SEC is an antiangiogenic virulence factor that promotes *Staphylococcus aureus* endocarditis independent of superantigen activity

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The superantigen staphylococcal enterotoxin C (SEC) is critical for *Staphylococcus aureus* infective endocarditis (SAIE) in rabbits. Superantigenicity, its hallmark function, was proposed to be a major underlying mechanism driving SAIE but was not directly tested. With the use of *S. aureus* MW2 expressing SEC toxoids, we show that superantigenicity does not sufficiently account for vegetation growth, myocardial inflammation, and acute kidney injury in the rabbit model of native valve SAIE. These results highlight the critical contribution of an alternative function of superantigens to SAIE. In support of this, we provide evidence that SEC exerts antiangiogenic effects by inhibiting branching microvessel formation in an ex vivo rabbit aortic ring model and by inhibiting endothelial cell expression of one of the most potent mediators of angiogenesis, VEGF-A. SEC’s ability to interfere with tissue revascularization and remodeling after injury serves as a mechanism to promote SAIE and its life-threatening systemic pathologies.

### INTRODUCTION

*Staphylococcus aureus* is the leading cause of infective endocarditis (IE) in high-income countries (1–3). *S. aureus* IE (SAIE) is an acute and invasive infection of the cardiac endothelium characterized by the appearance of vegetative lesions that form predominantly on heart valves (4). The pathognomonic vegetations are a meshwork of bacterial aggregates and host factors such as fibrin, fibrinogen, platelets, and erythrocytes (5). SAIE results in substantial damage to cardiac structures, in particular the valves and myocardium, due to tissue toxicity and necropurulent inflammation (6). Once established, SAIE can lead to severe complications, most notably congestive heart failure, stroke, acute kidney injury, and septic shock (3, 6, 7). Treatment of SAIE is challenging, requiring prolonged antibiotic therapy or surgery to remove infected valves (3, 6). Even with treatment, SAIE has a high rate of recurrence and a 22 to 66% mortality rate (1, 3). Infections are frequently associated with methicillin-resistant *S. aureus* (MRSA), which complicate treatment and increase risk of mortality (8). Furthermore, life-saving medical interventions (e.g., valve replacement, cardiac devices, and hemodialysis), an increasing population with underlying conditions (e.g., diabetes mellitus and immunosuppression), and advanced age also increase the risk of acquiring SAIE (1, 3). As a result, the incidence of SAIE has continued to increase (3). Unfortunately, the advances in cardiovascular medicine achieved in the past decade have failed to improve SAIE outcomes. Thus, understanding the mechanisms driving SAIE pathophysiology is not only of fundamental interest, particularly as it relates to bacterial factors critical for vegetation formation and secondary complications, but also of utmost importance to develop effective intervention strategies.

Epidemiological studies demonstrate a strong association between SAIE and a select group of superantigen (SAg) genes, where 18 to 25% of SAIE clinical isolates encode entC [staphylococcal enterotoxin C (SEC)], 9 to 20% encode tsstH [toxic shock syndrome toxin (TSST1)], and 58 to 90% encode the enterotoxin gene cluster (egc) (9). Consistent with these studies, SEC, TSST1, and the egc SAg SEI, SEM, SEO, and SE like (l)-U all contribute to IE and metastatic infection in experimental IE (10, 11). However, the underlying mechanism by which SAgS contribute to SAIE pathogenesis remains speculative. Classically, SAgS are known for their potent T cell mitogenic activity resulting in dysregulated activation and cytokine production, leading to inflammatory syndromes and, in extreme cases, toxic shock (12). Superantigenicity results from toxin cross-linking of the Vβ chain of the T cell receptor (TCR) to the major histocompatibility complex class II (MHC-II) receptor on antigen-presenting cells (12). Of relevance to SAIE, endothelial cells also express MHC-II and function as conditional antigen-presenting cells capable of cross-linking Vβ-TCR, resulting in endothelium-mediated superantigenicity (13).

The dysregulated immune activation caused by SAgS distracts and diverts the immune system (14). It is also the etiology of multiple diseases including atopic dermatitis, pneumonia, extreme pyrexia, purpura fulminans, and toxic shock syndrome (12). The commonly accepted model of the role of SAgS in SAIE includes localized or systemic superantigenicity that causes dysregulation of the immune system preventing clearance of *S. aureus* from the infected heart endothelium. SAgS also cause capillary leak and hypotension, which changes the hemodynamics of the vascular system (12). This alteration of blood flow may enhance vegetation formation. However, the requirement of superantigenicity in the pathogenesis and pathophysiology of SAIE has not been directly tested. In this study, we addressed the hypothesis that superantigenicity promotes SAIE and disease sequelae.

We used the rabbit model of native valve IE with the well-characterized MRSA strain MW2 (SEC) and MW2 stably expressing SEC toxoids (TCR or MHC-II/TCR inactivated) to provide evidence that SEC critically contributes to vegetation growth, the magnitude
of myocardial inflammation, and injury to the renal and hepatic systems independent of superantigenicity. We demonstrate that development of septic vegetations is a prerequisite for decreased renal function, while superantigenicity resulting from SAIE exacerbates hepatocellular injury and systemic toxicity. With the use of the ex vivo rabbit aortic ring model of angiogenesis and human aortic endothelial cells, we provide evidence that SEC inhibits revascularization and inhibits expression of vascular endothelial growth factor–A (VEGF-A), a potent proangiogenic factor. These results demonstrate that SEC is an antiangiogenic virulence factor that directly modifies endothelial cell function by a mechanism that can promote SAIE and its associated systemic pathologies.

**RESULTS**

*S. aureus* MW2 expressing SEC toxoids are deficient in SAg activity in vitro and in vivo

SAg activity is the most potent biological function of staphylococcal enterotoxins described to date and causes lethal pathologies. To establish whether superantigenicity promotes development of SEC-mediated SAIE, we complemented *S. aureus* MW2Δsec to produce SEC with an inactive TCR-binding site (SEC N23A) or to produce SEC with dual inactivation of the TCR- and MHC-II–binding sites (SEC N23A/F44A/L45A) (tables S1 and S2). Because of its biological inactivation and its associated systemic pathologies.

### Superantigenicity is not required for vegetation formation and high lethality

*S. aureus* SEC**, S. aureus** SEC**RO, S. aureus** SECN23A, and S. aureus** SECN23A/F44A/L45A** were tested in the rabbit native valve IE model. Strains were inoculated intravenously with 2 × 10⁷ to 4 × 10⁷ total colony-forming units (CFUs) after mechanical damage to the aortic valve and monitored for a period of 4 days. During that period, rabbits infected with *S. aureus* SEC**RO** had a ~66% decrease in overall vegetation formation, where 8 of 12 rabbits had no vegetation

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**Fig. 1. SEC toxoids are deficient in SAg activity.** (A) CD4⁺ T cell replication index of peripheral blood mononuclear cells (PBMCs) stimulated for 6 days with 0.3, 2, and 10 μl of cell-free supernates from overnight cultures of *S. aureus* SEC**RO**, SEC**, SECN23A, and SECN23A/F44A/L45A**. PBMCs ± 10 μg of purified SEC were used as controls. (B and C) Serum levels of interleukin-6 (IL-6) and lactate dehydrogenase (LDH) 48 to 96 hours after infection with indicated *S. aureus* strains in experimental IE. *S. aureus* SEC** (n = 12), S. aureus SEC**RO (n = 15), S. aureus SECN23A (n = 17), and S. aureus SECN23A/F44A/L45A (n = 17). Data are means ± SEM. The dashed line is the average analyte value of all rabbits before infection. Statistical significance was determined by one-way ANOVA with Holm–Šidák’s multiple comparisons test with each SEC-producing strain compared to *S. aureus* SEC**RO**. All experimental groups were tested against preinfection values and were statistically significant, *P < 0.0001. (B) *P = 0.03. (C) *P = 0.02.
(Fig. 2, A and B) and 4 of 12 had very small vegetations (sent to pathology) with an estimated weight of 5 to 15 mg. In stark contrast, *S. aureus* SEC+ formed vegetations in nearly all rabbits (14 of 15). Vegetation size ranged from 11 to 116 mg, with most vegetations weighing >25 mg (Fig. 2, A and B). Six vegetations were sent for histopathology by a board-certified veterinary pathologist. Unexpectedly, complementation with SEC_N23A or SEC_N23A/F44A/L45A restored vegetation formation to wild-type levels in most rabbits, with vegetation sizes ranging from 12 to 103 mg for SEC_N23A (in 13 of 17 rabbits) and 3 to 107 mg for SEC_N23A/F44A/L45A in 12 of 17 rabbits (Fig. 2, A and B). Together, infection with *S. aureus* expressing toxoids produced vegetations in 74% of rabbits (25 of 34), compared to 93% (14 of 15) in rabbits infected with *S. aureus* SEC+. Vegetations formed by *S. aureus* SEC_N23A and *S. aureus* SEC_N23A/F44A/L45A also had bacterial counts comparable to the wild-type strain (Fig. 2C). Consistent with the contribution of vegetation formation to lethal systemic complications, infection with either *S. aureus* SEC_N23A or *S. aureus* SEC_N23A/F44A/L45A led to high lethality, with ~50% of rabbits succumbing to infection during the experimental period compared to 73% of rabbits infected with *S. aureus* SEC+ (Fig. 2D). Rabbits infected with *S. aureus* expressing toxoids also exhibited higher degrees of splenomegaly (Fig. 2E) and, on average, higher bacteremia (>1 × 10^3 CFU/ml) than those infected with *S. aureus* SEC^KO^, although only those infected with SEC_N23A/F44A/L45A exhibited significantly higher bacteremia (Fig. 2F). Overall, there is an ~20% decrease in vegetation formation and lethality in rabbits infected with strains producing toxoids. Thus, superantigenicity alone does not fully account for the high lethal outcomes associated with SEC production in experimental SAIE. These results highlight a critical requirement of SEC in SAIE that is independent of superantigenicity and MHC-II interactions.

**Superantigenicity-independent effects drive myocardial inflammation in SAIE**

SAIE presents as rapidly growing vegetative lesions resulting in the quick destruction of valvar leaflets and progression of the infection into the myocardium and adjacent structures (5). We then asked whether SEC superantigenicity promotes extension of the vegetative lesion into the surrounding tissue changing the overall cardiac pathology. To address this, we performed histopathological analyses on transverse sections of hearts containing vegetations (figs. S3 and S4 and table S3). Hearts from rabbits infected with *S. aureus* SEC^+, *S. aureus* SEC_N23A, or *S. aureus* SEC_N23A/F44A/L45A were selected randomly based on the presence of vegetations (mean size, 6.6 ± 2 mm^2; n = 15). We processed all hearts of *S. aureus* SEC^KO^-infected rabbits with visible vegetations (mean size, 2.7 ± 1 mm^2; n = 4) and three hearts without vegetations. We recognize that there is selection bias in this case (and the data will reflect that), but we wanted to learn

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**Fig. 2. SEC is required for vegetation formation independent of superantigenic activity in the rabbit model of native valve IE and sepsis.** (A) Representative images of aortic vegetations (areas indicated with dashed lines). (B) Total weights of vegetations dissected from aortic valves from infected rabbits. (C) Bacterial counts recovered from aortic vegetations. (A to C) Bars represent median value. Statistical significance, one-way ANOVA with Holm–Šidák’s multiple comparisons test compared to SEC^KO^. (D) Percent survival of rabbits infected intravenously with 2 × 10^7 to 4 × 10^7 CFU of indicated strain over 4 days. *P* = 0.03, **P* = 0.006, ***P* = 0.0007, log-rank Mantel–Cox test. (E) Enlargement of the spleen resulting from *S. aureus* infection. (Dashed lines) Average spleen size of uninfected rabbit controls (0.86 ± 0.1, n = 3). (F) Bacterial counts per milliliter of blood recovered from rabbits postmortem. (D to F) Bars represent median value. *P* = 0.03, **P* = 0.003, ns = not significant, one-way ANOVA with Holm–Šidák’s multiple comparisons test. Statistical significance compared to SEC^KO^. 

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pathologies associated with vegetation formation, including the 33% that forms in the absence of SEC production. Consistent with histopathology of SAIE described in humans, *S. aureus* vegetations in rabbits were composed of large aggregates of bacterial colonies interspersed in a fibrinous meshwork of host factors and cell debris (figs. S3 and S4). However, the vegetative lesions were heterogeneous across infection groups in presentation (location and size of bacterial clusters) and in the magnitude of suppurative intracardial complications (myocardial inflammation and septic coronary arterial emboli) (Fig. 3). In rabbits infected with *S. aureus* SEC⁺, most vegetations were located on aortic valve cusps and valve leaflets, with large clusters of bacteria present on the leaflets and intermixed within the central core of the vegetation adjacent to the aorta. A few vegetations formed on the aortic wall, were transmural (across the entire wall), and extended into the adjacent adipose tissue (fig. S4A). Rabbits infected with *S. aureus* producing SEC toxoids (SECN23A or SECN23A/F44A/L45A) exhibited very similar histologic presentation to each other and to those infected with *S. aureus* SEC⁺ (fig. S4, B and C).

Most vegetative lesions presented with inflammation adjacent to the vegetations that was almost exclusively heterophilic (neutrophilic)

Fig. 3. Vegetative lesions present with bacterial infiltration, inflammation, and necrosis. (A) Examples of histopathologic assessment of myocardial inflammation during rabbit model of native valve IE (graded 0 to 3). Inflammation was graded on the basis of the amount of inflammatory cell infiltrate noted within the myocardium. 0 = no inflammation, 1 = rare, scattered infiltrate (arrows), 2 = multifocal bands of infiltrate, 3 = large coalescing bands of diffuse infiltrate with necrosis. Scale bars, 100 μm; inset scale bars, 20 μm. (B) Scoring of myocardial inflammation from H&E-stained images 48 to 96 hour after infection. No statistical significance among wild-type and toxoid groups, Fisher’s exact test with categorical pathology grades 0 and 1 and grades 2 and 3 against SECKO. (C) Examples of a fibrinonecrotic focus (dotted outline; note the myocardial necrosis within the outline, which is a lighter pink color), (*) a centrally located septic thrombus, and (arrows) intravascular bacteria (deeply basophilic/blue material). Scale bars, 100 μm (left) and 20 μm (right). (D) Examples of an epicardial lesion with saponification (necrosis) of epicardial fat (arrows) and a locally extensive zone of myocardial mineralization (dotted rectangle). Scale bar, 100 μm. (E) Histopathology scoring for the presence or absence of cardiac pathological findings: bacterial thrombi and associated necrosis, intravascular (I.V.) bacteria, and epicardial fibrin and inflammation.
FCN N23A/SEC KO infected kidneys, while exhibiting high-grade inflammation (Fig. 4F), failed to exhibit infarction, thrombi, and intravenous bacteria (all signs of embolic events) (Fig. 4G). Hence, SEC-mediated acute kidney injury is largely a result of vegetation formation and fragmentation rather than kidney failure that typically results from toxic shock syndrome (12).

Liver pathology presented grossly as pale, spleen-shaped, ischemic lesions that were focal, multifocal, or widespread (Fig. 5A and table S3). Livers from S. aureus SEC+–infected rabbits presented with high-grade pathology in 80% of the animals (Fig. 5B). Similar liver pathology developed in ~70% of rabbits infected with S. aureus producing SEC toxoids (Fig. 5B). Rabbids infected with S. aureus SEC+ had significantly increased mean levels of serum aspartate aminotransferase (AST; 236 U/liter) compared to the AST concentration from rabbits infected with S. aureus producing SEC toxoids or S. aureus SEC KO (AST < 73 U/liter) (Fig. 5C). Correspondingly, the ratio of AST to alanine aminotransferase (ALT) was significantly higher in rabbits infected with S. aureus SEC+ (Fig. 5D). These increases in liver aminotransferase enzymes are consistent with what is observed in humans during acute liver injury and ischemic hepatitis (26, 27). As with the kidneys, ischemic liver lesions were commonly associated with embolic events. Histologic analyses showed ischemic necrosis (well-demarcated zones of coagulation necrosis), often surrounded by necropurulent inflammation and/or hemorrhage, and more diffuse centrilobular vacuolation and/or necrosis (indicative of ischemia on a more global scale of the organ) (Fig. 5, E and F). Together, the data indicate that, in addition to the embolic events resulting in localized tissue injury, liver sensitivity to the toxic effects of SAg activity significantly increases hepatocellular injury.

**SEC inhibits neovessel formation in rabbit aortic explants**

SEC promoting SAIE and lethal systemic complications largely independent of superantigenicity centers its pathophysiologic effects directly on the endothelium. In acute IE, vegetative lesions develop rapidly with no evidence of repair (28). This is followed by fragmentation and septic embolization of cardiac vegetations leading to tissue injury and organ dysfunction, increasing mortality in patients with IE (29). Therefore, SEC targeting of endothelial cell function in vascular repair and revascularization of injured/ischemic tissues may be central to both localized and systemic pathologies characteristic of SAIE. Angiogenesis is the complex, coordinated process of vascular formation, critical to capillary development and tissue remodeling after injury (30, 31). To assess SEC effects on vascular regeneration in a physiologically relevant context, we used the ex vivo rabbit aortic ring model, which preserves the vessel microenvironment. Ring sections dissected from excised thoracic and abdominal aortas were embedded into a basement membrane matrix to induce angiogenic sprouting along severed edges. Rings were cultured under toxin conditions (SEC N23A), antiangiogenic conditions (axitinib), or proangiogenic conditions (basal medium with growth supplements) for 14 days. SEC N23A was used in our in vitro studies because (i) the endothelium does not express TCR and (ii) SEC is a select agent toxin that we can only modify in the SEC N23A background for follow-up studies. Axitinib is an antiangiogenic molecule due to its ability to inhibit signaling via VEGF-A receptors 1, 2, and 3, thereby preventing VEGF-A overexpression in endothelial cells (32). VEGF-A is the most potent proangiogenic growth factor (33, 34).
In the rabbit aortic ring model of angiogenesis, evidence of microvessel sprouting was observed between days 3 and 5 in rings maintained in supplemented basal medium, with sprouts elongating and branching over 14 days (Fig. 6A, top). Of interest, abdominal rings displayed denser microvessel networks with a sprout surface area of 24 ± 1.8 mm² compared to thoracic rings that exhibited a surface area of 15 ± 2.1 mm² (Fig. 6A, top). As expected, rings treated with the antiangiogenic molecule axitinib failed to sprout (fig. S5B).

For SEC N23A-treated rings, the dose-response analysis showed significant inhibition of sprouting from both thoracic and abdominal aortic rings (Fig. 6B). At the lowest treatment dose of 2.5 μg, SEC N23A significantly decreased the thoracic ring sprouting area by ~93% over time and the abdominal ring sprouting area by ~72% (Fig. 6B). Regardless of aortic origin, sprouts that formed on SEC N23A-treated rings were detected by day 7, but their further development was stunted (Fig. 6, C and D, and fig. S5, A and C). These results provide evidence that SEC is an antiangiogenic virulence factor with the potential to effectively inhibit vascular and tissue regeneration by interfering with neovessel formation.

SEC inhibits expression of the proangiogenic factor VEGF-A in human aortic endothelial cells

Revascularization and vascular injury repair mechanisms are driven by proangiogenic factors released by injured tissues (30, 31). Hence, we sought to establish whether SEC modifies the angiogenic response in endothelial cells by altering expression of angiogenesis-related genes. To do so, we profiled the expression of 24 genes important in driving multiple steps of the angiogenic response by quantitative reverse transcription polymerase chain reaction (RT-qPCR) (table S4). Expression was assessed in immortalized human aortic endothelial cell (iHAEC) at 2, 4, and 6 hours under toxin conditions [lipopolysaccharide (LPS) or SEC N23A], angiogenesis-enhancing conditions (VEGF-A), or angiogenesis-inhibiting conditions (axitinib). The gene expression profiles were expressed as fold change from medium control, with a cutoff of ≥2-fold or ≤−2-fold as thresholds for 100% increases or decreases in expression levels, respectively. iHAECs grown in standard endothelial cell culture conditions have been shown to produce an angiogenesis protein profile consistent with cells that...
are triggered to sprout and are therefore already under proangiogenic conditions (35). We therefore stimulated iHAECs with bacterial LPS (25 ng/ml) to induce an expression profile characteristic of proinflammatory and pathological angiogenesis (36). As expected, LPS-treated cells exhibited a highly proinflammatory profile, inducing expression of genes encoding for monocyte chemoattractant protein-1 (MCP-1), granulocyte-monocyte colony-stimulating factor-2 (GM-CSF2), and IL-8 (Fig. 7A). The effect of this proinflammatory profile is reflected on the increased gene expression for factors important in extracellular matrix degradation (Serpin E1), growth and survival [heparin-binding epidermal-like growth factor (HB-EGF), VEGF-A, and VEGF-C], and migration, proliferation, and vessel sprouting (pentraxin 3 and angiopoietin-2) (Fig. 7B). VEGF-A stimulation of iHAECs under already proangiogenic conditions resulted in induction of its own gene expression while inhibiting genes involved in inflammation, growth, migration, and proliferation (Fig. 7). These results are consistent with an endothelial cell monolayer in homeostasis with no need for further growth. The VEGF-A receptor inhibitor axitinib induced a highly suppressive profile, exhibiting a ≥5-fold down-regulation of genes important in extracellular matrix (ECM) degradation [a disintegrin and metalloproteinase with thrombospondin motifs 1 (ADAMTS-1)], growth and survival [acidic fibroblast growth factor (FGF) and VEGF-A], proliferation and migration [tissue inhibitor of matrix metalloproteinase 4 (TIMP-4) and thrombospondin 1], and inflammation (GM-CSF2) (Fig. 7). In stark contrast, iHAEC treatment with SEC N23A at nontoxic concentrations (20 µg/ml; fig. S6) induces marked inhibition of VEGF-A gene expression, also observed in early time points of axitinib-treated cells (Fig. 7). These results indicate that SEC can exert antiangiogenic effects by inhibiting VEGF-A expression in endothelial cells and help explain SEC N23A suppression of microvessel formation in the aortic ring model.

**DISCUSSION**

*S. aureus* SAgs are ubiquitous among human clinical isolates and have been implicated in both colonization and pathogenic mechanisms (37). Although their significance as virulence factors has been established, how these multifunctional SAgs specifically contribute to
S. aureus pathogenesis has become a pressing question (38–40). Current evidence indicates that the SAg SEC, TSST1, and select eGc toxins play a previously unidentified and essential role in the etiology of SAIE by yet uncharacterized mechanisms (10, 11). Still, in SAg-mediated illnesses, superantigenicity is placed as a triggering event of SAIE by yet uncharacterized mechanisms (10). It has been proposed, albeit not directly tested, that superantigenicity leading to hypotension and immune dysregulation allows bacterial immune evasion and persistence, while direct interaction with the heart endothelium promotes endothelial dysfunction and disease progression (10, 11, 38). With the use of S. aureus producing SEC inactivated in MHC-II and/or TCR binding, we provide evidence that superantigenicity is not the primary mechanism by which SEC promotes vegetations formation, cardiac toxicity, and extracellular complications such as acute kidney injury. Our results are consistent with published studies noting that not all SAgS promote SAIE to the same extent (11). SEG and SEN have comparable T cell mitogenic activity as SEA, SEB, and TSST1 (41, 42), yet they exhibit decreased efficiency in promoting SAIE. Our studies highlight the critical contribution of SAgS in life-threatening pathologies by at least two novel mechanisms: (i) interference with vascular injury repair and (ii) inhibition of revascularization of injured or ischemic tissues.

During SAIE, vascular injury results from mechanical damage and/or toxin-mediated damage of the endothelium. Reendothelialization of injured vessels restores their integrity and is essential to limit the progressive loss of the endothelial layer and continual deposition of host factors (e.g., platelets, fibrin, and leukocytes) that leads to mural thrombus formation (37). In the case of SAIE, reendothelialization defects would promote expansion of the vegetative lesion and growth of the vegetation. We have previously shown that the SAg TSST1 directly causes dysregulated activation of iHAECs and disrupts reendothelialization in an in vitro wound-healing assay, suggesting a role of SAgS in endothelial dysfunction and a possible explanation for their role in SAIE (38). Both reendothelialization and sprout elongation are a function of cell proliferation and migration, and their impairment results in angiogenesis defects (43). Angiogenesis is a highly complex multistep process requiring endothelial cell activation, basement membrane degradation, sprout formation and elongation, and capillary network stabilization (30, 31). These processes are also supported by endothelium-associated cells, such as pericytes, fibroblasts, monocytes, and smooth muscle cells (44, 45). These cell types are not present in standard monoculture in vitro wound-healing assays. For the study of angiogenesis, the ex vivo rabbit aortic ring model is a more physiologically relevant model (30, 46). With this model, we provide conclusive evidence that SEC disturbs branching microvessel formation, effectively inhibiting angiogenesis.

We propose that SEC inhibition of angiogenesis occurs via a mechanism dependent on down-regulation of one of the most potent inducers of angiogenesis, VEGF-A (33, 34, 47, 48). VEGF-A is a proangiogenic growth factor produced in virtually all vascularized tissues (33, 48). It promotes endothelial cell survival and mitogenic stimuli (49, 50), reendothelialization of denuded arterial walls to restore the integrity of the endothelium (51), and growth of new capillaries to restore perfusion to ischemic tissues (52, 53). Physiological levels of VEGF-A maintain vascular homeostasis and are vasculoprotective, while VEGF-A up-regulation is critical to trigger
sprouting angiogenesis and wound repair mechanisms (54–56). We provide evidence that SEC inhibits expression of VEGFA in iHAECs. This inhibition peaks between 2 and 6 hours after SEC exposure. Similar inhibition is observed in cells exposed to axitinib, a potent inhibitor of the VEGF-A receptor (32). Inhibition of the VEGF-A signaling pathway, regardless of the mechanism (e.g., VEGF-A neutralization or VEGF-receptor inhibition), results in endothelial dysfunction and vascular toxicity, including thrombosis, inappropriate pruning of the arterial vasculature, and revascularization defects during ischemic injury (57–59). These events can help explain the pathophysiology typical of SAIE.

SAIE has a high mortality rate owing to the high incidence of both intracardiac complications arising from the rapid local spread of the infection and the high incidence of embolization of septic vegetation fragments (1, 3, 60). Lodging of septic emboli within terminal blood vessels causes localized ischemia and infarction in multiple organ systems. In humans and in experimental rabbits, these complications can be manifested as myocardial infarction, kidney and/or liver injury, and strokes (60). Of these, kidney injury leading to acute renal insufficiency with progression to acute renal failure is tightly associated with SAIE severity, development of septic shock, and IE lethality (7). Our study demonstrates that SEC-associated kidney injury is a consequence of its role in vegetation formation and embolization of cardiac vegetations. The kidney lesions observed upon examination ranged from ischemic or hemorrhagic to necrotic. In this context, VEGF-A deficits and

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**Fig. 7. SEC inhibits VEGF-A expression in immortalized human aortic endothelial cells.** Expression analysis of 24 angiogenesis-related genes by RT-qPCR in iHAECs treated with LPS (25 ng ml⁻¹), VEGF-A (20 nM), VEGF-A receptor inhibitor (axitinib, 25 nM), or SECN23A (20 μg ml⁻¹) for 2, 4, and 6 hours. (A) Gene products that play significant roles in inflammatory responses. (B) Gene products that contribute to various angiogenesis processes. * denotes genes with twofold increases or decreases compared to untreated controls.
angiogenesis defects, in general, can lead to persistent vascular injury and insufficient revascularization, overall impairing the wound healing process, allowing renal deterioration and bacterial infection. The liver has central roles in clearing circulating bacteria and their toxins and is key in initiating or amplifying inflammatory responses during systemic infections (7). Yet, literature on the effects of SAIE on liver injury is scarce (3). In our study, increases in serum aminotransferases correlate with increases in IL-6 and LDH in rabbits infected with S. aureus SECC. Therefore, in contrast to the kidneys, in the liver, superantigenicity significantly contributes to cellular injury. It is intriguing that liver lesions from rabbits infected with S. aureus SECKO exhibited high prevalence of infarction and necrosis, when heart and kidney lesions were mostly devoid of embolic complications. It is almost as if the liver was the primary site of infection.

Our current understanding of SAIE involvement in the pathogenesis of many life-threatening S. aureus infections such as bacteraemia, pneumonia, and IE has been tied to complications arising from the extreme effects of SAAg activity. While adaptive immune system activation is the hallmark of staphylococcal SAgs, we provide evidence that this is not their only biological function involved in promoting life-threatening disease pathologies. It is clear that SAgs do much more than previously anticipated or expected. Therefore, it becomes fundamental to further understand the involvement of superantigenicity-independent mechanisms in other invasive and life-threatening diseases as well. Recently, the SAg SEI-X was found to inhibit neutrophil function via a sialic acid-binding motif uniquely present in this SAg (40). Meanwhile, TSST1 was reported to induce activation of epithelial cells through a dodecapeptide close to the base of the central α helix of the molecule (39). The dodecapeptide sequence is found in all staphylococcal SAgs, yet its effects on non-hematopoietic cells are poorly characterized (12, 39, 61). The relevance of the SEC dodecapeptide in endothelial cell function and S. aureus diseases such as IE is not known, but it is at the center of current studies.

Together, our study demonstrates that superantigenicity independent effects of SEC are essential to the pathogenesis and pathophysiology of SAIE. The data further show that SEC is an antiangiogenic toxin that inhibits sprouting of new capillaries from the activated endothelium, which would effectively impair revascularization of injured tissues. It is becoming evident that SAgs like TSST1 and SEC promote IE by interfering with vascular repair and the angiogenic process. One of the mechanisms by which SEC may exert its antiangiogenic effects is through the inhibition of endothelial cell expression of one of the key inducers of angiogenesis, VEGF-A. If so, deficient angiogenesis may be at the center of many pathologies associated with SAg-producing S. aureus infections. Given the reach of the vascular endothelium into every organ system and the prevalence of SAgs among both methicillin-susceptible and methicillin-resistant S. aureus strains, SAgs have the potential to promote pathologies in every organ that they encounter.

**MATERIALS AND METHODS**

**Bacterial strains and growth conditions**

Community-associated, methicillin-resistant USA400 strain MW2 was originally obtained in the Upper Midwest from a young patient who succumbed to necrotizing pneumonia. S. aureus MW2 encodes for the SAgs SEC, SEA, SEH, SEK, SEL, SEQ, and SEI-X. SEC is produced at 60 to 100 μg/ml in liquid culture, while the rest are produced at 0.0001 to 0.03 μg/ml. Staphylococcal strains were used from low-passage number stocks. All staphylococcal strains were grown in Bacto Todd Hewitt (TH) (Becton Dickinson) broth at 37°C with aeration (225 rpm) unless otherwise stated. Strains and plasmids used in this study are listed in Table S1. Plasmids used for complementation were maintained using carbenicillin (100 μg/ml) in Escherichia coli DH5α. For endocarditis experiments, strains were grown overnight, diluted, and washed in phosphate-buffered saline [PBS; 2 mM NaH2PO4, 5.7 mM Na2HPO4, and 0.1 M NaCl (pH 7.4)].

**Selection of mutations inactivating the SEC TCR-binding site (N23A) or the MHC-II-binding site (F44A/L45A)**

Asn23 is a highly conserved, surface-exposed residue located in a cleft between the O/B fold and β-grasp domain of SAgs (62). It forms hydrogen bonds with the backbone atoms of complementarity-determining region (CDR) 2 of VB-TCR (63). Hence, Asn23 contact with VB-TCR has one of the greatest energetic contributions of the complex. Mutations in this position, such as N23A or N23S, greatly destabilize the VB-TCR-SAgs interaction with profound effects in SAg activity (63). SECN23A has no detectable binding to VB-TCR as measured by surface plasmon resonance, no proliferative T cell responses in thymidine incorporation assays at concentrations up to 30 μg/ml (63), no lethality or signs of TSS in rabbits vaccinated subcutaneously with 25 μg three times every 2 weeks (16), and no lethality in rabbits after intravenous injection at 3000 μg/kg (15). Phe44 and Leu45, conserved among all enterotoxins, are located on a protruding hydrophobic loop that directly contacts MHC-II and forms strong electrostatic interactions with the α chain (20, 64). Leu45 is the most extensively buried amino acid residue in the SEC:MHC-II interface, but mutations in either residue (Phe44 or Leu45) effectively inactivate SEC binding (18–21, 64). F44A alone is 1000-fold less efficient in MHC-II binding, resulting in a concomitant reduction of IL-2 in T cell proliferation assays (19, 65).

**Construction of chromosomally complemented toxoid strains**

SEC is a CDC designated select agent. Hence, we are not allowed to use the wild-type copy of the gene in recombinant studies. For this reason, all PCR products generated in the making of the toxoid complement strains either included the permissible N23A mutation or were only a partial amplification of sec missing either the TCR or MHC-II domain. Each step of plasmid construction was verified by Sanger sequencing to contain the N23A TCR mutation. PCR amplification was performed using Phusion polymerase [New England Biolabs (NEB)] unless otherwise noted. S. aureus expressing SECN23A or SECN23AF44A/L45A were made by markerless chromosomal complementation in MW2sec with the genes expressed under the control of the native promoter and terminator (66). The SECN23A gene sequence was created by amplifying two fragments from MW2 with primer sets pJB38xN23AпромF/promN23R and termN23F/pJB38xN23AtermR. The chromosomal complementation plasmid, pJB38-NWMN29-30 [research resource identified (RRID): Addgene_84457], was digested with Eco RV, and PCR products were inserted by Gibson Assembly (NEB) as previously described, creating pKK29 (67). SECN23A was amplified from pKK29 with the primer set pUC19SECN23AaptF/pUC19SECN23AaptR and inserted by Gibson Assembly into pUC19 (RRID: Addgene_50005), linearized with Kpn I and Eco RI, to create pKK33. The MHC-II–binding site mutations were introduced into pKK33 by site-directed mutagenesis (QuikChange II,
Agilent Technologies) using the primer set SECF44A/L45Afor/SECF44A/L45Arev, creating pKk39. The SEC_{N23A/F44A/L45A} gene sequence was amplified from pKk39 with primer set pJB38xN23ApromF/pJB38xN23AtermR and inserted into pJB38-NWMN29-30 as described above, creating pKk42. pKk29 and pKk42 were electroporated into S. aureus RN4220 and moved into MW2Asec by generalized transduction with φ11 (68). S. aureus strains containing plasmids were selected with chloramphenicol (20 μg/ml) at 30°C. Allelic exchange was performed as previously described (66), and chromosomal insertions were detected by PCR with primer set XNWMN2930F/XNWMN2930R and verified by Sanger sequencing. Primers were purchased from Integrated DNA Technologies (table S2).

**SEC purification and T cell proliferation assay**

SEC was purified from S. aureus strain FR1913 in its native form by ethanol precipitation and thin-layer isoelectric focusing as previously described (69). Preparation of SEC resulted in a single band by Coomassie blue stain. Toxin preparations were tested for LPS contamination with the Toxin Sensor Chromogenic LAL Endotoxin Assay following the manufacturer’s instructions (GenScript). SEC preparations had <0.02 ng of LPS per 20 μg of SEC*. For analysis of SAg activity in S. aureus expressing SEC_{N23A}, SEC_{N23A/F44A/L45A}, SEC, or SEC^KO, cell-free supernates were collected from bacteria grown overnight, centrifuged, and filtered (0.22-μm pore size, MilliporeSigma). Mononuclear cells were isolated from human peripheral blood samples (PBMCs) originating from three donors by density gradient centrifugation using Ficoll–Paque PLUS (UW-Madison Minimal Risk IRB Protocol 2018-0304-CR004). PBMCs were labeled with 5 μM CellTrace Violet (CTV) dye (Thermo Fisher Scientific) following the manufacturer’s protocol. CTV-labeled cells were plated in 96-well plates at a concentration of 2 × 10^4 cells per well in RPMI 1640 medium supplemented with 10% heat-inactivated fetal bovine serum (FBS), 5% heat-inactivated bovine calf serum, 3% human AB serum, 1% L-glutamine, and 1% penicillin-streptomycin. Purified SEC (10 μg) or cell-free supernates at 0.3-, 2-, or 10-μl volumes were added to the wells. After 6 days, cell division of CD4+ T cells was assessed flow cytometrically by measuring dilution of CTV fluorescence intensity. Proliferation response was quantified using the FlowJ proliferation platform analysis tool, and data were presented as replication index or percent of CD4+ T cell divided.

**Rabbit model of SAIE**

The rabbit model of IE was performed as previously described with some modifications (10). New Zealand White rabbits (2 to 3 kg) were obtained from Bakkom Rabbity (Red Wing, MN) and anesthetized with ketamine (dose range, 10 to 50 mg/kg) and xylazine (dose range, 2.5 to 10 mg/kg). Mechanical damage to the aortic valve was done by introducing a hard plastic catheter via the left carotid artery, left to pulse against the valve for 2 hours, and removed, and the incision was closed. Rabbits were inoculated via the left carotid artery with 2 × 10^7 to 4 × 10^7 total CFU in PBS and monitored four times daily for a period of 4 days. For pain management, rabbits received buprenorphine (dose range, 0.01 to 0.05 mg/kg) twice daily. At the conclusion of each experiment, bacterial counts were obtained from heparinized blood (50 USP units/ml). Rabbits were euthanized with Euthasol (Virbac), and necropsies were performed to assess overall health. Spleens were weighed and used as an infection control, kidney and liver gross pathology was graded using gross lesion pathology scale (table S3), aortic valves were exposed to assess vegetation growth, and vegetations that formed were excised, weighed, and suspended in PBS for CFU counts. A minimum of four rabbit hearts from each infection group were placed in 10% neutral-buffered formalin and further processed by the Comparative Pathology Laboratory at the University of Iowa for histopathological analyses. Vegetation weight and bacterial counts cannot be obtained from hearts prepared for histology. Experiments were performed according to established guidelines and the protocol approved by the University of Iowa Institutional Animal Care and Use Committee (protocol 6121907). Numbers of rabbits per infection group were as follows: S. aureus SEC^+ (n = 12), S. aureus SEC^KO (n = 15), S. aureus SEC_{N23A} (n = 17), and S. aureus SEC_{N23A/F44A/L45A} (n = 17). Data from rabbit experimental groups are a result of at least three independent experiments.

**Serum analysis**

Rabbit serum was obtained from heparinized blood (50 USP units/ml) collected before infection and at 48, 72, and 96 hours after infection. Blood was centrifuged at room temperature at 5000g for 10 min. The collected supernatant was centrifuged for an additional 5 min, filter-sterilized using a 0.2-μm filter, and stored at −80°C for further analysis. Serum samples were sent to the University of Iowa Diagnostic Laboratories and evaluated for the following serum analytes: AST (U/liter), ALT (U/liter), BUN (mg/dl), creatinine (mg/dl), and LDH (mg/dl). IL-6 was quantified from serum samples using the R&D Systems DuoSet Rabbit IL-6 ELISA Kit according to the manufacturer’s instructions. Serum samples were diluted 1:10 in reagent diluent before use. The optical density (OD) was determined using a TECAN M200 plate reader (Tecan Group Ltd.) set to 450 nm with wavelength correction set to 540 nm. A standard curve was created by linear regression analysis of the IL-6 concentration versus OD, log-transformed (GraphPad Prism 9.3.1).

**Histopathology scoring**

Fixed tissues were routinely processed, cut at 5 μm, and hematoxylin and eosin (H&E)–stained or Gram-stained. The slides were scored in a semi-blinded fashion, with the pathologist only made aware of control animals. Histopathologic scoring was developed specifically for this animal model, as outlined by Gibson-Corley et al. (70) and explained in table S3. Rabbits infected with S. aureus SEC^KO with no vegetations did not exhibit pathology at the end of experimentation (n = 3). Hence, all hearts of S. aureus SEC^KO–infected rabbits with visible vegetations were processed (mean size, 2.7 ± 1 mm²; n = 4). Heart, kidney, and liver sections from rabbits infected with S. aureus SEC^+, S. aureus SEC_{N23A}, and S. aureus SEC_{N23A/F44A/L45A} were selected randomly based on the presence of vegetations (mean size, 6.6 ± 2 mm²; n = 15) and gross pathology lesions in the kidney and liver.

**SEC_{N23A} expression and purification**

SEC_{N23A} (C terminus 6×His-tagged) was expressed in E. coli BL21 (DE3) Star strain harboring pET25bHSVDelTEV. Following ethanol precipitation, toxins were purified through Np^2×-affinity chromatography using a HiTrap Heparin HP column (GE Healthcare). Protein capture was verified via visualization of a single band on SDS–polyacrylamide gel electrophoresis following Coomassie blue staining. Detoxi-Gel endotoxin removing resin (Thermo Fisher Scientific) was used to reduce LPS contamination to <0.001 ng of LPS.
per 30 μg of toxin. Endotoxin levels were quantified using the ToxinSensor Chromogenic LAL Endotoxin Kit (GenScript) to ensure toxin purity.

**Ex vivo rabbit aortic ring model of angiogenesis**

Mixed-sex New Zealand White rabbits, 2 to 3 kg, were purchased from Charles River Laboratories (Massachusetts) and maintained at Charmany Instructional Facility at the School of Veterinary Medicine of the University of Wisconsin-Madison according to established guidelines and the protocol approved by the Institutional Animal Care and Use Committee (protocol V006222-A02). Aortic ring explants were cultured by modifying the thin-layer method (71). Briefly, the thoracic and abdominal aortas were excised immediately after euthanasia, cleaned to remove excess fascia and connective tissue, dissected into 1- to 1.5-mm cross-sections, and embedded into GFR-Matrigel-coated wells [200 μl of phenol red–free Corning Matrigel Growth Factor Reduced (GFR) Basement Membrane Matrix] in a 48-well plate format. After 10 min of polymerization at 37°C, 5% CO₂, 200 μl of supplemented Medium 200 was added and plates were incubated at 37°C, 5% CO₂ for up to 14 days. Medium 200 contained low-serum growth supplement (LSGS) (see the next section), penicillin-streptomycin (100 U ml⁻¹), amphotericin B (2.5 μg ml⁻¹), and designated treatment. Medium (± treatments) was changed every 3 to 5 days. Rings were treated with SEC/N23A at 2.5, 5, 10, and 20 μg and axitinib at 10 μM. Rings treated initially with 20 μg were switched to 10 μg following the first medium change to maintain toxin pressure while limiting potential cytotoxic effects. Equal numbers of thoracic and abdominal aortic rings were used for each treatment condition as follows: medium control (25 ng ml⁻¹), VEGF-A (20 nM), axitinib (25 nM), or SEC (20 μg ml⁻¹) and designated treatment. Medium (± treatments) was changed every 3 to 5 days. Rings were treated with SEC/N23A at 2.5, 5, 10, and 20 μg and axitinib at 10 μM. Rings treated initially with 20 μg were switched to 10 μg following the first medium change to maintain toxin pressure while limiting potential cytotoxic effects. Equal numbers of thoracic and abdominal aortas were used for each treatment condition as follows: medium control (n = 24), axitinib (n = 8), SEC/N23A 2.5 μg (n = 6), SEC/N23A 5 μg (n = 9), SEC/N23A 10 μg (n = 9), and SEC/N23A 10 μg (n = 8). All treatment conditions were assessed from three separate rabbits with the exception of SEC/N23A at 2.5 μg, which was completed in two rabbits. Phase-contrast images were captured on days 0, 7, and 14 with Leica DMi8 equipped with a Tokai Hit stage-top incubator set to 37°C, 5% CO₂ using an HC PL FLUOTAR 4×/0.13 objective lens. Sprouting was assessed using ImageJ (72).

**Human aortic endothelial cell culture and RNA extraction**

hiAECs are a recently established cell line shown to retain phenotypic and functional characteristics of primary cells (38). iHAECs were cultured at 37°C, 5% CO₂ in phenol red–free, endothelial cell basal medium (Medium 200) supplemented with LSGS (final concentrations of the following: FBS, 2%; hydrocortisone, 1 μg ml⁻¹; human epidermal growth factor, 10 ng ml⁻¹; basic fibroblast growth factor, 3 ng ml⁻¹; heparin, 10 μg ml⁻¹, both from Gibco Life Technologies), as previously described (38). LSGS contains the final concentrations of the following: FBS, 2%; hydrocortisone, 1 μg ml⁻¹; human epidermal growth factor, 10 ng ml⁻¹; basic fibroblast growth factor, 3 ng ml⁻¹; heparin, 10 μg ml⁻¹. All experiments were conducted with iHAECs at passages 4 to 10 from a single clone seeded at 5 × 10⁴ per well in a six-well plate. iHAECs were treated with LPS (25 ng ml⁻¹), VEGF-A (20 nM), axitinib (25 nM), or SEC (20 μg ml⁻¹) for 2, 4, and 6 hours. Data are the result of three biological replicates. RNA was extracted at the indicated time points using the miRNeasy Mini Kit (Qiagen), according to the manufacturer’s instructions, and stored at −80°C. Contaminating genomic DNA (gDNA) was removed using the Turbo DNA-Free Kit (Invitrogen). RNA quantity and purity were assessed using NanoDrop 1000 (Thermo Fisher Scientific) with median A260/280 values of 1.93 and A260/230 of 1.55.

**cDNA synthesis and real-time RT-PCR**

Complementary DNA (cDNA) synthesis was carried out using DNA-free RNA with the RT First Strand Kit according to the manufacturer’s instructions (Qiagen). Each cDNA sample was used at 120 ng per well in 96-well plates. qPCR was conducted using the PowerUP SYBR Green Master Mix (Thermo Fisher Scientific, USA), with each reaction having 5 μl of PowerUP SYBR Green Master Mix, 2 μl of 1 μM forward and reverse gene-specific primers (table S4), and 3 μl of diluted cDNA experimental sample in a final reaction volume of 10 μl. Reactions were run with the following cycle profile: initial denaturation, 95°C 10 min; 40-cycle amplification/elongation, 95°C 15 s followed by 60°C 1 min. gDNA contamination from each sample was assessed using a total of 120 ng of RNA (no reverse transcriptase) with reaction conditions as listed above, with all samples having Ct values of >35. No-template controls were run with each qPCR plate and had Ct values of >38. All qPCRs were run and analyzed on the QuantStudio 3 Real-Time PCR System with analysis software v1.5.1 (Applied Biosystems). Each time point was collected in biological triplicates, with qPCRs from each experimental sample done in technical triplicates.

**Statistical analyses**

The log-rank Mantel-Cox test was used for statistical significance of survival curves. Normality was assessed using the D’Agostino and Pearson test along with associated Q-Q plots for data distribution. For comparison across means, log-transformed data were used and statistical significance was determined by using one-way analysis of variance (ANOVA) with Holm-Šídák’s multiple comparisons test for the following data sets: vegetation size, vegetation CFU, blood CFU, spleen size, BUN, creatinine, AST, ALT, LDH, and IL-6. Statistical significance for gross pathology data was determined using Fisher’s exact test along with calculated odds ratios and 95% confidence intervals. Statistical significance for sprouting area in the aortic ring model was determined by two-way ANOVA with Holm-Šídák’s multiple comparisons test. Statistical significance of virulence factor production was determined using the nonparametric Kruskal-Wallis test. α = 0.05 (GraphPad Prism 9.3.1).

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at https://science.org/doi/10.1126/sciadv.abo1072

View/request a protocol for this paper from Bio-protocol.

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Acknowledgments: We thank the University of Iowa Comparative Pathology Laboratory and University of Iowa Diagnostic Laboratories for histology and serum analysis services. *Funding:* This work was supported by NIH grants R01AI34692-01 (to W.S.-P.), R01AI136500 (to J.E.G.), ST32AI007511-23 (to P.M.T.), and T32GM008365 (to K.J.K.). *Author contributions:* K.J.K., S.S.T., X.-J.W. carried out qPCR analyses. K.N.G.-C. provided intellectual and technical support on histopathological analysis and gross pathological grading. S.S.T. carried out ex vivo experiments. K.J.K., P.M.T., A.N.F., K.K., and W.S.-P. carried out in vivo rabbit experiments. K.N.G.-C. provided intellectual and technical support on histopathological analysis and gross pathological grading. S.S.T. carried out ex vivo experiments. X.-J.W. carried out qPCR analyses. N.S.B. carried out T cell proliferation assays and analyzed the data. All authors reviewed the manuscript. *Competing interests:* K.J.K. is currently an employee of Integrated DNA Technologies, which sells reagents used or like those used here. *Data and materials availability:* All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials.

Submitted 9 February 2022
Accepted 24 March 2022
Published 11 May 2022
10.1126/sciadv.abo1072