Suppressor mutations in the Glutamine Dumper1 protein dissociate disturbance in amino acid transport from other characteristics of the Gdu1D phenotype

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Intracellular amino acid transport across plant membranes is critical for metabolic pathways which are often split between different organelles. In addition, transport of amino acids across the plasma membrane enables the distribution of organic nitrogen through the saps between leaves and developing organs. Amino acid importers have been studied for more than two decades, and their role in this process is well-documented. While equally important, amino acid exporters are not well-characterized. The over-expression of GDU1, encoding a small membrane protein with one transmembrane domain, leads to enhancement of amino acid export by Arabidopsis cells, glutamine secretion at the leaf margin, early senescence and size reduction of the plant, possibly caused by the stimulation of amino acid exporter(s).

Previous work reported the identification of suppressor mutations of the GDU1 over-expression phenotype, which affected the GDU1 and LOG2 genes, the latter encoding a membrane-bound ubiquitin ligase interacting with GDU1. The present study focuses on the characterization of three additional suppressor mutations affecting GDU1. Size, phenotype, glutamine transport and amino acid tolerance were recorded for recapitulation plants and over-expressors of mutagenized GDU1 proteins.

Unexpectedly, the over-expression of most mutated GDU1 led to plants with enhanced amino acid export, but failing to display secretion of glutamine and size reduction. The results show that the various effects triggered by GDU1 over-expression can be dissociated from one another by mutagenizing specific residues. The fact that these residues are not necessarily conserved suggests that the diverse biochemical properties of the GDU1 protein are not only born by the characterized transmembrane and VIMAG domains. These data provide a better understanding of the structure/function relationships of GDU1 and may enable modifying amino acid export in plants without detrimental effects on plant fitness.

Keywords: Arabidopsis, suppressor screening, ethyl methanesulfonate, structure–function, amino acid transporter, glutamine transport
## Introduction

Amino acids are critical metabolites in plants which fulfill several roles, in addition to being the constituting blocks of proteins. Amino acids are used as precursors for the synthesis of many secondary metabolites, like flavonoids (Falcone Ferreyra et al., 2012), alkaloids (Ziegler and Facchini, 2008), and glucosinolates (de Kraker and Gershenzon, 2011), which are metabolites critical for interaction of the plant with the environment (attraction, defense, and protection). Amino acids, especially Gln and Asn, also serve as essential carriers for organic nitrogen throughout the plant, being transported through the xylem and phloem saps between leaves, roots, storage organs, and meristems (Tegeder, 2014). Transport in the plant is mediated at the cell level by membrane proteins with specialized functions. Amino acid importers and exporters mediate transport of amino acids in opposite directions: importers mediate transport toward the cytosol, exporters mediate transport out from the cytosol, and respectively from or to the apoplasm, vacuoles or intracellular vesicles.

Characterized amino acid importers belong to the APC superfamily (Amino Acid Polyamine transporter; Vastermark et al., 2014), composed of 63 members in Arabidopsis (Tegeder and Rentsch, 2010). Importers utilize energy of the proton gradient across membranes to import amino acids against their concentration gradient, and are involved in many roles, like uptake from the soil, import into the phloem, phloem–xylem exchange, and transport into the embryo (Tegeder, 2014). Much less is known about amino acid exchangers and exporters. One member of the CAT subfamily has recently been described as an amino acid exchanger in tomato (Snowden et al., 2015). Two APC members have been described as possible amino acid exporters: CAT8 (Yang et al., 2010) and BAT1/GABP1 (Dundar and Bush, 2009; Michaeli et al., 2011). Finally, one gene belonging to the Drug and Metabolite Transporter superfamily (Jack et al., 2001), SIAR1, has been shown to unequivocally mediate amino acid export from plant cells (Ladwig et al., 2012). The family SIAR1 belongs to contains 47 members with only one other gene characterized, the auxin transporter WAT1 (Ranocha et al., 2013).

The existence of processes controlling the activity of amino acid transporters was evidenced by the discovery of the GDU1 gene, encoding an 158 amino acid protein, with a single transmembrane domain (Pilot et al., 2004). Two domains are conserved among GDU proteins: a membrane domain, and a cytosolic 19 amino acid-long region, called the VIMAG domain (Pilot et al., 2004; see Figure 1). Over-expression of GDU1, for instance in the gdu1-1D mutant, leads to a complex phenotype characterized by reduced plant size; early leaf senescence; crystallization of Gln at the leaf margins; increased amino acid content in leaves, apoplasm, and xylem and phloem saps; tolerance to exogenously supplied amino acids; and notably enhanced amino acid export from cells, while amino acid import remains unaffected (Pilot et al., 2004; Pratelli and Pilot, 2007; Pratelli et al., 2010). This complex phenotype (called Gdu1D) can almost entirely be explained by enhanced amino acid export from cells: this phenomenon would increase amino acid content in the apoplasm and phloem and xylem saps, and prevent absorption of Gln from the xylem in the leaf, which then is excreted by the hydathodes (Pilot et al., 2004). Size reduction likely comes from the induced disturbance in nitrogen metabolism. The fact that over-expression of GDU1 in Nicotiana tabacum and GDU1-homologs in Arabidopsis leads to a Gdu1D-similar phenotype suggests that the proteins of this family have a conserved function in plants, related to the regulation of amino acid export (Pratelli and Pilot, 2006; Pratelli et al., 2010). Nevertheless, the precise function of GDU1 in this process remains unknown.

Attempts to understand the function of GDU1 in the plant led to the identification of the ubiquitin ligase LOG2 in a yeast-two-hybrid screening. GDU1 and LOG2 localize at the plasma membrane and are able to co-immunoprecipitate when expressed in Nicotiana benthamiana (Pratelli et al., 2012). Decrease in LOG2 expression suppressed the Gdu1D phenotype, indicating that LOG2 activity is necessary for the development of the phenotype (Pratelli et al., 2012). A mammalian homolog of LOG2 is the mahogunin protein (MGRN1; 45% similarity at the protein level), involved in the regulation of the activity and trafficking of membrane proteins and in degrading aggregated proteins (Jiao et al., 2009; Perez-Oliva et al., 2009; Gunn et al., 2013; Chhangani et al., 2014). Despite coming from different organisms, MGRN1 and LOG2 were found to have several overlapping functional properties, notably the ability of the mammalian protein to partially complement the loss of the plant LOG2 in Arabidopsis (Guerra et al., 2013). It is hypothesized that LOG2 and GDU1 are involved in the regulation of the activity or trafficking of amino acid exporters, such as, when GDU1 is over-expressed, the exporters are more active at the plasma membrane (Pratelli et al., 2012; Guerra et al., 2013).

An ethyl methanesulfonate (EMS) screening was previously performed to isolate suppressor mutations of the Gdu1D phenotype. For this purpose, two GDU1 over-expressing lines (gdu1-5D and gdu1-6D, created by introducing a GDU1-over-expressing construct into wild type plants) were EMS-mutagenized, and screened for progenies that grew similarly to the wild type. These suppressor lines hence over-express a mutant GDU1 protein in addition to the endogenous GDU1 protein. Two suppressor mutants, loss of gdu1-1 and 2-1 (log1-1 and log2-1) were isolated and characterized (Pratelli and Pilot, 2006; Pratelli et al., 2012). log1-1 carries a G100R substitution in the conserved VIMAG domain of GDU1, which abolishes the interaction with LOG2 (Pratelli et al., 2012). The log2-1 is a R12K substitution in LOG2, whose effect has not been determined at the biochemical level (Pratelli et al., 2012). Another mutation, log2-3, was isolated from the same screening, and is a nonsense mutation in LOG2 (R15stop; Pratelli and Pilot, unpublished data). These three log mutations led to plants that were phenotypically indistinguishable from the wild type when grown on soil and on amino acid-containing media. From the same screening, we isolated three additional log1 mutations, whose characterization is reported here.
Results

Identification of Three New Mutations Suppressing the Gdu1D Phenotype

The pipeline previously described by Pratelli and Pilot (2006) was used to isolate new Gdu1D suppressor mutants: visual screening of M2 plants from the EMS mutagenesis of the GDU1 over-expressors gdu1-5D and gdu1-6D, and confirmation of continued GDU1 mRNA over-accumulation. Out of a total of 110,000 M2 seeds screened, three additional suppressor mutations were isolated that nearly restored the wild type size of the plants (Supplementary Figure S1), and suppressed the early senescence and Gln secretion. Analysis of the phenotype of F2 plants from crosses of the mutants with gdu1-6D and the wild type Col-7 showed that the three new mutations were recessive and segregated in accordance with an intragenic mutation in GDU1 (data not shown). Similar to gdu1-5D and segregated in accordance with an intragenic mutation in Col-7 showed that the three new mutations were recessive and segregated in accordance with an intragenic mutation in GDU1 (data not shown). Similar to log1-1, the corresponding mutants came from the mutagenesis of the over-expressor gdu1-6D (Pratelli and Pilot, 2006), and were named log1-2, log1-3, and log1-4. Sequencing of the GDU1 CDS in the T-DNA construct leading to its over-expression revealed that the mutations in log1-2, log1-3, and log1-4 corresponded to G202A, G119A, and C95T mutations (numbered from the ATG), respectively, in the GDU1 DNA sequence. These mutations led to E68K, G40D, and S32L substitutions, respectively, in the GDU1 protein sequence (Figure 1). Because these mutations suppressed the Gdu1D visual phenotype (plant size, Gln secretion, and early senescence), it was hypothesized that they affected GDU1 protein function. The log1 mutants were hence hypothesized to over-express inactive GDU1 proteins and the characterization of the mutations was thus a way to better understand GDU1’s structure–function relationships.

Amino acid uptake of GDU1 over-expressing lines was shown to be reduced, while efflux was enhanced (Pratelli et al., 2010). It was expected that the suppressor lines would display similar uptake and efflux as wild type plants, because of their wild type phenotype on soil. To test this hypothesis, Gln uptake of the suppressor lines was analyzed and compared to the wild type and the parental line gdu1-6D. Since this assay was not performed in the log1-1 plants (Pratelli and Pilot, 2006), they were included in the present study. The log1-1, log1-2, log1-3, and log1-4 suppressor mutants showed uptake and efflux similar to one another, but surprisingly different from both the wild type and gdu1-6D: the four log1 mutants had an uptake about 60% lower than the wild type (Figure 2A) and an efflux twice as large as the wild type (Figure 2B). As a comparison, Gln uptake of gdu1-6D was 75% lower and the efflux was three times larger than the wild type. The intermediate phenotype was clearly visible when uptake and efflux were plotted on the same graph (Figure 2C). This result suggests that these recessive suppressor mutations decrease GDU1 overall activity and thus lead to a milder phenotype. It has indeed been shown that the strength of the Gdu1D phenotype depends on the GDU1 expression level and that the plants are small and display early senescing leaves only when GDU1 is over-expressed 10 times or more than the wild type (Pilot et al., 2004).

Characterization of Plants Over-Expressing the GDU1 Variant Proteins

Despite two different attempts, no suitable antibody could be raised against GDU1. The gdu1-5D and gdu1-6D plants and the corresponding log suppressor mutants express an un-tagged GDU1 protein, preventing any quantitation of the GDU1 protein accumulation. In order to test for any effect of the suppressor mutations on the accumulation of the GDU1 protein, wild type GDU1, the three GDU1 variants, G100R GDU1 (corresponding to the log1-1 suppressor mutant; Pratelli and Pilot, 2006) and a GDU1 protein lacking the VIMAG domain (ΔVIMAG) were fused with the HA tag, placed under the control of the CaMV 35S promoter, and expressed in N. benthamiana. The ΔVIMAG, G100R, E68K, and S32L mutations did not affect protein accumulation, while the G40D mutation led to a consistent reduction of about 30% in protein accumulation.
in this system (Figure 3). The sub-cellular localization of these five GDU1 variants was determined by expression of the GFP-tagged proteins (GFP positioned in C-terminal) in \textit{N. benthamiana} leaves and observed by confocal microscopy. Similar to GDU1, the proteins localized to the plasma membrane and in compartments that could correspond to endosomes, similar to the wild type GDU1 protein (Pratelli et al., 2012). Interestingly, the fluorescence of these compartments was stronger for the ΔVIMAG, G100R and G40D GDU1 proteins, suggesting that the proteins located more in these structures than the other GDU1 proteins (Supplementary Figure S2), possibly revealing some effect of the mutations on the protein properties.

To study the ability of the proteins to lead to the Gdu1D phenotype, the HA-fusion constructs were used to transform \textit{Arabidopsis} and create recapitulation lines. The accumulation of the proteins in lines that segregated 3:1 for the kanamycin resistance was tested by western blot (Figure 4B). G40D GDU1-HA accumulated at a lower level than GDU1-HA, in good agreement with the \textit{N. benthamiana} results. Surprisingly, S32L GDU1-HA did not accumulate to the GDU1-HA levels in both tested lines, but was still present at high levels. In all cases, the rosette sizes of the plants expressing the GDU1 suppressor variants were identical to the wild type. The plants over-expressing GDU1-HA showed a ∼45% reduction in rosette diameter compared to the wild type and the empty vector-transformed plants (Figure 4A). Only the GDU1-HA over-expressors displayed Gln secretion crystals, typical of the Gdu1D phenotype. The size reduction of the GDU1-HA lines was less than for the original gdu1-1D over-expressor (∼60%; Figure 4A), and could be attributed to the difference in the construct used or the presence of the HA tag, which might slightly interfere with the protein stability or activity.

Since the log1 mutants showed intermediate Gln uptake and efflux between the wild type and the gdu1-6D mutant, Gln transport by the recapitulation lines was studied. G40D and S32L GDU1-HA over-expressing plants behaved similarly to the corresponding log1-3 and log1-4 suppressor mutants: the uptake was reduced by ∼50% while the efflux was increased by ∼50–100%. On the contrary, E68K GDU1-HA over-expressors transported Gln similarly to the GDU1-HA plants, differently from the log1-2 plants that they were supposed to recapitulate (Figures 5A,B). This discrepancy between the Gln uptake and efflux of the log1-2 suppressor mutant and...
E68K GDU1-HA over-expressors was unexpected and is not completely explained. It is possible that the presence of the HA tag interferes with the function of the mutated protein, similar to what was observed with the wild type GDU1 (the GDU1-HA over-expression line exhibits a weaker phenotype than the original gdu1-1D; Figure 4A). These data prove that the intermediate phenotype of the log1-3 and log1-4 mutants is caused by the over-expression of variant GDU1 proteins endowed with reduced functionality.

**Testing the GDU1 Variants for Interaction with LOG2**

The suppression of the Gdu1D phenotype by the G100R mutation (in the log1-1 mutant; Pratelli and Pilot, 2006) was explained by the loss of interaction with LOG2, the ubiquitin ligase necessary for the development of the Gdu1D phenotype (Pratelli et al., 2012). The G100R mutation affects the Gly100 residue of the VIMAG domain that is conserved in all GDU proteins examined so far. This Gly to Arg mutation (Figure 1) is supposed to either affect the folding of the VIMAG domain or create a steric clash at the interface surface between GDU1 and LOG2 caused by changes in residue charge and/or size. Because the three other log1 mutants displayed a similar phenotype as log1-1, we tested if any of the corresponding mutations would
affect the GDU1-LOG2 interaction, which would explain the suppression of the Gdu1D phenotype.

The LOG2 and GDU1 proteins were fused with the Myc or the HA tag respectively, expressed in N. benthamiana leaves and immunoprecipitated using cMyc agarose beads. The ΔVIMAG GDU1 protein, shown to be unable to interact with LOG2 (Pratelli et al., 2012) and to lead to the Gdu1D phenotype when over-expressed (Pratelli and Pilot, 2006), was used as a negative control (Figure 6). GDU1-HA, E68K GDU1-HA, and S32L GDU1-HA could be co-purified with LOG2-Myc, but G40D GDU1-HA did not co-immunoprecipitate (Figure 6). The G40D mutation, in addition to decreasing protein abundance (Figure 3), thus seems to prevent the interaction of GDU1 with LOG2, which could explain the Gdu1D suppressor effect. On the contrary, the Gdu1D suppressor effect of the other variants cannot be explained by an inability to interact with LOG2.

Site Directed Mutagenesis to Understand the S32L Mutation

Ser32 is highly conserved among GDU proteins, being sometimes replaced by Thr only, a residue with similar chemical properties (in 13% of the ~100 analyzed sequences from 20 plant species, including lower plants, conifers, monocots, and dicots; data not shown). Running prediction algorithms such as PhosPhAT (Heazlewood et al., 2008; Durek et al., 2010) and PlantPhos (Lee et al., 2011) suggested that Ser32 can be phosphorylated (Supplementary Figure S3) despite the fact that this part of the protein is supposed to be extra-cellular. The PhosPhAT tool also predicted that Ser28, next to Ser32, can be phosphorylated. We hypothesized that these two Ser can be phosphorylated, and that this phosphorylation is important for GDU1 function. Ser32 was mutagenized to Thr, Asp, and Ala to test for (1) the importance of the Ser vs. Thr in GDU1 function, the effect of (2) mimicking and (3) suppressing phosphorylation at this site respectively. Ser28 and Ser32 were also mutagenized to Ala at the same time to ensure that lack of Ser32 phosphorylation could not be complemented by phosphorylation of Ser28. The mutagenized GDU1 proteins were stably expressed in Arabidopsis in fusion with the HA tag. The size of the plants, protein accumulation, and Gln transport were determined as above.

Western blotting confirmed that all lines expressed the GDU1 protein variants (Figure 7B). The size and the phenotype of

![Figure 6](image_url)

**FIGURE 6** | Co-immunoprecipitation of the GDU1 variants and mLOG2 expressed in N. benthamiana. GDU1 variants were expressed as C-terminal HA fusion (H), mLOG2 (ubiquitination defective) as C-terminal Myc fusions (M) in N. benthamiana leaves. Proteins were extracted and immunoprecipitated (IP) with an anti-cMyc antibody, and detected by western blot (WB). Numbers on the right indicate molecular weight in kDa. In this particular experiment, LOG2-Myc (lane 4) did not accumulate at the same level as in the other samples.

![Figure 7](image_url)

**FIGURE 7** | Analysis of plants over-expressing the GDU1 variants.
(A) Rosette diameter of 4 week-old Arabidopsis. Plants were grown at the same time as plants in Figure 1 (the values for gdu1-1D, GDU1-HA and Col-7 are the same as in Figure 1). Error bars are SEM of 6–8 plants; statistical significance determined by ANOVA using Tukey’s HSD p < 0.01.
(B) Accumulation of the HA-tagged GDU1 proteins in each line was estimated by western blot with an anti-HA antibody. The wild type Col-7 and gdu1-1D lines are indicated as white and black bars; these lines do not express any tagged protein and were not tested by western blotting.
the plants over-expressing S32T GDU1 was identical to the
GDU1-HA over-expressors, in terms of size, Gln secretion
and leaf senescence (Figures 7A and 8), suggesting that S32T
GDU1 is fully functional. Over-expression of the S32D, S32A,
and S28A+S32A mutant proteins led to plants with slightly
reduced size (down to 80% of the wild type), but never as
much as the GDU1 or S32T GDU1 over-expressors. On the
other hand, Gln transport analyses showed that the plants over-
expressing the S32D, S32A, and S32T variants displayed uptake
and efflux similar to the plants over-expressing the wild type
GDU1 (Figure 5C). It is worth noting plants over-expressing the
S32T variant displayed a higher efflux than the GDU1-HA over-
expressor (Figure 5B), suggesting that this protein is more active
than GDU1. Interestingly, when the GDU1 protein bearing the
double mutation S28A+S32A was over-expressed, Gln uptake
and efflux were very similar to the S32L mutation, present in the
log1-4 mutant. No difference between the effect of the S32A and
S32D mutations was detected by this assay.

**A E68R Mutation Leads to an Almost Fully Functional GDU1**

We tested whether the suppressor effect of the E68K mutation
came from the change in charge (negative to positive) or from
the specific substitution to a Lys residue. For this purpose, Glu68 was mutagenized to Arg (a positively charged amino
acid), and the variant protein expressed in *Arabidopsis*. This
protein accumulated at the same level as the wild type protein
in the over-expression lines (Figure 7B), and the corresponding
plants had a size similar to one over-expressing the wild type
GDU1 (Figure 7A) but did not display the Gdu1D phenotype,
i.e., secretion and early senescence (Figure 8). Transport assays
showed that the plants over-expressing E68R behaved essentially
similar to a GDU1 over-expressor (Figure 5), showing that the
Arg at this position has little effect on the protein function in this
assay.

**Analysis of Amino Acid Tolerance of the Suppressed Mutants and of Over-Expressors of the GDU1 Variants**

Some amino acids, when supplied at high concentration, have
been shown to inhibit cell and plant growth (Bonner et al., 1992;
Lee et al., 2007; Pratelli and Pilot, 2007).

Another characteristic of the Gdu1D phenotype is the
tolerance of GDU1 over-expressors to toxic concentrations of
amino acids (Pratelli and Pilot, 2007; Pratelli et al., 2010). The
effect of high concentration of amino acids was used as a
supplemental assay to characterize the phenotype of the
log1 suppressors, the recapitulation lines and the over-expressors of
the GDU1 variants. The presence, absence or strength of the
Gdu1D phenotype was tested by growing the plants on 10 mM
Leu and Phe, shown to be particularly toxic to wild type plants
(Pratelli and Pilot, 2007). All lines grew equally on the control
medium (Figure 8 and Supplementary Figure S4), but showed
remarkable differences on the Leu- and Phe-supplemented
media. The log1-1, log1-3, and log1-4 mutants behaved similarly
to the wild type, while log1-2 behaved intermediate between
gdu1-1D and the wild type, with strong tolerance on Phe but
to Leu. Both E68K and E68R GDU1 over-expressors were
fairly tolerant to Leu and Phe, in good accordance with the
Gln transport assays, reminiscent of the Gdu1D phenotype.
The recapitulation line over-expressing G40D GDU1 was as
sensitive as the wild type. The S32L and S28A+S32A GDU1
over-expressors behaved similarly to the wild type; the S32T
GDU1 over-expressor looked like the gdu1-1D mutant; and the
S32D and S32A mutants showed an intermediate phenotype
between the wild type and the gdu1-1D mutant (Figure 8 and
Supplementary Figure S4).

**Discussion**

**Suppressed Mutants Show Only Partially Abolished Gln Uptake**

A total of four mutants were found (log1-1, log1-2, log1-3, and
log1-4), all carrying recessive mutations, hence expected
to behave as loss-of-functions, that suppressed the Gdu1D
phenotype caused by the over-expression of the GDU1 protein.
These suppressor lines over-express a mutant GDU1 protein
in addition to the endogenous GDU1 protein. The thorough
characterization of these lines presented in this study led to

| Transport group | Plant size (mm) | GDU1 phenotype | Amino acid tolerance | Leu | Phe | Ø |
|-----------------|----------------|----------------|---------------------|-----|-----|---|
| Col1 WT         | 68             | GDU1           | yes                 | 1   | 1   | 4 |
| Col1 HA        | 74             | GDU1           | yes                 | 1   | 1   | 4 |
| gdu1-1D GDU1    | 26             | yes            | 4                   | 4   | 4   | 4 |
| GDU1-HA        | 43             | yes            | 4                   | 4   | 4   | 4 |
| log1-1         | -              | yes            | 1                   | 1   | 1   | 4 |
| log1-2         | -              | yes            | 2.5                 | 4   | 4   | 4 |
| E68K GDU1-HA   | 65             | yes            | 3                   | 4   | 4   | 4 |
| E68R GDU1-HA   | 47             | yes            | 3                   | 3   | 3   | 4 |
| log1-3         | -              | yes            | 2                   | 1.5 | 4   | 4 |
| G40D GDU1-HA   | 74             | yes            | 1                   | 1   | 1   | 4 |
| log1-4         | -              | yes            | 2                   | 2   | 4   | 4 |
| S32L GDU1-HA   | 73             | yes            | 1                   | 1   | 1   | 4 |
| S32T GDU1-HA   | 40             | yes            | 4                   | 4   | 4   | 4 |
| S32D GDU1-HA   | 58             | yes            | 2                   | 2.5 | 4   | 4 |
| S32A GDU1-HA   | 64             | yes            | 2.5                 | 4   | 4   | 4 |
| S28S2A GDU1-HA | 57             | yes            | 2.5                 | 4   | 4   | 4 |

FIGURE 8 | Summary of uptake, size, phenotype, and amino acid tolerance of the log1 suppressor mutants and of lines over-expressing the GDU1 variants. WT: uptake and efflux similar to the wild type; GDU1: uptake and efflux similar to gdu1-1D; log1: intermediate between gdu1-1D and the wild type (see Figures 2 and 5). Value is the average of two lines (see Figures 4 and 7). “-“ indicates that the line was not grown in this experiment. Gln secretion and early senescence. Growth was scored according to the growth of the wild type (set to 1) and the gdu1-1D mutant (set to 4) in the corresponding growth condition; wild type and gdu1-1D plants grew equally well on the medium lacking amino acids (Ø); average of two experiments (the pictures of the plants from one experiment are displayed in Supplementary Figure S4).
unexpected observations. The size of all the mutants was similar to the wild type and the early senescence observed in the leaves of gdu1-1D and gdu1-6D was absent. However, analysis of Gln uptake showed that the mutations did not suppress the Gdu1D phenotype completely (Figure 2), with uptake and efflux of the mutants being intermediate between the wild type and the parental line gdu1-6D. Interestingly, the suppressor mutation in log2-1, which affects the ubiquitin ligase LOG2, interactor of GDU1 (Pratelli et al., 2012), similarly does not completely suppress all characteristics of the Gdu1D phenotype: the wild type size is restored, the amino acid susceptibility is abolished, the plants do not secrete Gln anymore, but amino acid export is lowered to an intermediate level between the wild type and the parent (Pratelli and Pilot, unpublished data).

These mutations were obtained from a visual screening which led to the isolation of non-secreting plants of wild type size, possibly introducing a bias toward finding mutations that do not necessarily affect transport, but rather toward ones that affect plant size and Gln secretion. The fact that the mutations in the log1-1, log1-3, and log2 suppressor mutants abolish the Gdu1D phenotype partially, in a similar way suggest that they suppress the same component of the phenotype, e.g., the one involving LOG2, leaving other components intact. This implies that the over-expression of GDU1 leads to several independent effects, some characterized by decreased size, induction of early senescence and amino acid tolerance, and others characterized by increased amino acid export.

The exact role of GDU1 is not known but two proteins interacting with it have been identified: LOG2 and its homolog LUL1, two membrane-associated ubiquitin ligases (Pratelli et al., 2012). It is proposed that GDU1 and LOG2/LUL1 form a complex stable enough to be co-immunoprecipititated, and involved in ubiquitination of yet unknown target(s) with a role in the regulation of amino acid transport (Pratelli et al., 2012). Loss of interaction with, or loss of expression of LOG2 is a simple explanation for the absence of the Gdu1D phenotype in the log1-1 and log1-3 mutants, over-expressors of a VIMAG domain-deleted GDU1 (Pratelli and Pilot, 2006) or the log2 mutants (Pratelli et al., 2012). The fact that the loss of this interaction leads to a similar phenotype as the log1-2 and log1-4 mutants is striking, since the corresponding mutations in GDU1 (S32L and E68K) do not appear to affect the interaction with LOG2 (Figure 6). However, it is possible that the functional properties of the GDU1-LOG2 complexes resulting from the interaction of S32L and E68K GDU1 and LOG2 are affected by these mutations, explaining the resulting phenotype of the corresponding plants.

The active structure of GDU1 is not known, and a possible structure could be a multimer, either with itself or other GDU proteins (seven GDU proteins are present in the Arabidopsis genome). In this case, the mutated proteins in the suppressor mutants might assemble with the endogenous wild type proteins to form hetero-multimers, which could have retained only some functions of the original multimers while being less active. This hypothesis could explain the attenuated phenotype of the plants, in terms of uptake, size or amino acid tolerance, and the diversity of the phenotypes of the plants over-expressing the various GDU1 variants (Figure 8). It has then to be postulated that GDU1 has diverse functions, and that each suppressor mutation affects them differently. These functions remain to be determined.

**The G40D Mutation Might Affect GDU1 Protein Stability and/or Function**

The G40D suppressor mutation in GDU1 has two effects on the protein’s properties: it does not accumulate at the same level as the other proteins when expressed in N. benthamiana leaves (Figure 3), and it does not co-immunoprecipitate with LOG2 (Figure 6). This mutation is located in the transmembrane domain, and affects one of the conserved residues of this domain. In about 100 GDU proteins from higher plants, Gly40 is sometimes replaced by Ala (8%), Ser (4%), Leu (1%), or Val (1%), but never by Asp (data not shown). Gly is a very common residue in membrane helices, involved in the formation of a glycine zipper in helix packing, thus enabling helix–helix interactions (Iavapour et al., 1999; Kim et al., 2005). Mutations of such important Gly residues led to abolition of transport (G114A in DcTA from Sinorhizobium meliloti; Trainer et al., 2007) or loss of dimer interaction in EmR from Escherichia coli (Elbaz et al., 2008), supposedly by disruption of the helix structure or ability to interact. The amount of sequence conservation of the GDU helix across different species is rare for a membrane protein, and one explanation could be that it is involved in interaction with membrane helices from other proteins. In this context, the G40D mutation likely affects such interaction, and may be the reason for the Gdu1D suppressor effect on the Gdu1D phenotype. Alternatively, the presence of a charged residue (Asp) inside the hydrophobic membrane helix could destabilize the protein, making it more difficult to be integrated in the lipid bilayer. This could explain the lower accumulation of the protein: membrane proteins that have folding problems are typically degraded by the cellular quality control system (Nagy and Sanders, 2004; Houck and Cyr, 2012). While the localization of the G40D GDU1 protein was not dramatically changed in N. benthamiana leaves, this does not preclude that GDU1 and its variants are addressed to different membrane subdomains. Such a hypothesis is supported by the fact that G40D GDU1 does not interact with LOG2: these proteins would not interact if G40D GDU1 and LOG2 are targeted to different membrane subdomains. Changes in subcellular distribution have indeed been observed for mutations affecting transmembrane Gly residues (Rosnoiblet et al., 2013). Nevertheless, the fact that the over-expression of G40D GDU1 in wild type plants affects amino acid transport (Figure 5C) supports the idea that the protein still retained at least one of its functions, the one affecting amino acid transport.

**Putative Role of the External Ser28 and Ser32**

The suppressor mutation in log1-4 affected Ser32, which is predicted to be phosphorylated, together with Ser28 (Supplementary Figure S3). It is not clear if these residues can be phosphorylated in vivo, because they are expected to lie in the extracellular region of the GDU1 protein. Based on the identification of bona fide extracellular phosphoproteins with no transmembrane domain by a proteomic analysis of the extracellular matrix of Arabidopsis cell suspension cultures, an extracellular phosphorylation network has been suggested.
ATP has been found in the extracellular medium, and is supposed to be involved in phosphorylation of extracellular proteins, as well as in intercellular signaling or during pathogen recognition (Ndimba et al., 2003; Chivasa et al., 2005). The fact that the mutation of the conserved Ser32 to Leu suppresses the Gdu1D phenotype led to the hypothesis that phosphorylation is important in GDU1 function. To test this hypothesis, Ser32 was mutagenized to Asp and Ala (to mimic or suppress phosphorylation, respectively) or Thr (another phosphorylation site), and over-expressed in plants. None of these mutations affected the functional properties of the GDU1 proteins as related to its effects on amino acid transport, since the over-expressors displayed similar Gln transport as GDU1 over-expressors (Figure 5). On the contrary, while S32T GDU1 over-expressors were small and tolerated amino acids, S32A and S32D over-expressors had near wild type size (Figure 7) and were somewhat susceptible to amino acids, suggesting that the S32T mutation does not alter GDU1 functional properties. S32A and S32D thus dissociated Gln uptake/export from the size and amino acid tolerance (Figure 8), showing that there is no direct relationship between these characteristics of the phenotype. The fact that their effect is similar is unexpected since these mutations typically lead to opposite phenotypes in terms of phosphorylation.

The double S28A+S32A mutation created a protein with functional properties very similar to the suppressor S32L mutation, and similar to the log1 mutants (Figures 2, 5, 7, and 8), different from the single S32A mutation. Ser28 and Ser32 thus seem to have redundant functions. In the hypothesis that these Ser are extra-cellular phosphorylation sites, mutagenesis of Ser32 individually would not have any effect, as observed; and only mutagenesis of the two Ser at the same time would lead to a phenotype. This hypothesis could be tested by mutagenizing Ser28 to Asp and Ala, which is expected to lead to a similar phenotype as the S32D and S32A over-expressing plants respectively. It can also be tested by mutagenizing S28 and S32 to Asp at the same time, the expected outcome of which would be a hyperactive GDU1 protein, and lead to plants with a phenotype reverse of the S28A+S32A over-expressing plants. In this hypothesis, the S32L mutation would prevent the phosphorylation of Ser28 and Ser32, possibly by affecting binding to a protein kinase, and would explain the observed similarity of the effect of S32L with S28A+S32A. If these Ser are not phosphorylation sites, these residues have a role in the function of the protein, which will require further investigation.

Substitution of Glu68 by a Positive Residue Affects GDU1 Protein Function
The effect of the suppressor mutation E68K has been investigated based on the hypothesis that it locally changes the charge of the protein from negative to positive. The E68R mutation was expected to lead to a non-functional GDU1, which, when expressed in plants, would lead to the same effect as the E68K GDU1 from the log1-2 mutant. Plants over-expressing E68R GDU1 were smaller, and Gln transport was similar to the GDU1 over-expressors, suggesting that this variant GDU1 protein retained much of GDU1 functionality. The main difference with the gdu1-1D over-expressor is that these plants did not secrete Gln or display early senescence.

Surprisingly, the phenotype of the recapitulation line E68K GDU1 did not recapitulate the log1-2 suppressor mutant. Amino acid transport in these plants was similar to the gdu1-1D mutant, and not intermediate between this mutant and the wild type as expected from the analysis of the log1-2 suppressor mutant (Figures 2 and 5). The E68K GDU1 over-expressing plants displayed a phenotype similar to the plants over-expressing the E68R mutant, except that the size of the plants was similar to the wild type. It is not explained why the E68K GDU1-HA over-expressors do not behave as the log1-2 plants and will need further investigation. As suggested earlier, the addition of an HA tag to the protein could affect the function of the mutant protein and explain this discrepancy. It has also to be noted that the log1-2 mutant is tolerant to external amino acids, similar to the gdu1-1D mutant (Figure 8), but does not display size reduction, Gln secretion or early senescence.

These results suggest that the mutation of Glu68 to Lys, a positively charged residue, affects protein function in a way that the over-expressing plants display enhanced Gln transport but no dramatic growth alterations. Glu68 is not conserved in GDU sequences, and is sometimes replaced by Arg in GDU2 and GDU3 (Figure 1). Yet GDU2 and GDU3 over-expressors display the typical Gdu1D phenotype (small plants secreting Gln; Pratelli et al., 2010), showing that the function of the GDU2 and GDU3 proteins is not very different from GDU1, despite this difference in sequence.

Conclusion and Opportunities for Crop Engineering
This work showed that all specific mutations in GDU1 can dissociate the various components of the Gdu1D phenotype. In some cases, the amino acid transport is affected in plants over-expressing mutant GDU1 proteins as in gdu1-1D, but these plants grow similarly to the wild type (S32A, E68K, and S32D). In the other cases, the transport is less affected, and the plants show a wild type phenotype (G40D, S32L) or close to wild type phenotype (S28A+S32A). The residues important for each function of GDU1 are not necessarily born by conserved residues, or domains, like the membrane or VIMAG domains. The use of the wild type GDU1 protein as a tool to modify amino acid export was impaired by the associated effects on plant fitness in Arabidopsis (Pilot et al., 2004) and tobacco (Pratelli and Pilot, 2006). The mutant proteins described here could be used to modify amino acid transport, and in particular export, in the plant without affecting plant fitness. Targeting amino acid export in plants would be another strategy to modify allocation of amino acids between the plant organs, and ultimately control protein content in reserve organs, like seeds and roots.

Materials and Methods
Plant Material and Growth
Arabidopsis thaliana (ecotype Col-7) lines were grown under 120 µE/m²/s, 22°C, 16 h light/8 h dark on soil (Mix of Sunshine
Mix from below with 300 mg/l Miracle-Gro Fertilizer (24/8/16% (w/w) N/P/K; Scotts, Marysville, OH, USA). Rosette diameter of about eight plants from each line was measured with a ruler about 4 weeks after sowing. Arabidopsis thaliana were transformed by the floral dip method (Clough and Bent, 1998) using Agrobacterium tumefaciens GV3101 (pMP90). For transient expression of proteins in N. benthamiana, young leaves of 5-week-old plants were infiltrated with a suspension of Agrobacterium tumefaciens carrying the constructs of interest and the silencing suppressor p19 (Voinnet et al., 2003) according to (Batoko et al., 2000), with the following modifications. The bacteria were grown overnight in LB supplemented with appropriate antibiotics, washed twice in 10 mM MgCl2, 100 µM acetosyringone, and diluted to final OD600 of 0.05 in the same solution before infiltration in N. benthamiana leaves. Amino acid tolerance experiments were performed as described (Pratelli and Pilot, 2006): plants were grown for 10 days in long days conditions, on half-strength MS medium, supplemented with 0.5% sucrose, and 10 mM of indicated amino acids.

Cloning and Constructs

Primer sequences used for cloning are listed in Supplementary Table S1. The log1 variants of the GDU1 gene were cloned by PCR from genomic DNA and cloned into pDONR Zeo by the Gateway technology. The Kunkel method was used for site directed mutagenesis (Kunkel et al., 1991), from sequences cloned by Gateway cloning in the pDONR Zeo Gateway vector, containing the f1 replication origin (Lalonde et al., 2010). Mutagenized inserts were sequenced, and transferred by Gateway cloning (Life Technologies) to the binary vector pPWHTKan and pPWGTkan, derivative of pHA212K (Yoo et al., 2004; Pratelli and Pilot, unpublished data). pPWHTkan carries in this order the CaMV 35S promoter, the Gateway cassette, a double HA tag sequence and the terminator of the small subunit of the Rubisco from pea (Pisum sativum; accession no. X00806). pPWGTkan carries similar parts except that the HA tag is replaced by the enhanced GFP sequence.

Western Blotting and Co-immunoprecipitation

Protein extraction and western blotting were performed as previously described (Yu and Pilot, 2014): Leaves from each line (selected on kanamycin for 7 days, and transferred to soil and grown for three more weeks) were collected for protein extraction and western blot. Five hundred mg of leaves were ground with 1 ml of extraction buffer composed of 50 mM Tris-HCl, pH 7.3, 150 mM NaCl, 10 mM MgCl2, 10 mM DTT, 0.5% Nonidet P-40, and 1X Complete Protease Inhibitors (Roche) on ice. Homogenates were centrifuged at 14,000 g at 4°C for 15 min. Protein concentration of the supernatant was quantified by Bradford reagent. Twenty µg of total proteins were analyzed by SDS-PAGE (4–12% polyacrylamide MES gel; Life Technologies) and western blotting. Proteins were transferred on a nitrocellulose membrane (GE Healthcare) and detected using anti-HA (clone 3F10; Roche Diagnostics; 1:5,000) primary antibody, antirat (Thermo Scientific) secondary antibody, and the ECL-Prime western-blotting detection system (GE Healthcare). Co-immunoprecipitation experiments were performed from N. benthamiana infiltrated leaves as described (Pratelli et al., 2012).

Amino Acid Uptake

Amino acid uptakes in plants were performed as described (Pratelli et al., 2010), from segregating T2 seeds selected on kanamycin before growing in liquid medium and uptake.

Author Contributions

RP and GP designed and performed the suppressor screening. SY and RP characterized the suppressor mutants. SY and CD characterized the recapitulation lines and the mutagenized GDU1 proteins. SY and GP designed the experiments and wrote the paper.

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Supplementary Material

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fpls.2015.00593

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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