya University) for confocal microscopy. We also thank Satoshi Naito, Hitoshi Onouchi, Shinji Wakuta (Hokkaido University), and Wolf B. Frommer (Nagoya University) for discussion and helpful advice. Molecular graphics and analyses were performed with UCSF Chimera, developed by the Resource for Biocomputing, Visualization, and Informatics at the University of California, San Francisco. Parts of this work were conducted at Nagoya University Live Imaging Center supported by Japan
Advanced Plant Science Research Network and the Electron Microscope Laboratory, Research Faculty of Agriculture, Hokkaido University.

Funding
A.Y. was supported by the Japanese Society for the Promotion of Science (JSPS; Grant reference number: 252799). This work was supported by the Ministry of Education, Culture, Sports, Science, and Technology of Japan (Grants-in-Aid for Scientific Research [grant nos. 19K16164 to A.Y., 26712007 and 19H05763 to J.T., 19H05637 to T.F.]).

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