Mutational Analysis of the Glycosylphosphatidylinositol (GPI) Anchor Pathway Demonstrates that GPI-Anchored Proteins Are Required for Cell Wall Biogenesis and Normal Hyphal Growth in *Neurospora crassa*

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Using mutational and proteomic approaches, we have demonstrated the importance of the glycosylphosphatidylinositol (GPI) anchor pathway for cell wall synthesis and integrity and for the overall morphology of the filamentous fungus *Neurospora crassa*. Mutants affected in the *gpi*-*1*, *gpi*-*2*, *gpi*-*3*, and *gpi*-*4* genes, which encode components of the *N. crassa* GPI anchor biosynthetic pathway, have been characterized. GPI anchor mutants exhibit colonial morphologies, significantly reduced rates of growth, altered hyphal growth patterns, considerable cellular lysis, and an abnormal “cell-within-a-cell” phenotype. The mutants are deficient in the production of GPI-anchored proteins, verifying the requirement of each altered gene for the process of GPI-anchoring. The mutant cell walls are abnormally weak, contain reduced amounts of protein, and have an altered carbohydrate composition. The mutant cell walls lack a number of GPI-anchored proteins, putatively involved in cell wall biogenesis and remodeling. From these studies, we conclude that the GPI anchor pathway is critical for proper cell wall structure and function in *N. crassa*.

In eukaryotic cells, a number of proteins are anchored to the outer leaflet of the plasma membrane via glycosylphosphatidylinositol (GPI) anchors. The presence of the GPI anchor is thought to play an important role in the trafficking of these proteins and providing them with an attachment to the plasma membrane, and in the case of fungi, to the cell wall as well (24, 41). Proteins destined to receive a GPI anchor are directed into the lumen of the endoplasmic reticulum (ER) by a typical signal peptide. The carboxyl termini of these proteins have a sequence motif that is recognized by a protein complex located in the ER, known as the GPI transamidase. The GPI transamidase complex cleaves the substrate protein at a position within this motif, termed the omega site, and transfers the GPI anchor en bloc to the newly generated C terminus of the protein.

The structures of the GPI anchor in the trypanosome, yeast, and mammalian systems have been determined. Although there are differences in the various substrates present on the GPI anchors produced by these organisms, all GPI anchors appear to share a common core structure (15, 19, 20). This core structure consists of a phosphatidylinositol (or inositol-containing sphingolipid) with an attached oligosaccharide chain that is terminated with a phosphoethanolamine residue. The linkages between the sugar units within the carbohydrate chain are conserved, and the amino group of the phosphoethanolamine moiety is used to attach the GPI anchor to the C terminus of the target protein. The organization of this basic GPI anchor structure is as follows: protein—phosphoethanolamine—1Mannose1—2Mannose1—6Mannose1—4Glucosamine1—inositol—phospholipid.

The process of GPI anchor production and attachment is mediated by the concerted actions of approximately 20 proteins, which are organized into biosynthetic complexes in the ER membrane. Seven primary steps have been identified in the GPI anchor pathway, beginning with anchor biosynthesis and concluding with the final attachment of the completed anchor structure to the recipient protein (15, 36). Some of the proteins involved in the biosynthesis and attachment of the GPI anchor catalyze the steps in the pathway, while others function as auxiliary factors.

In our analysis of the GPI anchor pathway in the filamentous fungus *Neurospora crassa*, we have focused on the functions and components of the phosphoethanolamine transferase and GPI transamidase complexes. The phosphoethanolamine transferase complex has been shown to consist of at least 4 components in mammals and *Saccharomyces cerevisiae*. The PIG-N/Mcd4p, hGPI7/Gpi7p, and PIG-O/Gpi13p proteins are involved in the addition of phosphoethanolamine substituents to the first, second, and third mannose residues in the mammalian/S. cerevisiae GPI-anchors, respectively (4, 26, 30, 31, 64, 66). The PIG-F/Gpi11p proteins serve as an auxiliary factor within the phosphoethanolamine transferase complex (31, 64, 66). The GPI transamidase complex is minimally composed of 5 proteins in both the mammalian and *Saccharomyces cerevisiae* systems. The GPI8/Pgi8p proteins likely function as the proteolytic activity of the complex, which cleaves the target protein at the omega site (48). The PIG-T/Gpi16p proteins have been shown to be important for the formation and stabilization of the GPI transamidase complex (22, 48, 49). The PIG-U/Cde91p proteins have been implicated in the recognition of either the GPI anchor attachment motif in the substrate.

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protein or the long chain fatty acids of the GPI anchor (32). The functions of the remaining GPI transamidase subunits, PIG-S/Gpl17p and GAA1/Gaa1p, are somewhat unclear, but the mammalian GAA1 has been shown to be involved in the binding of the GPI anchor (68). Homologs of each of these mammalian and S. cerevisiae phosphoethanolamine transferase and GPI transamidase complex components exist in N. crassa.

We have identified and characterized N. crassa mutants affected in three genes encoding components of the phosphoethanolamine transferase complex and one gene encoding a component of the GPI transamidase complex. We have also further characterized a previously identified N. crassa mutant affected in the enzymatic activity of the N-acetylglucosamine transferase complex, which catalyzes the transfer of N-acetylglucosamine to phosphatidylinositol during the first step in the GPI anchor biosynthetic pathway. Mutants affected in these genes are unable to make normal hypphae and fail to produce many of the typical cell types found in the N. crassa life cycle. The mutants have a vastly reduced rate of growth and grow in a tight colonial manner. Functional studies demonstrate that these mutants produce a weaker, altered cell wall. Electron micrographs of mutant cells illustrate an unusual “cell-within-a-cell” morphology, which we attribute to a defective cell wall. In addition, we show that the mutants generate a cell wall that differs extensively from the normal hyphal cell wall in carbohydrate and protein components. We conclude that GPI anchoring plays an important role in the biosynthesis, structure, and function of the N. crassa cell wall.

**MATERIALS AND METHODS**

**Strains and culturing conditions.** The arg-12 (FGSC 1527) and GTH-16 strains of N. crassa were used as the wild-type parental strains for the isolation of mutants affected in the gpip-1, gpip-2, gpip-3, and gpi1 genes, respectively. The arg-12 strain was obtained from the Fungal Genetics Stock Center (Kansas City, Kansas). The GTH-16 strain has an al-2 arg-9 inv qa-2 genotype (37). All cells were grown on supplemented Vogel’s medium as described by Davis and DeSerres (12). Gene mapping experiments were conducted by mating strains on a corn meal agar medium supplemented with needed amino acids, vitamins, and 0.4% glucose and using standard mapping procedures (12). The gpip-1 mutant, a temperature-sensitive mutant affected in the catalytic subunit of the N-acetylglucosamine transferase complex, was obtained from Seiler and Plamann (60).

**Isolation of the MSA-7 mutant.** The MSA-7 mutant, which contains a mutation in the gpip-1 gene, was isolated in a mutant screening experiment designed to identify mutants affected in the process of cell fusion. Conidia from the arg-12 strain were harvested into 10 ml of sterile water, transferred to petri dishes, and mutagenized with a 10-min exposure to a UV light source held at a distance of 10 cm above the petri dish. The UV mutagenesis resulted in a 99.7% killing of the cells. Mutants affected in three genes encoding components of total cellular extracts, the gpip-3 mutant and the GTH-16 wild-type parental strain were grown on cellophane on standard agar medium. The cells were harvested, ground on liquid nitrogen to a fine powder, and resuspended in a solution of 50 mM Tris–Cl (pH 7.5) and 1% sodium dodecyl sulfate (SDS). Each extract was briefly sonicated, boiled, and centrifuged at 1,000 × g to pellet cell wall debris. The protein concentrations of the soluble fractions were determined using the Bio-Rad DC protein assay kit (Bio-Rad Laboratories, Hercules, CA). Forty micrograms of total protein from each extract was separated on a 4 to 12% Bis-Tris NuPAGE gel (Invitrogen Life Technologies, Carlsbad, CA) and transferred to a polyvinylidene difluoride membrane.

**Western blot analysis of the GPIP-3 protein in a wild-type membrane preparation.** The GTH-16 strain was grown, harvested, and ground as before. The frozen, powdered mycelia were then resuspended in an ice-cold solution containing 50 mM Tris–Cl (pH 7.5), 200 mM NaCl, 10 mM dithiothreitol (DTT), 1 mM phenylmethylsulfonyl fluoride, and a protease inhibitor cocktail (product number P8215 from Sigma Aldrich, St. Louis, MO) at a 1/15,000 dilution.

For Western blot analysis of the GPIP-3 protein in a wild-type membrane preparation, the GTH-16 strain was grown, harvested, and ground as before. The frozen, powdered mycelia were then resuspended in an ice-cold solution containing 50 mM Tris–Cl (pH 7.5), 200 mM NaCl, 10 mM dithiothreitol (DTT), 1 mM phenylmethylsulfonyl fluoride, and a protease inhibitor cocktail (product number P8215 from Sigma Aldrich, St. Louis, MO). The extract was preclarified by centrifugation at 100,000 × g for 1 h. The 100,000 × g supernatant was discarded, and the centripetal fraction containing the total cellular membrane was resuspended in 50 mM Tris–Cl (pH 7.5) and 1% SDS. The determination of the protein concentration, SDS–polyacrylamide gel electrophoresis (PAGE), and Western blot analysis of the sample were as described above.

**Assessment of gross and hyphal morphologies.** To assess the gross colony morphology of each strain, small inocula of the gpip-1, gpip-2, gpip-3, and gpi1 mutants and the wild-type parents were made in the center of petri dishes containing standard agar growth medium. All strains, with the exception of the gpip-1 temperature-sensitive mutant, which was placed at 39°C, were allowed to grow at room temperature for the times indicated in Fig. 1. Images of the plates were then captured using a digital scanner.

**PCR analysis and sequencing of MSA-7 candidate genes.** All PCR experiments were carried out using primer oligonucleotides designed to amplify genes of interest in the short genomic region identified as containing the mutant gene. The sequences for the oligonucleotides were derived from the published genomic DNA sequence provided by the Neurospora genome project at the Broad Institute/MIT Center for Genome Research. Genomic DNA was isolated from all strains using the Trizol reagent as described in the manufacturer’s instructions (Invitrogen, Carlsbad, CA). The amplified genomic DNA regions were sequenced at Retrogen, Inc. (San Diego, CA).

**Use of RIP to isolate gpip-2, gpip-3, and gpi1 mutants.** Mutants affected in the gpip-2, gpip-3, and gpi1 genes were obtained using the Neurospora RIP (repeat-induced point mutation) phenomenon. RIP is a process in which multiple point mutations (C to T and G to A mutations) occur in DNA regions that are found in two or more copies in the haploid Neurospora genome during the premeiotic phase of one mating process (37). These mutations are generated in both copies of the duplicated DNA sequences. To produce mutations in the gpip-2, gpip-3, and gpi1 genes, PCR-amplified sequences from each gene were subcloned into the prAL1 vector (1) and the gene/prAL1 constructs were used to transform the N. crassa GTH-16 strain. The prAL1 vector includes a copy of the N. crassa qa-2 gene and can be used to select for transformants (37). The resulting transformants were then mated with the inl strain (FGSC 1453), and mutant progeny were identified by virtue of the colonial growth phenotype. The mutations present in the gpip-2, gpip-3, and gpi1 genes from these RIP-generated mutants were then identified by PCR amplification and sequencing of the gene as described above.

**GPIP-3 and ACW-1 antibody production and Western blot analyses.** Peptides representing amino acid numbers 388 to 407 (NH2-PPKVPFGTKPEVTPATAPK-COOH) of the predicted GPIP-3 protein and 186 to 208 (NH2-HQANGLDMEVFGPNLIWMNMAI-COOH) of the predicted ACW-1 protein were synthesized and used to immunize rabbits (Proteintech Group, Inc., Chicago, IL).

For Western blot analysis of total cellular extracts, the gpip-3 mutant and the GTH-16 wild-type parental strain were grown on cellophane on standard agar medium. The cells were harvested, ground on liquid nitrogen to a fine powder, and resuspended in a solution of 50 mM Tris–Cl (pH 7.5) and 1% sodium dodecyl sulfate (SDS). Each extract was briefly sonicated, boiled, and centrifuged at 1,000 × g to pellet cell wall debris. The protein concentrations of the soluble fractions were determined using the Bio-Rad DC protein assay kit (Bio-Rad Laboratories, Hercules, CA). Forty micrograms of total protein from each extract was separated on a 4 to 12% Bis-Tris NuPAGE gel (Invitrogen Life Technologies, Carlsbad, CA) and transferred to a polyvinylidene difluoride membrane.

**GPIP-3 and ACW-1 proteins were detected using the GPIP-3 polyclonal antibody at a 1/25,000 dilution and the ACW-1 polyclonal antibody at a 1/5,000 dilution, respectively.** Immunoreactive bands were visualized using an anti-rabbit alkaline phosphatase-conjugated secondary antibody (Sigma Aldrich, St. Louis, MO) at a 1/15,000 dilution.

For Western blot analysis of the GPIP-3 protein in a wild-type membrane preparation, the GTH-16 strain was grown, harvested, and ground as before. The frozen, powdered mycelia were then resuspended in an ice-cold solution containing 50 mM Tris–Cl (pH 7.5), 200 mM NaCl, 10 mM dithiothreitol (DTT), 1 mM phenylmethylsulfonyl fluoride, and a protease inhibitor cocktail (product number P8215 from Sigma Aldrich, St. Louis, MO). The extract was preclarified by centrifugation at 100,000 × g for 1 h. The 100,000 × g supernatant was discarded, and the centripetal fraction containing the total cellular membrane was resuspended in 50 mM Tris–Cl (pH 7.5) and 1% SDS. The determination of the protein concentration, SDS–polyacrylamide gel electrophoresis (PAGE), and Western blot analysis of the sample were as described above.

**Assessment of gross and hyphal morphologies.** To assess the gross colony morphology of each strain, small inocula of the gpip-1, gpip-2, gpip-3, and gpi1 mutants and the wild-type parents were made in the center of petri dishes containing standard agar growth medium. All strains, with the exception of the gpip-1 temperature-sensitive mutant, which was placed at 39°C, were allowed to grow at room temperature for the times indicated in Fig. 1. Images of the plates were then captured using a digital scanner.

To analyze hyphal morphology, inocula of all mutant and wild-type strains were placed between two cellophane sheets on an agar medium in a petri dish. The two sheets of cellophane were cut from the same larger sheet of cellophane and oriented at 90° to one another as determined by their original orientation within the larger cellophane sheet. The gpip-1 mutant was initially grown at room temperature for 24 h before being shifted to 39°C for an additional 6 h to induce the mutant phenotype. Other strains were grown at room temperature for the times indicated in Fig. 1. A region of cellophane containing the growing edge of the colony was then cut from each dish and placed on a droplet of water, and the cells were photographed using a differential interference contrast microscope.

GTH-16 was chosen as the representative wild-type strain in Fig. 1, as it is the parental strain from which the majority of the mutants were derived (see “Use of RIP to isolate gpip-2, gpip-3, and gpi1 mutants” above).
FIG. 1. GPI anchor mutants have altered gross and hyphal morphologies. Panels A to F are photographs of strains that were inoculated on agar growth medium in standard petri dishes. Panels G to L are pictures of the same strains that were inoculated between two cellophane sheets on agar growth medium, and the growing edge of each colony was photographed at a magnification of ×400. All cultures were incubated at room temperature, with the exception of the 34-15 (gpig-1) temperature-sensitive mutant, which was grown at 39°C to induce the mutant phenotype. Colonies of the wild-type (GTH-16) strain (A and G), MSA-7 (gpip-1) mutant (B and H), 34-15 (gpig-1) mutant (C and I), gpip-2 mutant (D and J), gpip-3 mutant (E and K), and gpi-1 mutant (F and L) are shown. The images in panels A to D, H, and J to L were captured at 48 h after inoculation. Panels E and F are shown at 10 days after inoculation. Panel G is shown at 24 h after inoculation. For the micrograph in panel I, the 34-15 (gpig-1) mutant was initially grown at room temperature for 24 h and then shifted to 39°C for an additional 6 h prior to examination. The scale bar in panel L represents a distance of 10 μm.
[3H]inositol labeling and incorporation into protein. As a means to verify that the gpip-1, gpip-2, gpig-3, and gpi-1 mutants were impaired in the process of GPI anchoring, mutant and wild-type parental cells were labeled with [2-3H]inositol (Perkin Elmer) and the total 1H incorporated into protein was determined. For labeling experiments with the gpip-1 temperature-sensitive mutant, mutant and wild-type strains were grown to approximately mid-log phase in 50 ml liquid shaker cultures at room temperature (the permissive temperature) and then shifted to 39°C (the restrictive temperature) for an additional 4 h prior to labeling. While being retained at the restrictive temperature, each culture was labeled with 15 μCi of [3H]inositol for a 15-min time period. All cells were then harvested over a Buchner funnel, quickly washed with fresh medium that had been prevarmed to 39°C, frozen in liquid nitrogen, and stored at −20°C.

Due to the problems associated with growing the gpip-1, gpip-2, gpig-3, and gpi-1 mutant cells in liquid culture, these mutants were grown on cellophane sheets atop agar medium for labeling experiments. The mutants and wild-type parental strains were grown on cellophane to a point best estimated as a healthy and comparable stage of growth. All cells were then labeled with 1 μCi of [3H]inositol by removing the entire cellophane sheet from each original culture plate and placing it into a new petri dish containing 1 ml of labeled liquid medium. After 1 h, the cellophane sheets were removed from the labeling dishes and blotted dry on Whatman 3MM paper to remove excess medium. Cells were then scraped from the cellophane sheets using a razor blade, frozen in liquid nitrogen, and stored at −20°C.

All labeled, frozen mycelia were Dounce homogenized in 2 ml of an ice-cold extraction buffer containing 50 mM Tris-Cl (pH 7.5), 200 mM NaCl, 10 mM DTT, and 1 mM phenylmethylsulfonyl fluoride. Each cellular extract was divided into two separate 1-ml aliquots, and SDS was added to a final concentration of 1% to each of the two cell extracts. These extracts contained 1% SDS and were then used for determination of 1H-labeled protein. These extracts were preclearced of cell wall debris by centrifugation at 3,000 × g for 5 min, and 150 μl of each supernatant was combined with 500 μl of chloroform and 500 μl of methanol (10:10:3 chloroform/methanol/extract) to precipitate and delipidate total protein for analysis. The samples were incubated at −20°C for 2 h, and the precipitated protein was collected by centrifugation at 17,000 × g for 10 min. The protein pellets were washed with chilled acetone (−20°C). The volume of chloroform/methanol was determined by spectrophotometric assay (260 nm), briefly dried, and resuspended by boiling in extraction buffer supplemented with 1% SDS. Total protein-associated counts were then determined using a scintillation counter.

The extracts devoid of SDS were used for a determination of 1H-labeled lipids as described by Hamburger et al. (29). The sum of the protein- and lipid-associated counts was taken as the total amount of [3H]inositol incorporated by each strain. The amount of [3H]inositol incorporated into precipitable protein was then expressed as a percentage of the total incorporation and compared among the mutant and wild-type strains.

Growth in the presence of cell wall perturbing reagents. To assess the sensitivity of the GPI anchor mutants to reagents that affect osmotic pressure or cell wall biosynthesis, cells were grown on standard Vogel's medium supplemented with 10% (v/v) calf serum, defined medium for C. albicans, or Congo red or calcofluor white or Congo red for 5.0) and shaken at 150 rpm at 37°C for 1 h. The cells were collected by centrifugation at 17,000 × g for 10 min, and aliquots of the supernatants were saved to assay for released protein. The cells were resuspended in 10 mM sodium acetate buffer (pH 5.0) containing 1% SDS and sonicated in 30-s bursts for a total of 3 min. The extracts were then precalced of debris by a second centrifugation, and aliquots of the supernatants were saved to assay for cellular protein. As described above, protein concentrations of all samples were determined using the Bio-Rad DC protein assay kit, and the amount of protein released into the hypo-osmotic buffer was expressed as a percentage of the total protein present in the system (the sum of the protein released to the buffer by lysis and the cellular protein released by sonication).

Electron microscopy. Electron micrographs of mutant and wild-type cells were prepared similar to the method described by Lenhard et al. (38). The gpip-3 and gpi-1 mutants were grown on cellophane on standard agar medium for 7 days. The GTH-16 wild-type strain was grown on cellophane on the same medium for 2 days. Samples of the mutant colonies and wild-type mycelia were floated off the cellophane into a solution of 2% glutaraldehyde, 100 mM sodium cacodylate, 800 mM sorbitol, and 0.1% dimethyl sulfoxide and fixed for 24 h at 4°C. The cells were then embedded in 2% agar and postfixed in a solution containing 1% osmium tetroxide and 100 mM sodium cacodylate. Following dehydration in a graded ethanol and acetone series, the samples were embedded in epon-araldite and thin sectioned. The sections were then stained with 2% uranyl acetate and Reynolds's lead citrate and viewed with the transmission electron microscope.

Western blot analysis of cell wall-associated proteins. To assess how the loss of GPI anchoring affects the synthesis of cell wall proteins, we performed a Western blot analysis on cell extracts from the temperature-sensitive gpip-1 mutant and wild-type strains using an “anti-cell wall” antibody. The “anti-cell wall” antibody is a polyclonal rabbit antibody raised against a crude preparation of GPI-anchored cell wall proteins (106). The wild-type and mutant strains were grown at both 22°C and 39°C and harvested, and cell extracts of each were prepared. The cell extracts were centrifuged at 10,000 × g and 30 μg aliquots of protein from the soluble fractions were analyzed by SDS-PAGE and Western blotting. The “anti-cell wall” antiserum was used at a 1:5,000 dilution, detected with a goat anti-rabbit horseradish peroxidase-conjugated secondary antibody (Promega, Madison, WI) at a 1:5,000 dilution, and subsequently visualized with the ECL detection system (Amersham Pharmacia Biotech, Piscataway, NJ).

Purification of cell walls. Mutant and wild-type strains were grown on cellophane sheets on standard agar growth medium. Cells were harvested and ground to a fine powder in a mortar and pestle while being maintained in a frozen state with liquid nitrogen. The ground material was resuspended in an extraction buffer containing 50 mM Tris-Cl (pH 7.5), 200 mM NaCl, 1% SDS, and 10 mM DTT and boiled for 20 min. All extracts were then centrifuged at 3,000 × g to obtain cell wall material. The isolated cell walls were washed once in extraction buffer, followed by five additional washes in ice-cold distilled H2O. Purified cell wall material was then lyophilized to complete dryness and used for aniline blue dye binding assays or trifluoromethanesulfonic (TFMS) acid digestions as described below.

Aniline blue binding assay. As a measure of the beta-1,3-glucan present in the mutant and wild-type cell walls, we assessed the ability of the cell wall to absorb aniline blue dye. Purified cell walls were prepared from the gpip-3 and gpi-1 mutants, the mnt-1 mutant, and their wild-type parental strain, GTH-16, as described above. The mnt-1 is a null mutant for the N. crassa alpha-1,2-mannosyltransferase gene and has been shown to contain reduced levels of galactomannan and elevated levels of glucose in its cell wall (6). All lyophilized cell wall preparations were resuspended in distilled H2O to a final concentration of 2 mg/ml. Aliquots containing increasing amounts of cell wall material from the mutant and wild-type cells were centrifuged at 3,000 × g for 10 min, and the supernatants were decanted. The collected cell walls were then resuspended in 1 ml of 0.002% aniline blue (Sigma Aldrich, St. Louis, MO) and agitated on a platform shaker at room temperature for 24 h. After pelleting the cell wall material by centrifugation, the amount of unabsorbed aniline blue dye was determined using a spectrophotometer (optical density at 595 nm [OD595]).

TFMS acid digestion of cell walls and identification of integral cell wall proteins. To analyze the proteins covalently linked to the mutant and wild-type cell walls, we digested purified cell walls with TFMS acid using a procedure modified from that described by Edge (14) for deglycosylation of glycoproteins. Cell walls of the gpip-3 and gnt-1 mutant and wild-type strains were prepared as described above. TFMS acid, anisole, and pyridine were obtained from Sigma Aldrich chemical company (St. Louis, MO). Prior to performing the acid digestions, 20 mg of each cell wall preparation were relyophilized overnight to ensure complete dryness of the samples. To maintain anhydrous conditions during the digestions, all glass tubes and syringes were used were dried under a vacuum and all steps throughout the procedure were performed in a chamber being continually purged with N2 gas. Initially, a solution of 16% anisole in TFMS acid was
prepared and 1.25 ml of this mixture was added to each of the cell wall preparations. The samples were then purged with N₂ gas, quickly covered with Parafilm, and placed in the N₂-filled chamber at 4°C for 5 h. During the course of the digestion, the samples were periodically mixed with a Pasteur pipette, purged, and recovered with Parafilm. After 5 h, 3.75 ml of a solution of pyridine/ methanol/H₂O (3:1:1) were added in a dropwise fashion to each of the digestes, which were continually swirled in a dry ice-ethanol bath. The samples were then placed on dry ice for 20 min, followed by an additional 20 min incubation at −20°C. All samples were removed from −20°C and allowed to thaw, and 1 ml of 5% ammonium bicarbonate solution was added to each. The released proteins were then precipitated in 12.5% TCA in acetone at 5% SDS. Protein concentrations of all samples were determined using the Bio-Rad DC protein assay kit. The amount of protein released from 1 mg of starting cell wall material was separated by SDS-PAGE on a 4% to 12% Bis-Tris NuPAGE gel and visualized using the SilverQuest silver staining kit (Invitrogen Life Technologies, Carlsbad, CA) or with Coomassie blue. Selected protein bands were then excised from the stained gels and sent to Midwest Bio Services (Overland Park, KS) for nano-liquid chromatography/mass spectrometry/mass spectrometry (LC/MS/MS)-based identification.

RESULTS

Isolation and identification of the MSA-7 mutant. The MSA-7 mutant was originally isolated in a screening experiment designed to generate mutants affected in the process of vegetative cell fusion. In addition to its cell fusion defect, the MSA-7 mutant grows in an extremely slow, spreading colonial manner, markedly different from that of its wild-type parent (Fig. 1A). The mutant cells do not readily extend across the medium and have a significantly reduced rate of growth (Fig. 1B). Microscopic examination of the MSA-7 mutant shows that the mutant has a very different hyphal morphology and branching pattern than does the wild-type strain. The characteristic vegetative hyphae of the wild-type strain are elongated and branch at points behind the growing hyphal tip (Fig. 1G). In contrast, the mutant hyphae are more bulbous in shape and highly branched (Fig. 1H). Clear evidence of cell lysis is also often found in MSA-7 mutant colonies.

Classical genetic mapping experiments were done to locate the mutation responsible for the MSA-7 phenotype on the N. crassa genome. Following the segregation of the colonial morphology among 1,000 total progeny from a number of crosses, it was determined that the relevant mutation mapped to a location 0.6 centimorgans to the left of the inl gene on linkage group V. From an analysis of the annotated genes at this locus, five genes were selected as likely candidates to contain the mutation of interest. Oligonucleotide primers were designed to amplify and sequence these genes from the mutant and wild-type parent genomes. Four of the five genes from the mutant were identical in sequence to those of the wild-type parent. However, the NCU06663 gene was found to contain a mutation in the normal stop codon (TGA to CGA) that results in a protein with high levels of sequence similarity with genes encoding known S. cerevisiae and mammalian GPI anchor pathway components. Several such genes were readily identified, and those encoding selected components of the phosphoethanolamine transferase and GPI transamidase complexes were targeted for disruption. Specifically, we attempted to create RIP mutations within the following N. crassa genes as defined by the Broad Institute/MIT Neurospora genome project (25): NCU07999, NCU06215, and NCU06508 (members of the phosphoethanolamine transferase and GPI transamidase complexes required for complex formation and stabilization). We also attempted to use the RIP technique to obtain null mutants for the gpi-1 (NCU06663) and gpi-1 genes (NCU09757). Each of the genes was PCR amplified and subcloned into the pRAL1 plasmid (1). The resultant constructs were then used to transform the N. crassa GTH-16 strain (37). A number of individual transformants for each construct were obtained and subsequently mated with the inl (FGSC 1453) strain to activate the RIP process. Multiple matings for three of the six chosen putative GPI anchor pathway genes yielded progeny with a colonial phenotype characteristic of the gpi-1 and gpi-1 mutants. Colonial mutants were isolated for the NCU07999, NCU06508, and NCU05644 genes, suggesting that the products of these genes function in the GPI anchor pathway.

The NCU07999-encoded protein is a homolog of the mammalian PIG-N and S. cerevisiae Med4p proteins. The PIG-N and Med4p proteins function in the addition of phosphoethanolamine to the first mannose residue during GPI anchor biosynthesis (26, 30, 34). The protein encoded by NCU06508 is a homolog of the mammalian PIG-O and S. cerevisiae Gpi13p proteins putatively function as the N-acetylgalactosamine transferase responsible for catalyzing the first step in the GPI anchor biosynthetic pathway (15, 59). In their study, Seiler and Plamann named this gene gpi-3 after its S. cerevisiae homolog. As the abbreviation of “gpi” has been previously used to designate N. crassa mutants lacking glucosephosphate isomerase and to develop a specific nomenclature for naming multiple components of the GPI anchor pathway, we have opted to refer to this gene as gpi-1 (glycosylphosphatidylinositol anchor N-acetylgalactosamine transferase gene-1). We sequenced the 34-15 (gpi-1) mutant allele and found a single missense mutation (TGG to TTG), which changes amino acid number 176 of the GPG-1 protein from a serine to a leucine residue.

Isolation of additional mutants in the GPI anchor pathway. As a means to further demonstrate the importance of GPI anchoring to the overall morphology of N. crassa, mutations in other genes in the GPI anchor pathway were generated via the RIP phenomenon. RIP is a process in which genes found in duplicate copies within the N. crassa haploid genome are mutated during mating. To generate these RIP mutants, a BLAST search of the N. crassa genome was first conducted to identify those genes having high levels of sequence similarity with genes encoding known S. cerevisiae and mammalian GPI anchor pathway components. Several such genes were readily identified, and those encoding selected components of the phosphoethanolamine transferase and GPI transamidase complexes were targeted for disruption. Specifically, we attempted to create RIP mutations within the following N. crassa genes as defined by the Broad Institute/MIT Neurospora genome project (25): NCU07999, NCU06215, and NCU06508 (members of the phosphoethanolamine transferase and GPI transamidase complexes required for complex formation and stabilization). We also attempted to use the RIP technique to obtain null mutants for the gpi-1 (NCU06663) and gpi-1 genes (NCU09757). Each of the genes was PCR amplified and subcloned into the pRAL1 plasmid (1). The resultant constructs were then used to transform the N. crassa GTH-16 strain (37). A number of individual transformants for each construct were obtained and subsequently mated with the inl (FGSC 1453) strain to activate the RIP process. Multiple matings for three of the six chosen putative GPI anchor pathway genes yielded progeny with a colonial phenotype characteristic of the gpi-1 and gpi-1 mutants. Colonial mutants were isolated for the NCU07999, NCU06508, and NCU05644 genes, suggesting that the products of these genes function in the GPI anchor pathway.
proteins. The PIG-O and Gpi13p proteins function in the addition of phosphoethanolamine to the third mannose residue within the GPI anchor (31, 66). We have named the NCU07999 and NCU06508 genes gpip-2 and gpip-3, respectively (glycosylphosphatidylinositol anchor phosphoethanolamine transferase gene-2 and -3). The NCU05644 gene encodes a mammalian PIG-T and S. cerevisiae Gpi16p homolog. The PIG-T and Gpi16p proteins are thought to be involved in the formation and stabilization of the GPI transamidase complex, which cleaves specified proteins at the omega site and attaches the GPI anchor to the newly generated C terminus (22, 48, 49). We have opted to call the NCU05644 gene gpi-1 (glycosylphosphatidylinositol anchor transamidase gene-1).

Genomic DNA was isolated from three mutant progeny for each of the successful RIP experiments and used for PCR amplification and sequencing of the mutant genes. In each instance, we found that the progeny having the RIP-induced phenotype contained multiple mutations within the endogenous copy of the transforming genes and that the majority of the progeny isolated contained nonsense mutations. One isolate that clearly represented a null mutant for each of the genes was selected for further phenotypic characterization. The mutations present in these representative null mutants were as follows: the gpip-2 mutant had a total of 147 mutations, including the introduction of 4 stop codons and the disruption of a putative 5′ splice site; the gpip-3 mutant contained 33 total mutations, 5 of which produced stop codons; and the gpi-1 mutant had 32 total mutations, including the introduction of 6 stop codons.

To verify the effectiveness of the mutations introduced by the RIP process and to demonstrate that the mutant lacked the relevant protein, antibodies were raised against a peptide representing amino acids 388 to 407 of the GPI-3 protein and used to probe cell extracts of the representative gpip-3 null mutant and wild-type parental strain (Fig. 2). The GPI-3 antiserum detected an approximately 100-kDa protein in the wild-type cell extract (lane 2) that was absent in the gpip-3 null mutant (lane 1). The apparent size of the GPI-3 protein is slightly smaller than that predicted for the translated gene product, most likely due to proteolytic processing. The same 100-kDa protein was found to be enriched in a wild-type membrane preparation (lane 3), consistent with the expected ER membrane localization of the GPI-3 protein.

We were unable to obtain RIP mutant progeny from matings of numerous transformants containing DNA sequences from the NCU06215 gene, or the gpip-1 and gpi2-1 genes (for which we have the MSA-7 mutant with the 57-amino-acid extension and the 34-15 temperature-sensitive mutant, respectively). The NCU06215 gene is a homolog of the mammalian hGPI7 and S. cerevisiae GPI7, which encode a phosphoethanolamine transferase that functions in the addition of phosphoethanolamine to the second mannose residue of the GPI anchor (4, 64). One possible explanation for the lack of mutant progeny from RIP experiments using the NCU06215, gpip-1, and gpip-2 genes might be that their gene products are vital for the biosynthesis of the N. crassa GPI anchor and that null mutants for these genes are inviable. A second line of evidence also suggests that null mutations in the NCU06215 and gpip-2 genes may potentially result in a lethal phenotype. As part of the Neurospora genome project (Dartmouth College, Hanover, NH), gene knockouts are being generated via homologous recombination-mediated gene replacement and made available to the Neurospora community. Gene replacement experiments were done for the NCU06215 and gpip-1 genes, and in both instances, viable knockouts could not be recovered. Efforts to obtain knockout mutants for these genes are continuing at the Neurospora genome project (G. Park, H. V. Colot, L. Litvinkova, S. Curilla, C. Ringelberg, K. A. Borkovich, and J. C. Dunlap, personal communication).

As previously mentioned, each of the mutants was disrupted in a homolog of a known component of the S. cerevisiae and mammalian GPI anchor pathways. As a means of verifying that the mutants are defective in the process of GPI anchoring, we assessed the ability of each to synthesize GPI-anchored proteins. To do so, all mutants and the corresponding wild-type parental strains were pulse labeled with [3H]inositol, and the amount of [3H]inositol incorporated into the protein was measured as a percentage of the total [3H]inositol incorporation. In each instance, there was a three- to fourfold reduction in the amount of [3H]inositol-containing proteins produced by the mutants compared to the wild-type strains, demonstrating that the mutants are impaired in the production of GPI-anchored proteins. As presented below, an examination of the protein composition of mutant and wild-type cell walls provides additional evidence that the mutants are lacking in the ability to generate GPI-anchored proteins.

The GPI anchor pathway is required for cell wall integrity. All of the null mutants obtained from the RIP experiments displayed a classical colonial growth phenotype, but the severity of the colonial phenotype differed among the individual mutants. The gpip-3 and gpi2-1 mutants had extremely slow growing, tight, colonial growth patterns, while the gpip-2 mutant displayed a phenotype in which the colonial colonies showed some minor spreading. However, these GPI anchor pathway mutants, as well as the gpip-1 and gpi2-1 mutants described earlier, all exhibited significantly reduced growth rates, altered gross and hyphal morphologies, and obvious points of cell swelling and lysis (Fig. 1). The mutant hyphae were extremely bulbous and apolar in shape and had many more septa than did the wild-type cells. Each of the mutants was impaired in the ability to undergo cell fusion events, as determined by a heterokaryon formation assay. In addition, the mutants were unable to produce either protoperithecia (the female mating structure) or conidia (asexual spores). These phenotypic defects were most apparent in the gpip-3 and

FIG. 2. The GPIP-3 protein is absent in the gpip-3 mutant. Samples of gpip-3 null mutant (lane 1) and wild-type (GTH-16) (lane 2) total cellular extracts and a wild-type (GTH-16) membrane preparation (lane 3) were separated by SDS-PAGE and analyzed by Western blotting using an anti-GPIP-3 antibody.
gpit-1 mutants, which were seemingly identical in phenotype. Since the gpig-3 and gpit-1 mutants were the most severely affected, these two mutants were used for the majority of our GPI anchor mutant analyses. One likely explanation for the variation among the mutants is that the severity of the phenotype is dependent upon the role of the various gene products in the GPI anchor pathway and the relative amount of GPI anchoring that may remain in the mutant cells.

In addition to these GPI anchor pathway mutants, several other *N. crassa* mutants having a colonial phenotype have previously been isolated. The altered gene in a few of these colonial mutants has been identified and shown to be involved in cell wall biosynthesis, structure, or function (6, 17, 71). Studies of *S. cerevisiae* and *Aspergillus* have demonstrated the importance of the GPI anchor biosynthetic pathway and the role of certain GPI-anchored proteins for cell wall integrity (4, 21, 45, 56, 70). Based upon these studies, the putative functions of several *N. crassa* proteins predicted to be GPI anchored, and the significant degree of cell lysis observed within the mutant colonies, we hypothesized that the *N. crassa* GPI anchor mutants would have altered cell walls.

As a way of examining the cell wall, we cultured the mutant and wild-type cells in a variety of growth conditions. Salt sensitivity has been found to be associated with mutations affecting cell wall biosynthesis (11, 47). We tested the ability of the mutant and wild-type strains to grow on solid medium supplemented with elevated levels of salt and found that all of the mutants were impaired. There was a clear correlation between the tightness of the colonial growth pattern displayed by the individual mutants and the salt concentration that fully inhibited their growth. The wild-type parental strains were unable to survive on agar medium containing 12% NaCl. In contrast, the very slow-spreadig colonial mutants, *gpip-1*, *gpig-2*, and *gpig-1*, were unable to grow on agar medium supplemented with 5% NaCl. The most severe, sickly mutants, *gpig-3* and *gpit-1*, were unable to survive on agar medium containing 2% NaCl.

The GPI anchor mutants were unable to readily grow in liquid media. This fact, coupled with the cell lysis frequently seen within the mutant colonies, suggested that the mutants were osmotically sensitive. Sorbitol is often used to stabilize *N. crassa* spheroplasts and other osmotically sensitive cell types (69). We found that the addition of 1 M sorbitol to either liquid or solid media caused an apparent increase in the growth rate of the mutants.

We initially used the temperature-sensitive *gpig-1* mutant to assess why the GPI anchor mutants might have difficulty growing in liquid medium. To assay for cell lysis, the *gpig-1* mutant and its wild-type counterpart were grown at room temperature to approximately mid-log phase and then shifted to the restrictive temperature for an additional 6 h. The cells and their respective culture media were then harvested and assayed for total protein. The *gpig-1* mutant was found to lose approximately 50% of its cellular protein to the medium at the restrictive temperature. Less than 5% of the protein from the wild-type parental strain is released under identical conditions. The extensive protein loss due to cell lysis is consistent with the slow, colonial growth pattern of the mutant and suggests that cell wall integrity is compromised when the GPI anchor pathway is disrupted.

Based on our previous microscopic analyses of these mutants and our direct examination of the cell lysis associated with the temperature-sensitive *gpig-1* mutant, we expected the *gpig-3* and *gpit-1* mutants would be susceptible to hypo-osmotic conditions. To test this hypothesis, mutant and wild-type cells were grown atop cellophane sheets, transferred to a hypo-osmotic solution of 10 mM sodium acetate (pH 5.0), and shaken at 150 rpm at 37°C. After 1 h of agitation, the cells and hypo-osmotic buffer were collected and assayed for total protein as described in Materials and Methods. It was determined that approximately 45% of the total cellular protein was released or secreted from the *gpig-3* and *gpit-1* mutant cells into the buffer. In contrast, the wild-type parent released or secreted less than 5% of its total cellular protein under the same conditions. These findings are consistent with the degree of lysis experienced by the *gpig-1* mutant at the restrictive temperature and further demonstrate the importance of the GPI anchor pathway for cell wall strength and stability.

The sensitivity of the *gpig-3* and *gpit-1* mutants to various cell wall-perturbing agents provides additional evidence for these mutants having fragile cell walls. We found that the mutants were hypersensitive to calcofluor white and Congo red, two reagents that bind to and affect the synthesis of the chitin component of the cell wall (57). Specifically, the *gpig-3* and *gpit-1* mutants were unable to grow on solid media containing calcofluor white or Congo red at a concentration of 1 mg/ml. In contrast, the wild-type parental strain was able to grow in the presence of either reagent up to a concentration of 30 mg/ml. Similar results have been reported for *S. cerevisiae* mutants defective in the GPI anchor and cell wall biosynthetic pathways (4, 5, 21, 54, 66). It has been shown that when cell wall integrity is compromised, fungal cells respond by increasing cell wall biosynthesis (35, 67).

We would interpret our results to mean that when chitin synthesis is compromised by the presence of calcofluor white or Congo red, the *gpig-3* and *gpit-1* mutant cells are less able to compensate for its loss than are wild-type cells.

Transmission electron micrographs further illustrate the importance of GPI anchoring in the synthesis of the *N. crassa* cell wall. In approximately 20% of the *gpig-3* and *gpit-1* mutant micrographs, we find cells with an abnormal “cell-within-a-cell” organization. These mutant cells have clearly defined cytosolic, plasma membrane, and cell wall constituents that are enclosed within what appears to be another cell or second region of surrounding cytosol, plasma membrane, and cell wall (Fig. 3B and C). This unusual cellular morphology was never found in micrographs of the wild-type parental cells (Fig. 3A). The “cell-within-a-cell” morphology has some apparent similarity to the previously described phenomenon of intrahyphal hyphae. Intrahyphal hyphae have been reported in several fungi and are predominantly thought to occur under certain conditions of growth or in response to cellular damage or various genetic mutations (18, 33, 39, 40, 43). In each of these instances, the intrahyphal hyphae, or “invading hyphae,” are thought to reside within the remnants of older, empty, or degenerating host hyphae. The host hyphae either lack cytosol or contain cytosol that is highly vacuolated and disorganized. This is distinct from the “cell-within-a-cell” organization observed in the GPI anchor mutant colonies, where we find no evidence indicating that either the inner or outer cell is degenerating. Given the tight, colonial growth phenotype of the
mutants, we have not yet been able to determine the mechanism by which the “cell-within-a-cell” structures are formed. However, we attribute the phenomenon to alterations in cell wall structure and function.

**GPI anchor mutants have alterations in cell wall carbohydrate composition.** Several GPI-anchored proteins have been shown to be involved in the biosynthesis and remodeling of the glucan layer of the fungal cell wall. The GPI-anchored *S. cerevisiae* Gas1p and the *Aspergillus* Gel1p and Gel2p have been shown to function as endoglucanases/glucanosyltransferases that cleave and rejoin molecules of beta-1,3-glucan (44, 45, 46, 53). The *N. crassa* genome contains three Gas1p/Gel1p/Gel2p homologs, which are predicted to be GPI anchored (16). If *N. crassa* GPI-anchored proteins were critical for the synthesis of beta-1,3-glucan, we would expect that disruptions of the GPI anchor pathway would result in alterations of the beta-1,3-glucan component of the cell wall. To assay for differences in cell wall carbohydrate composition, we used the beta-1,3-glucan-specific dye, aniline blue (63). Cell walls from the *gpip-3* and *gpit-1* mutant and wild-type cells were purified, and the amount of aniline blue absorbed by each was determined. We found that the mutant cell walls bound less aniline blue than the wild-type cell wall on a per milligram of cell wall basis (Fig. 4). At low cell wall concentrations, the mutant cell walls bound only between 25% and 33% as much aniline blue as the wild-type cell wall. The data strongly suggest that the GPI anchor mutants have reduced levels of beta-1,3-glucan in their cell walls.

As a control for the aniline blue binding experiment, we used cell wall from the colonial *mnt-1* mutant, which is affected in the production of the galactomannan component of the *N. crassa* cell wall (6). The *mnt-1* mutant cell wall bound much more aniline blue than either the GPI anchor mutant or wild-type cell walls. This is not unexpected, given the fact that the *mnt-1* mutant cell wall has a higher percentage of glucose than that of the wild type. It is possible that the *mnt-1* mutant compensates for the loss of galactomannan by increasing the amount of beta-1,3-glucan in their cell wall.

**GPI anchor mutants have alterations in integral cell wall proteins.** Based upon the importance of GPI anchoring in targeting proteins to the cell surface, we decided to look for differences in the proteins associated with the mutant and wild-type cell walls. Polyclonal antibodies directed against a crude cell wall fraction were used to look at the pattern of cell wall-associated proteins in the temperature-sensitive *gpig-1* mutant and wild-type cells at both the permissive and restrictive temperatures. The mutant and wild-type parent were grown at the permissive and restrictive temperatures, and cellular extracts of each were subjected to SDS-PAGE and Western blot analysis with the polyclonal antibody (Fig. 5). As is clear from the Western blot, at the restrictive temperature, the *gpig-1* mutant extract lacked some larger-sized proteins (>83 kDa) that were found in extracts from the mutant at the permissive temperature and from wild-type cells at both temperatures. We would conclude that when GPI anchoring is impaired, some cell wall-associated proteins are lost. Because the

![FIG. 3.](image-url) The *gpip-3* and *gpit-1* mutants have an abnormal “cell-within-a-cell” morphology. The *gpip-3* mutant, *gpit-1* mutant, and wild-type (GTH-16) strains were grown atop cellophane sheets on standard agar growth medium and prepared for electron microscopy as described in Materials and Methods. Representative electron micrographs of a wild-type cell (A), the *gpip-3* mutant (B), and the *gpit-1* mutant (C) are shown. Note the “cell-within-a-cell” morphology characteristic of the mutant cells. The scale bar in panel C represents a distance of 1 μm.

![FIG. 4.](image-url) Cell wall absorption of aniline blue dye. Purified cell walls were prepared from the *gpip-3* and *gpit-1* mutants, the *mnt-1* mutant, and the wild-type (GTH-16) parental strain as described in Materials and Methods. Increasing amounts of the mutant and wild-type cell wall preparations were incubated with a solution of 0.002% aniline blue dye, and the amount of aniline blue dye absorbed by each was determined as described in Materials and Methods. Graphed values are the means ± standard deviations of the results from three independent determinations.
polyclonal antibody is directed against a constellation of cell wall proteins, we were unable to identify specific GPI-anchored proteins from this analysis.

The proteins detected in Fig. 5 represent either detergent extractable “cell wall-associated” proteins or “integral cell wall” proteins that were captured in transit to the cell surface. Those proteins which are lost in the mutant at the restrictive temperature are likely to be GPI anchored. Given the importance of GPI-anchored proteins to cell wall formation and stability, we decided to examine the “integral cell wall” protein content and composition of the mutant and wild-type cell walls. Among the “integral cell wall” proteins, some may have GPI anchors, while others may be transmembrane proteins or secreted nonanchored proteins. We expected these proteins to be glycoproteins, with long carbohydrate chains that are often covalently cross-linked into the beta-1,3-glucan of the cell wall (7).

To characterize “integral cell wall” proteins, they have to be released by enzymatic or chemical hydrolysis of the cell wall carbohydrates. We opted to use a chemical digestion to free these proteins from the cell wall. TFMS acid has been previously used to remove sugars from purified glycoproteins and is able to hydrolyze glycosidic linkages without hydrolyzing peptide bonds if used in a temperature-controlled, water-free environment (14). Cell walls from the gpip-3 and gpit-1 mutant and wild-type cells were purified and treated with the TFMS acid as described in Materials and Methods. The released proteins were then recovered by TCA precipitation, and the relative amount of protein released from each cell wall was determined. On a mass basis, we found that the gpip-3 and gpit-1 cell walls contained 2.4% and 3.8% protein, respectively. This is to be compared with the wild-type cell wall, which we found to be 15.3% protein by mass. Clearly, the loss of GPI anchoring has a dramatic effect on the incorporation of protein into the cell wall. The protein released from 1 mg of purified cell wall material was subjected to SDS-PAGE and visualized by silver staining (Fig. 6). As can be seen from the figure, the total amount of protein recovered from equivalent amounts of

FIG. 5. The gpig-1 mutant lacks a number of “cell wall-associated” proteins at the restrictive temperature. The gpig-1 temperature-sensitive mutant and wild-type strain were grown at the permissive (22°C) and restrictive (39°C) temperatures and harvested, and cell extracts of each were prepared. Aliquots of the SDS-soluble material from each cell extract were separated by SDS-PAGE and analyzed by Western blotting using an “anti-cell wall” antibody. Samples of the gpig-1 mutant at 22°C (lane 1) and 39°C (lane 2) and the wild-type strain at 22°C (lane 3) and 39°C (lane 4) are shown. The molecular masses indicated at the right are in kilodaltons.

FIG. 6. The gpip-3 and gpit-1 mutants have altered “integral cell wall” protein compositions. Purified cell walls from the gpip-3 and gpit-1 mutants and the wild-type (GTH-16) strain were digested with TFMS acid as described in Materials and Methods. The total protein released from 1 mg of starting cell wall material from the gpip-3 mutant (lane 1), gpit-1 mutant (lane 2), and wild-type strain (lane 3) were separated by SDS-PAGE and visualized by silver staining. The 10 major protein bands from the wild-type cell wall are indicated with arrows. Those bands containing GPI-anchored proteins are highlighted with thick arrows. See Table 1 for a listing of both the GPI-anchored and nonanchored proteins detected in the major protein bands.
Table 1. “Integral cell wall” proteins from wild-type Neurospora crassa hyphae

<table>
<thead>
<tr>
<th>Protein band no. (predicted molecular mass [kDa])</th>
<th>Identified GPI-anchored protein(s) (gene)</th>
<th>Identified nonanchored protein(s) (gene)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (135)</td>
<td>Endochitinase (NCU02184)</td>
<td>None</td>
</tr>
<tr>
<td>2 (80)</td>
<td>Mixed-linked glucanase (NCU01353)</td>
<td>Catalase-3 (NCU00355)</td>
</tr>
<tr>
<td>3 (74)</td>
<td>None</td>
<td>NCW-1 (NCU05137)</td>
</tr>
<tr>
<td>4 (63)</td>
<td>None</td>
<td>Beta-glucosidase (NCU09326), NCW-2 (NCU01752)</td>
</tr>
<tr>
<td>5 (49)</td>
<td>ACW-1 (NCU08936), glucan/beta-glucanase (NCU0175)</td>
<td>None</td>
</tr>
<tr>
<td>6 (37)</td>
<td>None</td>
<td>Non</td>
</tr>
<tr>
<td>7 (28)</td>
<td>ACW-2 (NCU00957), ACW-3 (NCU05667)</td>
<td>Non</td>
</tr>
<tr>
<td>8 (25)</td>
<td>None</td>
<td>NCW-3 (NCU07817)</td>
</tr>
<tr>
<td>9 (17)</td>
<td>None</td>
<td>Cellular proteins</td>
</tr>
<tr>
<td>10 (11)</td>
<td>None</td>
<td>Cellular proteins</td>
</tr>
</tbody>
</table>

The ChiA protein has been shown to be required for proper growth and morphogenesis of A. nidulans (65). ChiA contains a probable GPI anchor addition site at its C terminus. Bands 5 and 7 each contained two major proteins, which are also predicted to be GPI anchored. Band 5 contained a glucan/beta-glucanase and a protein which we have designated ACW-1 (anchored cell wall protein-1). The ACW-1 is a homolog of the S. cerevisiae GPI-anchored Sps2p/Ecm33p (50, 51), which has been shown to be important for the integrity of the S. cerevisiae cell wall. The glucan/beta-glucanase protein and the ACW-1 (band 5) were reduced in the gpip-3 and gpit-1 mutant cell walls compared to that of the wild type. We have designated the two major proteins detected in band 7 ACW-2 and ACW-3. ACW-2 and ACW-3 are putative GPI-anchored proteins, which are also expressed at reduced levels in the mutant cell walls. The ACW-2 is a homolog of the Fusarium oxysporum FEM1p. FEM1p has been shown to be a GPI-anchored protein covalently linked to the cell wall of F. oxysporum (58). The ACW-3 is a serine- and threonine-rich protein without homology to other previously identified cell wall proteins. The final putative N. crassa GPI-anchored protein identified was a mixed-linked glucanase. This mixed-linked glucanase is a homolog of the Mlg1a and Mlg1b proteins from Cochliobolus carbonum, which serve as bifunctional beta-1,3-1,4/beta-1,3-glucanases (28). The mixed-linked glucanase was detected as a minor component of band 2. The major protein present in band 2 was catalase-3, a nonanchored protein. Thus, it is not unexpected that the intensity of band 2 was not diminished in the gpip-3 and gpit-1 mutant lanes.

To analyze the expression of a single GPI-anchored protein, antibodies were raised against a peptide representing amino acids 186 to 208 of ACW-1. The ACW-1 protein, the product of the NCU08936 gene, was chosen as the representative GPI-anchored protein because its sequence is highly similar to that of a known GPI-anchored protein (the S. cerevisiae Sps2p/Ecm33p and it is highly expressed (based upon its abundance in the vegetative hyphal cell wall and the high number of known expressed sequence tags for the gene). The ACW-1 antisera detected three major bands at 59, 50, and 45 kDa in the wild-type cell extract (Fig. 7, lane 2) that were greatly reduced or absent in the gpip-3 mutant (lane 1). This, along with the data showing that ACW-1 is lost from the mutant cell...
wall (Fig. 6), suggests that ACW-1 is rapidly degraded in the absence of GPI anchoring. The 45-kDa band is consistent with the predicted molecular mass of the ACW-1 proprotein. The upper two bands most likely represent different species of the protein, which might be expected to differ from the proprotein in GPI anchor and/or glycosylation status. The detection of such species is not unexpected, since the ACW-1 protein seen in Fig. 7 is presumably in transit to the cell surface and may be found in multiple modification states. The small amount of ACW-1 protein detected in the gpip-3 null mutant might indicate that a minimal level of GPI anchoring persists in the mutant cells or might simply represent the steady-state level of the ACW-1 protein that is being rapidly degraded in the absence of the GPI anchor.

As shown in Table 1, we also identified five cell wall proteins whose amino acid sequences indicate they are secreted without a GPI anchor. These nonanchored cell wall proteins were represented in bands 2, 3, 4, and 8 of Fig. 6. As mentioned, catalase-3 (CAT-3) was the major protein comprising band 2. The N. crassa CAT-3 is a secreted protein which has been identified as the major catalase within vegetative hyphae and is inducible in response to different conditions of environmental stress (10, 42). It is interesting that the CAT-3 protein appeared in relatively equal amounts in the gpip-3 and gpit-1 mutant and wild-type cell walls. We determined that band 4 included two cell wall proteins, a beta-glucosidase and a protein that we have designated NCW-2 (nonanchored cell wall protein-2). The beta-glucosidase is a homolog of the S. cerevisiae Sco11p. Sco11p is a member of a family of cell wall glucanases involved in the mating process of S. cerevisiae (9, 62). NCW-2, as well as NCW-1 and NCW-3 found in bands 3 and 8, respectively, are serine- and threonine-rich proteins without known homologs. Such serine- and threonine-rich regions are often characteristic of cell wall proteins and may serve as O-linked glycosylation sites that might function to direct and covalently link them to the fungal cell wall (23, 24).

The TFMS acid digestion leaves a single sugar from the N-linked and O-linked oligosaccharides attached to the protein. The presence of these sugars interferes with the identification of peptide fragments containing them in the analysis. For this reason, it is possible we may have been unable to identify some of the smaller cell wall proteins in our analysis (for example, from band 6).

It should also be noted that, in our proteomic analyses, we detected a number of small cytosolic, ribosomal, and mitochondrial proteins in bands 5 through 10. These proteins, which we have referred to as “cellular proteins” in Table 1, were the primary constituents of bands 9 and 10. Such small “intracellular” proteins are often found in fungal cell wall preparations, and there is some evidence suggesting they are cross-linked into the cell wall (2, 3, 27).

**DISCUSSION**

Here we report the characterization of five mutants, each affected in a different gene in the GPI anchor pathway of Neurospora crassa. All of the mutants have a colonial morphology and altered hyphal growth pattern and frequently undergo cell lysis. We attribute these abnormalities to structural and functional defects in the cell wall that result from the absence of a number of GPI-anchored proteins.

Although the GPI anchor mutants shared a general phenotype, the severity of the phenotype varied among the mutants. The gpig-1 mutant (containing an amino acid substitution in the N-acetylglucosamine transferase), gpip-1 mutant (containing a 57-amino-acid extension to an auxiliary factor of the phosphoethanolamine transferase complex), and gpit-2 mutant (a null mutant for the phosphoethanolamine transferase involved in the modification of the first mannose) most closely resembled one another and were less impaired than the gpip-3 mutant (a null mutant for the phosphoethanolamine transferase involved in the modification of the third mannose) and gpit-1 mutant (a null mutant for a subunit of the GPI transamidase complex involved in complex stabilization), which were phenotypically indistinguishable. Such variation could be explained by the type of mutation(s) present in each mutant gene, the potential role of the mutant gene product in the process of GPI anchoring, and the relative amount of anchoring that may persist with the alteration or abolition of that gene product.

Null mutants were obtained for the gpip-2, gpip-3, and gpit-1 genes yet could not be recovered for the gpig-1, gpip-1, or NCU06215 genes, despite repeated attempts. One likely explanation for this finding is that the gpip-1, gpit-1, and/or NCU06215 gene products may be absolutely essential for biosynthesis of the GPI anchor and survival of the cell. This explanation presumes that the mutants we have isolated experience a greatly reduced amount of GPI anchoring but are not completely devoid of the process. The gpip-1 temperature-sensitive (34-15) and gpit-1 (MSA-7) mutants were generated by the introduction of missense mutations in their respective GPI genes, which perhaps allows for a minimal level of product function and subsequent GPI anchoring.

Microscopic examination and functional assays indicated that all of the N. crassa GPI anchor mutants had altered hyphal morphologies and experienced a significant degree of cell lysis. Subsequent analyses demonstrated that each of the GPI anchor mutants was defective in the production of GPI-anchored proteins. We focused on the gpip-3 and gpit-1 mutants because they had the most severely impaired phenotype and sought to associate this phenotype with their defect in the process of GPI anchoring. The gpip-3 and gpit-1 mutants had defects in cell wall strength, as determined by their hypersensitivity to hypotonic conditions and the cell wall-perturbing agents calcofluor white and Congo red. Aniline blue binding assays and TFMS acid digestions illustrated obvious alterations in both carbohydrate and protein composition of the gpip-3 and gpit-1 mutant cell walls. Proteomic analyses of “integral cell wall” proteins demonstrated the abolition or reduction of a number of putative GPI-anchored proteins from the gpip-3 and gpit-1 mutant cell walls. Western blot analysis of an individual GPI-anchored protein, ACW-1, showed that its synthesis was affected in the gpip-3 mutant. In completing these studies, we have demonstrated clear defects in the gpip-3 and gpit-1 mutant cell walls and established a connection between these cell wall alterations and the deficiency of GPI-anchored proteins.

Another interesting observation was the unusual “cell-within-a-cell” organization found in electron micrographs of the gpip-3

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Vol. 5, 2006 GPI-ANCHORED PROTEINS IN N. CRASSA 597
and gpiI mutants. The biogenesis of this abnormal cellular morphology remains to be elucidated.

Consistent with the findings for other fungi (8, 72), we have shown that GPI-anchored proteins represent a significant fraction of the N. crassa cell wall protein. Specifically, we have identified 6 “integral cell wall” proteins from vegetative hyphae which are predicted to contain a GPI anchor based upon their primary sequence and sequence similarity to other known GPI-anchored proteins. The N. crassa GPI-anchored proteins were likely integrated into the cell wall via covalent cross-linking of N- and O-linked glycosylation and/or the GPI anchor present on the proteins to the carbohydrate component of the cell wall. These proteins, as well as other “nonanchored integral cell wall” proteins, were released from the cell wall by chemical hydrolysis with TFMS acid. To the best of our knowledge, this is the first report of the use of TFMS acid to digest a fungal cell wall. Using this procedure, we were able to isolate and identify a number of N. crassa cell wall proteins.

A consideration of the potential roles of the GPI-anchored proteins detected in the wild-type cell wall (Table 1) provides a likely explanation for the phenotype of the GPI anchor mutants. Some of these GPI-anchored proteins might be expected to have important roles in cell wall biogenesis and remodeling (the identified endochitinase, beta-glucanase, and mixed-linked glucanase might clip cell wall polymers to allow for wall remodeling during growth or function as cross-linking enzymes), while others may serve as major structural components of the wall (ACW-1, ACW-2, and ACW-3). The loss of such proteins would not only contribute to the decreased protein content of the mutant cell wall but would also lead to an overall decline in cell wall stability and function. Including those identified here, the N. crassa genome encodes approximately 90 proteins predicted to contain a GPI anchor (13, 16). Many of these proteins putatively serve as glycosyl hydrolases, glycosyl transferases, or peptidases, which would also be required for proper cell wall synthesis, structure, and function. Among the predicted N. crassa GPI-anchored proteins are homologs of the S. cerevisiae Gas1p and Aspergillus Gell1p and Gel2p, Gas1p, Gel1p, and Gel2p function in the remodeling of cell wall glucans (44, 45, 46). The absence of these N. crassa homologs could account for the reduction of beta-1,3-glucan in the cell walls of the GPI anchor mutants.

The GPI anchor mutants had decreased amounts of two major cell wall components, protein (Fig. 6) and beta-1,3-glucan (Fig. 4). It is unclear what component(s) are found in increased levels in the mutant cell walls as measured on a per-mg-cell wall basis. One likely possibility is that the mutant cell walls could have a higher percentage of chitin than the wild-type cell wall. Elevated levels of chitin have been associated with defects in cell wall integrity and result from the activation of the cell wall salvage pathway (52, 54, 55). In addition, our calcfluor white and Congo red sensitivity assays demonstrate that the mutants are extremely sensitive to the loss of chitin synthesis. It is also possible that the mutant cells compensate for the lack of beta-1,3-glucan by producing other polymers, such as beta-1,6-glucan. Although the exact composition of the mutant cell walls remains a subject for further study, it is clear that, when GPI anchoring is lost, the cell wall lacks key components and has an abnormal organization.

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