The *Arabidopsis thaliana* nucleotide sugar transporter GONST2 is a functional homolog of GONST1

Beibei Jing1,2 | Toshiki Ishikawa3 | Nicole Soltis4 | Noriko Inada5 | Yan Liang1,2 | Gosia Murawska1,2| Lin Fang1,2 | Fekadu Andeberhan1,2 | Ramana Pidatala1,2 | Xiaolan Yu6 | Edward Baidoo1,2 | Maki Kawai-Yamada3 | Dominique Loque1,2 | Daniel J. Kliebenstein4 | Paul Dupree6 | Jenny C. Mortimer1,2,7

1 Joint BioEnergy Institute, Emeryville, CA, USA
2 Environmental Genomics and Systems Biology Division, Lawrence Berkeley National Laboratory, Berkeley, CA, USA
3 Graduate School of Science and Engineering, Saitama University, Japan
4 Plant Sciences Department, UC Davis, Davis, CA, USA
5 Graduate School of Biological Sciences, NAIST, Nara, Japan
6 Department of Biochemistry, University of Cambridge, Cambridge, UK
7 School of Agriculture, Food and Wine, University of Adelaide, Adelaide, SA, Australia

Correspondence
Jenny C. Mortimer, Biosciences Area, Lawrence Berkeley National Laboratory, Berkeley, CA 94720, USA.
Email: jcmortimer@lbl.gov

Present address
Noriko Inada, Graduate School of Life and Environmental Sciences, Osaka Prefecture University, Osaka, Japan
Gosia Murawska, Chemistry Department, Basel, Switzerland
Lin Fang, Guangdong Provincial Key Laboratory of Applied Botany, South China Botanical Garden, Chinese Academy of Sciences, Guangzhou, China

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Abstract
Glycosylinositolphosphorylceramides (GIPCs) are the predominant lipid in the outer leaflet of the plasma membrane. Characterized GIPC glycosylation mutants have severe or lethal plant phenotypes. However, the function of the glycosylation is unclear. Previously, we characterized *Arabidopsis thaliana* GONST1 and showed that it was a nucleotide sugar transporter which provides GDP-mannose for GIPC glycosylation. *gonst1* has a severe growth phenotype, as well as a constitutive defense response. Here, we characterize a mutant in GONST1’s closest homolog, GONST2. The *gonst2-1* allele has a minor change to GIPC headgroup glycosylation. Like other reported GIPC glycosylation mutants, *gonst1-1gonst2-1* has reduced cellulose, a cell wall polymer that is synthesized at the plasma membrane. The *gonst2-1* allele has increased resistance to a biotrophic pathogen *Golovinomyces orontii* but not the necrotrophic pathogen *Botrytis cinerea*. Expression of GONST2 under the GONST1 promoter can rescue the *gonst1* phenotype, indicating that GONST2 has a similar function to GONST1 in providing GDP-D-Man for GIPC mannosylation.
1 | INTRODUCTION

The plant plasma membrane is an asymmetric lipid bilayer which acts as both a selective barrier and a point of contact between the interior and exterior of the cell. Glycosylinositolphosphoryleramides (GIPCs) are a glycosylated form of sphingolipid, that comprise an estimated 64% of plant sphingolipids and ~25% of the total lipids in the Arabidopsis thaliana (Arabidopsis) leaf (Bure et al. 2011; Cacas et al. 2013, 2016; Markham & Jaworski, 2007; Markham et al. 2006, 2013). GIPCs are found predominantly in the outer leaflet of the plasma membrane.

GIPCs are a highly diverse class of lipids, comprising a long chain base (LCB) linked via an amide group to a fatty acid (FA) to form a ceramide, and a polar glycan head group. The diversity results from variation in the length, degree and position of unsaturation and hydroxylation of the FA and LCB, as well the structure and identity of the glycan head group. The ceramide is synthesized in the endoplasmic reticulum (ER), where it is either glucosylated to produce glucosylceramides, or it is then trafficked to the Golgi for GIPC biosynthesis. The first GIPC-specific step is the addition of an inositol phosphate to the ceramide via headgroup exchange with a phospholipid phosphatidylinositol by inositolphosphorylceramide (IPC) synthase (IPCS) (Wang et al. 2008). The IPC core is first glucosylated with a glucuronic acid (GlcA) by the Carbohydrate Active enYzeme (CAzy) family 8 glycosyltransferase (GT8) INOSITOL PHOSPHORYLKERAMIDE GLUCURONOSYLTRANSFERASE (IPUT1) (Rennie et al. 2014). The GlcA-IPC core is further glucosylated to form mature GIPCs (Bure et al. 2011; Cacas et al. 2016; Fang et al. 2016; Ishikawa et al. ,2016, 2018; Markham & Jaworski, 2007; Tellier et al. 2014). In Arabidopsis vegetative tissues, the dominant GIPC carries a mannose (Man) on the GlcA-IPC, which is added by GIPC MANNOSYL-TRANSFERASE1 (GMT1), from CAZy family GT64 (Fang et al. 2016). However, other glucosylated forms of GIPCs have also been identified. For example, in Arabidopsis seeds and pollen, rice, and tobacco leaves, the major GIPC glycosylation is a GlcN(Ac) linked to the GlcA, which can then be extensively decorated (Bure et al. 2011; Cacas et al. 2016; Carter et al. 1958; Hsieh et al. 1978, 1981; Ishikawa et al. 2018; Kaul & Lester, 1975, 1978; Luttegehrm et al. 2015; Tellier et al. 2014).

Recent evidence supports a role for GIPC glycosylation in plant-microbe interactions. For example, a peptide (NLP) which determines pathogenicity in many plant pathogens, including oomycetes, was shown to bind to the GIPC headgroup (Lenarcic et al. 2017). The degree of GIPC glycosylation was important in determining the degree of NLP cytotoxicity (Lenarcic et al. 2017). In Medicago, the type of GIPC glycosylation is important for the successful formation of root-microbial symbioses, both with nodulating bacteria and arbuscular mycorrhizal fungi (Moore et al. submitted), and in Arabidopsis, plants with mutated GIPC glycosylation display a constitutive hypersensitive response, including elevated salicylic acid (SA) and reactive oxygen species (ROS; Fang et al. 2016; Mortimer et al. 2013).

In addition to GIPC glycosylation, many other glycosylation reactions occur in the lumen of the Golgi, including the synthesis of polysaccharides, and glycoproteins. Nucleotide sugars are the universal sugar donors for these processes. In plants, the majority of nucleotide sugars are UDP-linked, but the GDP-linked sugars GDP-D-Mannose (GDP-Man), GDP-D-Glucose (GDP-Glc), GDP-L-Fucose (GDP-Fuc), and GDP-L-Galactose (GDP-Gal) are also critical (Barel & O’Neill, 2011). Most nucleotide sugars required in the Golgi, including all of the GDP-sugars, are synthesized in the cytosol and therefore need to be translocated into the Golgi lumen via nucleotide sugar transporters (NSTs). Many of the Arabidopsis NSTs have now been heterologously characterized (Bakker et al. 2005; Baldwin et al. 2001; Ebert et al. 2015; Handford et al. 2004; Mortimer et al. 2013; Niemann et al. 2015; Norambuena et al. 2002, 2005; Rautengarten et al. .2014, 2016, 2017; Reyes et al. 2010; Rollwitz et al. 2006; Saez-Aguayo et al. 2017), although in vivo functionality is described for far fewer. Arabidopsis NSTs belong to the NST/triose phosphate translocator (TPT) superfamily which has 51 members that are distributed in six clades (Rautengarten et al. 2014). From this superfamily, only four members, the GOLGI LOCALIZED NUCLEOTIDE SUGAR TRANSPORTER (GONST) subclade, are predicted to transport GDP-sugars due to the presence of the conserved GX[LV]NK motif (Baldwin et al. 2001; Gao et al. 2001; Handford et al. 2004).

The substrate for GMT1 is provided, at least in part, by GONST1, and indeed both gonst1 and gmt1 have very similar phenotypes (Fang et al. 2016; Mortimer et al. 2013). GONST1 was initially identified based on sequence similarity to Saccharomyces cerevisiae Vrg4p and Leishmania donovani LPG2 GDP-Man transporters (Baldwin et al. 2001). GONST1 can complement the Saccharomyces cerevisiae vrg4-2 mutant and was the first biochemically characterized plant NST (Baldwin et al. 2001). GONST1 can transport all four plant GDP-sugars in vitro (Mortimer et al. 2013). However, analysis of gonst1 plants revealed a specific role in vivo as a GDP-Man transporter which provides essential substrate for GIPC glycosylation (Figure 1) (Mortimer et al. 2013). GONST2 to GONST4 were identified as GONST1 homologues on the basis of their sequence similarity to GONST1 (Handford et al. 2004). GONST4 has now been characterized as the Golgi GDP-Fuc transporter and has therefore been renamed GDP-FUCOSE TRANSPORTER1 (GFT1) (Rautengarten et al. 2016). GONST3 has recently been shown to be responsible for GDP-Gal transport and has been renamed GOLGI GDP-L-GALACTOSE TRANSPORTER1 (GGLT1) (Sechet et al. 2018). GONST2 was also able to complement vrg4-2 (Handford et al. 2004) and able to transport all four GDP-linked sugars in vitro (Rautengarten et al. 2016), but its function in planta, as well as its specificity, remains unknown.
function to GONST1 in providing GDP-Man for GIPC mannosylation. Demonstrate that even Arabidopsis plants with minor modifications gonst1 rescue the phenotype, indicating that GONST2 has a similar phenotype, as described in Liang et al. (2019). Gonst2-2 and gonst2-3 alleles as described above. Arabidopsis seeds were surface sterilized and sown on solid medium containing 0.5x Murashige and Skoog salts including vitamins and 1% (w/v) sucrose. Following stratification (48 hr, 4°C, in the dark), plates were transferred to a growth room (22°C, 100–200 µmol/m² s⁻¹, 14 hr light/10 hr dark, 60% humidity). After 2–3 weeks, plants were transferred to soil or Magenta boxes under the same conditions. For G. orontii experiments, plants were grown under a 12 hr light/12 hr dark photoperiod for 4–5 weeks before inoculation. Liquid callus cultures were derived from Arabidopsis roots and maintained as described previously (Prime et al. 2000). 2.4 | Plant material and growth conditions The T-DNA line gonst2-1 (FLAG_406C01; ecotype Ws; insertion into AT1G07290) as well as gonst1-1 (FLAG_164D07; insertion into AT2G13650) were previously described in Mortimer et al. (2013). A second independent null GONST2 T-DNA insertion was not available, so two additional gonst2 alleles were generated using CRISPR/Cas9 gene editing technology as described below. Arabidopsis seeds were surface sterilized and sown on solid medium containing 0.5x Murashige and Skoog salts including vitamins and 1% (w/v) sucrose. Following stratification (48 hr, 4°C, in the dark), plates were transferred to a growth room (22°C, 100–200 µmol/m² s⁻¹, 14 hr light/10 hr dark, 60% humidity). After 2–3 weeks, plants were transferred to soil or Magenta boxes under the same conditions. For G. orontii experiments, plants were grown under a 12 hr light/12 hr dark photoperiod for 4–5 weeks before inoculation. Liquid callus cultures were derived from Arabidopsis roots and maintained as described previously (Prime et al. 2000). 2.5 | Subcellular localization Agrobacterium tumefaciens (GV3101) transformed with either 35S promoter (GONST2-YFP) or the Golgi marker Man49-GFP (Nelson et al. 2007) were co-infiltrated into 4-week-old tobacco leaves. An additional A. tumefaciens strain carrying the p19 plasmid was also co-infiltrated to stabilize the transgene expression. Forty-eight hours after infiltration, the epidermal cells were removed from the tobacco leaves, fixed with formaldehyde and imaged using a Zeiss LSM 710 (Carl Zeiss, http://www.zeiss.com/) as previously outlined (Parsons et al. 2012). Image analysis and processing (scale bar, brightness, and contrast) were performed using IMAGEJ (Version 1.6r) (Schneider et al. 2012). 2.6 | Histochemical detection of H₂O₂ Detection of H₂O₂ was by endogenous peroxidase-dependent histochemical staining using 3,3'-diaminobenzidine (DAB) as described in Mortimer et al. (2013). Leaves of 15-day-old agar-grown plants were submerged in 1 ml buffer (100 mM HEPES-KOH, pH 6.8) or 1 mg/ml

### 2.3 | Phylogenetics

Arabidopsis protein sequences were downloaded from TAIR (www.arabidopsis.org) and used in a BLASTp search (standard parameters) in NCBI. Sequences were aligned with Clustal Omega (www.ebi.org) (standard parameters). The Phylib program set (v3.95) was used to build the tree, using standard parameters except where stated, as follows: seqboot (2000 replicates), proml (not rough analysis), consense, and drawgram. All bootstrap probabilities were 1.0 with 2000 replicates.

### FIGURE 1  Phylogenetic characterization of Arabidopsis GONST family. Protein sequences of Arabidopsis GONST 1–5 were downloaded from TAIR (www.arabidopsis.org) and (a) aligned in Clustal Omega (standard parameters) in EMBL-EBI to obtain the sequence identity. (b) The Phylib program set (v3.95) was used to build the tree, using standard parameters except where stated, as follows: seqboot (2000 replicates), proml (not rough analysis), consense, and drawgram. All bootstrap probabilities were 1.0 with 2000 replicates.

|       | GONST1 | GONST2 | GONST3/GGLT1 | GONST4/GFT1 | GONST5 |
|-------|--------|--------|--------------|-------------|--------|
| Identity | 100%   | 100%   | 100%         | 100%        | 100%   |
| Alignment | 63.64% | 22.22% | 20.47%       | 21.55%      | 18.56% |
| Consense |        |        | 48.67%       | 16.29%      | 19.05% |

| GONST1 | GONST2 | GONST3/GGLT1 | GONST4/GFT1 | GONST5 |
|--------|--------|--------------|-------------|--------|
| AT1G07290 (GONST1) |
| AT2G13650 (GONST2) |
| AT5G19980 (GONST4/GFT1) |
| AT1G76340 (GONST3/GGLT1) |
| AT1G21870 (GONST5) |

Here, we characterize the function of GONST2 in planta and demonstrate that even Arabidopsis plants with minor modifications to their GIPCs have increased resistance to a biotrophic pathogen (Golovinomyces orontii), but not to a necrotrophic pathogen (Botrytis cinerea). Expression of GONST2 under the GONST1 promoter can rescue the gonst1 phenotype, indicating that GONST2 has a similar function to GONST1 in providing GDP-Man for GIPC mannosylation.

### 2 | MATERIALS AND METHODS

#### 2.1 | Materials

All constructs described in this publication are available upon request from the Joint BioEnergy Institute (JBEI)’s Inventory of Composable Elements (ICE) (registry.jbei.org). All chemicals are from Sigma Aldrich, unless otherwise noted. The GONST2 C-terminal YFP construct pEarleygate101 GONST2 was a generous gift from Dr. Carsten Rautengarten, Lawrence Berkeley National Laboratory under the 35S promoter, and was described previously (Rautengarten et al. 2016).

#### 2.2 | Samples

All experiments were performed on at least three independently grown biological replicates unless otherwise stated.

#### 2.3 | Phylogenetics

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#### 2.4 | Plant material and growth conditions

The T-DNA line gonst2-1 (FLAG_406C01; ecotype Ws; insertion into AT1G07290) as well as gonst1-1 (FLAG_164D07; insertion into AT2G13650) were previously described in Mortimer et al. (2013). A second independent null GONST2 T-DNA insertion was not available, so two additional gonst2 alleles were generated using CRISPR/Cas9 gene editing technology as described below. Arabidopsis seeds were surface sterilized and sown on solid medium containing 0.5x Murashige and Skoog salts including vitamins and 1% (w/v) sucrose. Following stratification (48 hr, 4°C, in the dark), plates were transferred to a growth room (22°C, 100–200 µmol/m² s⁻¹, 14 hr light/10 hr dark, 60% humidity). After 2–3 weeks, plants were transferred to soil or Magenta boxes under the same conditions. For G. orontii experiments, plants were grown under a 12 hr light/12 hr dark photoperiod for 4–5 weeks before inoculation. Liquid callus cultures were derived from Arabidopsis roots and maintained as described previously (Prime et al. 2000). gonst2-2 and gonst2-3 alleles are as described in Liang et al. (2019).

#### 2.5 | Subcellular localization

Agrobacterium tumefaciens (GV3101) transformed with either 35S promoter (GONST2-YFP) or the Golgi marker Man49-GFP (Nelson et al. 2007) were co-infiltrated into 4-week-old tobacco leaves. An additional A. tumefaciens strain carrying the p19 plasmid was also co-infiltrated to stabilize the transgene expression. Forty-eight hours after infiltration, the epidermal cells were removed from the tobacco leaves, fixed with formaldehyde and imaged using a Zeiss LSM 710 (Carl Zeiss, http://www.zeiss.com/) as previously outlined (Parsons et al. 2012). Image analysis and processing (scale bar, brightness, and contrast) were performed using IMAGEJ (Version 1.6r) (Schneider et al. 2012).

#### 2.6 | Histochemical detection of H₂O₂

Detection of H₂O₂ was by endogenous peroxidase-dependent histochemical staining using 3,3'-diaminobenzidine (DAB) as described in Mortimer et al. (2013). Leaves of 15-day-old agar-grown plants were submerged in 1 ml buffer (100 mM HEPES-KOH, pH 6.8) or 1 mg/ml
DAB in buffer. After 4 min of vacuum infiltration, leaves were incubated at 22°C under a light intensity of 100–200 μmol/m² s⁻¹ for 6 hr. Leaves were cleared for 30 min in 96% (v/v) ethanol solution at 70°C, and examined using a light microscope. Leaves were visually assessed as having either “no staining,” “light staining,” or “heavy staining.”

2.7 | Quantitation of SA

For total SA determination, 500 mg leaves were frozen and ground in liquid nitrogen. The powder obtained was mixed with 1 ml 80% (v/v) methanol and incubated for 15 min at 70°C. This step was repeated four times. Pooled extracts were centrifuged and filtered through Amicon Ultra centrifugal filters (10,000 Da MW cutoff, EMD Millipore, Billerica, MA). The conjugated SA in the filtered extracts was dried and the hydrolyzed in 1 N HCl at 95°C for 3 hr. The mixture was subjected to three ethyl acetate partitioning steps. Ethyl acetate fractions were pooled, dried in vacuo, and resuspended in 50% (v/v) methanol. SA was quantified using HPLC-electrospray ionization (ESI)-time-of-flight (TOF) MS. Details of the running condition were described previously (Eudes et al. 2013).

2.8 | G. orontii infection assay

G. orontii MGH was maintained on pad4 leaves (Inada et al. 2016), and WT or gonst2-1 leaves were inoculated using a settling tower method, as previously described (Plotnikova et al. 1998). Five days after inoculation, leaves were collected and cleared in 99% ethanol, stained with Trypan Blue (250 μg/ml Trypan Blue in 1:1:1 glycerol:lactic acid:water) for ~15 min at room temperature, destained (1:1:1 glycerol:lactic acid:water), and visualized under a light microscope. Three independent inoculation experiments were performed, and 12–30 leaves were used to count numbers of conidiophores per colony, for each genotype, in each experiment.

2.9 | B. cinerea infection assay

B. cinerea inoculation followed a previously described protocol (Denby et al. 2004; Klieberstein et al. 2005) using the following four strains: MEAP6G, 1.02.01, 1.03.01, and NobleRot. We grew seedlings in a randomized complete block design on soil (SunGro Horticulture, Agawam, MA) growth chambers in 20°C, short-day (8h photoperiod) conditions. Spores were collected from mature B. cinerea cultures grown on canned peach plates and diluted to 10 spores/μL in filter-sterilized 50% organic grape juice. At 7 weeks of age, we detached leaves from plants and arrayed them on 1% phytoagar by their order in the planting flats. We inoculated 4 μL spore solution droplets onto each leaf in a randomized complete block design, then incubated under 12 hr light/12 hr dark at room temperature for 96 hr. The spore solution was continuously agitated to ensure equal distribution of spores. Digital images were taken at 48, 72, and 96 hr post inoculation. Lesion area was measured using custom R scripts along with the EBImage and CRImage packages (Failmezger et al. 2012; Pau et al. 2010; RDevelopment CORE TEAM 2008).

2.10 | GIPC analysis by TLC

Powdered, lyophilized liquid-grown callus (200 mg) was added to 5 ml of the lower layer of isopropanol:hexane:water (55:20:25) and incubated at 50°C for 15 min. Following centrifugation (500 x g, 10 min), the supernatant was transferred to a fresh tube, and the pellet was re-extracted with a further 5 ml of the lower layer of isopropanol:hexane:water (55:20:25). The supernatants were combined, dried under N2, and de-esterified by incubation with 33% (v/v) methanolamine in ethanol:water (7:3) at 50°C for 1 hr. After centrifugation (500 g, 10 min), the supernatant was retained, dried under N2, and incubated in 1 ml of chloroform:methanol:ammonia:water (10:60:6:24) overnight at 21°C with agitation. Samples were subjected to weak anion exchange chromatography as described in Mortimer et al. (2013), and following elution from the cartridge were resuspended in chloroform:methanol:1.8 M ammonium hydroxide (4 M ammonium acetate) (9:7:2) and separated by thin layer chromatography (TLC) using high-performance-TLC Silica gel on glass plates (Merck) developed in the same buffer. GIPCs were visualized using primuline (Skipski, 1975).

2.11 | GIPC analysis by LC/MS

Total lipid for sphingolipidomics was prepared from lyophilized tissues (5–10 mg dry weight) using a methanol/butanol-based extraction coupled with weak alkaline hydrolysis and HCl treatment to remove glycerolipids and polysaccharides, respectively, according to the previous report (Ishikawa et al. 2018). Each sphingolipid species was quantified using LC-MS/MS (LCMS-8030, Shimadzu, Kyoto, Japan) with the MRM mode targeting glucosylceramides, free ceramides, and GIPCs with 0, 1, and 2 hexoses on GlcA-IPCs. The contents of Hex-GIPCs and ceramides were absolutely quantified by an internal standard-based calculation method, and GlcA-IPCs and Hex-Hex-GIPCs (for which we lack standards) were relatively quantified using the calculation factors as for Hex-GIPCs as previously described (Fang et al. 2016; Ishikawa et al. 2016).

2.12 | Cell wall monosaccharide analysis

AIR was prepared according to Mortimer et al. (2010) and 5 mg was hydrolyzed with fresh 2 M trifluoroacetic acid (TFA; 400 μL, 1 hr, 121°C). The supernatant was removed, and the pellet washed twice with water (400 μL). The supernatant and washings were combined,
dried in vacuo, and analyzed by HPAEC-PAD as previously described (Fang et al. 2016). The TFA-insoluble pellet was subjected to Saeman hydrolysis. Briefly, following incubation in 72% (v/v) sulfuric acid (63 µl, 21°C 1 hr), water was added to each sample to give a final sulfuric acid concentration of 1 M and incubated at 100°C for 3 hr. The samples were then neutralized with barium carbonate, to precipitate the sulfate ions, and the Glc content measured by HPAEC-PAD as above.

2.13 | Mannan structural analysis using PACE

PACE was performed according to Goubet et al. (2009) with slight modifications. Briefly, AIR (500 µg) was incubated with concentrated NH3 for 30 min at 21°C, and then dried in vacuo. Following resuspension in ammonium acetate buffer (0.1 M, 500 µl, pH 6.0), samples were incubated for 14 hr at 21°C with an excess of the mannanases CJMan5A and CJMan26A (a kind gift from Professor Harry Gilbert, University of Newcastle, UK). The released oligosaccharides were derivatized with 8-aminonaphthalene-1,3,6-trisulfonic acid (Invitrogen) with 2-picoline-borane as the reducing agent, and separated by electrophoresis in large-format polyacrylamide gels. Gels were visualized using a Syngene G:BOX gel doc system (Synoptics), equipped with long-wave UV transilluminator bulbs and appropriate filters.

2.14 | Promoter swap

The GONST1 promoter (1.3 kb upstream of the start codon) and GONST2 promoter (1.0 kb upstream of the start codon) were amplified by PCR from Col-0 genomic DNA, and cloned into the binary vector pCAMBIA1305 to obtain pCAMBIA1305 GONST1pro and pCAMBIA1305 GONST2pro. Full-length CDNA of GONST2 were amplified by PCR and cloned into pCAMBIA1305 GONST1pro and pCAMBIA1305 GONST2pro to obtain pCAMBIA1305 GONST1pro:GONST2 (Fusion 1) and pCAMBIA1305 GONST2pro:GONST2 (Fusion 2). Constructs were transformed into Agrobacterium tumefaciens strain GV3101 and used to transform gonst1-1 with the floral dip method. T3 plants, which were confirmed to be homozygous for the gonst1-1 T-DNA insertion (Mortimer et al. 2013), were analyzed.

3 | RESULTS

3.1 | GONST2 is a close homolog of GONST1

Arabidopsis nucleotide-sugar transporters containing a conserved GDP-binding motif (GX[L/V]NK) were first identified by Baldwin et al. (2001) and named GONST, and consist of a clade of four proteins (GONST1-2, GONST3/GGLT1, GONST4/GFT1) (Handford et al. 2004) (Figure 1). A fifth transporter (GONST5) is found in a distinct clade from GONST1-4 and lacks the GXLNK (Handford et al. 2004; Rautengarten et al. 2014). GONST2 shares 61% identity with GONST1 at the amino acid level, as compared to only 19% with GFT1 (Figure 1), and is expressed at a low level in most tissues (Figure S1). We also confirmed that GONST2 is localized to the Golgi, as previously reported (Rautengarten et al. 2016) (Figure S2).

3.2 | Use of CRISPR to generate new gonst2 alleles

Previously, we isolated and partially characterized a homozygous gonst2-1 allele (Ws ecotype) which lacked detectable GONST2 transcript by RT-PCR but did not have a visible phenotype (Mortimer et al. 2013). Since no further T-DNA lines were available, we used CRISPR/Cas9 gene editing to create two further gonst2 alleles, gonst2-2 and gonst2-3, in the Col-0 ecotype, as described in Liang et al. (2019) (Figure S3). As was the case for gonst2-1, gonst2-2, and gonst2-3 did not show a visible phenotype compared to WT.

3.3 | Loss of GONST2 enhances the gonst1 constitutive hypersensitive response

gonst1-1gonst2-1 has a more severe growth phenotype, as compared to gonst1 alone (Figure S4) (Mortimer et al. 2013). gonst1 has biochemical phenotypes consistent with the constitutive activation of plant defense responses, including elevated salicylic acid (SA) and reactive oxygen species (ROS) (Mortimer et al. 2013). To test whether loss of GONST2 also resulted in the constitutive activation of plant defense responses, we measured in situ H2O2 production using 2-aminobenzidine +staining (DAB, as a proxy for ROS) and SA in gonst2-1. gonst2-1 did not show a significant change in either SA or H2O2 production compared to WT (Figure 2). However, the gonst1-1gonst2-1 double mutant showed increased frequency of heavier DAB staining and significantly higher salicylic acid as compared to WT and gonst1-1 alone (Figure 2).

3.4 | gonst2-1 has increased resistance to a biotrophic pathogen but not to a necrotrophic pathogen

Plant pathogens can be divided into two major groups depending on their lifestyle strategies: necrotrophy and biotrophy. Nectrophic pathogens kill host cells and extract nutrition from the dead host, while biotrophic pathogens colonize living cells and obtain nutrition from living hosts (Hammond-Kosack & Jones, 1997). Classically, SA signaling triggers resistance against biotrophic pathogens, whereas a combination of jasmonic acid (JA) and ethylene (ET) signaling activates resistance against necrotrophic pathogens and these two pathways are mostly antagonistic (Robert-Seilaniantz et al. 2011). We wanted to test whether gonst1 or gonst2 plants show increased pathogen resistance, and whether this was generic or specific to biotrophic pathogens. However, gonst1-1 and gonst1-1gonst2-1 rosette
leaves are not suitable for pathogen assays, as they are fully senesced by ~20 days under normal conditions. Therefore, we tested gonzst2-1, despite the lack of a detectably significant increase in SA or ROS (Figure 2), since its rosette leaves are healthy. Indeed, gonzst2-1 showed a significant increase in resistance to the biotrophic pathogen G. gronitii MGH (an Arabidopsis-adapted powdery mildew) (Figure 3), as measured by conidiophores/colony. This was in contrast to pathoassays with the necrotrophic pathogen Botrytis cinerea. Four different isolates were tested, but there was no significant difference in susceptibility between WT and gonzst2-1 (Figure 3; Table 1).

3.5 | gonzst2-1 and gonzst1-1gonst2-1 have altered GIPC glycosylation

Previously, we had shown that gonzst1 has reduced GIPC mannosylation (Mortimer et al. 2013). To explore GIPC glycosylation in gonzst2, as well as gonzst1gonst2, we developed a simple thin layer chromatography (TLC) method which separates GIPCs primarily due to the nature and the degree of glycosylation. Due to the small stature and tissue death in gonzst1 and gonzst1gonst2, it was not possible to isolate GIPCs from whole plants. Therefore, we generated root-derived callus from all of the genotypes, isolated an enriched GIPC fraction, and performed TLC. The plates were stained with primuline and visualized under UV light (Figure 4a). gonzst1 was used as a control, and it shows a large shift in the mobility of the GIPCs, as compared to the Ws WT due to the loss of mannosylation as previously reported (Mortimer et al. 2013). A small fraction of the lower pool remains (marked with an arrow head), gonzst2 has a profile similar to WT with a small upper fraction of GIPCs (marked with an arrow head), and gonzst1-1gonst2-1 had essentially all GIPCs in the faster moving upper fraction.

To explore this further, we then used LC-MS/MS multiple reaction monitoring (MRM) to perform sphingolipidomics. No overall significant difference was detected in the glucosylceramides, hydroxyceramides or ceramides (Figure S5, Dataset S1). Plant GIPCs are enormously complex, due to the possible variations in FA, LCB, and glycan structure. Since no changes in the total amount of GIPCs nor the ceramide composition were detected (Figure S5). Following the nomenclature described in Fang et al. (2016) (Figure S6), the data have been aggregated to show the relative amount of GIPCs containing either 0, 1, or 2 hexoses terminal to GlcA-IPC (Figure 4b). While gonzst2-1 did not show a significantly different GIPC headgroup profile from the WT, there was significantly less Man-GlcA-GIPCs in gonzst1-1gonst2-1 than gonzst1-1 (t-test, p = 0.04, Figure 4b, Dataset S1).

3.6 | gonzst2-1 and gonzst1-1gonst2-1 Golgi-synthesized cell wall polysaccharides are unaffected

Since GONST2 is a Golgi-localized nucleotide sugar transporter, we next tested whether the loss of GONST2 could impact other glycosylation processes in the Golgi, in addition to GIPCs. The majority of cell wall polysaccharide biosynthesis, with the exception of cellulose and callose, occurs in the Golgi, so we investigated the monosaccharide composition of the non-cellulosic polysaccharides of callus, leaves, and stems by hydrolyzing an alcohol insoluble residue (AIR) cell wall preparation with trifluoroacetic acid (TFA) (Figure S7). No significant difference was detected between gonzst2-1, gonzst1-1gonst2-1, and the WT.
3.7 | gontst2-1 and gontst1-1gontst2-1 glucomannan structure and quantity are unchanged

Glucomannan is a Golgi-synthesized cell wall polysaccharide composed of β(1,4)-Man and α-Glc, which is synthesized by the CSLA family of GTs. CSLA9 (the dominant mannan synthase in Arabidopsis vegetative tissue) requires GDP-Man and GDP-Glc for glucomannan synthesis (Dhugga et al. 2004; Goubet et al. 2009; Liepmann et al. 2005), and it has been proposed that it has a luminal active site (Davis et al. 2010). However, no NST responsible for providing these substrates to the Golgi lumen has yet been identified. Loss of GONST1 does not affect glucomannan biosynthesis (Mortimer et al. 2013), GFT1 is a GDP-Fuc transporter (Rautengarten et al. 2016), and GGLT1 is a GDP-Gal transporter (Sechet et al. 2018). Since glucomannan is a relatively minor component of the cell wall (Handford et al. 2003), the monosaccharide analysis may not reveal alterations to its quantity or the Glc:Man ratio of the glucomannan backbone. Therefore, we used Polysaccharide Analysis by Carbohydrate gel Electrophoresis (PACE) to investigate glucomannan quantity and structure (Handford et al. 2003). No difference was seen either in the type or quantity of the oligosaccharides released by hydrolysis of gontst-1, gontst1-1gontst2-1, and WT AIR by mannanases (Figure 5). Therefore, this suggests that GONST1 and GONST2 are not providing substrate in the Golgi lumen for mannan biosynthesis.

3.8 | gontst1-1gontst2-1 has less cellulose

Previously, we showed that a mutant in GIPC mannosylation (GMT1) has reduced cellulose (Fang et al. 2016). To test whether this phenotype is common to plants with altered GIPC mannosylation, we hydrolyzed the TFA-insoluble AIR fraction with sulfuric acid to release glucose derived from cellulose (Figure 6). gontst1-1 had a significant reduction in upper and lower stem cellulose content as compared to WT, whereas callus and seedling were unaffected. gontst2-1 did not show a significant difference in any tissue type analyzed compared to WT. However, gontst1-1gontst2-1 mutants showed a significant decrease in callus cellulose content (a tissue rich in primary cell wall), compared to the WT or single mutants, but not in other tissue types. These data are consistent with a specific role for GIPC mannosylation in determining cell wall cellulose content.

3.9 | Expression of GONST1pro:GONST2 in gontst1-1 rescues growth and GIPC glycosylation

To test whether the phenotypic differences observed between gontst1 and gontst2 are due to differences in the protein function, or whether they are due to differences in expression level, we expressed the GONST2 coding sequence (CDS) driven by either the GONST1 promoter (GONST1pro:GONST2) or the GONST2 promoter (GONST2pro:GONST2) in the gontst1-1 background. Multiple independently transformed lines were selected for analysis. Analysis of T3 segregants revealed that some of the homozygous gontst1-1 plants had a restored growth phenotype (Figure 7). GONST2 expression was analyzed by real-time RT-PCR (Figure 7). The suppression of the gontst1-1 growth phenotype was only apparent in those lines in which GONST2 expression was driven by the GONST1 promoter (Figure 7). The rescue of the growth phenotype was reflected in the biochemical characterization of GIPC headgroup composition (Figure 7). This result supports the view that GONST2 has the same function as GONST1, and that the function is cell type specific and/or dose dependent.

![Figure 3](image-url) **Figure 3** Susceptibility of gontst2-1 plants to biotrophic and necrotrophic pathogens. (a) 5 days after inoculation with the biotroph G. orontii, leaves were harvested, stained with trypan blue and conidiophores per colony counted. The data represent the mean of 12–30 leaves per genotype per experiment, scored in three independent experiments, ±SD (Student’s t-test, * p < 0.05, *** p < 0.001). (b) 72 or 96 hr after inoculation with four phenotypically diverse B. cinerea isolates (1.02, 1.03, MEAP6G, and NobleRot), lesion size was measured. The data represent the mean of 105–166 leaves per plant genotype per experiment, ±SE. No significant difference was detected between WT and gontst2-1 (F-test; Table 1)
Resistance to biotrophic pathogens, such as the powdery mildew-causing G. orontii, is regulated by SA signaling (Wildermuth et al. 2001). Activation of SA signaling is often correlated with accumulation of reactive oxygen species including H₂O₂ (Herrera-Vasquez et al. 2015). Resistance to necrotrophic pathogens such as B. cinerea requires JA/ET signaling, which mostly function antagonistically with SA (Robert-Seilaniantz et al. 2011). We found that the gnost1-gnost2-1 double mutant contains significantly increased SA and enhanced H₂O₂ accumulation. While uninfected gnost2-1 did not show an increased SA level, it is possible that the SA level is enhanced in gnost2-1 after G. orontii infection, contributing to the enhanced resistance to G. orontii. It has been reported that altered ceramide profiles are associated with altered phytohormone levels, and thus with an altered response to pathogens (Magnin-Robert et al. 2015). In this case, ceramide functions as a signaling component. While gnost2 does not show a significant change to the ceramide pool, changes to the GIPC glycosylation may be enough to affect SA signaling and thus the response to G. orontii. Alternatively, a defect in membrane trafficking in gnost2-1 may negatively impact G. orontii infection. G. orontii forms a specialized infection hypha called the haustorium in the host apoplast to establish infection. The haustorium is surrounded by host-derived membrane called the extrahaustorial membrane, which has modified endosomal characteristics (Inada, Betsuyaku, et al., 2016). It has been shown that GIPCs are important for secretory sorting of proteins (Markham et al. 2011; Wattelet-Boyer et al. 2016), and therefore, it may be that changes to a minor class of GIPC are enough to disrupt these processes, thereby negatively affecting G. orontii infection. GIPC glycan engineering therefore offers a promising approach for developing plants with increased disease resistance.

Cellulose content is decreased in gnost1-1 and gnost1-1-gnost2-1 plants. Cellulose is synthesized at the plasma membrane by rosettes of CESA proteins which move through the plane of the plasma membrane (McFarlane et al. 2014). The rosettes are assembled in the Golgi and are delivered to the plasma membrane via the secretory system (Wightman & Turner, 2010). The reduced cellulose

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### TABLE 1

ANOVA results for the various factors in the Botrytis cinerea infection experiment on WT and gnost2 plants at 72 hr post inoculation

| Sources of variation | df | SS   | LRT  | p     |
|----------------------|----|------|------|-------|
| Botrytis isolates    | 3  | 1.623| <.001|       |
| WT versus. gnost2    | 1  | 0.00963| .3784|       |
| Time                 | 1  | 0.04897| .1171|       |
| Isolate x plant genotype | 3  | 0.2676| .8487|       |
| Random effects       |    |      |      |       |
| Tray/flat            | 1  | 1.3844| .2394|       |
| Tray                 | 1  | 9.1055| .002548|    |      |

Abbreviations: df, degrees of freedoms; p, estimated p-value; SS, Type III Sums-of-Squares.

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**FIGURE 4** Glycan headgroup composition of gnost GIPCs (a) TLC of a GIPC-enriched membrane fraction which has been stained with primuline. Bands discussed in the text are marked with a red arrow head. (b) An enriched GIPC fraction was analyzed by LC-MS/MS MRM. The data here are collapsed to describe only the number of hexoses on the GIPC headgroup. All data are mean ± SD of three independently grown replicates of liquid grown cell culture; asterisk indicates significant difference from the wild type (Student’s t-test, *** p < .001). The full dataset is shown in Figure S5 and Dataset S1.
phenotype was also reported for \textit{gmt1}, which has the same biochemical GIPC phenotype as \textit{gonst1gonst2} (Fang et al. 2016). The reasons for this decrease are not clear. It is possible that the altered GIPC glycosylation affects trafficking of the rosettes to the plasma membrane, or alternatively, the change to plasma membrane composition affects CESA function. CESA proteins are S-acylated, and it has been suggested that this decoration may either localize proteins to lipid microdomains (which are rich in GIPCs) or even facilitate their formation (Konrad & Ott, 2015; Kumar et al. 2016). COBRA and COBRA-like proteins which are also essential for normal cellulose biosynthesis are glycosylinositolphosphatidylinositol (GPI) anchored (Roudier et al. 2005). GPI anchored proteins are targeted to the outer leaflet of the plasma membrane, and to lipid microdomains (Borner et al. 2005). Therefore, correct GIPC glycosylation may be necessary for either CESA activity or localization and retention of GPI-anchored proteins in the plasma membrane. More recently, a role for GIPCs in modulating the salt-dependent activation of a plasma membrane calcium channel (Jiang et al. 2019) and for plasmodesmata function (Yan et al. 2019) suggesting that GIPCs may have a broad role in regulating plasma membrane functionality.

It should be noted that alterations to cellulose content can affect susceptibility to some pathogens (Hernandez-Blanco et al. 2007; Malinovsky et al. 2014). However, some difference has been observed between whether primary wall or secondary wall cellulose is impaired, and the type of immune response that is induced (Bacete et al. 2018). For example, CESA3 mutants (a primary cell wall CESA) are more resistant to powdery mildews (Cano-Delgado et al. 2003; Ellis & Turner, 2001) and mutants in secondary cell wall CESAs and secondary cell wall deposition are more resistant to necrotrophs (Hernandez-Blanco et al. 2007; Ramirez et al. 2011).

Mannan content is unchanged in the \textit{gonst1gonst2} plants. It had been reported that CSLA9, unlike related GT2 proteins (CSLC4, CESAs), has a topology which results in a luminal active site (Davis et al. 2010). This would necessarily require a nucleotide transporter to provide GDP-sugars for mannan biosynthesis. However, none of the predicted GDP-sugar transporters seem to have this function in planta (Mortimer et al. 2013; Rautengarten et al. 2016; Sechet et al. 2018). This implies that either the mannann synthases do not require a nucleotide sugar transporter, or that the transporter does not have a canonical GDP-binding motif.

Future work will be required to confirm these data, making use of the additional \textit{gonst2} alleles now available (Liang et al. 2019). It will also be important to establish how GIPC glycosylation affects these assorted membrane-based processes. For example, molecular dynamics could be applied to model the plant plasma membrane and understand how the GIPC glycan headgroup structure affects protein movement within the membrane. It will also be interesting to understand what drives the differences in functionality of the NSTs in in vitro assays versus in planta function. Both GONST1 and GONST2 can transport all GDP-sugars when tested in liposome-based assays (Mortimer et al. 2013; Rautengarten et al. 2016), but it is clear that they are highly specific in vivo. This could be mediated by substrate concentration, interaction with non-catalytic proteins, or interactions with the GT that utilizes the substrate (in this case GMT1 (Fang et al. 2016)). The recent crystal structure of the yeast Vrg4 NST provided new insights into how NST function is regulated.
(Parker & Newstead, 2017). To our knowledge, no plant NSTs have yet been structurally characterized, but we expect that this information will be critical for understanding NST specificity.

FIGURE 7 Expression of GONST2 under the GONST1 promoter in the gns1 background restores the dwarfed phenotype of gns1. (a) Schematic of the two introduced constructs. (b) Top row: 15-day-old, agar grown WT, gns1-1, Fusion1 T-1, Fusion1 T-2, Fusion2 T-3, and Fusion2 T-4 seedlings. Scale bar = 1 cm. Bottom row: 6-week-old WT, gns1-1, Fusion1 T-1, Fusion1 T-2, Fusion2 T-3, and Fusion2 T-4. Plants were first grown on agar for 10 days, and then transplanted onto soil. Scale bar = 3 cm. (c) Gene expression analysis of GONST2 relative to WTWs and normalized against TUBULIN using Q-PCR. Values represent average of three biological replicates ±SD. (d) An enriched GIPC fraction was analyzed by LC-MS/MS MRM. The data here are collapsed to describe only the number of hexoses on the GIPC headgroup. All data are mean ± SD of three independently grown replicates of liquid grown cell culture. Asterisk indicates significant difference from the wild type (Student’s t-test, * p < .05, *** p < .001).

ACCESSION NUMBERS
GONST1 (Q941R4.2) and GONST2 (AEE28103.2).

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CONFLICTS OF INTEREST
No conflicts of interest to declare.

AUTHOR CONTRIBUTIONS
JCM and BJ designed the research. BJ, JCM, FA, GM, LF, and RP carried out the research. TI performed spherolitomics; NS and NI performed pathogen assays; EB analyzed salicylic acid quantities; YL generated CRISPR lines; XY analyzed mannan structure by PACE. JCM, BJ, TI, NS, NI, MKY, DL, and PD analyzed data.

ORCID
Gosia Murawska https://orcid.org/0000-0003-1822-9607
Daniel J. Kliebenstein https://orcid.org/0000-0001-5759-3175
Paul Dupree https://orcid.org/0000-0001-9270-6286
Jenny C. Mortimer https://orcid.org/0000-0001-6624-636X

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SUPPORTING INFORMATION

Additional Supporting Information may be found online in the Supporting Information section.

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