**Introduction**

*Streptococcus pyogenes*, which is commonly referred as *Group A Streptococcus* (GAS), is a major human pathogen that causes both relatively mild common infections, such as pharyngitis and superficial skin infections, and potentially lethal, severe invasive infections, including soft-tissue infections, pneumonia, and necrotizing fasciitis [1, 2]. These invasive infections often lead to systemic GAS dissemination, resulting in septic shock and streptococcal toxic shock syndrome. Progress has recently made on pathogenesis in severe invasive GAS infections that include the understanding of the role of CovRS mutations in hypervirulence, in vivo selection of GAS CovRS mutants, and innate immune evasion and vascular invasion by hypervirulent GAS CovRS mutants. This mini review focuses on the progress in these areas. Because of the limited scope, we do not intend to provide a thorough review on the subject but emphasize more on our own contributions.

**Hypervirulent Group A Streptococcus Variants**

Severe GAS infections were frequent and often fatal in the 19th century and reemerged in the 1980s. The reemergence of severe invasive GAS infections in the 1980s is associated with the emergence of the virulent MIT1 clone of genotype *emm*1 GAS and virulent *emm*3 GAS. The MIT1 clone of *emm*1 GAS has been evolved by the acquisition of DNase Sda1- and superantigen SpeA-encoding prophages and the replacement of a 36-kb chromosomal region of pre-1980 *emm*1 GAS with that of *emm*12 GAS that contains the NADase and streptolysin O genes [3]. Contemporary M3 GAS acquired a prophage that encodes the superantigen SpeK and phospholipase A2 SlaA [4]. Since 2000, M89 GAS with the loss of the genes for synthesis of the hyaluronic acid capsule has also emerged to cause severe invasive infections [5]. The 5 most prevalent *emm* genotypes of recent pharyngeal and invasive GAS isolates are *emm*1 (MIT1 clone), *emm*3, *emm*12, *emm*28, and *emm*89 [2, 6]. Invasive *emm*3 GAS causes a higher mortality rate than invasive strains of other genotypes [7].

Invasive GAS isolates are usually more virulent than pharyngeal isolates in experimental animal infections, displaying greater tissue invasion, innate immune evasion, and systemic dissemination than pharyngeal isolates [8, 9]. Invasive GAS isolates frequently carry mutations of the two-component regulator system CsrRS or CovRS [10, 11]. CovRS negatively regulates many virulence factors, including many of those that are...
involved in the evasion of innate immunity [12]. Natural CovRS mutations enhance virulence gene expression, innate immune evasion, systemic dissemination, leading to hypervirulence [8, 9, 12-14]. Natural nonsense mutations of the orphan kinase RocA in emm3 and emm18 GAS also contribute to enhanced virulence [15, 16]. Invasive emm3 GAS isolates also have prevalent mutations in RopB, the activator that is essential for the expression of the protease SpeB [10, 11].

**In vivo Selection of Hypervirulent GAS Mutants**

The emergence of CovS mutations during infection has been demonstrated in M1T1 and emm12 GAS during experimental mouse infections, and CovS mutants have higher expression of virulence genes and virulence [6, 13]. *In vivo* selected CovS mutants of M1T1 GAS has downregulated expression of SpeB, lacking detectable levels of the SpeB activity in culture supernatant (SpeB activity-negative or SpeB− phenotype) [14]. The SpeB− phenotype is a validated marker for selected CovS mutants of CovS of M1T1 and M12 GAS in mouse infection [6, 20].

No SpeB− variants can be detected after cutaneous infection with SpeB− emm3 isolates, and the emergence of emm3 GAS CovS mutants cannot be demonstrated in mice infection by screening for variants with the SpeB− phenotype [6]. These results are surprising because CovS mutations are frequent in clinical invasive emm3 isolates [10, 11]. However, the failure to demonstrate the emergence of emm3 GAS CovS mutants in mouse infection may be just a technical difficulty that arises in a possible distinction between emm1 and emm3 GAS in regulation of speB. The natural CovSΔG576V point mutation of invasive emm3 isolate MGAS315, like in M1T1 GAS, enhances expression of virulence genes and critically contribute to its virulence; however, this CovS missense mutation does not cause a SpeB− phenotype in MGAS315, and CovSΔG576V mutant cannot confer the SpeB− phenotype of covS deletion mutant of M1T1 GAS [9]. Apparently, screening with the SpeB activity assay is unable to identify certain emm3 GAS CovS mutations in mouse infection.

The DNase sda1 has been reported to be required for the *in vivo* selection of SpeB− variants of M1T1 GAS, leading to a proposal that Sda1 helps GAS avoid killing by neutrophil extracellular traps in the absence of SpeB production and thereby provides pressure for selection of CovRS mutations with the SpeB− phenotype [17]. However, the role of Sda1 in the selection of M1T1 GAS SpeB− variants cannot be confirmed, and deletion of all three DNase genes has no effect on the selection of M1T1 GAS CovS mutants in mouse infection [18]. The capsule synthase hasA and M protein emm genes are reported to be required for the selection of CovRS mutants of M1T1 GAS [19]. The first sequenced M1 GAS strain SF370, which has functional hasA and emm but does not belong to the M1T1 GAS clone, rarely switch to SpeB− during experimental mouse infection [18]. Thus, the exact switch for *in vivo* selection of hypervirulent M1T1 GAS CovS mutants remains elusive.

*In vivo* selection of M1T1 GAS CovS mutants is diminished in neutropenic mice that are obtained by depletion of neutrophils through the treatment of mice with anti-Ly6G monoclonal antibodies 1A8 and RB6-8C5 [20]. RB6-8C5 also depletes a subset of monocytes that differentiate into recruited macrophages during infections. However, the role of recruited macrophages in selection of GAS CovS mutants is ruled out based on the phagocyte responses and CovS mutant selection in CXCR2−/− mice. CXCR2 is a receptor for CXC chemokines and is involved in neutrophil recruitment. *In vivo* selection of M1T1 GAS CovS mutants is significantly reduced in CXCR2−/− mice. CXCR2−/− mice display impaired neutrophil recruitment but have higher recruited macrophages at infection sites in air-sac infection of mice with M1T1 GAS [20]. The decrease in the selection of M1T1 GAS CovS mutants in CXCR2−/− mice is apparently correlated with the impaired neutrophil recruitment but not with the increased recruitment of macrophages. The findings support that neutrophils are required for *in vivo* selection of GAS CovS mutants.

CovS mutant outcompetes wild-type bacteria in mice with normal neutrophil responses but is outcompeted by wild-type bacteria in neutropenic mice [20]. The finding demonstrates that CovS mutants can survive better than the parent GAS in the presence of neutrophils.

M1T1 GAS mouse infection also selects nonsense and missense mutations of RocA [21]. Selected RocA mutations enhance the expression of many CovRS-controlled virulence genes *in vitro* and *in vivo* but, unlike CovS mutations, do not downregulate speB transcription at stationary growth phase and in subcutaneous infection of mice.

The current model for the occurrence of hypervirulent GAS mutants during infection is presented in Figure 1. In this model, spontaneous covS and rocA mutations are selected by neutrophils, and enhance virulence gene expression, enhancing innate immune evasion, tissue invasion, and systemic dissemination and resulting in hypervirulence.

**Role of SpeB in an Apparent Source-Sink Dynamics of Wild-Type GAS and Hypervirulent GAS Variants**

Natural M1T1 GAS CovS mutations down-regulate SpeB production *in vitro* and *in vivo*. RopB mutations also down-regulate SpeB expression. M1T1 GAS RocA and CovS mutants have similar enhancement in many CovRS-controlled virulence factors [16] but RocA mutants do not downregulate SpeB production *in vivo* [21]. RocA mutants rank between wild type M1T1 GAS and CovS mutants in skin invasion, inhibition of neutrophil recruitment, and virulence in subcutaneous infection of mice [21]. RopB deletion of an emm3 isolate, which has a natural RocA mutation, enhances virulence [10]. These findings indicate that, in addition to high expres-
sion of CovRS-controlled virulence genes, the downregulation of SpeB is a significant factor for the hypervirulence of M1T1 GAS CovRS mutants and emm3 CovS/RocA/RopB and RocA/RopB mutants. It has been proposed that the downregulation of SpeB preserves virulence factors to enhance virulence [22]. This notion is supported by the phenotype of covR deletion mutants. Deletions of covR and covS enhance expression of many virulence genes similarly; however, covR deletion enhances SpeB expression. Deletion of covR enhances skin invasion but reduces systemic dissemination and virulence [23]. SpeB critically contributes to dermal ulceration caused by GAS covR deletion mutant [24]. Apparently, high levels of SpeB expression enhance localized infection and reduces systemic dissemination in soft tissue infections.

The majority of pharyngeal and invasive GAS isolates are SpeB+ [25], suggesting that CovRS mutations arise during human infection with GAS carrying wild-type CovRS and are not transmissible. The source-sink dynamics, a model in ecology that describe how variation in habitat quality affects the population growth or decline of organisms, can be borrowed for a proposal on the relationship of wild-type GAS and its hypervirulent variants and the role of SpeB in this population dynamics (Figure 2). GAS with wild-type CovRS and RopB produces high levels of SpeB, which is critical for causing dermal and mucosal purulent ulceration, degrades virulence factors for localized infection, and may also confer a survival advantage in pharyngeal infection. Thus, SpeB plays a significant role for localized, contagious

Figure 1. A model for in vivo selection of hypervirulent GAS mutants. M1T1 GAS CovRS mutants. Neutrophils select spontaneous covRS mutations of M1T1 GAS. M1T1 and M12 CovRS mutants and CovRS/RopB mutants enhance expression of multiple virulence genes and downregulate SpeB production, resulting in mutants with enhanced innate evasion, in vivo survival, skin invasion, systemic dissemination, and virulence. The figure was modified from one in reference 20.

Figure 2. A proposal for the role of SpeB in a model of source-sink dynamics of wild-type GAS and hypervirulent variants. The high expression of SpeB is critical for localized, contagious infection at the source habitat, non-invasive throat and skin surface whereas the downregulation of SpeB preserve virulence factors to facilitate systemic, less contagious but potentially lethal infections at the sink habitat, the tissues and blood that are usually germ-free.
infection at non-invasive throat and skin surface, which serves as the source habitat of GAS. GAS with CovRS and RopB mutations downregulates SpeB expression, which preserves virulence factors and indirectly enhances innate immune evasion and tissue and vascular invasion. Thus, the GAS variants cause systemic, less contagious, and potentially lethal infections at usually germ-free tissues and blood, which serves as the sink habitat of GAS.

**Evasion of Neutrophil Response by Hypervirulent GAS CovS Mutants**

Some necrotizing fasciitis (NF) patients have numerous bacteria but few or no neutrophilic responses at infection sites, which is classified as stage III NF [26], and other histopathologic types include a moderate-to-severe neutrophilic response and a positive Gram staining (stage II) and an intense neutrophilic response with the absence of bacteria (stage I) in infected tissues. Patients with stage III NF have a higher mortality rate than patients with stage I and II NF. Animals in a murine model of NF caused by hypervirulent GAS CovS mutants display stage III histopathologic features of few or no neutrophils at sites of bacterial infection [27, 28]. Correction of CovS mutations in hypervirulent GAS CovS mutants enhances the neutrophil response and lead to the Stage II histological type whereas covS deletion of wild-type M1T1 GAS reduces neutrophil recruitment and leads to the Stage II histological type in mouse infection [8,9]. Thus, CovS mutations enhance innate immune evasion.

CovS null mutations enhance the expression of the sse, spyCEP, and scpA genes. Sse, SpyCEP, and SscpA degrade the neutrophil chemoattractant molecules platelet-activating factor (PAF), interleukin 8/CXC chemokine, and C5a peptide, respectively. SsE, but not SpyCEP or SscpA, is required for the inhibition of neutrophil recruitment by hypervirulent M1T1 GAS CovS mutants [8]. Streptolysin S (SLS) delays the exodus of the neutrophils from the vessel lumen into the tissue at the early stage of GAS skin infection in mice [29]. SLS is also cytotoxic to neutrophils [30]. In mouse infection with hypervirulent CovS mutants, Sse and SscpA are both required and have synergistic effects for inhibition of neutrophil recruitment and systemic infection. Figure 3 illustrates the critical role of the covS mutation, Sse, and SLS in the inhibition of neutrophil recruitment by hypervirulent GAS CovS mutants in mouse model of skin infection.

**Vascular Invasion by Hypervirulent GAS CovS Mutants**

In a mouse model of pulmonary infection, hypervirulent GAS MGAS315 bacteria are sparsely scattered along the alveolar ducts and alveolar septa right after intratracheal inoculation, and mice are moribund at 24-48 h after inoculation. Alveolar ducts are often severely damaged and contain large numbers of bacteria, bacteria are associated with the septa of alveoli, and, more strikingly, intensive Gram staining for GAS are present in the peribronchovascular and perivascular interstitium (Figure 4) [31]. The peribronchovascular interstitium is the connective-tissue sheath that encloses the bronchi, pulmonary arteries, veins, and lymphatic vessels, whereas the perivascular interstitium extended from the peribronchovascular interstitium protects blood vessels that connect the microvasculature system in the alveolar region with the arteries and veins inside the peribronchovascular interstitium. Thus, MGAS315 has the capacity to invade the peribronchovascular interstitium.

MGAS315 has a natural covS G1370T mutation, which results in a CovS G457V missense mutation [9]. This CovS G457V mutation contributes to the hypervirulence of MGAS315 in subcutaneous infection of mice [9]. An isogenic strain derived from MGAS315 that carries the wild-type covS gene (MGAS315wtcovS) can still cause infections in the alveolar area but cannot invade the peribronchovascular and perivascular interstitium (Figure 4). Deletion of the covS gene in MGAS315wtcovS restored the infection of the peribronchovascular and perivascular interstitium (Figure 4). CovS mutation of MGAS315 is required for the peribronchovascular and perivascular invasion.

As MGAS315 invades the perivascular interstitium, the integrity of the smooth muscle and endothelial layers lining the blood vessels became disrupted and separated [31]. As segments of the smooth muscle and endothelial layers are further degraded, the bacteria can enter the lumen of the blood vessels. In contrast, blood vessels are intact in MGAS315wtcovS infection because bacteria infect the alveolar region but do not enter the connective-tissue sheath around blood vessels. MGAS315 loads in the liver and spleen are >1000-fold higher than MGAS315wtcovS in the infections. These data support a critical role of the vascular invasion in systemic MGAS315 dissemination.

The virulence factors that are directly involved in the vascular invasion are not known. Depletion of neutrophils and inflammatory monocytes does not lead to the perivascular invasion in MGAS315wtcovS infection. Thus, the role of CovS mutations in the perivascular invasion appears not to be directly linked to GAS evasion of neutrophil and inflammatory monocyte responses even though the innate immune evasion should be a prerequisite for the perivascular invasion. Hypervirulent GAS may cross the barrier between the alveoli and the peribronchovascular and perivascular interstitium and/or enter the peribronchovascular interstitium by crossing the epithelium in bronchioles. The damages on the smooth muscle and endothelial layers may involve CovRS-controlled virulence factors. Invasion of the vascular system by hypervirulent GAS likely involves coordinated actions of multiple virulence factors that are controlled by CovRS. Elucidation of the invasion route and virulence factors critical for understanding the process of systemic dissemination and developing novel therapies to treat severe invasive GAS infections.
Figure 3. The critical role of CovS mutation and PAF acetylhydrolase Sse and streptolysin S in inhibition of neutrophil recruitment by hypervirulent GAS CovS mutants. In mouse model of subcutaneous infection, hypervirulent GAS CovS mutant severely inhibits neutrophil recruitment, and correction of CovS mutation increases neutrophil recruitment. Deletion of sse increases neutrophils (PMN) recruitment that are largely in necrotic form, and further deletion of sagA, which encodes the peptide component of streptolysin S, leads to neutrophils full of the infection site and reduces neutrophil necrosis. Shown images are microscopic images of infection site sections with Gram stain. GAS and neutrophils were stained in blue and pink colors, respectively. The figure was derived from reference 28.

Figure 4. CovS mutations leads to the peribronchovascular and perivascular invasion the vascular attack in murine pulmonary infection. In pulmonary infection of mice from intratracheal GAS inoculation, hypervirulent GAS CovS mutant invades peribronchovascular and vascular invasion, and correction of CovS mutations leads the loss of perivascular and vascular invasions, which are restored by the deletion of the corrected sse gene. Shown are microscopic images of lung sections with Gram stain. Letters: A, alveoli; AS, alveolar septa; AD, alveolar duct; B, bronchiole; BT, bronchial terminus; PBVI, peribronchovascular interstitium; PVI, perivascular interstitium; and V, blood vessel. The figure was derived from reference 31.
In summary, natural mutations of virulence regulators CovRS, RocA, and RopB enhance GAS virulence. In vivo selection of CovRS mutants requires neutrophils, and CovRS mutants survive better than GAS with wild-type CovRS against neutrophil responses. Downregulation of SpeB due to CovS and RopB mutations contribute to GAS virulence in invasive infections. CovS mutations lead to inhibition of neutrophil recruitment at infection sites, and PAF acetylhydrolase Sse and streptolysin S are both required for and have synergistic contribution to the inhibition of neutrophil recruitment by hypervirulent GAS CovS mutants. CovS mutations also enable GAS to invade the perivascular interstitium and directly attack the vascular system for systemic dissemination. Despite the progress, the molecular mechanisms for in vivo selection of hypervirulent GAS mutants and their vascular invasion remain to be elucidated. Efforts in the elucidation of these mechanisms are expected to provide targets for the prevention and treatment of severe GAS infections.

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References
