A Temperature-sensitive Mutation in the Arabidopsis thaliana Phosphomannomutase Gene Disrupts Protein Glycosylation and Triggers Cell Death*

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Eukaryotic phosphomannomutases (PMMs) catalyze the interconversion of mannose 6-phosphate to mannose 1-phosphate and are essential to the biosynthesis of GDP-mannose. As such, plant PMMs are involved in ascorbic acid (AsA) biosynthesis and N-glycosylation. We report on the conditional phenotype of the temperature-sensitive Arabidopsis thaliana pmm-12 mutant. Mutant seedlings were phenotypically similar to wild type seedlings when grown at 16–18 °C but died within several days after transfer to 28 °C. This phenotype was observed throughout both vegetative and reproductive development. Protein extracts derived from pmm-12 plants had lower PMM protein and enzyme activity levels. In vitro biochemical analysis of recombinant proteins showed that the mutant PMM protein was compromised in its catalytic efficiency ($K_{cat}/K_m$). Despite significantly decreased AsA levels in pmm-12 plants, AsA deficiency could not account for the observed phenotype. Since, at restrictive temperature, total glycoprotein patterns were altered and glycosylation of protein-disulfide isomerase was perturbed, we propose that a deficiency in protein glycosylation is responsible for the observed cell death phenotype.

Phosphomannomutases (PMMs) catalyze the interconversion of mannose 6-phosphate and mannose 1-phosphate and are required for the synthesis of GDP-mannose (Fig. 1). Both prokaryotes and eukaryotes utilize GDP-mannose in the synthesis of complex structural carbohydrates. Accordingly, this sugar nucleotide contributes to the synthesis of different structural carbohydrates in plant cell walls (1) and fulfills a key role in the biosynthesis of ascorbic acid (AsA) in plants (2, 3). GDP-mannose is also essential for post-translational modifications, such as protein glycosylation and glycosylphosphatidylinositol (GPI) anchoring in eukaryotes, because mannose, derived from GDP-mannose, is a crucial buliding block of the core glycan chain attached to the modified proteins (4, 5).

Because of their involvement in such fundamental processes, mutations in PMM genes often lead to severe and pleiotropic phenotypes. In the yeast Saccharomyces cerevisiae, a temperature-sensitive (ts) mutant affected in the PMM homolog SEC33 is arrested in growth at restrictive temperature. In fact, SEC33 has been classified as an essential gene because of the lethal phenotype of its null mutant (6, 7). Furthermore, PMM gene deletion mutants of the protozoan parasite Leishmania mexicana remain viable but are no longer able to establish an infection (8). In mouse, disruption of the PMM2 gene causes early embryonic lethality (9), and in humans mutations in PMM2 lie at the basis of Jaeken syndrome (also termed carbohydrate-deficient glycoprotein syndrome type Ia), a severe clinical disorder that provokes impaired neurological development and increased childhood mortality (10). In both S. cerevisiae and humans, defects in protein glycosylation due to lack of GDP-mannose have been pinpointed as the underlying cause for the observed phenotypes (7, 10).

The AsA deficiency of the Arabidopsis thaliana vitamin c1 (vtc1) mutant demonstrated that GDP-mannose is also involved in the biosynthesis of AsA (11). The VTC1 locus encodes a GDP-mannose pyrophosphorylase (GMPP), the enzyme that produces GDP-mannose from mannose 1-phos-

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8 The abbreviations used are: PMM, phosphomannomutase; AFLP, amplified fragment length polymorphism; AsA, ascorbic acid; ConA, concanavalin A; EMS, ethyl methanesulfonate; GMPP, GDP-ß-mannose pyrophosphorylase; GPI, glycosylphosphatidylinositol; Ndel, insertion/deletion; MS, Murashige and Skoog; PDI, protein-disulfide isomerase; ts, temperature-sensitive; BC, back-crossed; Mb, megabase(s).
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A mutation in the Arabidopsis PMM gene triggers cell death. PMM is required for the synthesis of GDP-mannose that is essential for AsA biosynthesis; posttranslational modifications, such as protein glycosylation and GPI anchoring; and the synthesis of different structural polysaccharides in the cell wall. PMM deficiency causes a marked drop in AsA content, developmental arrest, and subsequent cell death. The pmm-12 mutant plants also had a lower AsA content, but it is demonstrated that this AsA deficit cannot account for the cell death phenotype. Instead, we propose that a pmm deficiency in protein glycosylation is responsible for the observed cell death phenotype.

EXPERIMENTAL PROCEDURES

Plant Material, Growth Conditions, and Screening Procedure—A. thaliana (L.) Heynh. (ecotype Col-0) plants were grown either on Murashige and Skoog (MS) medium, including 10 g/liter sucrose or in soil at continuous light (≈60 μmol of photons m⁻² s⁻¹), unless otherwise stated. The permissive temperature used was either 16 or 18 °C, and the restrictive temperature was 28 °C.

The pmm-12 line was generated by treating ~200,000 seeds with 15 μM EMS for 24 h at 24 °C. Mutagenized seeds (M₁) were subdivided into batches consisting of ~1000 individual seeds (25 mg), grown at 18 °C, and allowed to self-pollinate, resulting in the corresponding M₂ seed batches. For 150 M₂ batches, aliquots of 25 mg of seeds were plated, grown at permissive temperature for 7 days, and subsequently screened for ts growth defects at restrictive temperature. Seedlings grown on MS medium containing 1 μM 1-naphthaleneacetic acid that showed growth defects within 4 days after transfer to restrictive temperature (compared with wild type) were considered putative mutants and subsequently rescued by transferring them back to the permissive temperature on fresh MS medium. Heritability of the observed phenotype was tested with M₃ plants produced by self-fertilization of M₂ plants of putative mutants. M₃ seeds were used to characterize the identified mutants.

For complementation, the AtPMM cDNA was first cloned into pDONR221 and subsequently cloned into the binary vector pK7WG2D (13) using Gateway technology (Invitrogen). M₃ plants were transformed at permissive temperature via the floral dip method (14).

DNA Extraction—Individual plants were ground and mixed with 400 μl of buffer composed of 200 mM Tris-HCl (pH 7.5), 2 M NaCl, 50 mM EDTA, and 2% N-cetyl-N,N,N-trimethylammonium bromide, followed by a 60-min incubation at 65 °C. After 15 min on ice, samples were extracted once with 250 μl of chloroform/isoamylalcohol (24:1). The aqueous phase was mixed with 200 μl of isopropyl alcohol, and samples were centrifuged. The DNA pellet was rinsed once with 70% ethanol, air-dried, and resuspended in 50 μl of 1 mM Tris-HCl (pH 8.0) and 0.1 mM EDTA.

Map-based Cloning and Fine Mapping—To obtain pmm-12 mutants with a more homogeneous genetic background, pmm-12 M₃ mutants were back-crossed (BC) into the Col-0 wild type genetic background. The resulting BC₁ plants were self-fertilized to obtain BC₁F₃ seeds, and the 3:1 (wild type/mutant) segregation ratio was tested at restrictive temperature.
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Subsequently, mutant BC₁F₂ plants were rescued by transfer to soil and growth at the permissive temperature. Self-pollination of these mutant plants led to lines whose genetic background was purified by 50%. Homogeneity of the resulting BC M₃ mutant lines was verified by growing at least 50 plants and checking the consistency of the mutant phenotype at restrictive temperature. To generate mapping populations, BC M₁ mutant lines were reciprocally crossed with Landsberg erecta (Ler) wild type.

In a first step, a genome-wide mapping procedure using the physical AFLP map described by Peters et al. (15, 16), was applied to 310 F₂ plants to determine the chromosomal location of the PMM gene. All AFLP markers, including the 85 Col-0/Ler framework AFLP markers resulting from a standard set of AFLP SacI/Msel primer combinations, were generated as described (17). AFLP markers were scored co-dominantly with the aid of the specific image analysis software AFLP-QuanTaqPro (Keygene Products, Wageningen, The Netherlands). For fine-mapping, a large population of 9,900 F₂ segregants were phenotyped, and 1,317 F₂ mutants (13.3%) were selected for further fine mapping of the mutant locus. In the case of a monogenic recessive trait, F₂ mutants are the individuals of choice. These 1,317 mutant F₂ segregants were subsequently genotyped for insertion/deletion (InDel) polymorphisms of the PMM frame of InDel markers flanking the genomic area containing the PMM gene. All AFLP markers, including the 85

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Sequence Analysis—PCR primers were designed based on the Arabidopsis information resource (TAIR) annotation (available on the World Wide Web) using Primer3 software (19) to amplify 500-bp exonic and intronic fragments having a minimal overlap of 100 bp. Amplicons were amplified by PCR using 50 ng of genomic DNA template, Silverstar DNA polymerase (Eurogentec, Seraing, Belgium), and a final concentration of 0.5 μM for each primer. The conditions used for the PCR amplification were as follows: 2 min at 95°C; 10 cycles of 30 s at 95°C, 30 s at Tₘ (depending on the primer sets), 2 min at 72°C. PCR products were purified with ExoSAP-IT (U.S. Biochemical Corp., Cleveland, Ohio) and transferred to a 96-well PCR plate (~10 ng/100 bp) together with 50 μM primer. After the plate had been dried at 65°C for 30–120 min, sequencing mix and buffer were added. The conditions for a second PCR were as follows: 2 min at 95°C; 40 cycles of 10 s at 95°C, 10 s at 48°C, 2 min at 60°C. Samples were sequenced with an automated sequencer (ABI3700 or ABI377; Applied Biosystems Inc., Foster City, CA).

Production of Recombinant Proteins—The open reading frame of PMM (at2g45790) and its mutant isoforms were PCR-amplified using cDNA obtained from wild type or mutant plants as a template. The pmm-1 mutation was introduced using the ATGGCGGCGAAAAATTCGCCAG primer, the pmm-2 mutation with cDNA from pmm-12 plants, and the above primer adapted to the wild type sequence. Fragments were cloned into pDONR221, verified by DNA sequence analysis, and then cloned into pDEST17 using Gateway technology (Invitrogen). Escherichia coli (strain BL21(DE3)pLysE) harboring the various constructs were, by means of a preculture, inoculated at OD 0.15 in 5 ml of Luria broth medium containing 100 μg/ml ampicillin and 25 μg/ml chloramphenicol. Cultures were subsequently grown for 1.5 h at 37°C and induced with 0.2 mM isopropyl-β-d-galactopyranoside. After 5 h of incubation at room temperature, 1 ml of bacteria was harvested. Recombinant proteins were purified with TALON metal affinity resin (Clontech), using 50 mM Hepes (pH 7.5), 300 mM NaCl, 10% glycerol, 1 mM phenylmethylsulfonyl fluoride (1:1.5, w/v) as equilibration buffer and equilibration buffer including 250 mM imidazole as elution buffer. Protein concentrations were determined according to Bradford (20).

Enzyme Activity Assays—PMM activity was determined via a coupled assay; the conversion of mannose 1-phosphate to mannose 6-phosphate was coupled to the reduction of NADP⁺ to NADPH (catalyzed by glucose 6-phosphate dehydrogenase) through the activities of phosphomannose isomerase and glucose-6-phosphate isomerase. The reaction was monitored through the increase in absorbance at 340 nm as a result of the increase in NADPH. In planta PMM activity was determined essentially as described (21). In short, leaf tissue ground in liquid nitrogen was resuspended in 50 mM Hepes (pH 7.6), 10 mM MgCl₂, 5 mM dithiothreitol, 1 mM EDTA, 1 mM ethylene glycol-bis(β-aminoethyl ether) N,N',N''-tetraacetic acid, 1 mM benzamidine hydrochloride, and 0.5 mM phenylmethylsulfonyl fluoride (1:1.5, w/v). Cell debris was removed by centrifugation, and sample supernatants were desalted on Sephrose G25 NAP5 columns (GE Healthcare). PMM activity was determined in a reaction mixture consisting of protein extract, 25 mM Hepes (pH 7.6), 10 mM MgCl₂, 0.1% bovine serum albumin, 1 mM NADP, 50 μM glucose 1,6-bisphosphate (as a hexose bisphosphate activator), 0.5 units/ml glucose-6-phosphate dehydrogenase, 0.5 units/ml phosphoglucone isomerase, and 0.5 units/ml phosphomannose isomerase. The reaction was initiated by the addition of the relevant concentration of d-mannose 1-phosphate, and the absorbance at 340 nm was recorded for 120 min. For each sample, material from at least 40 seedlings was pooled. In vitro activity of purified recombinant PMM was measured similarly in the same reaction mixture. The PMM content of each recombinant protein fraction was verified by protein gel blot analysis using penta-His antibody (Qiagen, Hilden, Germany).

Ascorbate Measurements and Protein Gel Blotting—The AsA content of control and mutant Arabidopsis leaves was determined as described previously (22, 23).

For concanavalin A (ConA) staining, 5 μg of total soluble (NAP5 column-purified) protein from Arabidopsis leaves was resolved on a 12.5% SDS-polyacrylamide gel and blotted onto a Hybond ECL membrane (GE Healthcare). The membrane was incubated for 1 h in PBSCT buffer (138 mM NaCl, 2.7 mM KCl, 10 mM Na₂HPO₄, 1.8 mM KH₂PO₄, 1 mM MgCl₂, 1 mM CaCl₂, 0.5% Tween 20, pH 7.2) and then for 1 h in PBSCT buffer con-
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Identification of a ts Cell Death Mutant—An EMS-mutagenized population of ~200,000 Arabidopsis thaliana (ecotype Col-0) seeds was screened for deficiencies in growth and development when grown at a restrictive temperature of 28 °C. Candidate mutants with severe growth-related phenotypes were rescued by transfer to permissive temperature (16 to 28 °C). Candidate mutants with severe growth-related phenotypes were rescued by transfer to permissive temperature (16 to 28 °C). Candidate mutants with severe growth-related phenotypes were rescued by transfer to permissive temperature (16 to 28 °C). Candidate mutants with severe growth-related phenotypes were rescued by transfer to permissive temperature (16 to 28 °C).

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son of wild type and mutant cDNAs and their deduced amino acid sequences revealed that point mutation \textit{pmm-1} resulted in a GGA to AGA codon change (codon 7), leading to a substitution of a neutral glycine residue with a basic arginine. The \textit{pmm-2} mutation resulted in a CGA to CAA codon change (codon 37), thereby replacing an arginine with a neutral glutamine (Fig. 3). PMMs are phosphotransferases belonging to the superfamily of haloalkanoic acid dehalogenases (27). Members of the phosphotransferase branch share a conserved N-terminal DXDX(T/V) motif. However, the \textit{pmm-2} mutation affected a highly conserved arginine residue, whereas the \textit{pmm-1} mutation involved the substitution of an apparently less conserved amino acid (Fig. 3).

**Complementation and Identification of a T-DNA Insertional Mutant**—To confirm that the mutations in the \textit{at2g45790} locus were responsible for the \textit{pmm-12} phenotype, homozygous \textit{pmm-12} plants were transformed with a construct containing the \textit{wild type} \textit{at2g45790} coding region under control of the CaMV 35 S promoter. T2 seeds with a 3:1 segregation ratio on selective medium, indicative of a single insertion locus, also unambiguously displayed a 3:1 phenotypic segregation ratio when grown on nonselective medium at restrictive temperature (data not shown). This indicated that the \textit{pmm-12} mutant phenotype could be complemented by over-expression of the \textit{at2g45790} locus.

We retrieved the T-DNA insertion line 045D04 from the GABI-Kat collection (29). This line contains an insertion 5 bp

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upstream from the start codon of the PMM gene, as verified via the standard GABI-Kat quality control on DNA extracted from T$_2$ seedlings (29). Unfortunately, none of the T$_2$ plants grown from the received seeds gave T$_3$ seeds that were 100% resistant to sulfadiazine. In other words, no transgenic lines homozygous for antibiotic resistance could be recovered. A closer look at T$_3$ plants revealed that the siliques of plants resistant to sulfadiazine lacked ~25% of the seeds, whereas siliques of nonresistant T$_3$ plants were undistinguishable from wild type siliques (data not shown), strongly suggesting that a knock-out mutation in at2g45790 causes embryo lethality.

The pmm-12 Mutant Has a ts Cell Death Phenotype—When grown in vitro at permissive temperature, 2-week-old pmm-12 mutant seedlings did not show any obvious phenotype when compared with wild type Col-0 seedlings. However, when transferred to 28 °C, growth was inhibited, and seedlings died within several days (Fig. 4A). The first signs of cell death visible to the naked eye consisted of tissue browning at the base of developing leaves and in the shoot meristem (Fig. 4B) and became apparent after 4 days at 28 °C. In 6-week-old plants grown in soil at 16 °C under a 10-h photoperiod, cell death became visible only after exposure to 28 °C for at least 2 weeks (Fig. 4C). In addition, mutant pmm-12 plants were germinated and grown in soil at 16 °C under continuous light conditions until flowering. These plants also displayed strong growth inhibition and completely wilted within 2 weeks after transfer to 28 °C (Fig. 4D).

Growth analysis of the primary root of germinating seedlings disclosed that, at permissive temperature, the growth rate of pmm-12 roots was slightly decreased compared with that of wild type roots (Fig. 5A). Upon transfer to restrictive temperature, 7 days after germination, root growth rates of wild type plants significantly increased until plants were transferred back to 18 °C, 4 days later. In contrast, root growth of 7-day-old pmm-12 seedlings was rapidly inhibited upon transfer to 28 °C. Root growth was completely arrested within 2 days and could not be rescued by placing the seedlings back to 18 °C after 4 days at 28 °C (Fig. 5B and C). When root explants were transferred to callus-inducing medium and placed at 28 °C, pmm-12 explants, in contrast to wild type explants, could not induce callus formation (data not shown). Taken together, we demonstrated that the ts cell death phenotype is conferred throughout both vegetative and reproductive development and that the progression of cell death depends on the plant's developmental stage.

pmm-12 Plants Exhibit Decreased PMM Enzyme Activity and Protein Levels—To determine the effect of the pmm-12 mutation on PMM activity in planta, total PMM activity was monitored in extracts from wild type and pmm-12 plants. Plants were first grown for 2 weeks at 16 °C and then transferred to 28 °C or kept at 16 °C. Shoot tissue was harvested 0 h, 24 h, 48 h, and 4 days after transfer, and PMM activity was monitored via a coupled assay. When grown at 16 °C, pmm-12 plants had PMM activity significantly lower than that of wild type plants. At the beginning of the experiment (0 h), the conversion velocity (V) measured in pmm-12 extracts was ~30% of that in wild type samples. After transfer to 28 °C, wild type plants showed a decrease in activity of ~25%, whereas activity in the mutant plants remained relatively stable (Fig. 6A).

Western blotting using a polyclonal AtPMM antibody revealed that pmm-12 plants had considerably decreased PMM protein levels, both at 16 and 28 °C. Furthermore, the temperature shift to 28 °C had a negative effect on protein levels in wild type and in mutant plants (Fig. 6B). We conclude that pmm-12 plants suffer from a considerable reduction in both total PMM activity and PMM protein levels.

The Different Mutant PMM Isoforms Have Altered Kinetic Properties—We assessed the impact of each of the identified point mutations on recombinant enzyme activity. Consequently, four different Arabidopsis PMM isoforms were tested: the wild type enzyme, two isoforms containing one of the identified mutations each (PMM-1 and PMM-2), and one isoform
containing both point mutations (PMM-12). Kinetic parameters of the His-tagged purified proteins were determined at both 18 and 28 °C. The different PMM isoforms displayed distinct kinetic properties. At 18 °C, the turnover numbers \( K_{\text{cat}} \) of wild type PMM and PMM-1 were nearly identical, whereas those of both PMM-2 and PMM-12 were markedly reduced, with \( K_{\text{cat}} \) values of 11 and 4% relative to wild type values, respectively. The \( K_m \) value of wild type PMM was 387 ± 28 μM, comparable with that of PMM-1 (475 ± 34 μM). Unexpectedly, the \( K_m \) of PMM-12 was roughly 2-fold lower. Nevertheless, the stacked mutations resulted in a catalytic efficiency \( (K_{\text{cat}}/K_m) \) that was only 8% of the wild type value. No differential effects of assay temperature on \( K_{\text{cat}}/K_m \) were observed (Table 1). Taken together, these data indicate that the decrease in catalytic activity of the mutant PMM-12 enzyme is largely due to the pmm-2 mutation. Second, the strong decrease in catalytic efficiency of PMM-12 observed at 28 °C is comparable with that at 18 °C.

The pmm-12 Cell Death Phenotype Is Not Caused by AsA Deficiency—Reduction of PMM expression levels through virus-induced gene silencing in Arabidopsis has been shown to result in decreased AsA levels (3). Therefore, we measured AsA levels in pmm-12 after transfer to restrictive temperature, using the ascorbate-deficient vtc1 mutant (11, 12) as a reference. In wild type plants, AsA levels increased significantly after transfer to 28 °C, whereas in pmm-12 plants they remained relatively stable and amounted to only 20–50% of wild type levels. In fact, AsA content in pmm-12 seedlings was very comparable with that in vtc1, either when expressed per g

**FIGURE 5.** Inhibition of pmm-12 root growth upon transfer to restrictive temperature. Primary root growth rates (in μm/h) of Col-0 wild type and pmm-12 mutant plants grown for 16 days at 18 °C (A) or for 7 days at 18 °C, 4 days at 28 °C, and finally placed back to 18 °C for 5 days (B). Each data point was calculated by dividing the average root length difference by the time. S.E. values are indicated (n = 8–14). C, typical plate with 7-day-old Col-0 wild type and pmm-12 mutant seedlings (grown at 18 °C), 4 days after transfer to 28 °C.

**FIGURE 6.** Reduction of PMM enzyme activity and protein levels in pmm-12 plants. A, PMM activity of total protein extracts isolated from both wild type (Col-0) and pmm-12 mutant seedlings, initially grown at 16 °C for 2 weeks and subsequently transferred to 28 °C for 24 h, 48 h, or 4 days. Proteins were isolated from at least 40 pooled seedlings, and equal amounts of total protein were used for each sample. As a substrate, 1 mM D-mannose 1-phosphate was used. Values are averages of three measurements. B, Western blot of protein extracts used in enzyme activity measurements probed with polyclonal Arabidopsis PMM antibody (α-PMM) detecting a protein of ~30 kDa. Equal amounts of protein (25 μg) were loaded in each lane. Error bars depict S.E.

The pmm-12 Cell Death Phenotype Is Not Caused by AsA Deficiency—Reduction of PMM expression levels through virus-induced gene silencing in N. benthamiana has been shown to result in decreased AsA levels (3). Therefore, we measured AsA levels in pmm-12 after transfer to restrictive temperature, using the ascorbate-deficient vtc1 mutant (11, 12) as a reference. In wild type plants, AsA levels increased significantly after transfer to 28 °C, whereas in pmm-12 plants they remained relatively stable and amounted to only 20–50% of wild type levels. In fact, AsA content in pmm-12 seedlings was very comparable with that in vtc1, either when expressed per g
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Table 1

| Protein | 18 °C $K_m$ (μM) | 28 °C $K_m$ (μM) | 18 °C $k_{cat}$ (s$^{-1}$) | 28 °C $k_{cat}$ (s$^{-1}$) | 18 °C $k_{cat}/K_m$ (μM$^{-1}$ s$^{-1}$) | 28 °C $k_{cat}/K_m$ (μM$^{-1}$ s$^{-1}$) |
|---------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| PMM-WT   | 387 ± 28        | 200 ± 15        | 4.13 ± 0.11     | 10.01 ± 0.23    | 10.7 $\times 10^3$ | 50.1 $\times 10^3$ |
| PMM-1 (G7R) | 475 ± 34      | 229 ± 18        | 4.18 ± 0.11     | 10.31 ± 0.25    | 8.8 $\times 10^3$ | 45.1 $\times 10^3$ |
| PMM-2 (R37Q) | 294 ± 21      | 111 ± 8         | 0.44 ± 0.01     | 0.67 ± 0.01     | 1.5 $\times 10^3$ | 6.0 $\times 10^3$  |
| PMM-12 (G7R/R37Q) | 204 ± 13    | 68 ± 5          | 0.18 ± 0.00     | 0.24 ± 0.00     | 0.9 $\times 10^3$ | 3.7 $\times 10^3$  |

**FIGURE 7.** Leaf AsA content per g fresh weight of 2-week-old wild type (Col-0), pmm-12, and vtc1 seedlings grown at 16 °C, after transfer to 28 °C. AsA levels were calculated from three independent observations. Error bars, S.E.

**FIGURE 8.** Effect of the pmm-12 mutation on protein glycosylation and GPI anchoring. Protein was extracted from both wild type (Col-0) and mutant seedlings, initially grown at 16 °C for 2 weeks and subsequently transferred to 28 °C for 6 h, 24 h, 48 h, 4 days, or 7 days. A, affinodetection of glycoproteins containing high mannose type N-glycans with ConA. Protein sizes (in kDa) are indicated on the right. B, N-glycosylation of PDI, visualized via protein gel blotting, in pmm-12 plants is affected by transfer to restrictive temperature. The arrows on the right point out the electrophoretic migration of the mono- and nonglycosylated forms of PDI, in contrast to the upper band corresponding to the fully (di)glycosylated protein. C, protein gel blot detection of SKU5 in wild type and pmm-12 plants after transfer to restrictive temperature.

Protein Glycosylation in the pmm-12 Mutant is Temperature-sensitive—GDP-mannose is required for the synthesis of the core glycan chain that is attached to N-linked glycoproteins (5). Therefore, we studied the effect of the temperature shift on total glycoproteins by affinodetection with ConA, a lectin that binds to the terminal mannose residue of N-linked oligosaccharides (30). Several differences in protein composition were observed in extracts from pmm-12 mutant plants transferred to 28 °C compared with extracts from plants kept at 16 °C or wild type plants. For example, there was a small but marked shift in the position of some bands at about 66-kDa protein, which moved to a lower molecular mass. Moreover, two new bands of ~30 kDa were detected following transfer to 28 °C (Fig. 8A).

f fresh weight (Fig. 7) or per g of chlorophyll (data not shown). However, despite their relatively low AsA content, vtc1 seedlings showed neither enhanced growth retardation nor cell death following transfer to 28 °C. In addition, we tried to complement the ts phenotype of pmm-12 with L-galactono-1,4-lactone, the immediate precursor of AsA. Although AsA levels increased in pmm-12 seedlings grown on MS medium supplemented with 6 mM L-galactono-1,4-lactone, cell death could not be prevented after transfer to 28 °C (data not shown). Taken together, these results indicate that the cell death phenotype observed in pmm-12 plants is not caused by AsA deficiency.

| FIGURE 8. Effect of the pmm-12 mutation on protein glycosylation and GPI anchoring. Protein was extracted from both wild type (Col-0) and mutant seedlings, initially grown at 16 °C for 2 weeks and subsequently transferred to 28 °C for 6 h, 24 h, 48 h, 4 days, or 7 days. A, affinodetection of glycoproteins containing high mannose type N-glycans with ConA. Protein sizes (in kDa) are indicated on the right. B, N-glycosylation of PDI, visualized via protein gel blotting, in pmm-12 plants is affected by transfer to restrictive temperature. The arrows on the right point out the electrophoretic migration of the mono- and nonglycosylated forms of PDI, in contrast to the upper band corresponding to the fully (di)glycosylated protein. C, protein gel blot detection of SKU5 in wild type and pmm-12 plants after transfer to restrictive temperature. |
A defect in N-glycosylation has been shown to result in the perturbed addition of N-glycans to PDI, a phenomenon that can be detected as a band shift on a protein gel blot (24, 31). PDI is an abundant protein of the endoplasmic reticulum that is posttranslationally modified with N-linked glycans (32, 33). We therefore analyzed the effect of the temperature shift on the mobility of PDI extracted from wild type and pmm-12 seedlings by using specific antibodies. PDI obtained from wild type seedlings, initially grown at 16 °C and subsequently transferred to 28 °C, had a stable molecular mass of ~62 kDa (Fig. 8 B). However, with PDI obtained from pmm-12 mutant seedlings, two bands of lower molecular mass appeared 48 h after transfer to 28 °C (Fig. 8B). On overexposed films, these bands were detectable 24 h after transfer to 28 °C (data not shown). Note that the cell death phenotype only became apparent 4 days after the temperature shift (Fig. 4).

Another posttranslational protein modification depending on GDP-mannose is the addition of a GPI membrane anchor (34). We used antibodies to Arabidopsis SKU5, a protein that has been shown to be degraded in the absence of a functional GPI anchor (35), to investigate whether in pmm-12 the temperature shift might also trigger a defect in GPI anchoring. SKU5 protein levels were decreased in mutant plants, 7 days after transfer to 28 °C (Fig. 8C) and 3 days after the appearance of the cell death phenotype (compare with Fig. 4B).

DISCUSSION

Map-based Cloning of the pmm-12 Mutation Locus—During a genetic screen, aimed at identifying EMS-induced ts mutations in Arabidopsis genes involved in growth and development, we isolated a ts mutant that died within several days upon transfer to restrictive temperature. The mutation was initially mapped to a 673-kb region on the lower arm of chromosome II by scoring a limited set of physically mapped AFLP markers segregating in 310 F2 individuals. AFLP markers segregating in the F2 population were co-dominantly scored (see “Experimental Procedures”). Co-dominant scoring of AFLP markers makes it possible to unequivocally distinguish homozygous from heterozygous plants based on quantitative measurements of the band intensities. This strategy allows the extraction of additional genetic information from the AFLP fingerprints resulting in a more precise delineation of the region of interest. Further fine mapping eventually revealed that the affected gene (at2g45790) encoded a PMM and, remarkably, that there were two point mutations present in the DNA sequence of the gene cloned from the pmm-12 mutant compared with the wild type gene present in the public data base.

The Temperature-sensitive pmm-12 Mutant Offers a Unique Tool for Functional Characterization of Plant PMM—Based on sequence similarity, the Arabidopsis genome contains a single PMM gene (at2g45790). During the preparation of this manuscript, it was reported that the encoded protein indeed showed PMM activity, effectively converting mannose 1-phosphate to mannose 6-phosphate (3). Overexpression and virus-induced gene silencing knockdown experiments in Arabidopsis and N. benthamiana, respectively, indicated a role for PMM in AsA biosynthesis. However, no pmm knock-out mutants were reported, and efforts to obtain knockdown lines via RNA interference were unsuccessful. The lack of such mutants impaired assessment of the possible involvement of PMM in processes other than AsA biosynthesis (3). Thus, the conditional phenotype of the ts pmm-12 mutant described here has enabled us to further characterize PMM functions in plants. The data presented show that the ts pmm-12 mutant is a unique tool with which to study the molecular function of the Arabidopsis PMM gene and to characterize the role of PMM and GDP-mannose in plant growth and development.

PMM Enzyme Activity Is Differentially Affected by the pmm-1 and pmm-2 Mutations—PMMs are members of the magnesium-dependent phosphotransferase superfamily of haloalkanoic acid dehalogenases that catalyze phosphoryl group transfer reactions (27). All members of the haloalkanoic acid dehalogenase superfamily, including PMMs, possess a tertiary structure consisting of a typical α/β core domain and a second cap domain, with the active site located at the domain-domain interface. Transfer of the phosphoryl group is mediated by a strongly conserved aspartate nucleophile in the core domain (28) that has a highly conserved tertiary structure, whereas the cap domain that determines substrate specificity is more divergent in sequence and length (36, 37). Both the pmm-1 and pmm-2 mutations are located in the core domain. However, by using the PSIREN tool (38), the GenTHREADER tool (39), and the crystal structure of human PMM1 as a template (37), we inferred that the affected amino acid residues are most probably not essential for catalysis or substrate binding (data not shown). Nevertheless, the identified point mutations might disrupt the highly conserved tertiary structure of the core domain, because both amino acid substitutions are rather drastic. The pmm-1 mutation changes a neutral aliphatic glycine residue into a positively charged basic arginine, whereas pmm-2 results in the substitution of an arginine into a negatively charged acidic glutamine. In addition, the arginine affected by pmm-2 is highly conserved among virtually all PMMs present in public data bases (data not shown), suggesting that it is important for adequate functioning of the protein.

In vitro assays with the different recombinant mutant PMM isoforms (performed at both 18 and 28 °C) showed that the pmm-2 mutation resulted in a significant reduction in PMM enzyme activity, whereas the pmm-1 mutation only marginally affected enzyme kinetics (Table 1). This result implies that, due to the location and/or nature of the amino acid substitution, the pmm-1 mutation alone would have few biological consequences. The activity assays also revealed that the observed decrease in catalytic efficiency of PMM-12 is independent of the assay temperature. When viewed together with the observed decreases in PMM protein and enzyme activity observed at 18 °C (Fig. 6), these results indicate that the pmm-12 mutant is not a typical ts mutant in the sense that the protein becomes inactive or unstable only at restrictive temperature. Rather, the data suggest that Arabidopsis plants growing at 18 °C can cope with reduced PMM enzymatic capacity, whereas in plants growing at 28 °C this becomes lethal.

Aberrant Protein Glycosylation Precedes the Cell Death Phenotype—Disturbed PMM enzyme activity will probably lead to GDP-mannose deficiency and have pleiotropic effects. PMM has been described to play an important role in the biosynthesis of N-linked glycoproteins (24, 31). Moreover, PMM activity is required for efficient AsA biosynthesis because AsA levels are only moderately decreased in pmm-12 mutant plants. Given that all the plants grown in the current study were pmm-1 pmm-2 double mutants, it is likely that the observed AsA deficiency is due to the pmm-2 mutation, and that PMM-1 is not essential in Arabidopsis for AsA biosynthesis.
of AsA (2, 3). Therefore, we first tested whether pmm-12 contained less AsA than wild type plants. Our measurements indicated that, after transfer to restrictive temperature, AsA levels in the pmm-12 mutant were only 20–50% of those in wild type plants and comparable with those in the AsA mutant vtc1 (Fig. 7). This result strengthens the genetic evidence for PMM involvement in AsA biosynthesis. During this experiment, it was noted that vtc1, unlike pmm-12, did not display cell death when transferred to 28 °C. Moreover, supplying the pmm-12 mutant with the immediate precursor of AsA, l-galactono-1,4-lactone, did not prevent the occurrence of cell death after transfer to 28 °C (data not shown). Taken together, these results demonstrate that the ts cell death phenotype observed in the pmm mutant is not caused by AsA deficiency.

Besides its role in AsA biosynthesis, GDP-mannose also plays a key role in posttranslational modifications, such as protein glycosylation. N-Glycosylation is essential for proper folding, targeting, and function of many secreted proteins. The core glycan attached to N-linked glycoproteins is assembled on a dolichol phosphate and contains nine mannose residues, which are transferred to it from GDP-mannose (4). The results presented here indicate that, shortly after the temperature shift, a deficiency in N-glycosylation occurs in pmm-12 plants. In addition, GPI anchoring, a second posttranslational protein modification depending on GDP-mannose (34), is affected several days later (Fig. 8). Based on these observations, we propose that the cell death phenotype of pmm-12 plants at restrictive temperature and the embryo lethality observed in the T-DNA insertion knock-out line are caused by strongly decreased PMM activity, eventually leading to a deficiency in N-glycosylation.

Various examples in the literature support the hypothesis that glycosylation defects can result in cell death and lethality. In both yeast and animals, aberrations in protein glycosylation have been linked to cell death. Mutations that block early steps in the assembly of core glycans (e.g. alg1, alg2, alg4, and vig9) are lethal (40, 41) as is the knock-out mutation of the yeast PMM gene SEC53 (6). Furthermore, glycosylation mutants with defects in the oligosaccharyltransferase protein complex were reported to undergo programmed cell death (42, 43). In Arabidopsis, mutant alleles of the gene encoding GCS1/KNF, a protein necessary for complex N-glycan formation, and the genes encoding DGL1 and STT3, two different subunits of the Arabidopsis oligosaccharyltransferase complex, have been described to result in aborted embryos or to produce nonviable seeds (31, 44–46).

The Arabidopsis cyt1 mutation abolishes GMPP gene function and results in embryo lethality by arresting embryonic development at a very early stage, providing strong evidence for the lethality of a severe GMPP deficiency. Characteristic aspects of the cyt1 phenotype can be mimicked with tunicamycin, an inhibitor of N-glycosylation. This observation led to the conclusion that a deficiency in this process can account for most of the phenotypic features of cyt1 (24). It is interesting to note that GMPP acts directly downstream of PMM in the GDP-mannose biosynthesis pathway (Fig. 1). However, similar to the pmm-12 mutant, cyt1 could not be rescued by exogenously applied AsA (24), suggesting that in both cases the lack of AsA is not sufficient to explain the developmental arrest and subsequent cell death. The cyt1 mutant is allelic to vtc1, a mutant deficient in its AsA biosynthesis (12) that was isolated by virtue of its sensitivity to oxidative stress (47). In marked contrast to the lethality caused by cyt1 mutations, vtc1 plants are viable and fertile, indicating that the vtc1 mutation is weak. Consistently, GMPP activity is reduced but not absent from vtc1 protein extracts (11).

The evidence described above demonstrates that the lethality of the pmm-12 mutation results from glycosylation defects rather than AsA depletion. It remains intriguing why the GDP-mannose depletion in pmm-12 is severe enough to result in cell death, whereas its AsA levels remain comparable to those in vtc1. The presence of alternative biosynthesis or recycling pathways for AsA that are independent of mannose would provide an explanation for this paradox (48). However, a recent analysis of the vtc2 vtc5 double mutants has revealed that the GDP-mannose pathway is the only significant source of AsA in Arabidopsis seedlings (49). Another potential explanation would be that pmm-12 prioritizes the flux through GDP-mannose toward AsA biosynthesis at the expense of glycosylation. More detailed comparisons of the regulation of AsA synthesis in the vtc mutants and pmm-12 are required in order to determine whether any form of prioritization exists in planta. Although this comparison is beyond the scope of the present paper, it is possible to consider factors that might influence the path of metabolite flow, such as toxic effects of elevated mannose 6-phosphate or a role for mannose 1-phosphate in cell survival.

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