Iron is an essential micronutrient that plays important roles as a redox cofactor in a variety of processes, many of which are related to DNA metabolism. The E2 ubiquitin conjugase UBC13, the only E2 protein that is capable of catalyzing the formation of non-canonical K63-linked ubiquitin chains, has been associated with the DNA damage tolerance pathway in eukaryotes, critical for maintenance of genome stability and integrity. We previously showed that UBC13 and an interacting E3 ubiquitin ligase, RGLG, affect the differentiation of root epidermal cells in Arabidopsis. When grown on iron-free media, Arabidopsis plants develops root hairs that are branched at their base, a response that has been interpreted as an adaption to reduced iron availability. Mutations in UBC13A abolished the branched root hair phenotype. Unexpectedly, mutations in RGLG genes caused constitutive root hair branching. Based on recent results that link endocytotic turnover of plasma membrane-bound PIN transporters to K63-linked ubiquitination, we reinterpreted our results in a context that classifies the root hair phenotype of iron-deficient plants as a consequence of altered auxin distribution. We show here that UBC13A/B and RGLG1/2 are involved in DNA damage repair and hypothesize that UBC13 protein becomes limited under iron-deficient conditions to prioritize DNA metabolism. The data suggest that genes involved in combating detrimental effects on genome stability may represent essential components in the plant’s stress response.

Keywords: auxin, DNA repair, endocytosis, iron deficiency, K63-linked polyubiquitination, root hairs

INTRODUCTION
Iron participates in a variety of vital processes as a redox cofactor and is an indispensable element for virtually all organisms. In plants, suboptimal iron availability results in decreased yield and reduced quality of edible plant parts, causing severe economic losses. When plants are the major source of dietary iron, a low iron concentration may pose severe health problems. Iron deficiency-induced anemia (IDA) is the most widespread nutritional disorder worldwide. In severe cases, IDA can affect infant development and increase the risk of maternal and child mortality. Due to the limited solubility of iron in aerated soils, mechanisms have evolved that aid in mobilizing otherwise sparingly soluble Fe(III)oxyhydroxides, which represent the prevailing form of iron in most soils at neutral or basic pH. Induction of proton-translocating P-type ATPases decreases the rhizospheric pH, thereby increasing the activity of Fe(III) by a factor of 1,000 for each unit the pH decreases (Santi et al., 2005; Santi and Schmidt, 2008, 2009). In addition, secretion of iron binding compounds (IBCs) facilitates the mobilization of iron particularly at high pH (Sussin et al., 1993; Jin et al., 2007; Fourcroy et al., 2013; Rodriguez-Celma et al., 2013). A further set of supposedly separately regulated responses comprise morphological changes such as the formation of additional root hairs and invaginations of secondary walls in the rhizodermis/hypodermis, responses that were suggested to improve iron acquisition by increasing the absorptive surface area (Schikora and Schmidt, 2001; Schmidt et al., 2003). In contrast to phosphate deficiency, which triggers a nutrient-specific and conserved set of developmental responses including denser and longer root hairs (Ma et al., 2001; Savage et al., 2013) and a dramatically altered root architecture where primary root growth is attenuated and lateral root formation is stimulated (Ticconi et al., 2009), the morphological responses to iron deficiency are less well studied and appear to be less conserved among species. While some plants such as sunflower and cucumber produce very dense root hairs in response to iron deficiency (Landsberg, 1996; Li and Schmidt, 2010), other species such as Plantago lanceolata and tomato show only a moderate increase in root hair frequency (Schmidt and Bartels, 1996; Schikora and Schmidt, 2002). Interestingly, both the induction of cell wall invaginations and the formation of extra root hairs are inducible by application of exogenous auxin, suggesting to us that auxin is involved in the induction of these responses (Landsberg, 1996; Schmidt et al., 2003).

In Arabidopsis, the morphological responses to iron deficiency are neither pronounced nor uniformly described. Branching of the hairs was described as a major response and interpreted as an alternative to longer and/or denser root hairs as a strategy to increase the surface area of the roots (Müller and Schmidt, 2004). Instead, formation of shorter and misshapen root hairs was described for iron-deficient Arabidopsis plants by Dünény et al.

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et al., 2009). In contrast to that of phosphate-deficient plants, which form longer and denser root hairs, the ubiquitin conjugase UBC13. UBC13 was identified in the root hair zone of iron-deficient cucumber plants by a proteomic approach and cloned using the CODEHOP strategy (Li and Schmidt, 2010). The sequence of UBC13 is highly conserved among eukaryotes, and its function has been related to the error tolerance branch of DNA repair in yeast and Arabidopsis and to the NF-kB signal transduction pathway in mammals (Hofmann and Pickart, 1999; Wen et al., 2006; Wu et al., 2009). In Arabidopsis, UBC13 is encoded by two close sequencologs, UBC13A (UBC35) and UBC35B (UBC36). UBC13 is the only known E2 ubiquitin conjugating enzyme that is capable of catalyzing the formation of ubiquitin chains linked to K63, a function that appears to be conserved in eukaryotes (Hofmann and Pickart, 1999). In contrast to the formation canonical of ubiquitin chains linked via K48, which target proteins for degradation via the 26S proteasome, proteins conjugated to K63-linked polyubiquitin chains are involved in signaling and in the coordination of cellular processes such as endocytotic trafficking and DNA repair. UBC13 acts in conjunction with E3 ligases such as the RING domain ligase protein RGLG2 that has been shown to interact with UBC13 in Arabidopsis (Yin et al., 2007). Non-canonical ubiquitination through K63-linked ubiquitin chains is required for DNA damage tolerance, a pathway that allows the bypass of lesions in the DNA template during replication. The AtUBC13 genes complemented the yeast ubc13 null mutant for sensitivity to DNA damaging agents and for spontaneous mutagenesis, suggesting that in Arabidopsis UBC13 proteins are involved in the error-free DNA damage tolerance pathway (Wen et al., 2006). The UBC enzyme variant Mms2 is an E2-like protein that interacts with Ubc13 in eukaryotic cells (ubiquitin conjugating enzyme variant, UEV; Hofmann and Pickart, 1999; VanVliet et al., 2001). The Arabidopsis genome harbors four Mms2 homologs, UEV1A to UEV1D, all of which can form stable complexes with UBC13 from yeast and Arabidopsis (Wen et al., 2008). For UEV1D a function in DNA damage response has been experimentally verified (Wen et al., 2008). We here present evidence that both UBC13 and RGLG2 are critical in DNA damage repair and hypothesize that the root hair phenotype of iron-deficient plants is caused by diminished availability of RGLG2 protein, probably due to stress-induced movement from the plasma membrane to the nucleus and subsequently changed auxin distribution in epidermal cells. We further speculate that under iron-deficient conditions different E2–E3 complexes are favored in the nucleus to prioritize genome stability.

THE Arabidopsis IRON DEFICIENCY ROOT HAIR PHENOTYPE: A CONSEQUENCE OF CHANGES IN AUXIN DISTRIBUTION? The branched root hair phenotype of iron-deficient plants differs from that of phosphate-deficient plants, which form longer and denser root hairs resulting from restricted longitudinal elongation of root epidermal cells and additional cell fate assignment by increased expression of the Myb-type transcription factor ETC1 (Ma et al., 2001; Müller and Schmidt, 2004; Savage et al., 2013). Branched root hairs have also been described for auxin-related mutants such as axr1, aux1, and axr2, and for mutants defective in actin filament organization or vesicle transport such as scf1 or tip1 (see Guimil and Dunand, 2007 for an overview). Plants harboring mutations in UBC13A do not respond to iron deficiency with the formation of branched root hairs. Moreover, ubc13A ubc13B double mutants have shorter root hairs compared with the wild type, a trait that was also observed in some auxin-deficient mutants such as trh1 (Vicente-Aguilera et al., 2004; Li and Schmidt, 2010). Transgenic plants overexpressing UBC13A (UBC13A OE) showed a phenotype that was essentially similar to that of the wild type when grown on iron-free media, with no further increase in the number of branched hairs. Iron-sufficient UBC13A OE plants are indistinguishable from Col-0 plants, indicating that an iron deficiency signal is required for the induction of root hair branching. Notably, double mutants defective in the expression of the E3 ligases RGLG1 and RGLG2 constitutively displayed the branched root hair phenotype (Li and Schmidt, 2010). This was an unexpected result since disruption of the E2/E3 cascade should result in similar phenotypes, regardless of the site of disruption. Interestingly, the phenotype of rglg1 rglg2 mutants could be rescued by omitting phosphate from the growth media (Li and Schmidt, 2010). Such a rescue by phosphate deficiency was described for the short root hair phenotype of the auxin transport mutant trh1 (Müller and Schmidt, 2004). TRH1 is a member of the AKT/AKUP/AHAK family of potassium carriers required for the correct distribution of auxin (Vicente-Aguilera et al., 2004). It seems reasonable to speculate that altered auxin metabolism is the cause of the variable phenotype of the mutants grown under the different conditions. In support of this assumption, DR5–GUS reporter lines showed reduced GUS expression under iron-deficient conditions (Lam et al., 2012), indicative of decreased levels of or a diminished responsiveness to auxin. In phosphate-deficient DR5–GUS plants, staining was more intense than under control conditions, indicating increased auxin levels in this growth type. Together these results suggest that compromised auxin transport or metabolism could be the cause for the branched root hair phenotype under iron-deficient conditions.

A MODEL FOR THE FUNCTION OF K63-LINKED UBIQUITINATION IN ROOT HAIR CELLS The following model would explain the results obtained by us (Li and Schmidt, 2010) and others (Figure 1). The model is based on observations made in Arabidopsis but may also apply to other species. Under iron-sufficient conditions, RGLG1/2 binds a protein X that, when not captured by RGLG1/2, acts as an inhibitor of proper root hair initiation, probably via a reduction of auxin responsiveness. Protein X may act directly or indirectly on auxin distribution. AtUBC13 is not induced by iron deficiency (Li and Schmidt, 2010) and not much affected by other stresses1 and thus

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1 https://www.genevestigator.com/
iron-deficient roots, RGLG1/2 is located to the nucleus where it likely to be present in iron-sufficient plants in equal amounts. In iron-deficient conditions, the expression of two genes encoding the iron storage proteins and their effects on root hair differentiation. (A) In wild-type roots, protein X is bound to RGLG proteins, preventing the formation of branched root hairs. UBC13 is present, but only under iron-deficient conditions UBC13 interacts with RGLG proteins which move to the nucleus. The interaction of UBC13 with RGLG1/2 releases protein X which leads to the formation of branched hairs. (B) In rglg mutants, protein X is free both in the presence and absence of iron, causing root hairs to branch independent of the iron supply. (C) In root hair cells of ubc13 plants, no interaction between UBC13 and RGLG1/2 can occur, leaving protein X inactive under both iron-sufficient and iron-deficient conditions. (D) Over-expression of UBC13 increases the level of UBC13 protein but does not affect UBC13-RGLG1/2 interaction, resulting in a phenotype similar to that of the wild type.

We can only speculate on the nature of protein X. A possible scenario implies that protein X post-translationally regulates auxin distribution or metabolism. The plasma membrane-localized auxin carrier protein PIN2 is constitutively recycled by endocytosis (Benjamins and Scheres, 2008; Grunewald and Friml, 2010). Recently, RGLG proteins were shown to be involved in the control of the proteolytic turnover of PIN2 via K63-linked ubiquitination (Leitner et al., 2012). Loss of PIN2 ubiquitination interferes with vacuolar targeting, stabilizes PIN2, and alters auxin availability in Arabidopsis roots. rglg1 rglg2 mutants had reduced auxin levels and transgenic rglg1 rglg2 plants carrying the DR5-reporter construct showed reduced auxin responsive ness (Yin et al., 2007). Translated to the model, PIN2 could represent protein X. A possible scenario is outlined in Figure 2. Under iron-sufficient conditions, RGLG proteins mediates K63-linked ubiquitination of PIN2, thereby controlling its proteolytic turnover (Figure 2A). Loss of PIN2 polyubiquitination, caused for example by a mutation in RGLG1/2, causes an arrest of endocytotic cycling of PIN2, decreased auxin levels and, ultimately, branching of root hairs. Such a scenario may apply to iron-deficient plants in which UBC13 is recruited by RGLG1/2 in the nucleus, compromising the ubiquitination of PIN2, which accumulates in the cell (Figure 2B). Recruitment of UBC13 may be facilitated by translocation of RGLG from the plasma membrane to the nucleus.

UBC13 was suggested to be involved in the error-free DNA damage repair pathway (Wen et al., 2006). In yeast, this function is fulfilled by an Ubc13-Mms2-Rad5-mediated polyubiquitination of the homotrimeric protein complex proliferating cell nuclear antigen (PCNA); in humans, PCNA ubiquitination is catalyzed by the Rad5 homolog HLF and SHPRH (Ulich, 2007; Chang and Cimprich, 2009). A role for AtRAD5A in DNA repair has been demonstrated (Chen et al., 2008; Mannuss et al., 2010; Wang et al., 2011) and an involvement of UBC13-UEV1 and AtRAD5A in this pathway in Arabidopsis represents a likely scenario.

In ubc13a mutant plants, several iron-responsive genes are less induced than in the wild type (Li and Schmidt, 2010), indicative of an involvement of UBC13A in processes that utilizes iron. For example, induction of the transcription factor HHLH38, an essential regulator of the iron deficiency response, was markedly less pronounced in roots of the ubc13a mutant. Notably, under iron-deficient conditions the expression of two genes encoding the iron storage proteins FER1 and FER2 was highest in ubc13a plants when compared with the wild type, suggesting a higher iron status of the mutant. These results may be interpreted in a sense that more iron...
FIGURE 2 | Experimental evidence for processes associated with the formation of K63-linked ubiquitin chains. (A) Under control conditions, RGLG mediates the endocytic turnover of PIN2, thereby regulating the distribution of auxin. UBC13-UEV1-RAD5 is involved in housekeeping DNA damage repair. (B) Under iron-deficient conditions, RGLG moves to the nucleus where it interacts with UBC13. UBC13-RGLG1/2 might be associated with DNA repair or with other processes related to genome stability. Loss of PIN2 ubiquitination cause a cessation of endocytic PIN2 cycling, thereby altering auxin distribution. N, nucleus; LV, lytic vacuole; EE, early endosome; RE, recycling endosome.

is available if the UBC13-RGLG pathway is not engaged. Based on the conserved function of UBC13 in DNA damage repair among eukaryotes, it can be assumed that the UBC13-RGLG1/2 complex participates in DNA damage repair.

To test our hypothesis, we germinated seeds from wild-type plants, rglg1 rglg2, and ubc13a ubc13b double mutants on media containing various concentrations of the DNA-damaging agent methyl methanesulfonate (MMS). While wild-type plants were largely unaffected both under iron-sufficient and iron-deficient conditions, the germination rate of ubc13a ubc13b double mutants showed a dramatic decrease in germination rate which was more pronounced under iron-deficient conditions (Figure 3). rglg1 rglg2 mutants showed a less pronounced and iron-independent decrease in seed germination (Figure 3). These data are consistent with the assumption that under iron-deficient conditions RGLG1/2 is translocated into the nucleus where it may associate with UBC13 and other E2 proteins, thereby diminishing the efficiency of UBC13 in DNA damage tolerance. UBC13 may have critical, currently unknown functions in DNA metabolism in conjunction with RGLG1/2, probably associated with genome stability (Figure 2B).

THE ROLE OF IRON IN DNA SYNTHESIS AND REPAIR

Iron is required for several DNA-related processes. For example, ribonucleotide reductases (RNRs) require iron as an essential cofactor and iron deficiency leads to reduced RNR activity (Cavanaugh et al., 1985; Saletta et al., 2011). RNRs activity is linked to DNA synthesis and tightly regulated in response to iron deficiency in yeast and mammals to assure accurate DNA replication (Furukawa et al., 1992). In yeast, a post-transcriptional regulatory mechanism promotes destabilization of mRNAs involved in non-essential pathways and degradation of such transcripts via
While in plants such information is not available, we speculate DNA-repair-related proteins (Gao et al., 2003; Guo et al., 2008) repair (Boulon et al., 2010; Antoniali et al., 2013), and for several recent, nucleoli have been associated with a function of sensing material, i.e., the nucleus, plastids, and mitochondria. More interestingly, many of the down-regulated transcripts are related to the tricarboxylic acid cycle or participate in mitochondrial electron transport, switching respiration to fermentative metabolism (Sanvisens et al., 2011) and also, storage of iron in the vacuole via CCC1 is inhibited by this pathway, a further parallel to Arabidopsis roots in which CCC1-like vacuolar iron transporters are down-regulated in response to iron deficiency (Gollhofer et al., 2011).

In humans, iron deficiency leads to G1/S cell cycle arrest and to the induction of several members of the growth arrest and DNA damage (GADD) gene family members, probably associated with DNA damage resulting from iron deficiency (Saletta et al., 2011). In addition, DNA replication and repair depends on Fe-S clusters (Netz et al., 2010; Pokharel and Campbell, 2012) the synthesis of which is impaired upon iron deficiency (Chen et al., 2004). Defects in Fe-S cluster biogenesis result in genome instability (Veatch et al., 2009). This is consistent with a competition for UBC13 protein in the nucleus due to stress-induced translocation of RGLG1/2 to the nucleus. The role of RGLG1/2 under iron-deficient conditions awaits clarification, but from our results a function for RGLG1/2 in genome stability appears to be plausible. Further experiments addressing the fitness of plants treated with DNA-damaging agents and grown with different iron supply may falsify or validate our hypothesis. The effects associated with adverse environmental condition on genome stability are generally understudied and, despite big steps forward in the past few years (reviewed by Waterworth et al., 2011), not fully understood. From the example outlined here for iron deficiency it might be inferred that mechanisms that are involved in combating the constant assault of environmental stresses on genome stability may be an important component of the plant stress response.

**MATERIALS AND METHODS**

Arabidopsis thaliana (L.) Heynh Columbia (Col-8) ecotype was used as the wild-type. Seeds of the wild-type, ubc13a and ubc13b mutants were obtained from the Arabidopsis Biological Resource Centre (Ohio State University), ubc13a ubc13b double mutants were generated by genetic crossing. The rgl1 rgl2 mutant lines were provided by Dr. Andreas Bachmeir (University of Vienna). Seeds were surface-sterilized and sown on media as described by Estelle and Somerville (1987), supplemented with 0%, 0.0066%, and 0.014% MMS. Data points indicate the percentage of seed germination after 5 days with SD (n = 36).

**CONCLUSION**

In summary, the re-interpretation of the data outlined above hints at a new facet of the iron deficiency response of Arabidopsis, which might also be important for other plant species and may thus represent a general aspect in the adaptation of plants to low iron availability. Under iron shortage, DNA-related processes such as replication and repair might be prioritized to secure essential housekeeping functions. While alternative scenarios such as a role of UBC13 in root development that is not linked to DNA repair cannot be ruled out at present, the fact that ubc13a mutants appear to have a higher, healthier iron status implies that activation of the DNA damage pathway occurs at the expense of other genes encoding iron-containing proteins. This situation is similar to what has been reported for yeast where iron deficiency limits iron utilization in energy-generating pathways via the action of the RNA-binding protein Chs2 to avoid that essential processes such as DNA synthesis and repair and thus genome integrity are compromised (Martínez-Pastor et al., 2013). While the evidence reported here is indirect and awaits experimental validation, it is tempting to assume that, similar to yeast, post-transcriptional mechanisms that bias the translation of messenger RNAs encoding proteins involved in essential iron-requiring processes such as DNA replication and repair also occur in Arabidopsis and other plants. Our initial experiments indicate that both UBC13 and RGLG are involved in DNA repair, but only for ubc13a ubc13b mutants iron deficiency severely the effect of MMS. This situation in nuclei/nucleoli is important for prioritizing DNA replication and repair when iron becomes limited.

**FIGURE 3** Analysis of DNA damage response during seed germination. Seeds of wild-type plants, rgl1 rgl2, and ubc13a ubc13b double mutants were placed on either iron-replete or iron-deplete media containing 0, 0.0066%, 0.014%, and 0.014% MMS. Data points indicate the percentage of seed germination after 5 days with SD (n = 36).
0.01% and 0.014% MMS either with or without 40 μM FeEDTA, and stored at 4°C for 2 days. A total of 36 seeds were placed on each plate, three plates were used for each treatment. After 5 days, seed germination was determined and normalized with the rate observed on iron-replete standard without MMS.

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