M1T1 group A streptococcal pili promote epithelial colonization but diminish systemic virulence through neutrophil extracellular entrapment

Laura E. Crotty Alexander · Heather C. Maisey · Anjuli M. Timmer · Suzan H. M. Rooijakkers · Richard L. Gallo · Maren von Köckritz-Blickwede · Victor Nizet

Received: 14 September 2009 / Revised: 22 October 2009 / Accepted: 10 November 2009 / Published online: 4 December 2009
© The Author(s) 2009. This article is published with open access at Springerlink.com

Abstract Group A Streptococcus is a leading human pathogen associated with a diverse array of mucosal and systemic infections. Cell wall anchored pili were recently described in several species of pathogenic streptococci, and in the case of GAS, these surface appendages were demonstrated to facilitate epithelial cell adherence. Here we use targeted mutagenesis to evaluate the contribution of pilus expression to virulence of the globally disseminated M1T1 GAS clone, the leading agent of both GAS pharyngitis and severe invasive infections. We confirm that pilus expression promotes GAS adherence to pharyngeal cells, keratinocytes, and skin. However, in contrast to findings reported for group B streptococcal and pneumococcal pili, we observe that pilus expression reduces GAS virulence in murine models of necrotizing fasciitis, pneumonia and sepsis, while decreasing GAS survival in human blood. Further analysis indicated the systemic virulence attenuation associated with pilus expression was not related to differences in phagocytic uptake, complement deposition or cathelicidin antimicrobial peptide sensitivity. Rather, GAS pili were found to induce neutrophil IL-8 production, promote neutrophil transcytosis of endothelial cells, and increase neutrophil release of DNA-based extracellular traps, ultimately promoting GAS entrapment and killing within these structures.

Keywords Group A Streptococcus · Pilus · Virulence factor · IL-8 · Neutrophil extracellular traps

LE Crotty Alexander and HC Maisey contributed equally to this work.

L. E. Crotty Alexander · H. C. Maisey · A. M. Timmer · S. H. M. Rooijakkers · M. von Köckritz-Blickwede · V. Nizet
Department of Pediatrics, University of California San Diego, 9500 Gilman Drive, La Jolla, CA 92093, USA
e-mail: vnizet@ucsd.edu

L. E. Crotty Alexander
Department of Medicine, Division of Pulmonary and Critical Care, Massachusetts General Hospital, 55 Fruit Street, Boston, MA 02114, USA

S. H. M. Rooijakkers
Medical Microbiology, University Medical Center Utrecht, Heidelbergraaf 100, 3584 CX Utrecht, Netherlands

R. L. Gallo
Division of Dermatology, University of California San Diego, 9500 Gilman Drive, La Jolla, CA 92093, USA

R. L. Gallo
VA San Diego Health Care System, San Diego, CA, USA

V. Nizet
Skaggs School of Pharmacy and Pharmaceutical Sciences, University of California San Diego, 9500 Gilman Drive, La Jolla, CA 92093, USA

V. Nizet
Rady Children’s Hospital, 3020 Children’s Way, San Diego, CA 92123, USA
Introduction

*Streptococcus pyogenes* (group A *Streptococcus* (GAS)) is a leading human pathogen estimated to cause over 500,000 deaths each year globally, with disproportionate effects upon children, the elderly, and those living in developing countries [1, 2]. While GAS can asymptomatically colonize the upper respiratory tract or skin of healthy individuals, it is also associated with a large spectrum of mucosal and invasive diseases, from simple pharyngitis or impetigo to the potentially life-threatening pneumonia, necrotizing fasciitis (NF) and toxic shock syndrome (TSS) [3]. A resurgence of severe GAS disease in recent decades has been mirrored by the emergence of a globally disseminated clone of the M1T1 serotype [4]. M1T1 strains are the most common cause of GAS pharyngitis and are strongly overrepresented in severe cases such as NF and TSS [5].

The ability of GAS to produce life-threatening infections even in previously healthy people reflects a diverse array of surface-associated and secreted virulence factors that together allow the bacterium to invade host cellular barriers, resist innate immune clearance, injure host tissues, and provoke harmful or dysregulated inflammatory responses [6, 7]. Remarkably, only in recent years was it recognized that major hair-like oligomeric surface organelles known as pili were present on the surface of Gram-positive bacteria including the pathogenic streptococci GAS, group B *Streptococcus* (GBS), and *Streptococcus pneumoniae* [8]. Targeted mutagenesis coupled with tissue culture assays and murine infection models have indicated that both GBS and pneumococcal pili each contribute to disease pathogenesis, but in different manners. Both pilus structures promote cell adherence and invasion [9, 10]; however, while GBS pili aid in resistance to host antimicrobial peptide and neutrophil killing mechanisms [11], pneumococcal pili stimulate exaggerated proinflammatory responses including tumor necrosis factor-α (TNF-α) release [9].

The role of the GAS pilus structure in disease pathogenesis has not been studied extensively in vivo. GAS pili are encoded by a highly variable locus known as the fibronectin-binding, collagen-binding T antigen (FCT) region. In M1 strains, this pathogenicity island encodes RofA, a transcriptional regulator, Cpa, a collagen-binding protein [12], Spy0128, the pilus backbone subunit, Spy0125 and Spy0130, two pilus accessory proteins, and Spy0129 (SrtC1), a sortase enzyme that polymerizes the GAS pilus proteins via covalent linkage. GAS pilus assembly requires expression of Spy0128 and SrtC1, but not the other genes. Interestingly, Spy0128 is now recognized to represent the classical Lancefield T1 antigen [13], a variable trypsin-resistant surface protein used for decades along with M protein for GAS typing schema based on serum recognition [14]. Deletion of Spy0128 suggests that GAS pili can contribute to biofilm formation [15] and bacterial aggregation in saliva [16], as well as adherence to human tonsil explants and some but not all human epithelial cell lines tested [15, 17]. In the M53 serotype background, full-length GAS pili were not required for disease establishment in a humanized mouse model of impetigo [18].

In this study, we generate a targeted deletion of the Spy0128 major pilus backbone subunit in a representative isolate of the M1T1 GAS clone, originally isolated from a patient with NF and TSS, to study the role of pili in disease pathogenesis using several in vitro and in vivo model systems. While pili are seen to promote GAS colonization phenotypes, they are found to restrict invasive disease pathogenesis, in direct contrast to findings reported for GBS and pneumococcal pili. A novel association with neutrophil extracellular killing mechanisms may help explain this unexpected finding.

Methods

Bacterial strains, mutagenesis, and complementation

WT GAS M1T1 strain 5448 was isolated from a patient with NF and TSS [19]. GAS were propagated in Todd–Hewitt broth (THB) or on THB agar (THA; Difco, BD, Franklin Lakes, NJ, USA). Precise allelic replacement of the spy0128 gene was performed using an established methodology [20]. Briefly, ~1,000 bp of flanking DNA immediately upstream and downstream of the spy0128 gene was amplified from the M1T1 chromosome by PCR using primers with 25 bp extensions corresponding to the 5′ and 3′ end of the chloramphenicol acetyltransferase (*cat*) gene. The upstream and downstream PCR products were then joined with the *cat* gene (from plasmid pACYC1094) by fusion PCR, and the resultant amplicon, containing an in-frame substitution of *spy0128* with *cat*, was subcloned into the temperature-sensitive suicide vector pHY304 to yield the knockout vector pSpy0128-KO. This vector was introduced into GAS M1T1 strain 5448 by electroporation and single and double crossover events identified by differential temperature and antibiotic sensitivity, using 5 μg/ml erythromycin (Erm) and 2 μg/ml chloramphenicol (Cm) for selection. The isogenic mutant 5448ΔPil was confirmed unambiguously by PCR and sequence analysis to contain a precise chromosomal replacement of *spy0128* with *cat*. For complementation analysis, the full-length *spy0128* gene was amplified from the GAS M1T1 5448 genome, subcloned into the streptococcal expression vector pDCerm [20], and introduced into the 5448ΔPil mutant using Erm selection; an empty vector control strain was
incubated at 37°C overnight. Then 2×10^6 colony forming cells/well in a tissue culture-treated 24-well plate and previously described [21]. Briefly, human pharyngeal Bacterial adherence to epithelial cells was determined as Epithelial cell adherence assays

Detection of pilus expression via immunofluorescent microscopy

Logarithmic-phase GAS were plated on Polysine microscope slides (Eerie Scientific) and fixed with 3% paraformaldehyde (PFA). Slides were blocked with 3% bovine serum albumin (BSA) in PBS containing 0.05% Tween-20 (blocking buffer) overnight at 4°C in a humidified chamber. Slides were incubated with goat serum diluted 1:200 in blocking buffer, followed by incubation with primary rabbit antiserum against the GAS T1 antigen, i.e., Spy1028 (obtained from Centers for Disease Control Streptococcal Laboratories, Atlanta, GA), diluted 1:1,000 in blocking buffer. The slides were then incubated with a secondary fluorescein isothiocyanate (FITC)-conjugated goat anti-rabbit IgG diluted 1:200 in blocking buffer. Mounted samples were viewed simultaneously with a ×60/1.42 PlanApo objective after laser excitation at 488 nm and differential interference contrast using a spectral deconvolution confocal microscope (Olympus FV1000).

Epithelial cell adherence assays

Bacterial adherence to epithelial cells was determined as previously described [21]. Briefly, human pharyngeal epithelial cells (HeP-2) and skin keratinocytes (HaCaT) obtained from ATCC were propagated as monolayers in RPMI-1640+10% fetal bovine serum (FBS), 2 mM L-glutamine, and 100 μM nonessential amino acids at 37°C+5% CO2. For adherence assays, cells were plated at 2×10^5 cells/well in a tissue culture-treated 24-well plate and incubated at 37°C overnight. Then 2×10^6 colony forming units (CFU) of logarithmic phase GAS (multiplicity of infection=MOI=10:1) in fresh RPMI-1640+2% FBS were added to each well. The plate was centrifuged at 1600 RPM for 5 min, then incubated at 37°C+5% CO2 for 30 min. Monolayers were washed with PBS to remove unattached bacteria, and 100 μl of trypsin and 400 μl of 0.025% Triton-X 100 were added to each well to facilitate cell detachment and lysis. Dilutions were plated on THB at 37°C overnight for enumeration of CFU.

Skin adherence assay

Eight-week-old female CD-1 mice (Charles River Laboratories, CA) were shaved and treated with Nair® (Church & Dwight Co., Inc.) 24 h prior to infection. Logarithmic phase WT and ΔPil mutant GAS were mixed 1:1 in PBS and 10 μL drops containing 1×10^6 total CFU plated on THA. Six-mm skin biopsy punches (Acuderm Inc.) were used to cut out disks of agar. Mice were sedated with ketamine/xylazine intraperitoneally (i.p.). Agar disks were placed onto mouse skin for 1 h to allow bacterial adherence; control agar disks were added to 2 mL Micro-tubes (Sarstedt) on ice. Skin around the periphery of the agar disks was marked and agar disks gently lifted off and discarded. Skin within the marked perimeter was harvested using 8-mm skin biopsy punches and placed in 2 mL Micro-tubes containing 1 mL of PBS on ice. Tubes containing skin were placed in the Mini Bead Beater (Biospec Products, Inc.) and gently shaken on “wash” setting for 1 min, supernatant was discarded, and 1 mL of fresh PBS was added. Skin was washed again for 1 min, supernatant was discarded, and 1 mL of PBS with 1 mm Zirconia/Silica beads (Biospec Products, Inc.) was added to these tubes and control agar tubes. Skin and agar controls were homogenized for 2 min, serially diluted, and replica plated on THA (for enumeration of all bacteria) and THA+2 μg/mL Cm (for enumeration of ΔPil alone).

Animal infection models

Eight- to ten-week-old female CD-1 mice were used for all experiments. Necrotizing subcutaneous infection was studied using a well-established model [21–23]. Briefly, logarithmic phase GAS were diluted 1:1 with sterile Cytodex beads (Sigma). Eight- to ten-week-old female CD-1 mice were shaved and Nair® applied 24 h prior to infection. Mice were injected subcutaneously in the left flank with 5×10^7 WT or ΔPil+Pil complemented mutant GAS and simultaneously in the right flank with an equivalent inoculum of the isogenic ΔPil mutant, allowing each mouse (N=8) to serve as its own control. Developing lesion sizes were measured daily, and on d4 mice were euthanized, skin lesions excised, homogenized, and serially diluted for plating on THA and enumeration of CFU. *Pneumonia* was modeled by introducing 1×10^6 CFU WT or isogenic ΔPil mutant GAS into the left nare of mice held in the upright position. At 24 or 48 h, mice were euthanized with inhaled CO2. The right lung was homogenized, serially diluted, and plated on THA for enumeration of total CFU. The left lung was inflated with 4% PFA for 5 min, resected, and placed in PFA for histology. Hematoxylin-eosin-stained lung sections were evaluated under light microscopy at ×10, ×40, and ×100 using a Zeiss Axiolab microscope. Systemic sepsis was modeled by injecting 2×10^8 CFU logarithmic phase WT or isogenic ΔPil mutant GAS in 100 μL of PBS via tail vein; mice were then monitored twice daily for mortality (N=11 or 12 animals per group total from two separate experiments).
Whole blood, macrophage, and antimicrobial peptide killing assays

For whole blood killing assays, a phlebotomy sample was obtained from a healthy human volunteer and 300 μL placed in each 2-mL siliconized tube (Fisher Scientific). Next, 100 μL of logarithmic phase GAS was added to reach a final concentration of 1×10^4 CFU/mL. Tubes were incubated on a tube rotator at 37°C. At 2 h, blood was serially diluted and plated on THA for enumeration of CFU. For J774 murine macrophage killing assays, cells were plated at 7.5×10^5 cells/well in RPMI-1640+2% FBS in a 24-well plate and incubated at 37°C+5% CO_2 overnight. Then 7×10^6 CFU of logarithmic phase GAS were added to each well and the plates were incubated for 30 min at 37°C+5% CO_2. Cells were rinsed with PBS, detached and lysed with 0.025% Triton X-100, serially diluted, and plated on THA for enumeration of total cell-associated CFU. For cathelicidin antimicrobial peptide killing assays, GAS were grown to logarithmic phase, brought up at 1×10^5 CFU/mL in 70% DMEM+10%FBS+20%THB, and 90 μL was added to 10 μL of varying concentrations of human (LL-37) or mouse (CRAMP) cathelicidin in triplicate wells in a 96-well plate, then incubated for 24 h at 37°C. Bacteria were plated on THA to determine the MIC for each peptide.

Complement deposition

Human serum was isolated from whole blood by coagulation in glass serum tubes (BD Biosciences) and stored at −80°C. Normal human serum (nhs) was generated by pooling sera from six different donors. To inactivate complement, nhs was heat-inactivated at 56°C for 30 min. GAS were grown to logarithmic phase and washed with HEPES-buffered saline containing 5 mM CaCl_2 and 2.5 mM MgCl_2 (HBS+++)+1% BSA (HBS+++−BSA). Bacteria were diluted ×200 and incubated with nhs at 0%, 2%, 5%, and 10% for 20 min at 37°C in a fast shaker. Bacteria were washed and incubated with FITC-conjugated anti-human C3 antibodies (Protos Immunoresearch) on ice for 30 min. Washed bacteria were taken up in PBS and analyzed on BD FACSCalibur flow cytometer, using BD CellQuest Pro, and data was analyzed with FlowJo (Tree Star, Inc.). Fluorescent mean, median, and geometric mean were obtained on all samples.

Neutrophil endothelial transcytosis assays

Human brain endothelial cells were propagated as monolayers in RPMI-1640+10% FBS and human lung endothelial cells (kind gift of Dr. Jeffrey Esko) were propagated as monolayers in DMEM (low glucose ×1; Mediatech) +20% FBS+1% nonessential amino acids+50 μg/mL heparin at 37°C+5% CO_2. Transwells (Costar) were placed inside tissue culture-treated 24-well plates, with 500 μL of media in the lower chambers. Cells were plated at 5×10^5 cells/upper chamber in 150 μL of media and incubated at 37°C+5% CO_2 until confluent monolayers formed. Next, 1×10^6 CFU of logarithmic phase WT, ΔPil+Pil complemented mutant, or ΔPil mutant were added to each lower chamber and 1×10^6 fresh human neutrophils were added to each upper chamber. At indicated time-points, the media in lower chambers was gently agitation, 30-μL samples removed, and the total number of neutrophils migrating to the lower chamber quantified with a hemocytometer under light microscopy.

Neutrophil IL-8 release and phagocytosis assays

To measure neutrophil interleukin-8 (IL-8) release, 1×10^5 human neutrophils were infected with 1×10^4 CFU of logarithmic phase GAS. At 3 h, 200 μL of media was removed and spun at 1,400 RPM, and the supernatant frozen at −20°C. Quantitative ELISA for IL-8 was performed using an IL-8 Quantikine kit (R&D Systems) according to the manufacturer’s instructions, using samples thawed on ice and diluted 1:4 in ELISA buffer. To assess neutrophil phagocytosis, GAS were labeled with FITC through a modified protocol derived from Goldmann et al., 2004 [24]. Logarithmic-phase GAS were concentrated in PBS to OD_600 of 1.0, FITC was added to a final concentration of 0.2 mg/mL, and samples incubated for 30 min, on ice, in the dark. Bacteria were washed with PBS until the supernatant was clear, then 1×10^6 CFU were added to 1×10^5 neutrophils (MOI 10:1) in RPMI-1640+2% FBS. Samples were incubated for 60 min at 37°C. Trypan blue was added for a final concentration of 0.5% to quench extracellular fluorescence. Samples were rinsed with PBS and then run on BD FACSCalibur flow cytometer, using BD CellQuest Pro, and data was analyzed with FlowJo (Tree Star, Inc.). Fluorescent mean, median, and geometric mean were obtained on all samples.

Neutrophil extracellular trap assays

To assess neutrophil extracellular trap (NET) induction, coverslips (1/2 oz, 12 mm; Fisher) were coated with Poly-D-lysine (Sigma) and placed in 24-well ultra low attachment surface plates (Costar) with 500-μL RPMI-1640+2% heat-inactivated human serum. Human neutrophils were plated at 5×10^5 per well, infected with 5×10^4 CFU logarithmic-phase GAS, and spun at 1,600 RPM for 10 min prior to incubation at 37°C+5% CO_2 for 20 min. PFA was added to final concentration of 4% and coverslips were inverted into ProLong® Gold with DAPI (Invitrogen). NETs were
visualized and quantified per visual field using a ×40/0.65 Achroplan objective with a Zeiss Axiolab fluorescent microscope. For NET adherence and killing assays, neutrophils were incubated with 25 nM phorbol myristate acetate (PMA) for 4 h to maximally induce NETs and eliminate phagocytosis. NETs were infected with GAS and spun at 1,600 RPM for 10 min. To quantify adherence, NETs and bacteria were incubated at 37°C+5% CO2 for 5 min. NETs were gently rinsed with PBS, then harvested with 0.025% Triton X-100, serially diluted, and plated on THA for enumeration of total NET-associated CFU. For killing assays, NETs and bacteria were incubated at 37°C+5% CO2 for 20 min. NETs and bacteria were harvested with 0.025% Triton X-100, serially diluted, and plated on THA for enumeration of total cell-associated CFU. In both adherence and killing assays, at the end of their respective incubations, two wells of each condition were treated with LIVE/DEAD BacLight staining (Molecular Probes), fixed with 1% PMA, mounted onto glass slides using ProLong® Gold with DAPI and immediately visualized under fluorescent microscopy.

Statistical analysis

Differences in bacterial counts were evaluated by Student’s \( t \) test. Significance was determined as \( P<0.05 \). Survival curves were evaluated by the Wilcoxon’s rank-sum test.

Assurances

Human blood collection and neutrophil isolation were approved by the University of California San Diego (UCSD) Human Research Protections Program. Animal studies were approved by the UCSD Institutional Animal Use and Care Committee.

Results

Mutagenesis and complementation of pilus expression in GAS strain M1 5448

To explore the functional role of pili in the pathogenesis of invasive M1T1 GAS disease, a knockout mutant (ΔPil) of strain 5448 was generated by precise allelic exchange mutagenesis of spy0128, then complemented by return of the spy0128 gene on an expression plasmid. To confirm functional changes in GAS pilus expression, we performed immunofluorescent labeling against Spy0128. Whereas surface pili were observed in the WT GAS parent strain and the complemented mutant, no pili were detected on the surface of the ΔPil mutant (Fig. 1a).

Pili contribute to GAS M1T1 epithelial cell adherence in vitro and skin adherence in vivo

Adherence phenotypes were assessed in cell lines representative of the two principal sites of GAS colonization. Compared to the WT GAS parent strain, the isogenic ΔPil mutant showed an 87% reduction in adherence to HaCaT human keratinocytes \( (P<0.001; \text{Fig. 1b}) \), a defect that was partially restored in the complemented mutant \( (P<0.005) \), and a 28% reduction in adherence to HEp-2 human pharyngeal epithelial cells \( (P<0.05) \); although the observed partial complementation (~12%) of the latter defect did not reach statistical significance. To assess the role of pili in GAS adherence to intact skin, a competition experiment was performed in which a 1:1 mixture of WT and ΔPil mutant GAS was inoculated on mouse skin for 1 h before harvesting and bacterial enumeration. Control sample mixtures were plated to confirm the initial inoculum. WT GAS outcompeted the isogenic ΔPil mutant, as demonstrated by a ratio of greater than 1 in all samples. The
median ratio of WT:ΔPil bacteria recovered was >6:1 in the excised skin samples (P<0.02; Fig. 1c).

Pilus expression blunts M1T1 GAS virulence in three different murine infection models

The contribution of pilus expression to GAS invasive disease pathogenesis was tested in three different model systems of infection, providing surprising but internally consistent results. First a subcutaneous injection model was used, in which GAS M1T1 produces necrotizing lesions or ulcers with histopathologic features resembling human NF. Compared to the WT GAS parent strain, the isogenic ΔPil mutant produced significantly larger lesions by d2 and d3 post-infection (P<0.02; Fig. 2a; representative mouse shown in Fig. 2c), while the complemented mutant (ΔPil+Pil) mutant had significantly smaller lesions on d3 post-infection than ΔPil harboring an empty vector control (P<0.02; Fig. 2b). Next, GAS pneumonia was established in mice by intranasal inoculation and lungs harvested at 24 or 48 h. We found significantly more bacteria were recovered from lungs of mice infected with the GAS ΔPil mutant than the WT parent strain at either time point (P<0.04; Fig. 3a). By 24 h, histopathology reflected the increased bacterial burden in the lungs of ΔPil-infected mice, and by 48 h, markedly more severe pneumonia with dense neutrophilic infiltration was evident in the ΔPil-infected mice vs. the WT GAS-infected mice (Fig. 3b). Finally, we modeled GAS sepsis by injecting the bacteria intravenously and monitoring mortality. In this study, the ΔPil mutant GAS demonstrated more rapid kinetics of lethality (50% dead by 72 h vs 96 h for the WT strain; 100% mortality in 96 h vs. 144 h for the WT strain, P=0.002, Fig. 3c). Thus in three distinct models of invasive M1T1 GAS infection, we observed pilus expression to diminish virulence potential, a finding opposite to those reported for the GBS or pneumococcal pili in systemic mouse infection models.

Pilus expression increases GAS susceptibility to human whole blood killing

To extend our animal observations to interactions with components of the human innate immune system, we assessed the contribution of pilus expression to M1T1 GAS survival in freshly drawn human blood. Consistent with diminished systemic virulence in the mouse studies, we found the ΔPil mutant had a 50% increase in blood survival compared with WT GAS parent strain (P<0.001); survival was reduced toward WT levels in the ΔPil+Pil complemented mutant (P<0.02; Fig. 4a). However, in contrast to findings reported for the GBS pili [11], GAS pilus expression did not alter the bacterium’s susceptibility to killing by the human cathelicidin antimicrobial peptide LL-37 (MIC=32 μM for both WT and ΔPil mutant) nor the murine cathelicidin antimicrobial peptide mCRAMP (MIC =16 μM for both strains). No difference was observed in WT vs. ΔPil mutant GAS susceptibility to killing by J774 murine macrophages (Fig. 4b), and similar degrees of complement C3b deposition were measured on the surface of the WT and ΔPil mutant bacteria after incubation in human serum (Fig. 4c). Consistent with equivalent levels of C3b-mediated opsonization, human neutrophil phagocytotic uptake of FITC-labeled WT, ΔPil mutant and ΔPil+Pil complemented mutant bacteria was similar as assessed by flow cytometry (Fig. 4d). Together these data indicate that the attenuating effect of pilus expression on M1T1 GAS virulence in the animal infection models or human blood killing studies was not linked to increased neutrophil opsonophagocytosis.

GAS pilus expression stimulates neutrophil IL-8 production and endothelial transcytosis

An important property reported for pneumococcal pili is the induction of increased levels of host pro-inflammatory
Fig. 3 Deletion of GAS pilus expression increases virulence in mouse models of pneumonia and sepsis. 

a Mice infected intranasally with GAS WT had significantly fewer bacteria in their lungs at 24 and 48 h.

b Representative hematoxylin–eosin-staining of lung sections demonstrates more histopathological damage in ΔPil lungs vs. WT-infected lungs at 24 and 48 h.

c Mice infected intravenously with the ΔPil mutant died an average of 72 h after infection compared with 96 h after infection for WT-infected mice (P=0.002). The survival curve was generated from pooling data from two experiments. At least three mice were used per group and all experiments were repeated three times.

Fig. 4 Pilus expression in GAS leads to increased susceptibility to whole blood killing and stimulates increased IL-8 release and transendothelial migration by human neutrophils.

a The ΔPil mutant had increased survival in human whole blood compared with WT GAS (P=0.0001) or the ΔPil+Pil complemented mutant (P=0.016).

b No difference in survival between pilated and non-pilated GAS strains in a macrophage killing assay.

c No difference in surface deposition of complement component C3b in piliated vs. non-piliated GAS.

d Human neutrophils phagocytose FITC labeled GAS WT, ΔPil mutant, and ΔPil+Pil complemented mutant strains with equivalent efficiency.

e Increased secretion of IL-8 by neutrophils exposed to WT GAS vs. the ΔPil mutant (P=0.009).

f Increased migration of neutrophils through an endothelial monolayer in response to WT GAS compared with the isogenic ΔPil mutant (P=0.03). All experiments were performed in triplicate and were repeated three times; representative experiment shown.
cytokine expression [9]. IL-8 is a multifunctional human innate defense protein with a key role in the recruitment and activation of neutrophils [25]. IL-8 is produced in large amounts by neutrophils upon detecting bacteria [26, 27] and acts in an autocrine fashion to stimulate bactericidal activities such as oxidative burst and release of granule enzymes [28, 29]. To evaluate the effect of MIT1 GAS pili on IL-8 production, freshly isolated human neutrophils were exposed to GAS and IL-8 quantified by ELISA. Neutrophils infected with the WT parent strain released ~35% more IL-8 than those infected with the isogenic ΔPil mutant (P<0.01; Fig. 4e). An established Transwell assay was utilized to examine how the presence of pili influenced human neutrophil endothelial transmigration toward the pathogen. Human microvascular endothelial cells were grown to confluence on a Transwell membrane, the lower chambers inoculated with WT, ΔPil mutant or ΔPil+Pil complemented mutant MIT1 GAS, and freshly isolated human neutrophils added to the upper chambers. As shown in Fig. 4f, the ΔPil mutant GAS elicited the transmigration of less than half as many neutrophils as the WT parent strain (P<0.04); genetic complementation of the mutant partially reversed the observed phenotype with a third more neutrophils transmigrating compared with the ΔPil mutant.

GAS pilus expression increases NET production and the entrapment of GAS within these structures

Release of extracellular DNA-based antimicrobial traps has recently been discovered as a new paradigm in the innate immune function of neutrophils [30, 31]. This process is morphologically distinct from other classical cell death processes including apoptosis and necrosis, and has been dubbed “NETosis” [32]. Binding of many species of bacteria and fungi within the DNA backbone has been demonstrated, and NETs contain cationic histones, granule proteases, and antimicrobial peptides that together kill the entrapped microbes [31, 33]. One of the proinflammatory stimuli known to strongly activate NET production is IL-8 [34, 35]. We found that human neutrophil exposure to the WT GAS parent strain resulted in a more than 3-fold increase in NET production as compared with treatment with the ΔPil mutant (P<0.0001); NET induction was restored to near WT levels in the ΔPil+Pil complemented mutant (P<0.0001; Fig. 5a). Representative fluorescent microscopy images showing increased NET production (red arrows) by human neutrophils when infected with WT and ΔPil+Pil GAS strains compared with infection with the ΔPil mutant GAS are shown in Fig. 5b. Because GAS pilus expression induced IL-8 production and NET release, and because pilus represent projecting structures that increase the surface area of the bacterium, we hypothesized that piliated GAS would have greater adherence to NETs and susceptibility to NET killing. For equalization, neutrophils were stimulated with PMA prior to allow maximal NET production prior to GAS exposure. Significantly more WT GAS and ΔPil+Pil complemented mutant GAS adhered to NETs than did ΔPil mutant GAS (P<0.003; Fig. 5c).

Representative images of FITC-labeled GAS entrapped within NETs, shown in Fig. 5d, demonstrate increased entrapment of WT and ΔPil+Pil GAS compared with the ΔPil mutant. Correspondingly, killing of GAS within the induced NETs was significantly greater for WT GAS (78% of inoculum) and ΔPil+Pil complemented mutant GAS (99%) than the ΔPil mutant strain (55%; P<0.05; Fig. 5e). Thus, we conclude that pilus expression enhances neutrophil extracellular clearance of GAS, through induction of IL-8, activation of NET formation, and enhanced entrapment of the bacterium within these structures.

Discussion

Through more than 50 years of research, the presence of long, pilus-like polymers extending from the surface of pathogenic streptococci went unrecognized. Now, a few years after genome analyses inspired their discovery [9, 13, 36], streptococcal pili are appreciated to influence host cell adherence, biofilm formation, interactions with the innate immune system, and virulence potential [9–11, 15, 17, 18, 37–39]. In the present study, we use targeted mutagenesis to examine the role of the major pilus subunit of GAS, now recognized to represent the classical T antigen of this species, in virulence phenotypes of the globally disseminated MIT1 clone that is the leading agent of pharyngitis and invasive infections. While extending the tissue culture results to demonstrate that GAS pili enhance adherence to intact mouse skin, we find a surprising negative effect of pilus on invasive disease potential, which contrasts the established virulence role of GBS and pneumococcal pili in similar mouse models. Thus, the functional roles of pili in streptococcal pathogenesis are not universal, but rather bear important species-specific attributes, such that generalizations should not be assumed. Although similarities are identified in the general genomic structure of streptococcal pilus loci, including the presence of sortases for covalent linkage of LP(X)TG sequence motifs in the major pilus subunits, significant sequence diversity in the pilus genes themselves would appear to allow for functional diversity of the encoded proteins [8].

A validated model for development of invasive MIT1 GAS infection shows that innate immune-mediated selection of mutations in the covR/S two-component regulator are a critical step in systemic dissemination of the pathogen [40–44]. Such mutations lead to upregulation of key MIT1 GAS virulence factors that promote resistance to neutrophil
defenses, while allowing accumulation and activation of host plasminogen on the bacterial surface, promoting tissue spreading [42, 44]. Upon in vivo selection, two of the most strongly upregulated GAS M1T1 genes, the IL-8 peptidase SpyCEP and the phage-encoded DNAse Sda1, promote resistance to NET-mediated killing, by blocking IL-8 stimulation of NET production and by degrading the DNA backbone of NETs, respectively [23, 35, 45]. Here we identify GAS M1T1 pilus expression as instead promoting susceptibility to NET-mediated killing, likely through increased IL-8-dependent NET production coupled with increased adherence to and physical entrapment within the generated NETs. The GAS M1T1 pilus thus plays an important role in virulence during establishment of infection by increasing epithelial adherence, but may simultaneously represent a pathogen-associated molecular pattern recognized by phagocytes, promoting increased killing of GAS upon arrival in blood or deeper tissues. Transcriptional analysis indicates the M1T1 GAS *spy1028* major pilus subunit gene is slightly downregulated (by 1.2 fold) upon the in vivo selected *cov*R/S mutations [40], which would reduce its contributions to NET generation and entrapment just as the above-mentioned NET-resistance genes are upregulated.

The observed activation of host cytokine release for GAS M1T1 was also reported for the pneumococcal pilus [9]; Similarly, the P pilus or type I pilus of uropathogenic *Escherichia coli* have been shown to stimulate IL-8 release and induce migration of neutrophils across cellular barriers [46, 47]. As we observed for the M1T1 GAS pilus, the surface expressed GAS M1 protein was recently recognized to stimulate NET production [48, 49], but M1 nevertheless promotes GAS survival within these structures because it reduces the bacterium’s susceptibility to human and murine cathelicidin antimicrobial peptides present in NETs [48]. In contrast, the GAS pilus was not observed to promote cathelicidin resistance, a feature reported for pili of certain strains of GBS [11].

Finally, though impeding invasive infection, our data confirm that pilus expression is important for M1T1 GAS adherence to human keratinocytes in vitro, as previously reported [17], and we show that this phenotype is robust enough to promote bacterial adherence to intact mouse skin. Similarly, our finding that pili promote adherence to human HEp-2 pharyngeal epithelial cells supports binding studies performed with tonsillar explants [17]. Because of the documented immunogenicity of the GAS pilus (T antigen) in natural infection and mouse vaccination studies [13, 15], the major pilus subunit Spy1028 continues to represent an outstanding candidate GAS vaccine antigen, with the potential to inhibit GAS colonization of the two major epithelial surfaces (skin and pharynx) where it is normally resident.
Acknowledgements This work was supported by NIH grant AI077780 to V.N. L.E.C.A. received a fellowship from The Hartwell Foundation. S.H.M.R. was supported by a grant from the European Molecular Biology Organization, H.C.M. by a National Science Foundation (NSF) Graduate Research Fellowship, A.M.T. by an A.P. Giannini Foundation Postdoctoral Fellowship and M.v.K.-B. through a fellowship from the Deutsche Akademie der Naturforscher Leopoldina (BMBF-LPD 9901/8-187). We thank Anna Cogen for sharing protocols for bacterial adherence to mouse skin.

Conflicts of interest The authors declare that they have no conflicts of interest.

Open Access This article is distributed under the terms of the Creative Commons Attribution Noncommercial License which permits any noncommercial use, distribution, and reproduction in any medium, provided the original author(s) and source are credited.

References

1. Carapetis JR, Steer AC, Mulholland EK, Weber M (2005) The global burden of group A streptococcal diseases. Lancet Infect Dis 5:685–694
2. Steer AC, Danchin MH, Carapetis JR (2007) Group A streptococcal infections in children. J Paediatr Child Health 43:203–213
3. Lamagni TL, Darenberg J, Luca-Harari B, Siljander T, Efstratiou AR, Schlievert PM (1992) Clonal basis for resurgence of serious Streptococcus pyogenes disease in Europe. J Clin Microbiol 46:2359–2367
4. Aziz RK, Korb M (2008) Rise and persistence of global M1T1 clone of Streptococcus pyogenes. Emerg Infect Dis 14:1511–1517
5. Cleyra PP, Kaplan EL, Hundle J, Wiazlo A, Kim MH, Hahser AR, Schliever PM (1992) Clonal basis for resurgence of serious Streptococcus pyogenes disease in the 1980s. Lancet 339:518–521
6. Cunningham MW (2000) Pathogenesis of group A streptococcal infections. Clin Microbiol Rev 13:470–511
7. Kwinn LA, Nizet V (2007) How group A Streptococcus circumvents host phagocyte defenses. Future Microbiol 2:75–84
8. Telford JL, Barocci MA, Margaret I, Rappuoli R, Grandi G (2006) Pili in gram-positive pathogens. Nat Rev Microbiol 4:509–519
9. Barocci MA, Ries J, Zogaj X, Hemsley C, Albiger B, Kanth A, Dahlberg S, Fernebro J, Moschioni M, Masignani V, Hultenby K, Taddei AR, Beiter K, Wartha F, von Euler A, Covacci A, Holden DW, Normark S, Rappuoli R, Henriques-Normark B (2006) A pneumococcal pili influences virulence and host inflammatory responses. Proc Natl Acad Sci U S A 103:2857–2862
10. Maisey HC, Hensler M, Nizet V, Doran KS (2007) Group B streptococcal pili proteins contribute to adherence to and invasion of brain microvascular endothelial cells. J Bacteriol 189:1464–1467
11. Maisey HC, Quach D, Hensler ME, Liu GY, Gallo RL, Nizet V, Doran KS (2008) A group B streptococcal pili protein promotes phagocyte resistance and systemic virulence. FASEB J 22:1715–1724
12. Kreikemeyer B, Nakata M, Oehmcke S, Ghoshwender C, Normann J, Podbielski A (2005) Streptococcus pyogenes collagens type I-binding Cpa surface protein. Expression profile, binding characteristics, biological functions, and potential clinical impact. J Biol Chem 280:33228–33239
13. Mora M, Bensi G, Capo S, Falugi F, Zingaretti C, Manetti AG, Maggi T, Taddei AR, Grandi G, Telford JL (2005) Group A Streptococcus produce pili-like structures containing protective antigens and Lancefield T antigens. Proc Natl Acad Sci U S A 102:15641–15646
14. Lancefield RC, Dole VP (1946) The properties of T antigens extracted from group A hemolytic streptococci. J Exp Med 84:449–471
15. Manetti AG, Zingaretti C, Falugi F, Capo S, Bombaci M, Bagioni F, Gambellini G, Bensi G, Mora M, Edwards AM, Musser JM, Graviss EA, Telford JL, Grandi G, Margarit I (2007) Streptococcus pyogenes pili promote pharyngeal cell adhesion and biofilm formation. Mol Microbiol 64:968–983
16. Edwards AM, Manetti AG, Falugi F, Zingaretti C, Capo S, Buccato S, Bensi G, Telford JL, Margarit I, Grandi G (2008) Scavenger receptor gp340 aggregates group A streptococci by binding pili. Mol Microbiol 68:1378–1394
17. Abbot EL, Smith WD, Siou GP, Chiriboga C, Smith RJ, Wilson JA, Hirst BH, Kehoe MA (2007) Pili mediate specific adhesion of Streptococcus pyogenes to human tonsil and skin. Cell Microbiol 9:1822–1833
18. Lizano S, Luo F, Bessen DE (2007) Role of streptococcal T antigens in superficial skin infection. J Bacteriol 189:1426–1434
19. Chatellier S, Ihendyane N, Kansal RG, Khambaty F, Basma H, Norrby-Teglund A, Low DE, McGee R, Korb M (2000) Genetic relatedness and superantigen expression in group A streptococcus serotype M1 isolates from patients with severe and non-severe diseases. Infect Immun 68:3532–3534
20. Jeng A, Sakota V, Li Z, Datta V, Beall B, Nizet V (2003) Molecular genetic analysis of a group A Streptococcus operon encoding serum opacity factor and a novel fibronectin-binding protein. SbX. J Bacteriol 185:1208–1217
21. Tommer AM, Kristian SA, Datta V, Jeng A, Gillen CM, Walker MJ, Beall B, Nizet V (2006) Serum opacity factor promotes group A streptococcal epithelial cell invasion and virulence. Mol Microbiol 62:15–25
22. Datta V, Myskowski SM, Kwinn LA, Chiem DN, Varik NL, Kansal RG, Korb M, Nizet V (2005) Mutational analysis of the group A streptococcal operon encoding streptolysin S and its virulence role in invasive infection. Mol Microbiol 56:681–695
23. Buchanan JT, Simpson AJ, Aziz RK, Liu GY, Kristian SA, Korb M, Feramisco J, Nizet V (2006) DNase expression allows the pathogen group A Streptococcus to escape killing in neutrophil extracellular traps. Curr Biol 16:396–400
24. Goldmann O, Rohde M, Chhatwal GS, Medina E (2004) Role of macrophages in host resistance to group A streptococci. Infect Immun 72:2956–2963
25. Kunkel SL, Standiford T, Kasahara K, Strierer RM (1991) Interleukin-8 (IL-8): the major neutrophil chemotactic factor in the lung. Exp Lung Res 17:17–23
26. Bazzoni F, Cassatella MA, Rossi F, Ceska M, Dewald B, Bagnioli M (1991) Phagocytosing neutrophils produce and release high amounts of the neutrophil-activating peptide 1/interleukin 8. J Exp Med 173:771–774
27. Strierer RM, Kasahara K, Allen RM, Standiford TJ, Rolf MW, Becker FS, Chesneau SW, Kunkel SL (1992) Cytokine-induced neutrophil-derived interleukin-8. Amer J Path 141:397–407
28. Simms HH, D’Amico R (1997) Studies on polymorphonuclear leukocyte bactericidal function: the role of exogenous cytokines. Shock 7:84–89
29. Thelen M, Peper P, Kernen P, von Tschamer V, Walz A, Bagnioli M (1998) Mechanism of neutrophil activation by NAF, a novel monocye-derived peptide agonist. FASEB J 2:2702–2706
30. Brinkmann V, Reichard U, Goossman C, Fauler B, Uhlemann Y, Weiss DS, Weinrauch Y, Zychlinsky A (2004) Neutrophil extracellular traps kill bacteria. Science 303:1532–1535
31. von Kockritz-Blickwede M, Nizet V (2009) Innate immunity turned inside-out: antimicrobial defense by phagocyte extracellular traps. J Mol Med 87:775–783

© Springer
32. Wartha F, Henriques-Normark B (2008) ETosis: a novel cell death pathway. Sci Signal 1:pe25
33. Fuchs TA, Abed U, Goosmann C, Hurwitz R, Schulze I, Wahn V, Weinrauch Y, Brinkmann V, Zychlinsky A (2007) Novel cell death program leads to neutrophil extracellular traps. J Cell Biol 176:231–241
34. Gupta AK, Hasler P, Holzgreve W, Gebhardt S, Hahn S (2005) Induction of neutrophil extracellular DNA lattices by placental microparticles and IL-8 and their presence in preeclampsia. Human Immunol 66:1146–1154
35. Zinkernagel AS, Timmer AM, Pence MA, Locke JB, Buchanan JT, Turner CE, Mishalain I, Sriskandan S, Hanski E, Nizet V (2008) The IL-8 protease SpyCEP/ScpC of group A Streptococcus promotes resistance to neutrophil killing. Cell Host Microbe 4:170–178
36. Lauer P, Rinaudo CD, Soriani M, Margarit I, Maione D, Rosini R, Taddei AR, Mora M, Rappuoli R, Grandi G, Telford JL (2005) Genome analysis reveals pili in Group B Streptococcus. Science 309:105
37. Dramsi S, Caliot E, Bonne I, Guadagnini S, Prevost MC, Kojadinovic M, Laliou I, Poyart C, Trieu-Cuot P (2006) Assembly and role of pili in group B streptococci. Mol Microbiol 60:1401–1413
38. Krishnan V, Gaspar AH, Ye N, Mandlik A, Ton-That H, Narayana SV (2007) An IgG-like domain in the minor pilin GBS52 of Streptococcus agalactiae mediates lung epithelial cell adhesion. Structure 15:893–903
39. Konto-Ghiorghi Y, Maire E, Dumenil G, Caliot E, Trieu-Cuot P, Dramsi S (2009) Dual role for pilus in adherence to epithelial cells and biofilm formation in Streptococcus agalactiae. PLoS Pathog 5:e1000422
40. Sumby P, Whitney AR, Graviss EA, DeLeo FR, Musser JM (2006) Genome-wide analysis of group a streptococci reveals a mutation that modulates global phenotype and disease specificity. PLoS Pathog 2:e5
41. Cole JN, McArthur JD, McKay FC, Sanderson-Smith ML, Cork AJ, Ranson M, Rohde M, Itzek A, Sun H, Ginsburg D, Kotb M, Nizet V, Chhatwal GS, Walker MJ (2006) Trigger for group A streptococcal MIT1 invasive disease. FASEB J 20:1745–1747
42. Walker MJ, Hollands A, Sanderson-Smith ML, Cole JN, Kirk JK, Henningham A, McArthur JD, Dinkla K, Aziz RK, Kansal RG, Simpson AJ, Buchanan JT, Chhatwal GS, Kotb M, Nizet V (2007) DNase Sda1 provides selection pressure for a switch to invasive group A streptococcal infection. Nat Med 13:981–985
43. Trevino J, Perez N, Ramirez-Pena E, Liu Z, Shelburne SA 3rd, Musser JM, Sumby P (2009) CovS simultaneously activates and inhibits the CovR-mediated repression of distinct subsets of group A Streptococcus virulence factor-encoding genes. Infect Immun 77:3141–3149
44. Musser JM, Shelburne SA 3rd (2009) A decade of molecular pathogenomic analysis of group A Streptococcus. J Clin Invest 119:2455–2463
45. Sumby P, Barbian KD, Gardner DJ, Whitney AR, Welty DM, Long RD, Bailey JR, Parnell MJ, Hoe NP, Adams GG, Deleo FR, Musser JM (2005) Extracellular deoxyribonuclease made by group A Streptococcus assists pathogenesis by enhancing evasion of the innate immune response. Proc Natl Acad Sci U S A 102:1679–1684
46. Sauer FG, Mulvey MA, Schilling JD, Martinez JJ, Hultgren SJ (2000) Bacterial pili: molecular mechanisms of pathogenesis. Curr Opin Microbiol 3:65–72
47. Godaly G, Otto G, Burdick MD, Strieter RM, Svanborg C (2007) Fimbrial lectins influence the chemokine repertoire in the urinary tract mucosa. Kidney Int 71:778–786
48. Lauth X, von Köckritz-Blickwede M, McNamara CW, Myskowski S, Zinkernagel AS, Beall B, Ghosh P, Gallo RL, Nizet V (2009) MI protein allows group A streptococcal survival in phagocyte extracellular traps through cathelicidin inhibition. J Innate Immun 1:202–214
49. Oehmcke S, Mörgelin M, Herwald H (2009) Activation of the human contact system on neutrophil extracellular traps. J Innate Immun 1:225–230