Expression of Hygromycin Phosphotransferase Alters Virulence of Histoplasma capsulatum

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The Escherichia coli hygromycin phosphotransferase (hph) gene, which confers hygromycin resistance, is commonly used as a dominant selectable marker in genetically modified bacteria, fungi, plants, insects, and mammalian cells. Expression of the hph gene has rarely been reported to induce effects other than those expected. Hygromycin B is the most common dominant selectable marker used in the molecular manipulation of Histoplasma capsulatum in the generation of knockout strains of H. capsulatum or as a marker in mutant strains. hph-expressing organisms appear to have no defect in long-term in vitro growth and survival and have been successfully used to exploit host-parasite interaction in short-term cell culture systems and animal experiments. We introduced the hph gene as a selectable marker together with the gene encoding green fluorescent protein into wild-type strains of H. capsulatum. Infection of mice with hph-expressing H. capsulatum yeasts at sublethal doses resulted in lethality. The lethality was not attributable to the site of integration of the hph construct into the genomes or to the method of integration and was not H. capsulatum strain related. Death of mice was not caused by altered cytokine profiles or an overwhelming fungal burden. The lethality was dependent on the kinase activity of hygromycin phosphotransferase. These results should raise awareness of the potential detrimental effects of the hph gene.

Hygromycin B is commonly used as a dominant selectable marker in selection of genetically manipulated organisms. Expression of an Escherichia coli hygromycin phosphotransferase (hph) gene results in hygromycin B resistance in bacteria, fungi, plants, insects, and mammalian cells (7, 8, 10–12, 16, 21, 27, 28). Expression of the hph gene has rarely been reported to result in effects other than those expected by the expression of hygromycin B resistance, insertional mutagenesis, or the expression of the recombinant DNA carrying the hph gene (3, 19). hph gene expression is generally well tolerated and has been used in the generation of transgenic mice, viable Drosofila, and infectious bacteria and fungi.

Histoplasma capsulatum is an ascomycetous fungus with worldwide distribution and is the causative agent of histoplasmosis. Hygromycin B is the most common dominant selectable marker used in the molecular manipulation of H. capsulatum (20, 25–27, 31). Recombinant molecular genetics is limited in H. capsulatum, and few selectable markers have been characterized (18). Several studies have used hygromycin as a selectable marker in the generation of knockout strains of H. capsulatum or as a marker in mutant strains (20, 25–27, 31). hph-expressing organisms appeared to have no defect in long-term in vitro growth and survival and have been successfully used to exploit host-parasite interaction in short-term cell culture systems and animal experiments. We sought to introduce the gene encoding green fluorescent protein (GFP) into wild-type strains of H. capsulatum, and we selected the hygromycin resistance gene as our selectable marker. This work was being conducted to examine compartmentalization and reactivation of H. capsulatum. Surprisingly, infection of mice with hph-expressing H. capsulatum at normally sublethal numbers resulted in lethality. Five independent hph-expressing strains in the H. capsulatum G217B background all displayed the unexpected lethality. The lethality was not related to the site of integration of the hph construct into the genomes, the method of integration, or the strain of H. capsulatum. Lethality was dependent on kinase activity of hygromycin phosphotransferase.

MATERIALS AND METHODS

Fungal and bacterial strains. Escherichia coli was grown in LB broth with appropriate antibiotics. H. capsulatum was grown in HMM medium with the addition of 200 µg/ml hygromycin, 200 µM uracil, or 150 µg/ml zeocin as appropriate (32). H. capsulatum strain WU 15 was obtained from William Goldman. Chemically competent E. coli strain TOP10 was used for plasmid cloning. Agrobacterium tumefaciens strains for H. capsulatum transformation were generated by electroporation of plasmid constructs generated in the pCB301 backbone into A. tumefaciens LBA1100 (15, 33). Agrobacterium-mediated transformation (AMT) of H. capsulatum was performed as previously described (20). Electroporation of H. capsulatum was performed as previously described with modifications due to the use of a square-wave electroporator (31). Electroporations were performed using a BTX 830 instrument at 375 kV for 5.5 ms using 1-mm cuvettes.

Generation of plasmid constructs. Restriction digestions and ligations were performed under standard conditions. A BglII-SalI fragment comprising the E. coli hph gene under control of the Agrophilus nidulans GPD regulator sequences ligated to the A. nidulans TrpC terminator sequence was ligated between the BglII and Xhol sites of pUG27, resulting in a hygromycin resistance cassette flanked by loxp sequences. A SalI-SaclI fragment containing the hygromycin resistance cassette flanked by loxp sequences was cloned into the polylinker of pCB301-BLE, the hygromycin open reading frame was replaced by the Streptocollotrichus hindustanus BLE gene. To generate pCB301-GFP-HYG, the GFP gene under control of the CBP regulatory sequence was excised from pSBB9.2 (14) and cloned downstream of the hygromycin resistance cassette within the pCB301 multiple cloning region. The vector pCB301-BLE-HYG was

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generated by insertion of the GPD promoter-hygromycin phosphotransferase-TrpC terminator cassette into pCB301-BLE between the Apal and the Kpnl sites. Site-directed mutagenesis was performed using a QuikChange mutagenesis kit according to the manufacturer’s instructions to generate pCB301-BLE-muthYHG, resulting in a D196A substitution within the hph gene product (Stratagene, La Jolla, CA). The plasmid pCR186 was obtained as a kind gift from Chad Rappleye.

Animal studies. C57BL/6 mice were purchased from The Jackson Laboratory. Animals were housed in microisolator cages and were maintained by the Department of Laboratory Animal Medicine (University of Cincinnati), which is accredited by the Association for Assessment and Accreditation of Laboratory Animal Care. All animal experiments were performed in accordance with the Animal Welfare Act guidelines of the National Institutes of Health, and all protocols were approved by the Institutional Animal Care and Use Committee of the University of Cincinnati. H. capsulatum was harvested in mid-log phase by centrifugation from a liquid culture in HMM medium. To produce infection, animals were inoculated intranasally with between 2 \times 10^4 and 2 \times 10^6 H. capsulatum yeast cells in a 30-μl volume of Hanks balanced salt solution (HBSS).

Quantitative organ culture. Mice were euthanized and examined for fungal burdens in lungs and spleens. Organs were harvested and homogenized in 5 ml of HBSS. Organ homogenates were plated on blood heart infusion agar plates at burdens in lungs and spleens. Organs were harvested and homogenized in 5 ml of HBSS. Organ homogenates were plated on blood heart infusion agar plates at multiple 10-fold dilutions and incubated at 30°C until colony growth could be measured. The fungal burden was expressed as mean log_{10} CFU per whole organ ± standard error of the mean. The limit of detection was 10^{4} CFU.

Cytokine measurement. Lungs from infected mice (n = 4 or 5) were removed, homogenized in 10 ml of HBSS, centrifuged at 1500 × g, filtered sterilized, and stored at −70°C until assayed. Commercially available enzyme-linked immunosorbent assay kits were used to measure gamma interferon (IFN-γ), interleukin-6 (IL-6), and tumor necrosis factor alpha (TNF-α) (Pierce, Rockford, IL) and IL-10 and IL-12 (R & D Systems, Minneapolis, MN).

Flow cytometry. Inflammatory cell infiltrates were analyzed by flow cytometry. Lung leukocytes and splenocytes were obtained by teasing apart lungs between the frosted ends of two glass slides. Mononuclear lung cells were further isolated by running over Lymphocyte M (Cedar Lane Laboratories, Burlington, NC). Cells were washed three times with HBSS. The following monoclonal antibodies were purchased from BD Biosciences: CD3-fluorescein isothiocyanate, CD4-peridinin chlorophyll protein (APC), CD8-APC, CD11c-APC, GR-1-APC, Mac3-phyceroerythrin, and B220-APC. A total of 2 × 10^6 cells were incubated with 0.5 μg of monoclonal antibody in staining buffer (1% bovine serum albumin in phosphate-buffered saline) for 10 min at 4°C. The cells were washed in staining buffer, and fluorescence was measured using a FACScalibur flow cytometer (BD Bio-sciences, San Jose, CA).

Lung histology. Histology was performed on formalin-fixed inflamed lung samples by the Comparative Pathology Laboratory of the University of Cincinnati.

Determination of site of T-DNA integration. To determine the site of Agrobacterium-mediated integration into the H. capsulatum genome, thermal asymmetrical PCR (TAIL-PCR) was performed as previously described (17). Briefly, H. capsulatum genomic DNA was isolated from transformed strains. Sequential rounds of TAIL-PCR were performed using three right- or left-border primers and a pool of degenerate primers (AD1 to AD4) as described. After three rounds of amplification, products were gel purified, cloned into pCR2.1-TOPO, and transformed into E. coli. Transforms were isolated, screened, and sequenced at the Cincinnati Children’s Hospital Medical Center Sequencing facility using vector primers. The sequence was compared to the H. capsulatum G217 genome sequence (www.wustl.edu) by BLAST analysis to identify the site of genomic integration.

Hygromycin B phosphotransferase activity assay. A semiquantitative hygromycin B phosphotransferase activity dot blot assay was performed as previously described with slight modification (24, 29). H. capsulatum strains UC18, UC19, UC24, and UC25 were grown in HMM medium to mid-log phase. Yeast cells from equal volumes of culture were lysed by bead beating at 4°C in 150 μl of ice-cold kinase lysis buffer (25 mM Tris-HCl [pH 7.5], 5% glycerol, 10 mM MgCl₂, 1 mM dithiothreitol, 1 mM phenylmethylsulfonyl fluoride, and 1× EDTA protease inhibitor cocktail [Roche Applied Sciences, Indianapolis, IN]), and cellular debris was removed by centrifugation. The cell culture supernatant obtained following centrifugation of yeast cells was concentrated 100-fold by vacuum centrifugation. Five microliters of cell lysate or concentrated cell culture supernatant was added to 50 μl of reaction buffer (13.4 mM Tris maleate [pH 7.1], 8.4 mM MgCl₂, 50 mM NH₄Cl, 60 mM hygromycin B, 15 mM ATP, 25 μM [γ-32P] ATP per ml) and incubated at room temperature for 1 h. The reaction samples were diluted with 150 μl of distilled water (ddH₂O), loaded into a dot blot apparatus fitted with a layer of nitrocellulose paper above a double layer of P81 phosphocellulose paper (Whatman, Florham Park, NJ), and allowed to drain by gravity. Each well was then washed with ddH₂O. The P81 filter was then washed three times in ddH₂O at 65°C. The filter was then exposed to a phosphor storage screen and visualized on a Storm PhosphorImager (GE Healthcare, Piscataway, NJ).

Statistics. One-way analysis of variance was used to compare groups, and the log rank test was used to analyze survival.

RESULTS

GFP-expressing H. capsulatum strains were generated by AMT of T-DNA containing a GFP expression cassette and a hygromycin selectable marker. The GFP/HYG cassette was introduced into the widely used H. capsulatum laboratory strains G217B and G186AR (14, 18). In addition, a GFP-expressing ura auxotrophic derivative of G217B was generated by the introduction of pCB301-GFP/HYG into strain WU 15 (20). Transformants with high levels of GFP expression were isolated. These strains demonstrated stable high levels of fluorescence after prolonged growth in the absence of hygromycin B selection. Growth curves of the GFP-expressing strains were similar to those of their parent G217B and G186AR strains (data not shown).

A G217B-derived strain, UC1, was then used for in vivo studies in mice. Studies with durations of 7 to 9 days were performed, with no unexpected deaths. During a long-term study examining compartmentalization and reactivation of H. capsulatum, 55% (33/60) of animals infected with 2 × 10⁴ yeast cells unexpectedly died in the first 3 weeks of the study. Because of this unusual occurrence, additional studies were performed to investigate the animal deaths. In the next series of studies, groups of C57BL/6 mice were infected with doses of UC1 organisms that varied between 2 × 10⁴ and 2 × 10⁶ yeast cells and followed for survival. Animals infected with >2 × 10⁶ yeast cells died between days 9 and 20. The hypervirulence of H. capsulatum UC1 compared to the parent strain H. capsulatum G217 was confirmed in survival studies (Fig. 1; Table 1). H. capsulatum UC1 was 100- to 1,000-fold more virulent than its parental H. capsulatum G217 strain.

Organ cultures were performed to determine if UC1-infected mice died of overwhelming H. capsulatum infection. Organ cultures of animals infected with G217B or UC1 showed no significant difference in CFU (P > 0.05) in the lungs of mice.
sacrificed on day 7 or 9. The fungal burden in spleens of animals infected with UC1 exceeded that of wild-type G217B organisms at days 7 ($P < 0.0001$) and 9 ($P < 0.05$) (Fig. 2). The number of CFU present in both the lung and the spleen declined between days 7 and 9. Of note, neither of the numbers of organism isolated is usually associated with the demise of animals. Most often, death is accompanied by a level of $\geq 10^6$ CFU per organ (1). Thus, it is unlikely that the apparent hypervirulence of UC1 could be attributed solely to an increased organism burden in animals infected with G217B-derived UC1 compared to the G217B strain of *H. capsulatum*.

Exuberant inflammatory responses may cause immunopathology and result in the death of animals independent of organism burden. The phenotype of cells infiltrating into the lungs of animals infected with G217B and UC1 at day 7 was determined by fluorescence-activated cell sorter analysis. The lungs of animals infected with G217B and UC1 showed no difference in the number or proportion of CD4$^+$ or CD8$^+$ T cells, dendritic cells, neutrophils, macrophages, or B cells present on day 7 (Fig. 3).

Cytokine analysis of lung and spleen homogenates provided additional evidence that the inflammatory response in animals infected with G217B was similar to that in animals infected with UC1. No significant differences in TNF-α, IFN-γ, IL-4, IL-10, or IL-12 levels were noted in the lungs or spleens at day 7 and day 9 despite infection with either G217B or UC1 (Fig. 3).

Histological analysis of lungs from C57BL/6 mice sacrificed at day 7 after inoculation with sublethal inocula of either G217B or UC1 revealed severe pyogranulomatous pneumonia in all animals, with no significant difference discerned between animals infected with either strain (data not shown).

We investigated if a correlation existed between the hypervirulence phenotype and the site of T-DNA integration into the *H. capsulatum* genome. A second independent isolate of GFP-expressing G217B, UC5, was studied. One hundred percent lethality between days 10 and 15 was similarly noted in C57BL/6 mice infected intranasally with a sublethal dose of $2 \times 10^6$ yeast cells of UC5. The sites of T-DNA integration in UC1 and UC5 were then determined by sequencing TAIL-PCR-generated products. Despite the similar hypervirulent phenotypes of UC1 and UC5 in mice, sequence analysis revealed single unique sites of integration within the *H. capsulatum* genome for each strain (upstream of putative open reading frame HCAG 08014 and downstream of HCAG 07026, respectively). The increased virulence and unexpected deaths were also noted when UC3, which had been rendered prototrophic by complementation with the *URA5*-carrying plasmid pCR186, was used for infection.

Studies were designed to determine the role of GFP expression, hygromycin B expression, and AMT in the lethality. *H. capsulatum* G217B strain UC17, generated by the introduction of only the hph resistance cassette by AMT, was then used to
infect C57BL/6 mice. Despite the lack of a GFP expression cassette, 100% of infected animals died between days 10 and 15, demonstrating that the presence of the \textit{hph} cassette was sufficient to cause the noted hypervirulence phenotype.

To determine if lethality was related to AMT, we generated a zeocin-resistant \textit{H. capsulatum} G217B strain, UC16, by introduction of the \textit{S. hindustanus} BLE gene under control of the \textit{A. nidulans} GPD regulatory sequences by AMT. A single zeocin-resistant transformant was characterized and used to infect C57BL/6 mice at a dose of $2 \times 10^6$ yeast cells via intranasal inoculation. All animals survived the challenge, and on sacrifice on day 21, organ culture of lungs and spleens demonstrated clearance of the infection. This suggested that the hypervirulence phenotype was related to expression of the \textit{hph} gene and hygromycin resistance.

To further examine the role of AMT, hygromycin resistance, and GFP expression in the lethality noted in mice, C57BL/6 mice were infected with \textit{H. capsulatum} strain WU13, a hygromycin-resistant GFP-expressing strain derived by \textit{A. tumifaciens}-mediated random insertion. This uracil auxotroph was rendered prototrophic by introduction of the \textit{Podospora anserina} ura5-carrying plasmid pCR186. As noted with other hygromycin-resistant strains, 85% of animals died between days 10 to 21 after infection (Table 1).

All studies described thus far had been performed on \textit{H. capsulatum} with a G217B strain background. Similar studies were performed with sublethal doses of a hygromycin-resistant strain of G186AR \textit{H. capsulatum}, UC2, generated by AMT. Eight of eight infected animals died by day 16 after inoculation, demonstrating that the hypervirulence was not restricted to the G217B strain of \textit{H. capsulatum} (Fig. 1; Table 1).

To explore the possibility that the virulence of \textit{H. capsulatum} was associated with functional activity of the hygromycin phosphotransferase, site-directed mutagenesis was performed, resulting in the substitution of an alanine for the essential aspartic acid residue within the kinase activation domain of the \textit{hph} gene product (2, 13). Introduction of the zeocin resistance cassette and a hygromycin resistance cassette containing the mutant \textit{hph} gene into \textit{H. capsulatum} G217B resulted in organisms (strains UC18 and UC19) that were resistant to zeocin but sensitive to hygromycin. A control strain was generated by the introduction of the zeocin resistance cassette and a hygromycin resistance cassette containing an active \textit{hph} gene in \textit{H. capsulatum} G217B (strains UC24 and UC25). UC24 and UC25 grew in the presence of hygromycin B concentrations of $>300 \mu g/ml$, while UC18 and UC19 were unable to grow in the presence of $>150 \mu g/ml$ of hygromycin B. Cellular lysates of UC24 and UC25 demonstrated hygromycin B phosphotransferase activity by dot blot assay, while no activity was detected in lysates of UC18 and UC19. No hygromycin B phosphotransferase activity was detected in culture supernatants of UC18, UC19, UC24, or UC25. These strains were then characterized at a molecular level. All strains demonstrated hybridization patterns consistent with a single site of integration by Southern blot analysis (data not shown). The sites of T-DNA integration in UC18, UC19, UC24, and UC25 were determined by sequencing TAIL-PCR-generated products. The strains integrated within four distinct putative open reading frames: HACG 06126, HACG 04373, HACG 00845, and HACG 01994, respectively. PCR analysis using primers located upstream or downstream of the site of integration paired with left- and right-border primers revealed no evidence of genomic rearrangement associated with T-DNA integration. C57BL/6 mice infected with $2 \times 10^6$ UC18 and UC19 organisms survived infection, demonstrating a course of disease seen following infection with wild-type G217B \textit{H. capsulatum} until sacrifice on day 21, whereas five of six mice infected with either UC24 or UC25 died with median times to death of 12 and 14 days, respectively (Fig. 5).

**DISCUSSION**

Generation of GFP-expressing \textit{H. capsulatum} in wild-type laboratory strains G217B and G186AR is an important tool for pathogenicity studies. GFP-positive \textit{ura5} gene deletions provide strains with better characterization than the UV-generated uracil auxotrophic strains, where the site of mutation(s) is unknown (20, 32). However, long-term studies in vivo with these strains unexpectedly led to lethality. The adverse outcomes were restricted to animal studies, and no differences were noted between wild-type and hygromycin-resistant strains in axenic or in vitro studies with macrophages.

Although small increases in organism burden were noted in animals infected with hygromycin-resistant \textit{H. capsulatum}
strains compared to wild-type strains, the number of organisms isolated is not sufficient to explain the deaths of mice (1). Typically, death from overwhelming infection is not observed with G217B in C57BL/6 mice until $\approx 10^8$ CFU are present. Moreover, the burden of infection in mice infected with a strain of *H. capsulatum* that contains the hygromycin resistance gene was not that dissimilar from that with the wild type at days 7 and 9. Another possible explanation for these findings is an altered pathological response. This consideration is highly unlikely, since the pathology did not differ between the two groups and the characters of the inflammatory response as assessed by flow cytometry were similar.

The hypervirulence in animals was related to integration of a hygromycin resistance cassette comprising the *A. nidulans* GPD regulatory sequence, the *E. coli* hygromycin phosphotransferase gene, and the *A. nidulans* TrpC terminator region into the *H. capsulatum* genome. It was not related to the method of integration, as strains generated by both *A. tumifaciens*-mediated and non-*A. tumifaciens*-mediated random integration had a similar phenotype. Similarly, the site of integration did not appear to be critical. AMT exhibits little bias in integration had a similar phenotype. Similarly, the site of integration did not appear to be critical. AMT exhibits little bias in genome. It was not related to the method of integration, as strains generated by both *A. tumifaciens*-mediated and non-*A. tumifaciens*-mediated random integration had a similar phenotype. Similarly, the site of integration did not appear to be critical. AMT exhibits little bias in genome. It was not related to the method of integration, as strains generated by both *A. tumifaciens*-mediated and non-*A. tumifaciens*-mediated random integration.
tion, while this phenomenon has not been recognized in other fungi, community awareness may lead to increased reporting of untoward effects.

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