

Genome-Wide Mapping of Cystitis Due to *Streptococcus agalactiae* and *Escherichia coli* in Mice Identifies a Unique Bladder Transcriptome That Signifies Pathogen-Specific Antimicrobial Defense against Urinary Tract Infection

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The most common causes of urinary tract infections (UTIs) are Gram-negative pathogens such as *Escherichia coli*; however, Gram-positive organisms, including *Streptococcus agalactiae*, or group B streptococcus (GBS), also cause UTI. In GBS infection, UTI progresses to cystitis once the bacteria colonize the bladder, but the host responses triggered in the bladder immediately following infection are largely unknown. Here, we used genome-wide expression profiling to map the bladder transcriptome of GBS UTI in mice infected transurethrally with uropathogenic GBS that was cultured from a 35-year-old women with cystitis. RNA from bladders was applied to Affymetrix Gene-1.0ST microarrays; quantitative reverse transcriptase PCR (qRT-PCR) was used to analyze selected gene responses identified in array data sets. A surprisingly small significant-gene list of 172 genes was identified at 24 h; this compared to 2,507 genes identified in a side-by-side comparison with uropathogenic *E. coli* (UPEC). No genes exhibited significantly altered expression at 2 h in GBS-infected mice according to arrays despite high bladder bacterial loads at this early time point. The absence of a marked early host response to GBS juxtaposed with broad-based bladder responses activated by UPEC at 2 h. Bioinformatics analyses, including integrative system-level network mapping, revealed multiple activated biological pathways in the GBS bladder transcriptome that regulate leukocyte activation, inflammation, apoptosis, and cytokine-chemokine biosynthesis. These findings define a novel, minimalistic type of bladder host response triggered by GBS UTI, which comprises collective antimicrobial pathways that differ dramatically from those activated by UPEC. Overall, this study emphasizes the unique nature of bladder immune activation mechanisms triggered by distinct uropathogens.

rinary tract infections (UTIs) are among the most common infectious diseases of humans, with up to 40% of healthy adult women experiencing at least one UTI episode in their lifetime (24). Gram-negative bacteria, most notably, uropathogenic Escherichia coli (UPEC), are the leading cause of UTIs. These organisms express a range of virulence factors that provoke inflammation in the bladder as part of the early steps of UTI pathogenesis (34, 47, 69, 91). Rapid host inflammatory cascades are initiated in the bladder following UPEC infection, and these encompass interleukins (ILs) (8, 51, 52, 68), tumor necrosis factor (TNF), nitric oxide (16), and CXC chemokine receptor 2 (50), among other host factors (7, 65-67). Notably, UPEC may also suppress innate immune activation as a virulence strategy (7, 41, 75), and roles for immune suppressive and regulatory elements, including Th17 cells and IL-10, were recently shown in the bladder response leading to UPEC UTI pathogenesis (13, 70).

Gram-positive pathogens are less frequently associated with UTIs than organisms such as *E. coli* but are nonetheless common causes of UTI (63). *Streptococcus agalactiae*, also known as group B streptococcus (GBS), has emerged in the past 10 years as an important Gram-positive uropathogen. GBS is generally associated with maternal cervicovaginal colonization and disease in newborns and older persons with chronic medical illness (3, 15); however, GBS also causes acute and subacute UTI. GBS UTIs include asymptomatic bacteriuria (ABU), cystitis, pyelonephritis, and urosepsis, which, clinically, are indistinguishable from UTIs

caused by other uropathogens (46). Urinary GBS is cultured from 1% to 2% of all cases of clinically suspected UTI (84), and GBS UTI complicates up to 7% of pregnancies and may account for up to 10% of pyelonephritis during pregnancy (46, 54). GBS UTI may also contribute to chorioamnionitis (1) and premature onset of labor (44) and has been associated with an increased risk of vertical transmission of the bacteria to infants (55, 90). While the incidence of GBS neonatal disease has decreased in the past decade, diseases such as UTI have increased in adults during this period (15).

In terms of the mechanisms of virulence and disease underlying GBS UTI, high rates of ABU among elderly adults (15, 19, 46) prompted a recent survey of the GBS serotypes that cause UTI,

Received 8 January 2012 Returned for modification 12 February 2012 Accepted 9 June 2012 Published ahead of print 25 June 2012 Editor: S. M. Payne Address correspondence to Glen C. Ulett, g.ulett@griffith.edu.au. * Present address: Kimberly B. Ulett, Royal Brisbane and Women's Hospital, Department of Medicine, Herston, Australia. Supplemental material for this article may be found at http://iai.asm.org/. Copyright © 2012, American Society for Microbiology. All Rights Reserved. doi:10.1128/IAI.00023-12 since distinct serotypes of GBS, most notably types III, Ia, and V, are associated with invasive disease in neonates and adults (84). In that study, a wide spectrum of serotypes were shown to cause UTI, with a disproportionately high distribution of serotype III isolates associated with acute disease. The pathogenesis of GBS UTI, however, and the immune responses that are initiated in the bladder during infection are largely unknown. GBS UTI triggers high levels of IL-1 α in the bladder (83) and also upregulates macrophage inflammatory protein-1 α (MIP-1 α), MIP-1 β , IL-9, and IL-10 (35). These findings suggest that, despite GBS UTI being clinically indistinguishable from UTI caused by other pathogens, divergent mechanisms of pathogenesis are associated with cystitis caused by GBS.

The lack of knowledge on the pathogenesis of acute GBS UTI prompted us to map the bladder transcriptome of GBS cystitis in mice to define the host pathogen interactions that occur immediately after infection. Comparing the bladder defense pathways initiated by GBS with those triggered by UPEC on a genome-wide scale revealed a unique bladder response triggered by GBS which is both more limited in scope and slower to develop than that seen with the prototypic Gram-negative uropathogen. This report provides new insight into defense strategies in the host bladder and uncovers a surprisingly minimal local transcriptional program in acute GBS UTI that develops in a focused, pathogen-specific manner. (This study was presented in part at the American Society for Microbiology Annual General Meeting, 21 to 24 May 2011, New Orleans, LA [76a]).

MATERIALS AND METHODS

Bacteria. The uropathogenic GBS used in this study was cultured from clean-catch voided urine of a 35-year-old nondiabetic woman with no identified risk factors who presented to the University of Alabama at Birmingham (UAB) Hospital with symptoms that were consistent with uncomplicated cystitis, including dysuria, single-organism bacteriuria (> 100,000 CFU/ml), and leukocyte esterase and pyuria (\geq 10 white blood cells [WBC]/µl; nonspun), noted on urinalysis. GBS was grown at 37°C on Todd-Hewitt (TH) agar or in Todd-Hewitt broth (THB). Capsular serotyping, which identified this isolate as serotype III, was performed as described elsewhere (84). Use of the GBS cultured from the patient in this study was enacted according to the principles of the Helsinki Declaration with approval from the Institutional Review Board (IRB) of UAB (X070722011) and the Human Ethics Committee (HEC) of Griffith University (MSC/11/10/HREC). The UAB IRB and the Griffith University HEC waived the need for specific informed consent. In some comparative assays, we used the prototype UPEC CFT073 strain (ATCC 700928), which was originally cultured from a patient with pyelonephritis (43). This strain has been used in multiple pathogenesis studies in mice, as previously described (32, 38, 82, 92).

Analysis of GBS adherence to bladder uroepithelium. Initially, we sought to determine whether the uropathogenic serotype III GBS isolate in this study was capable of binding to bladder cells *in vitro*. For this, we performed binding assays using T24 and 5637 human bladder uroepithelial cells (ATCC; HTB-4, HTB-9). Thirty thousand bladder cells were grown in poly-D-lysine-coated multiwell chamber slides (BD BioCoat) as previously described (83). Fluorescein isothiocyanate (FITC)-stained GBS (0.25 mg ml⁻¹ in phosphate-buffered saline [PBS], 15 min, 37°C) were added (multiplicity of infection [MOI], 100 CFU cell⁻¹), and after 2 h, monolayers were rinsed with PBS and fixed for 45 min with 3% paraformaldehyde and F-actin was labeled with phalloidin Alexa Fluor 594 (Molecular Probes) to colocalize GBS with host cell surfaces. DNA was counterstained with the nuclear dye Hoechst 33258. Cells were visualized with a Leica TCS SP2 confocal microscope. We also performed quantitative measurements of GBS binding to 5637 bladder uroepithelial cells by

the use of antibiotic protection assays at 30 min (initial binding), 2 h (initial binding and invasion), and 24 h (intracellular survival), as described elsewhere (80). Briefly, 5×10^4 to 8×10^4 cells were seeded into wells of a 24-well tissue culture-treated plate (Nunc), grown for 24 h at 37°C in 5% CO₂, and challenged at a multiplicity of infection of 10 bacteria per uroepithelial cell. After 30 min or 2 h, monolayers were rinsed with PBS (five times) and either the monolayers were processed for colony counts to determine numbers of adherent GBS or fresh medium with 100 U/ml penicillin, streptomycin, and gentamicin was added for subsequent processing at 24 h to measure intracellular GBS (78, 81). For some assays, supernatants from quadruplicate cultures were collected at 30 min, 2 h, and 24 h for analysis using an 8-target multiplex protein assay (Bio-Rad Laboratories, Australia) to assess the role of the uroepithelial cells in host responses to GBS.

Murine model of GBS cystitis. Female C57BL/6 mice (8 to 10 weeks of age) were purchased from the Animal Resources Center (Australia) and Jackson Laboratories (United States). Urine was collected 24 h prior to challenge and examined microscopically and by culture to exclude mice with a preexisting condition as described elsewhere (85). Mice were anesthetized by isoflurane, and the periurethral area was sterilized with 10% povidone-iodine. Mice were catheterized using a sterile catheter, and 40 µl of PBS containing the challenge inoculum was instilled into the bladder in a transurethral manner. Control mice received PBS only. For comparisons with UPEC, another group of mice received an equivalent dose of strain CFT073 (based on equal cell numbers) that was grown at 37°C in lysogeny broth as previously described (13). Urine was collected at 22 to 24 h for colony counts, after which mice were euthanized and bladders removed and homogenized for counts or processed for microscopy to visualize GBS binding to the uroepithelial surface. Escalating-dose experiments were used to determine the 90% infectious dose (ID₉₀) of the GBS isolate based on the methods of Reed and Muench (61). A challenge dose of approximately 3×10^9 CFU was used in most assays unless otherwise indicated. In some experiments, bladders were collected from mice 30 min after challenge and were sectioned and washed in PBS (five washes of 5 min each wash, in 1 ml, with rotation) and then processed as normal to determine the numbers of initially adherent GBS in the bladder. Immunohistochemistry (IHC) analysis was also performed on duplicate bladders for histological assessment of local cellular infiltrates. This processing was based on standard IHC methods performed with rat anti-mouse Ly-6g (GR-1) (eBiosciences, San Diego, CA) and goat anti-rat IgG-FITC conjugate (Southern Biotech, Birmingham, AL). To visualize the distribution of FITC-stained GBS on the bladder uroepithelium, fluorescence dissection stereomicroscopy was performed using an Olympus SZX16 microscope fitted with a charge-coupled-device (CCD) DP72 camera. Additional bright-field and epifluorescence microscopy was performed using an AxioImager M2 microscope fitted with AxioCam MRm Rev. 3 and MRc 5 cameras (Carl Zeiss, Australia). All animal studies were performed with approval from and in accordance with the ethical standards of the Animal Ethics Committee of Griffith University (MSC/14/08/AEC) and the Institutional Animal Care and Use Committee of UAB (080708186).

Urine growth assays. To determine whether GBS growth in urine could affect the colonization dynamics of the organisms in the bladder, we undertook *in vitro* urine growth assays. Human urine was collected from six healthy adult volunteers who had not had UTI or undergone any antibiotic treatment in the 2 weeks prior. Equal volumes of urine were pooled, filter sterilized (0.45- μ m-pore-size filters), and stored at 4°C until use (within 48 h). Growth assays were also performed using mouse urine. Duplicate 200- μ l aliquots were inoculated with approximately 10³ CFU of GBS and grown in 96-well microtiter plates at 37°C with shaking (200 rpm), and the optical density at 600 nm (OD₆₀₀) was recorded between 0 and 72 h. The growth experiments were repeated three times, and data from one representative experiment are shown. Ethics approval for use of human volunteer subjects was granted by Griffith University (Human Research Ethics Committee approval MSC/11/10/HREC).

		Mean $C_T \pm$ SEM (<i>n</i>	= 5)			
Target gene ^a	Time (h)	Control	Infected	ΔC_T	Fold change	P^b
CXCL10	2	34.111 ± 1.29	33.351 ± 0.53	0.75	1.411	0.110
	24	35.208 ± 3.14	28.019 ± 6.82	7.189	30.75	0.001
iNOS	2	36.73 ± 1.25	36.75 ± 2.02	-0.02	-1.082	0.817
	24	35.73 ± 0.44	32.26 ± 2.81	3.47	11.92	0.001
CCL5	2	28.37 ± 1.40	28.425 ± 1.02	-0.055	1.4	0.153
	24	29.192 ± 1.15	28.015 ± 1.21	1.177	2.525	0.003
CXCL5	2	35.34 ± 0.87	35.21 ± 2.45	0.13	1.13	0.717
	24	35.64 ± 0.56	33.73 ± 1.80	1.91	4.96	0.001
CXCL9	2	35.573 ± 0.81	34.213 ± 3.57	1.36	1.777	0.075
	24	36.753 ± 1.32	31.950 ± 2.78	4.803	6.464	0.008
IL-1α	2	31.717 ± 1.05	30.737 ± 3.19	0.98	2.788	0.001
	24	32.655 ± 0.95	29.012 ± 2.05	3.643	22.11	0.001
IL-1β	2	27.766 ± 1.11	25.343 ± 1.87	2.423	7.58	0.001
	24	29.455 ± 2.35	24.013 ± 3.15	5.442	21.096	0.001
IL-6	2	33.47 ± 5.70	30.77 ± 10.9	2.5	7.913	0.001
	24	37.03 ± 1.34	33.07 ± 6.22	3.14	17.94	0.001
EMR1	2	28.06 ± 0.12	28.40 ± 0.38	0.4	1.072	0.824
	24	34.48 ± 2.68	33.14 ± 1.02	1.34	2.521	0.124
CSF1R	2	22.16 ± 0.06	22.60 ± 0.02	0.44	1.51	0.396
	24	28.02 ± 1.3	27.22 ± 0.69	0.8	1.73	0.199
INFGr1	2	33.097 ± 0.33	33.110 ± 0.62	-0.013	1.103	0.539
	24	35.340 ± 3.68	33.477 ± 0.62	1.863	1.12	0.480
Caspase-3	2	30.91 ± 0.63	31.25 ± 1.17	-0.34	-1.089	0.724
	24	30.92 ± 1.27	30.64 ± 1.11	0.28	1.293	0.3
β-Actin	2	19.938 ± 0.22	20.256 ± 0.13	-0.318	0.815	0.113
	24	20.26 ± 0.31	20.36 ± 0.15	-0.1	0.937	0.127
GAPDH	2	20.162 ± 0.3	20.662 ± 0.43	-0.5	-1.18	0.5
	24	20.531 ± 0.23	20.352 ± 0.25	0.179	1.132	0.658

TABLE 1 qRT-PCR data for selected genes identified as significantly changed or unchanged in expression at 2 and 24 h postinfection in bladder during GBS cystitis according to microarrays

^a Primer sequences are listed in Table S1 in the supplemental material.

^b P values compare GBS-infected group with control (PBS) group within each time point per Materials and Methods.

Electron microscopy. For scanning electron microscopy (SEM), whole bladders were collected from euthanized mice at 2 h after transurethral challenge and immediately fixed in 3% glutaraldehyde–0.1 M cacodylate buffer (pH 7.4) and stored at 4°C until processing. After being washed in fresh buffer, bladders were pinned onto wax sheets to prevent curling, postfixed in 1% osmium tetroxide, and then dehydrated through a graded ethanol series and critical point dried. Samples were mounted on SEM stubs and sputter coated with platinum for viewing using a JEOL 6300F SEM operated at 8 kV.

RNA isolation and microarrays. Total RNA was isolated from GBSinfected, UPEC-infected, and PBS (control) bladders at 2 h and 24 h after challenge using group sizes of five mice per treatment group per time point. Total RNA was isolated using TRIzol reagent (Gibco) according to the manufacturer's instructions, treated with RNase-free DNase, and analyzed using a Bioanalyzer 2100 instrument (Agilent). RNA that passed Bioanalyzer analysis was quantified, and 100 ng was amplified into cDNA using SuperScript III reverse transcriptase (Invitrogen) with random hexamers tagged with a T7 promoter sequence. Microarray analyses were carried out in quintuplicate for each group and time point using one array per mouse (each array representing one single bladder). This approach provided a high degree of statistical power based on nonpooled samples to incorporate variances between mice within each treatment group. We regarded the use of five biological replicates (mice) in each treatment group and time point preferable to technical replicate experiments, which were not performed.

qRT-PCR. To interrogate array data sets, quantitative reverse transcriptase PCR (qRT-PCR) was carried out for selected genes (listed in Table 1) that were identified as differentially regulated or unchanged following the treatment according to array analysis. Amplification of cDNA (500 ng RNA amplified) was performed using a GeneAmp 7700 system (Applied Biosystems). Target genes were amplified using thermal cycling conditions previously described (13, 83). Glyceraldehyde-3-phosphate

dehydrogenase (GAPDH) and β -actin were used as reference genes. Primer sequences for each target gene are shown in Table S1 in the supplemental material. Separate reactions were carried out to ensure that the efficiency of amplification of the reference gene was approximately equal to that of the target gene. Relative expression levels of target genes were determined by normalizing reaction threshold cycle (C_T) values to the housekeeper genes, GAPDH, and β -actin. ΔC_T values were used in the formula 2.0 – (ΔC_T) for calculation of the relative mRNA expression levels of target genes as previously described using PCR efficiencies and mean crossing-point deviations between samples and controls (56, 57).

Statistical analysis and bioinformatics. For data preprocessing, raw array data were normalized using quantile normalization and summarized using RMA in the affy package in Bioconductor (http://www .bioconductor.org), as described elsewhere (13). Differentially expressed genes were identified by analyzing data using the MAANOVA package (http://www.bioconductor.org/packages/bioc/1.8/html/maanova. html). To identify significant changes in expression induced by GBS, we used a shrinkage-based t test together with permutation (9, 10). A false discovery rate (FDR) of 0.1 was used to generate the significant-gene list. Gene class testing was performed using the SAFE package in Bioconductor by applying thresholds to select significant pathways (4). Both the Kyoto Encyclopedia of Genes and Genomes (KEGG) (29, 48) and Gene Ontology (GO) (2) databases were used to generate a signaling network from the gene class analysis. An FDR of 0.1 was used as a cutoff for both tools. GoMiner was used to analyze functional groups of genes activated in biological pathways and to generate Venn diagrams of functionally related responses. InnateDB (http://innatedb.ca) was used to provide an integrative system-level analysis of functional groups activated within biological pathways. Overrepresentation analysis (ORA) of pathways in innateDB was performed using significant-gene lists, the hypergeometric algorithm, and the Benjamini Hochberg correction method with the pathway ORA P value significance set at 0.05.

Microarray data accession number. The raw data sets for array comparisons have been deposited in the Gene Expression Omnibus database under accession no. GSE27575 (http://www.ncbi.nlm.nih.gov/geo/).

RESULTS

Serotype III GBS colonizes the bladder uroepithelium in vitro and in vivo. Initial binding assays using human T24 and 5637 bladder uroepithelial cells showed adherence of the serotype III GBS isolate to these cells within 30 min after infection (Fig. 1A and B; T24 cells shown). These results are consistent with results previously reported for serotype II GBS (83). Changes in cellular architecture were noted in some cells to which GBS had bound (compare Fig. 1A and C). Similar results were observed in 5637 bladder cells (not shown). In the murine model, the serotype III GBS isolate was also observed bound directly to the surface of the bladder uroepithelium as early as 2 h following transurethral infection. Colonization was observed as focal areas of adhesion across the bladder surface, which was visualized using fluorescence dissection stereomicroscopy to detect FITC-GBS (Fig. 1D and E). Resected areas of bladder processed and imaged using high-resolution SEM confirmed binding of serotype III GBS to the uroepithelium *in vivo* at 2 h following challenge (Fig. 1F and G). Histological assessment of control (Fig. 1H to J) and GBS-infected (Fig. 1K to M) bladders using hematoxylin and eosin (H&E) and IHC illustrated a local cellular infiltrate at 24 h comprised of polymorphonuclear leukocytes (PMNs), as shown by IHC (compare controls [I and J] with GBS-infected panels [L and M]).

Dynamics of early GBS uroepithelium binding and dose response. Initial quantitation of early binding of GBS to 5637 bladder uroepithelial cells after infection *in vitro* showed that only 0.15% of the total bacteria used for challenge bound to bladder cells after 30 min (Fig. 2). This amount of adherence was higher than the UPEC CFT073 adherence at 30 min; however, this difference was not statistically significant. A notable difference after 2 h in which total GBS adherence remained at only 0.15% (adherence/ invasion) but UPEC adherence had increased to 2.5%, which represented a statistically significant difference, was observed (Fig. 2). Intracellular survival data at 24 h after antibiotic protection showed that GBS was recovered from the intracellular compartment of host cells in numbers comparable to that of UPEC (Fig. 2). Thus, GBS binds to bladder uroepithelial cells rapidly (within 30 min) and efficiently compared to UPEC *in vitro*, is able to invade these cells, and is able to survive under intracellular conditions for at least 24 h.

In escalating challenge dose assays in mice, we calculated the ID₅₀ and ID₉₀ values of the GBS isolate in this study to be 1.2 and 2.9×10^9 CFU, respectively. We based these calculations on the number of mice exhibiting positive bladder cultures at 24 h postinfection (Fig. 3A). We then quantitated early GBS binding to the bladders in mice that received an ID₉₀ challenge dose of 3 \times 10⁹ CFU, which was used in subsequent assays. In assays based on extensive washing of bladders immediately after infection, we observed efficient binding of GBS to the surface of the bladder uroepithelium in vivo after 30 min (Fig. 3B). The majority of GBS present in the bladders of mice at 30 min represented bound bacteria, since there was no difference in total bladder load between tissues that were unwashed (U/W) and those that were extensively washed (W). The differences between U/W and W bladders from mice infected with UPEC CFT073, on the other hand, were significant (P = 0.015; Fig. 3B). These data demonstrate at least equally efficient binding of GBS to the bladder uroepithelium surface compared to UPEC. The average bladder GBS burden after 2 h was 10⁶ CFU/0.1 g tissue, which decreased 100-fold after 24 h (Fig. **3C**). These decreases in GBS load between 2 h, 24 h, and 5 days were statistically significant (P < 0.05), demonstrating a decline in bacterial burden over time to a point at 5 days when most mice were bladder culture negative. A few mice exhibited positive kidney cultures at 2 h, 24 h, and 5 days (Fig. 3D); however, there were no significant differences in the mean recovery of GBS from kidneys between any of these time points. High numbers of GBS were detected in the urine at 2 h that also subsequently declined by 24 h but then exhibited equivalent mean bacteriuria levels between 24 h and 5 days (Fig. 3E).

Some uropathogens, including E. coli, are able to utilize urine for growth (33, 64, 89), and we next sought to determine whether GBS might replicate in urine in the bladder to gain insight into whether this might affect the dynamics of uroepithelium colonization. In vitro growth assays performed using GBS and fresh human urine demonstrated virtually no bacterial replication over a time course of 72 h (Fig. 4A). Similar data were observed for growth curves determined using fresh urine from C57BL/6 mice (not shown). In contrast, we observed rapid growth for two control E. coli strains that were used in these assays, UPEC CFT073 and E. coli 83972 (Fig. 4A). To determine whether a general growth defect might be present in this GBS isolate, we also performed parallel growth curve assays using rich THB. However, there was no general growth defect in the uropathogenic serotype III GBS isolate in THB (Fig. 4B). Together, these data show that GBS is unable to utilize urine for growth; hence, GBS growth in urine is unlikely to influence the dynamics of uroepithelium colonization during infection in mice.

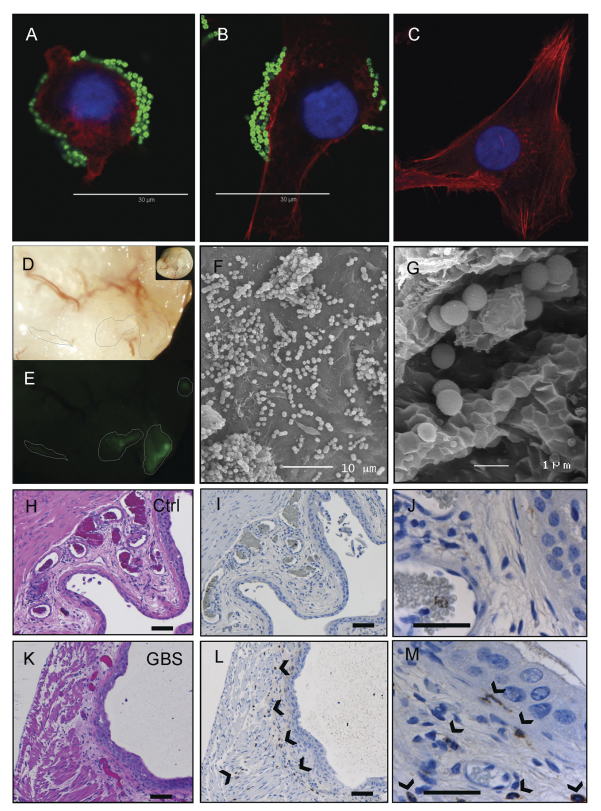


FIG 1 Attachment of uropathogenic serotype III GBS to human bladder uroepithelial cells *in vitro* (A, 30 min; B, 2 h; C, control) and colonization of the surface of bladder uroepithelium in murine GBS UTI as analyzed by fluorescence microscopy of whole tissue, illustrating focal areas of adhered GBS (D and E) alongside images of pockets of bound GBS between folds of uroepithelium (F and G) as analyzed with high-resolution SEM. Histological assessment of control (H to J) and GBS-infected (K to M) bladders illustrates a PMN infiltrate at 24 h (arrowheads), as demonstrated in serial sections stained using either H&E (H and K) or PMN-specific Gr-1 antibody and IHC techniques (I and J, L and M). Bars, 50 μM (small bars) and 180 μM (large bars).

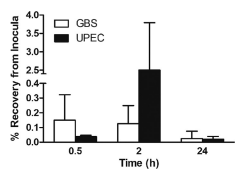


FIG 2 Quantitation of adherence and intracellular survival of serotype III GBS in human bladder uroepithelial cells *in vitro*. The percentage of bacteria bound to 5637 cells after 30 min (representing initial adherence) is shown for GBS, alongside the percentage bound and invaded at 2 h and the percentage of intracellular bacteria recovered from bladder cells after 24 h after antibiotic protection. The equivalent data for UPEC CFT073 are shown for comparison.

GBS bladder transcriptome. Initial genome-wide expression profiling of bladder failed to identify any genes with significantly altered expression immediately after GBS infection at 2 h. This finding was unexpected given the high GBS bladder burdens that were detected at this early time point (Fig. 3C) and the overall lower variance in total bladder-urine bacterial loads between mice within the 2-h group compared to later time points. In contrast to the array data sets at 2 h, comparison of infected and control mice at 24 h revealed a diverse set of 172 genes with significant changes in expression (FDR ≤ 0.1). Heat maps for arrays illustrating overall gene signal intensities in the samples at 24 h are shown in Fig. 5A alongside a volcano plot illustrating the overall breadth of the GBS bladder transcriptome at 24 h in Fig. 5B. The complete significant-gene list at 24 h is presented in Table 2. A summary of the gene expression changes plotted according to FDR for the top 75 genes (FDR \leq 0.05) is shown in Fig. S1 in the supplemental material alongside a Venn diagram derived using GoMiner for responses related to defense against bacteria, cytokines, and inflammation. The raw data sets for array comparisons have been deposited on the Gene Expression Omnibus website (see Materials and Methods for accession number).

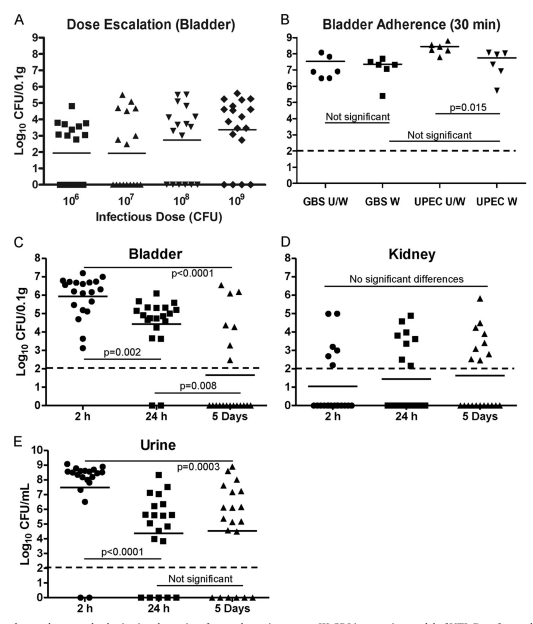
Overall, the GBS bladder transcriptome comprised genes encoding multiple mediators of defense, including those known to be responsive to GBS in other models of infection, including IL-1 α (3.7-fold change, FDR = 0.035), IL-1 β (10.1, 0.011), IL-6 (3.3, 0.093), Stat1 (4.5, 0.03), Stat2 (3.11, 0.04), Ccl5 (2.10, 0.044), and multiple poly(ADP-ribose) polymerases (PARPs) (5, 20, 23, 31, 49, 58, 77, 81, 83). Other transcripts that encode ILs and their receptors or receptor antagonists, including IL-1 family member 9 (4.7, 0.033), IL-1r antagonist (1.8, 0.082), IL-2rg (3.4, 0.048), IL-13ra1 (3.0, 0.088), and IL-17rd (0.7, 0.066), were also identified in the response. Chemokine and chemokine receptor gene expression, including that of Cxcl10 (25.4, 0.006), Cxcl5 (4.27, 0.071), Cxcl9 (3.99, 0.075), and Ccl12 (2.9, 0.078), was also prominent. Other responses of note were related to apoptosis regulators and effectors, including PARP family members 9 (2.8, 0.058), 12 (2.5, 0.053), and 14 (3.1, 0.088), CARD 5 (3.0, 0.035), and neutrophil apoptosis (3.3, 0.093), given the ability of GBS to induce apoptosis and PARP cleavage (79). Unexpectedly, several key defense molecules associated with GBS in other (non-UTI) infection models such as inducible nitric oxide synthase (iNOS) (78) and TNF- α

(27) were not detected as significantly altered in expression in the bladder transcriptome.

qRT-PCR. qRT-PCR showed that expression levels of selected target genes were consistent with individual gene responses identified using arrays. For example, Cxcl10, which was the most strongly activated gene induced by GBS according to arrays at 24 h (25.40-fold increase [P = 0.006]; Table 2), was also highly expressed according to qRT-PCR (Table 1; 30.75-fold increase P =0.001]). Similarly, multiple other genes, including those that encode iNOS, CCL5, CXCL5, CXCL9, IL-6, IL-1α, and IL-1β, were upregulated according to both qRT-PCR and arrays. As expected, there were a number of genes that were tested by qRT-PCR that exhibited higher fold increases compared to arrays because PCR is more sensitive than the arrays (Table 1 and data not shown). All of these transcripts were detected at several cycles above background levels, which validates the detection of genes with low C_T values (e.g., IL-6). The housekeeping genes β -actin and GAPDH also showed no significant changes according to qRT-PCR, which was consistent with findings in arrays. This was also consistent with findings determined for several other genes that showed no significant changes in expression according to arrays and qRT-PCR (e.g., genes for EMR1, CSFR1, INFGr1, and Caspase-3).

Bioinformatics analysis of biological pathways in the transcriptome. Initial analysis of biological pathways within the GBS bladder transcriptome by gene class testing using the KEGG database identified no significant interactions with *P* values < 0.25 for comparisons between infected and control mice at 2 h. In contrast, there were 34 significantly enriched KEGG pathways at 24 h (Table 3). The most strongly represented pathways at 24 h were those related to host responses to bacterial infection; for example, pathways for metabolism (KEGG pathways 561, 562, and 630), ABC transport (2010), focal adhesion (4510), extracellular matrix receptor interactions (4512), Toll-like receptors (TLRs) (4620), NK cell-mediated cytotoxicity (4650), and B cell receptor signaling (4662), which were all significantly overrepresented. Others of interest were those for mitogen-activated protein kinase (MAPK) signaling (KEGG pathway 4010), apoptosis (4210), T cell receptor signaling (4660), antigen processing and presentation (4612), and cytokine-cytokine receptor interactions (4060).

Subsequent GO biological process analyses of the array data for GBS-infected mice at 2 h and 24 h, shown in Table S2 in the supplemental material, revealed larger numbers of overrepresented pathways compared to those derived using KEGG as described above: 40 and 278, at 2 h and 24 h (P < 0.05), respectively. Interestingly, several pathways identified at 2 h using GO indicated that GBS might suppress early host responses in the bladder following infection. These pathways related to downregulation of the defense response (pathway 0031348), downregulation of the response to stimulus (0048585), and downregulation of the inflammatory response (0050728) (see Table S2 in the supplemental material). Pathways related to immune activation and upregulation of antimicrobial defense were highlighted at 2 h but were more prominent at 24 h. These included pathways for T-lymphocyte-mediated immunity (0002456), nitric oxide synthesis (0006809), TNF- α production (0032640), and upregulation of the innate immune response (0045089). Concomitant activation of immune-suppressive pathways for adaptive responses (0002820) and cytokine synthesis (0042036) was also identified at 24 h, illustrating a balance of pro- and anti-inflammatory responses.



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FIG 3 Infectious dose and temporal colonization dynamics of uropathogenic serotype III GBS in a murine model of UTI. Data for escalating dose assays (A) and measures of early bladder adherence of GBS in mice challenged with the ID_{90} after 30 min (B) are shown alongside persistence data for GBS in the bladder (C) and kidneys (D) and bacteriuria levels (E). Broken lines indicate the detection limit for each sample type based on the volume plated. Initial binding of GBS to bladder uroepithelium as represented in panel B was determined by comparing unwashed (U/W) bladders (bound GBS plus those present in bladder lumen) (n = 6) with washed (W) bladders (representing only adherent bacteria after 30 min), and the results are shown alongside data for UPEC for comparison. Temporal colonization data (C, D, and E) represent pooled data from mice challenged with the ID_{90} in two separate experiments with at least 9 mice per group per experiment. Statistical comparisons between groups are based on Mann-Whitney *U* tests with *P* values set for significance at 0.05.

Integrated system-level analysis. We next used the InnateDB database as a complementary bioinformatics tool to enable integrative system-level analyses of the array data. This identified 293 distinct biological pathways at 24 h, which included an additional 15 pathways compared to those that were identified using GO. Mining of the data within InnateDB by ORA identified 168 overrepresented pathways at 24 h (P < 0.05; see Table S2 in the supplemental material). The most significant of these with the highest number of activated genes were those for cytokine receptor interactions (pathway 604) and chemokine signaling (4351), shown in

Fig. 6. Many of the interactors in the pathways that were identified exhibited a large number of interactions such as TRAF6 (525 interactors), although others, such as TNF (39 interactors) and TLR-9 (16 interactors), exhibited fewer interactors.

Common and GBS-unique elements of the bladder transcriptome. Direct comparisons of gene expression responses in bladders of mice challenged with GBS and UPEC enabled us to ascertain the degree of response conservation in the bladder transcriptome. These comparisons used bladders from GBS-infected, UPEC-infected, and PBS control mice in a single experiment per-

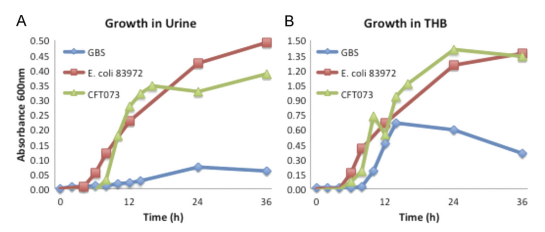


FIG 4 *In vitro* growth curve assays of uropathogenic serotype III GBS in pooled human urine (A) and in THB (B) as measured at OD₆₀₀. The equivalent growth curve data derived from control *E. coli* strains UPEC CFT073 and 83972 in parallel assays are shown for comparison.

formed with the one control group to map global gene expression responses. An absence of significant changes in gene expression induced by GBS in the bladder at 2 h according to array analysis contrasted strikingly with data obtained from mice infected with UPEC for 2 h. These data, which were recently published (13), showed that mice develop a 2-h bladder transcriptome of UPEC UTI that comprises 1,564 transcripts with significant fold differences compared to controls. Microarray analysis at 24 h in the current study showed that, in contrast to the 172 significant changes in gene expression triggered by GBS (Table 2), UPEC induced 15-fold-more significant changes in gene expression at this time point. The 24-h bladder transcriptome of UPEC UTI in C57BL/6 mice revealed a significant-gene list of 2,507 genes (FDR \leq 0.05), which is shown in Table S3 in the supplemental material. Raw array data for UPEC-infected mice at 24 h have been deposited in the Gene Expression Omnibus database (see Materials and Methods for accession number), and data defining bacterial loads in these mice, along with 2-h array data sets, are described elsewhere (13).

An overall comparison of the bladder transcriptomes triggered by GBS and UPEC UTI at 2 h and 24 h is illustrated by volcano plots in Fig. 7. These plots highlight the total number of shared gene expression responses that we identified as commonly induced/repressed by both pathogens at 24 h (black stars in panels B to D; FDR < 0.05) and also illustrate the much larger number of microbe-specific gene expression patterns that were induced in the bladder at 24 h (red diamonds in panels A to C; FDR < 0.05). In total, the common bladder gene expression responses that were induced/repressed by both GBS and UPEC models at 24 h comprised only 68 genes, which are listed in detail in Table S4 in the supplemental material. The plots in Fig. 7 also illustrate the notably slower innate bladder response to GBS (Fig. 7A and B) compared to the more rapid and broader response induced by UPEC (Fig. 7C and D).

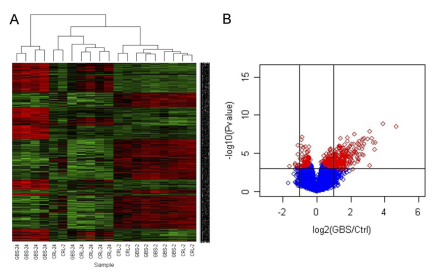


FIG 5 Global gene expression in bladder during GBS cystitis in mice comprises a significant list of 172 genes (FDR < 0.1) derived from comparisons of infected and control (Ctrl) mice at 24 h (n = 5). Gene signal intensities are illustrated in the heat maps for individual arrays (A). The complete significant-gene list is provided in Table 2. The summary response is shown using a volcano scatterplot (B) where significant genes are displayed as shrinkage *t* test results [$-\log_{10}(P \text{ value})$] (*y* axis) versus expression ratio [$\log_2(\text{mean GBS infected/control})$] (*x* axis), with vertical lines representing a 2-fold difference; red diamonds represent FDR < 0.1.

TABLE 2 Global gene expression in bladder during GBS cystitis in mice comprises a significant-gene list of 172 genes (FDR < 0.1) derived from

TABLE	2 ((Continued)
	- 1	Commuca	

	nt-gene list of 172 genes (s of infected mice with co			Gene assignment	Target gene product	Fold change	FDR'
ostinfection	s of finected finec with et	model mee (n - s)) at 24 II	NM_008390	Irf1	3.37	0.047
	TT (1 (F 11 1	FDDd	NM_013563	Il2rg	3.35	0.048
Gene assignment	Target gene product	Fold change	FDR ^a	NM_008491	Lcn2	3.34	0.079
JM_021274	Cxcl10	25.4	0.006	NM_031168	Il6	3.31	0.094
JM_008324	Indo	14.87	0.006	NM_008332	Ifit2	3.3	0.051
JM_011579	Tgtp	10.94	0.019	EF660528	AW112010	3.27	0.046
JM_021792	Iigp1	10.55	0.015	NM_029509	5830443L24Rik	3.26	0.068
VM_008361	Il1b	10.12	0.011	NM_009251	Serpina3g	3.19	0.069
NM_011410	Slfn4	9.42	0.015	NM_011157	Srgn	3.15	0.053
3C034256	Tgtp	8.75	0.011	NM_001040005	Rnf213	3.15	0.077
NM_011260	Reg3g	8.7	0.075	NM_145227	Oas2	3.13	0.058
KM_905096	Irgb10	7.92	0.015	NM_019963	Stat2	3.11	0.035
.38281	Irg1	7.03	0.011	NM_001039530	Parp14	3.07	0.088
NM_008620	Gbp4	6.89	0.022	NM_139198	Plac8	3.06	0.094
NM_008331	Ifit1	6.81	0.031	NM_011347	Selp	3.05	0.075
NM_008326	Irgm	6.54	0.015	NM_133990	Il13ra1	3.04	0.088
3C112328	LOC626578	6.51	0.022	NM_183201	Slfn5	3.03	0.091
NM_011315	Saa3	6.44	0.046	NR_003508	Mx2	3.01	0.081
NM_009114	S100a9	6.38	0.035	AK220210	Nlrc5	2.98	0.035
NM_018738	Igtp	6.35	0.018	NM_018851	Samhd1	2.96	0.057
NM_144559	Fcgr4	5.94	0.053	NM_011408	Slfn2	2.93	0.069
NM_011407	Slfn1	5.83	0.026	NM_013683	Tap1	2.87	0.075
VM_010999	Olfr56	5.75	0.025	NM_011909	Usp18	2.86	0.046
NM_020498	Ly6i Cha5	5.71	0.035	NM_011331	Ccl12	2.86	0.078
NM_153564	Gbp5	5.48	0.022	ENSMUST051264	Cd300lf	2.85	0.082
VM_177162	9930032O22Rik	5.43	0.047	NM_008848	Pira6	2.84	0.078
VM_010260 VM_029499	Gbp2 Ms4a4c	5.42 5.41	0.022 0.079	NM_030253	Parp9	2.83	0.058
ENSMUST031549	EG240327	5.3	0.079	NM_001013371	Dtx3l	2.8	0.046
NM_021394	Zbp1	5.04	0.051	NM_027835	Ifih1	2.8	0.09
VM_011346	Sell	4.96	0.031	NM_001038604	Clec5a	2.78	0.083
VM_010927	Nos2	4.74	0.015	NM_001033308	BC013712	2.74	0.066
NM_153511	Illif9	4.7	0.033	NM_010259	Gbp1	2.67	0.029
VM_133871	Ifi44	4.65	0.033	NM_011852	Oas1g	2.65	0.079
VM_009099	Trim30	4.62	0.035	NM_010553	Il18rap	2.59	0.066
VM_009283	Stat1	4.5	0.035	NM_029084	Slamf8	2.58	0.044
M_172777	BC057170	4.45	0.046	NM_172893	Parp12	2.51 2.5	0.053 0.049
NM 010501	Ifit3	4.33	0.035	NM_181754	Gpr141		0.049
NM 009141	Cxcl5	4.27	0.071	NM_013673 ENSMUST047498	Sp100 AA467197	2.48 2.47	0.09
VM_023137	Ubd	4.22	0.035	NM_145209	Oasl1	2.38	0.043
NM 011472	Sprr2f	4.09	0.057	NM_008152	Gpr65	2.32	0.098
VM_021384	Rsad2	4.07	0.037	NM_008152 NM_008348	Il10ra	2.32	0.085
VM_018734	Gbp3	4.05	0.044	NM_010747	Lyn	2.29	0.078
NM_145545	Gbp6	4.03	0.047	NM_172689	Ddx58	2.20	0.040
NM_016850	Irf7	4.02	0.056	NM_001024134	Trim15	2.22	0.069
JM_010819	Clec4d	4	0.077	NM_145133	T2bp	2.19	0.009
JM_008599	Cxcl9	3.99	0.075	NM_009890	Ch25h	2.19	0.078
JM_020557	Tyki	3.96	0.025	NM_019401	Nmi	2.17	0.078
VM_008230	Hdc	3.91	0.068	BC117807	Ibrdc3	2.1	0.070
JM_008462	Klra2	3.87	0.068	NM_013653	Ccl5	2.1	0.034
JM_013532	Lilrb4	3.87	0.083	NM_026913	Mitd1	2.1	0.056
M_008329	Ifi204	3.83	0.068	NM_001013817	Sp140	2.02	0.030
JM_011854	Oasl2	3.78	0.035	NM_009277	Trim21	1.99	0.078
M_010554	Illa	3.72	0.035	NM_008217	Has3	1.88	0.083
M_023386	Rtp4	3.7	0.041	NM_030701	Gpr109a	1.84	0.089
JM_145226	Oas3	3.57	0.022	AK034303	9330175E14Rik	1.82	0.044
B112025	Sirpb1	3.52	0.088	NM_145391	Tapbpl	1.82	0.078
JM_021893	Cd274	3.49	0.015	NM_031167	Illrn	1.82	0.070
VM_028595	Ms4a6c	3.43	0.078	NM_207648	H2-Q6	1.8	0.002
KR_033405	LOC546714	3.43	0.094	NM_030720	Gpr84	1.73	0.092
M_025777	Duoxa2	3.41	0.046	NM_001001495	Tnip3	1.73	0.092
M_023044	Slc15a3	3.38	0.047	NM_011530	Tap2	1.71	0.075

(Continued on following page)

TABLE 2 (Continued)

Gene assignment	Target gene product	Fold change	FDR ^a
NM_183162	BC006779	1.71	0.078
NM_145373	Sectm1a	1.66	0.058
NM_175397	Sp110	1.59	0.099
NM_001033339	Mmp25	1.5	0.068
NM_024495	Car13	1.43	0.098
NM_010380	H2-D1	1.35	0.083
NM_198013	Cuedc1	0.79	0.1
NM_001085492	Rere	0.78	0.083
NM_181039	Lphn1	0.77	0.058
NM_173186	Tbc1d24	0.76	0.082
NM_172619	Adamts10	0.74	0.04
 NM_007405	Adcy6	0.74	0.046
NM_173788	Npr2	0.74	0.046
NM_198300	Cpeb3	0.74	0.079
NM_032418	Dmpk	0.74	0.091
NM_177030	Dock6	0.73	0.035
NM_029967	Adamtsl1	0.72	0.025
NM_011242	Rasgrp2	0.72	0.023
NM_053097	Cml3	0.72	0.083
NM_011066	Per2	0.72	0.005
XM 990451	AU042671		0.071
NM_173749	E430002G05Rik	0.71	0.088
—		0.71	
NM_134437	Il17rd	0.7	0.066
NM_001042779	Sema3b	0.7	0.088
NM_011218	Ptprs	0.69	0.051
BC057903	Col6a3	0.69	0.064
NM_022814	Svep1	0.69	0.081
NM_007496	Zfhx3	0.68	0.035
NM_175473	Frasl	0.67	0.1
NM_177030	Dock6	0.66	0.058
NM_008483	Lamb2	0.66	0.083
NM_053185	Col4a6	0.65	0.015
NM_031176	Tnxb	0.65	0.078
NM_027890	Susd2	0.64	0.05
NM_001033323	Igsf9b	0.64	0.071
NM_007463	Speg	0.64	0.078
NM_001012765	Adcy5	0.63	0.056
NM_001033476	Ahnak2	0.62	0.029
NM_013630	Pkd1	0.62	0.046
NM_012040	Pnck	0.62	0.088
NM_010570	Irs1	0.6	0.027
NM_013568	Kcna6	0.6	0.09
NM_175750	Plxna4	0.59	0.015
NM_139001	Cspg4	0.58	0.09
NM_008742	Ntf3	0.57	0.071
XR_034431	LOC100048255	0.56	0.058
XR_034431	LOC100048255	0.55	0.035
NM_052835	Rpl10	0.55	0.084
NM_009154	Sema5a	0.54	0.014
NM_009204	Slc2a4	0.54	0.035
NM_176922	Itga11	0.54	0.083
^{<i>a</i>} FDR, false discovery ra	8		

TABLE 3 Activation of diverse KEGG regulatory networks in bladder
during GBS cystitis

KEGG no.	Pathway	P
00561	Glycerolipid metabolism	0.01
00562	Inositol phosphate metabolism	0.01
00630	Glyoxylate and dicarboxylate metabolism	0.01
02010	ABC transporters	0.01
04510	Focal cell matrix adhesion	0.01
04512	Extracellular matrix-receptor interaction	0.01
04620	Toll-like receptor signaling	0.01
04650	Natural killer cell-mediated cytotoxicity	0.01
04662	B cell receptor signaling	0.01
04810	Regulation of actin cytoskeleton	0.01
04920	Adipocytokine signaling	0.01
04010	MAPK signaling	0.014
04930	Type II diabetes mellitus	0.014
05060	Unknown	0.017
00620	Pyruvate metabolism	0.018
03320	Peroxisome proliferator-activated receptors	0.018
04210	Apoptosis	0.02
00970	Aminoacyl-tRNA biosynthesis	0.023
04670	Leukocyte transendothelial migration	0.023
05020	Unknown	0.023
03450	Nonhomologous end joining	0.028
00480	Glutathione metabolism	0.03
00220	Urea cycle and metabolism of amino groups	0.033
00980	Metabolism of xenobiotics by cytochrome P450	0.033
00982	Drug metabolism—cytochrome P450	0.033
05222	Small cell lung cancer	0.033
04060	Cytokine-cytokine receptor interaction	0.035
04742	Taste transduction	0.035
04612	Antigen processing and presentation	0.036
04630	Jak-STAT signaling	0.039
04660	T cell receptor signaling	0.04
04912	Gonadotropin-releasing hormone signaling	0.04
00790	Folate biosynthesis	0.042
04664	Fc epsilon RI signaling	0.048

nificant gene expression responses in the bladder transcriptome and cellular infiltrate data. These experiments were performed in parallel with UPEC experiments for direct comparisons. The results, shown in Fig. 8, illustrate that uroepithelial cells contribute to host defense against GBS through the production of several cytokines, including IL-1 β and IL-1ra. Several of these responses were consistent with gene expression patterns observed in mice. Increased production of the chemokine IL-8 by uroepithelial cells at 24 h in response to GBS (Fig. 8) also correlated with the PMN infiltrate observed in the bladder of mice at 24 h (Fig. 1). Comparing the overall responses induced by GBS with those triggered by UPEC showed that several but not all target proteins were induced more strongly by UPEC. These included IL-8 and granulocytemacrophage colony-stimulating factor (GM-CSF) but not IL-1B or IL-1ra. Other target proteins, including IL-6, IL-13, IL-17, and CCL5, were not significantly induced by GBS over the time course assessed in the human cell culture model, even though these or closely related factors (e.g., IL-13ra1, IL-17rd) were significantly induced/repressed in the GBS bladder transcriptome. Thus, uroepithelial cells contribute to defense against GBS, and several (but not all) responses of human bladder cells are consistent with transcriptional responses in the murine GBS bladder transcriptome.

^{*a*} FDR, false discovery rate.

Bladder uroepithelial cells provide innate defense against GBS. We next undertook an analysis of GBS-infected human bladder 5637 uroepithelial cells *in vitro* to assess whether uroepithelial cells might contribute to the responses that we observed in the bladder transcriptome in mice. For this, we used a multiplex bead-based analysis of cell culture supernatants and compared the response of infected cells to that of control (noninfected) cells for a panel of 8 protein targets that were selected on the basis of sig-

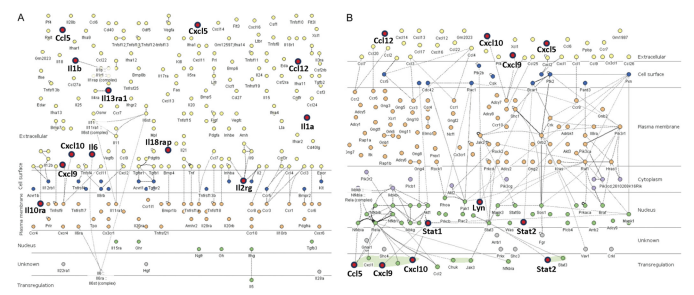


FIG 6 Immune activation in the GBS UTI transcriptome comprises prominent cytokine-cytokine receptor signaling (innateDB pathway number 604) (A) in which 12 genes were highly active (red circles; expression and significance cutoffs, 2.0 and 0.1). The pathway is illustrated according to subcellular localization and includes the total of 255 genes annotated in the pathway. The bladder transcriptome also includes chemokine signaling (innateDB pathway 4351) (B), with activation of the pathway occurring via 8 genes (red circles; Cxcl10, Cxcl9, and Stat2 are shown twice) among a total of 175 genes in the pathway.

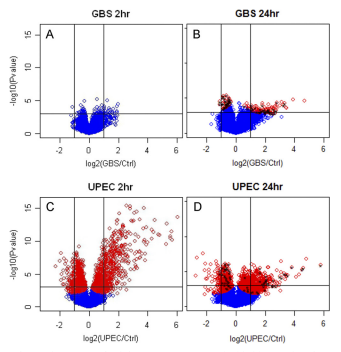


FIG 7 Pathogen-specific defense against GBS in the bladder involves fewer gene expression responses, and these are slower to develop than responses elicited by UPEC. Volcano scatterplots (A and B) illustrate significant genes identified by shrinkage *t* test results at 2 h and 24 h for GBS, compared to those for UPEC (C and D). Red diamonds represent genes at FDR < 0.05 for both pathogens. Significant changes in gene expression for those genes identified as common (i.e., induced/repressed by both GBS and UPEC after 24 h) are denoted by black stars in panels B and D and are listed in detail in Table S4 in the supplemental material. These common responses illustrate the exact combination of shared and pathogen-specific responses in the bladder transcriptomes due to these two organisms.

DISCUSSION

Cystitis due to bacterial infection stems from microbial adherence to uroepithelial cells, which facilitates resistance to the natural hydrodynamic forces of urine flow that remove potential pathogens from the bladder. Uropathogenic GBS was shown to adhere to uroepithelial cells more efficiently than nonuropathogenic GBS in a prior study (83), and the organisms' hemolysin-cytolysin was shown to contribute to the pathogenesis of UTI in mice (35). However, the early events in bladder colonization and the mechanisms of host defense activated immediately following infection with GBS are largely unknown. The results of the current study show that GBS is able to bind to mouse and human bladder uroepithelium rapidly, within minutes of infection, and relatively efficiently compared to the best-adapted bacterial uropathogen, UPEC. This study also shows for the first time that GBS is able to invade human bladder uroepithelial cells and is able to survive inside these cells for at least 24 h, which may contribute to UTI pathogenesis. Adherence to and invasion of bladder uroepithelium set the stage for progression to cystitis once the bacteria are detected by innate immune surveillance mechanisms.

Innate immune signaling cascades are initiated within hours of contact with bacterial pathogens in various infectious settings (17, 18, 71, 95, 96) and are induced in the murine bladder within 2 h of UPEC UTI (13). Such rapid inflammatory responses in the bladder in response to UPEC are exemplified by a 1,000-fold increase in TNF- α expression at 1 h postinfection, which precedes high bladder bacterial loads at 1 day (28), and induction of CXCL1, CXCL6, and MIP-2 within 4 h (7, 25). Surprisingly, despite substantial GBS bladder burdens early in the course of UTI in this study, GBS failed to trigger robust inflammation within 2 h of infection in mice according to genome-wide transcriptomic analysis. Thus, GBS UTI contrasts with UPEC UTI in terms of presentation of the higher GBS bladder burdens very early during infection which precede the bulk of inflammatory responses that occur

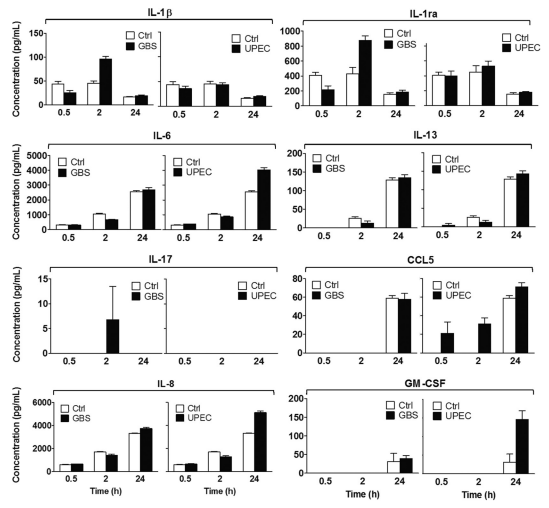


FIG 8 Bladder uroepithelial cells contribute to defense against GBS through production of chemokines and cytokines. Human 5637 uroepithelial cells were infected with GBS at an MOI of 10 as described in Materials and Methods, and culture supernatants at 0.5 h, 2 h, and 24 h were used to measure a panel of 8 protein targets identified from array data and histopathology as part of the bladder response to GBS. Data derived from multiplex protein assay are shown for each target and represent the means \pm standard errors of the means of the results determined for quadruplicate samples versus control (noninfected) bladder cells.

at 24 h postinfection. Our data also indicate that GBS bladder burdens are not influenced by the presence of urine because the bacteria could not utilize this for growth *in vitro*, in contrast to UPEC. High GBS loads observed at 2 h in this study are consistent with recent findings of high bladder bacterial loads in C3H/HeN mice at 3 h (35) that progressed with relatively mild immune responses in the bladder. Taken together, these findings suggest that the dynamics of early GBS bladder colonization in these two strains of mice are similar.

This study also showed that innate bladder responses to GBS unfold slowly compared to responses to UPEC and that the breadth of the responses to GBS is more limited in scope. The nature of the pathogen-specific response during cystitis in mice due to these two organisms was unmistakable because of the comparative analyses that were undertaken in parallel across the entire genome. We chose to administer the same infectious dose for both organisms in terms of cell number for our array experimental design in order to precisely control this comparison. However, it would also be interesting to compare gene expression responses in mice given lower doses of UPEC, which, based on prior observations, would be closer to the ID₉₀ for this pathogen. Our comparisons identified only 68 shared gene expression responses induced by both GBS and UPEC, which illustrates that the majority of gene expression changes in the bladder triggered by these organisms are pathogen specific. The bioinformatics findings in this study are of particular interest, because they reveal a marked induction of innate defense mechanisms against GBS in UTI that are essentially unknown in host responses to GBS in general. Some of these, such as ABC transporters, have prominent roles in infectious disease pathogenesis in other settings, and these would be of interest in future GBS host-pathogen studies. Other biological pathways identified by KEGG as enriched in the GBS bladder transcriptome such as TLRs are well-established elements of defense against GBS in non-UTI models both *in vitro* and *in vivo*.

A relatively slower and milder innate immune response triggered by GBS compared to UPEC could influence the survival of GBS in the bladder immediately after colonization of the uroepithelium. Interestingly, human blood cultures exposed to GBS also exhibit a delayed pattern of induction of proinflammatory cytokines, including IL-1 β and IL-6, which are not produced until 24 h (86, 87). Comparisons of bladder responses to GBS with non-UTI models reveal other commonalities in the transcriptome reported here: IL-6 in GBS sepsis (5, 20, 49), for example, and CCL5, Stat1, Parp9, IL-1 β , and Nos2 (23, 31, 58, 77, 81, 83). Factors that might contribute to tissue damage in the bladder during GBS cystitis, including multiple CXC chemokines such as CXCL10 (36) as well as IL-1 β (11) and Nos2 (45), were also shown to be induced. However, it should be noted that these factors mediate antimicrobial defense in some infections and thus may be important for disease control in GBS UTI. CXCL10 and Nos2, for example, aid in controlling Gram-negative uropathogens in the bladder (53). The functional contribution of many of these factors in the GBS bladder transcriptome to the progression of UTI can now be investigated based on their expression kinetics as shown in this study.

In terms of UTI pathogenesis in general, different microbes have been strongly linked to host features relevant to disease (76), which underscores the importance of host factors in both controlling and contributing to disease. Several studies have described pathogen-specific immune responses in the urinary tract, and these offer valuable insight into the results of the current study. In a recent study on upper UPEC UTI, a new TLR-4/interferon regulatory factor 3 (IRF3)-dependent pathway of pathogen discrimination was defined (21). Pathogen-specific responses to UPEC were previously associated with mucosal TLR responses involving TLR-4-dependent, lipopolysaccharide (LPS)/CD14-independent defense (22, 26), as reviewed elsewhere (60). In the newly described model, commensal E. coli selectively engaged receptors that drive IRF3 activation by UPEC, leading to epithelial gene transcription in a pathogen-specific manner (21). In the current study, GBS did not trigger IRF3; however, IRF1 and IRF7 were activated transcriptionally. This implies that mechanisms involving IRF1 and IRF7, which are involved in prominent signaling events in the GBS bladder transcriptome such as that associated with CXCL10 (88, 94), may drive immune engagement in the bladder early during GBS UTI. Thus, further studies would be of interest to address the potential role of interferon regulatory factors in GBS UTI.

TLR signaling genes were identified as overrepresented in several biological pathways in the GBS bladder transcriptome according to innateDB, which was used to gain insight into networks relevant to innate immunity beyond KEGG and GO (39, 40). While there remains a debate concerning how GBS initiates innate immune responses, there is a consensus with respect to an important role for TLR-2. In systemic GBS infection, TLR-2 deficiency is associated with impaired host resistance (59); in peritoneal macrophages, GBS induces TLR-2-dependent gene activation (12), and in a neonatal infection model, TLR-2 was necessary for GBS clearance (42). There is little published data on TLR-2 signaling in the bladder; however, a body of recent evidence suggests that TLR-4 is central to the induction of local innate immunity (60, 72, 73, 93). TLR-4 and CXCR1 deficiency predisposes to asymptomatic carriage of E. coli and possibly other uropathogens (60). In this study, GBS upregulated the genes for TLR-2, -11, -1, and -6 marginally but none of these responses were statistically significant. InnateDB identified interactions involving TLR-9, suggesting recognition of unmethylated CpG GBS DNA. Upregulation of many IL-1 family members in the GBS bladder transcriptome also suggests potential involvement of TLR-2, especially considering recent reports on the upregulation of IL-1 through TLR-2 in response to intracellular infection (37) and *Borrelia* (6). It is also valuable to interpret these data for GBS against those described for enterococci in mice, which revealed TLR-2-independent inflammation during pyelonephritis (30). In GBS UTI, upregulation of TLR-2 might occur later during infection (after 24 h) or may act in signal transduction in the absence of upregulation. However, an absence of significant upregulation of TLR-2 early during UTI may also explain the relatively slow bladder immune response to GBS and the high incidence of GBS ABU that is observed clinically (84).

In this study, we observed a number of important correlations between the GBS bladder transcriptome in mice and protein production in human bladder uroepithelial cells that were challenged with GBS in vitro. This is the first report, to our knowledge, that human bladder cells contribute to defense against GBS through the production of cytokines and chemokines, including IL-1β, IL-1ra, and IL-8. Some of these findings, such as that for IL-8, support our histopathology observations of a notable PMN inflammatory infiltrate in the bladder of mice after 1 day of GBS UTI. Other findings in the human bladder cell culture model, however, such as an absence of response for CCL5, and other factors related to genes identified as significant in the bladder transcriptome (e.g., IL-13 and IL-17), suggest limitations of this in vitro model. Here, it is important to underscore the limitations of epithelial cell monoculture models for the host response studies; cell-cell interactions, for example, are often necessary to generate effective antibacterial responses, as reviewed elsewhere (14). The timing and duration of infection in such models can also influence bacterial adherence to (and invasion of) host cells, as demonstrated in the present work for UPEC (0.5 h versus 2 h), and this could alter cytokine responses. In these assays, we tested culture supernatants at 0.5 h, 2 h, and 24 h, but it is plausible that these responses could differ depending on the timing of infection. Moreover, many cell types that would be present in the infected bladder in vivo such as resident macrophages, lymphocytes, and PMNs as well as other infiltrating cells were excluded from the human cell culture model. Thus, the contribution of these cells to defense, and the role of cytokines produced by human bladder cells in UTI, requires further investigation. Importantly, however, several pathogen (e.g., IL-1β, IL-1ra)-specific responses were also evident in the human bladder cell response, which supports the finding of distinct bladder transcriptomes for GBS and UPEC observed in mice. Finally, one must consider species differences between mouse and human when extrapolating protein expression data from human bladder cells in culture to data derived from mouse infection assays. The lack of IL-1ß protein detected at 24 h in the human cell culture model in this study, for example, contrasts with high levels of this cytokine induced by both GBS and UPEC in mice at 18 to 24 h during cystitis (83). However, these distinctions may reflect any of the multiple limitations inherent in these models as described, and such comparisons are therefore valid only in view of these various limitations.

Several recent studies on immune responses to bacterial uropathogens have investigated these responses in relation to the degree of immune activation in the urinary tract and the control of disease. In *Ureaplasma parvum* UTI, for example, complications of infection were shown to depend on host factors related to exaggerated, as opposed to minimal, immune responses (62). In that study, animals that developed asymptomatic, rather than acute, UTI exhibited divergent non-PMN-related responses in the urinary tract. More minimalistic immune responses led to a higher rate of pyelonephritis in infected animals. These findings support the notion that the degree of host response contributes to the pathogenesis of bacterial UTI, as discussed elsewhere (74). In the current study, only a few mice exhibited positive kidney culture results following challenge with GBS despite all mice harboring high GBS bladder burdens at 2 h and most exhibiting moderate bladder and urine burdens at 24 h. This suggests that that the early immune responses activated by GBS in the bladder are mostly effective in preventing ascension of the bacteria to the kidneys. Note, however, that GBS kidney colonization patterns may differ between mouse strains and that pyelonephritis is more prevalent in C3H/HeN mice, which exhibit inflammation and a cellular infiltrate (35). These data imply that array mapping of GBS pyelonephritis in mice would be useful to define immune activation under this condition; this would ideally be undertaken using a more consistent model of upper UTI than that described in the current work.

Finally, our data support the observations of a recent study that showed that GBS simultaneously stimulates and suppresses immune activation, through distinct pro- and anti-inflammatory mechanisms in the bladder (35). Our bioinformatics findings based on GO showed that GBS activates biological pathways related to downregulation of defense and to downregulation of inflammation early during infection. However, GBS also triggered pathways of T-lymphocyte-mediated immunity, nitric oxide synthesis, and TNF- α production. Collectively, these results confirm that GBS simultaneously suppresses and activates bladder inflammation.

In summary, this report defines a minimalistic, pathogen-specific bladder transcriptome of GBS cystitis. Many of the factors identified within this response require future investigation to define their potential roles in defense against GBS UTI. Overall, this report offers a critical new perspective on the uniqueness of bladder responses to different uropathogens and, more broadly, highlights the distinct mechanisms of UTI pathogenesis induced by different bacteria.

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