Necrotizing fasciitis of the abdominal wall caused by *Serratia marcescens*

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**Abstract**

In this article, we present the first case of necrotizing fasciitis affecting the abdominal wall caused by *Serratia marcescens* and share results of a focused review of *S. marcescens* induced necrotizing fasciitis. Our patient underwent aorto-femoral bypass grafting for advanced peripheral vascular disease and presented 3 weeks postoperatively with pain, erythema and discharge from the incision site in the left lower abdominal wall and underwent multiple debridements of the affected area. Pathology of debrided tissue indicated extensive necrosis involving the adipose tissue, fascia and skeletal muscle. Wound cultures were positive for *Serratia marcescens*. She was successfully treated with antibiotics and multiple surgical debridements. Since necrotizing fasciitis is a medical and surgical emergency, it is critical to examine infectivity trends, clinical characteristics in its causative spectrum. Using PubMed we found 17 published cases of necrotizing fasciitis caused by *Serratia marcescens*, and then analyzed patterns among those cases. *Serratia marcescens* is prominent in the community and hospital settings, and information on infection presentations, risk factors, characteristics, treatment, course, and complications as provided through this study can help identify cases earlier and mitigate poor outcomes. Patients with positive blood cultures and those patients where surgical intervention was not provided or delayed had a higher mortality. Surgical intervention is a definitive way to establish the diagnosis of necrotizing infection and differentiate it from other entities.

**Introduction**

Necrotizing fasciitis, also known as flesh-eating disease, is a rare infection of the deeper layers of skin and subcutaneous tissues, and easily spreads across the fascial plane within those tissues. As bacterial toxins and the immune response cause vasoconstriction of the vasculature, the fascial spaces become avascular resulting in necrosis, which also prevents penetration of antibiotics into the tissues.1 Common causes are Group A streptococcus (GAS) (*Streptococcus pyogenes*), *Staphylococcus aureus*, *Vibrio vulnificus*, *Clostridium perfringens*, and *Bacteroides fragilis*. Mortality ranges from 4.2 to 38% with improving prognosis as time to treatment decreases.2 *Serratia marcescens*, a motile bacillus, gram-negative, facultative anaerobe, is an opportunistic pathogen of increasing importance. It is part of normal colon flora, and is also found on skin, sewage, and water. This Enterobacteriaceae organism also typically colonizes the respiratory and urinary tracts and causes infections in those organ systems.3 *S. marcescens* is often an opportunistic infection and may cause osteomyelitis, septic arthritis, endocarditis, and, rarely cellulitis or necrotizing fasciitis.4 Soft tissue infections due to gram-negative organisms are relatively uncommon, and typical predisposing factors include: a history of trauma, alcoholism, peripheral vascular disease, systemic lupus erythematosus, immunosuppression, diabetes mellitus, urinary tract infection (UTI), bactereemia, pneumonia, infective arthritis, burns, and renal failure.5,6 Other predisposing factors include: antibiotic use (most often first generation cephalosporins), steroid use, surgical instrumentation, urinary catheters, respiratory equipment, intravenous lines, injections, lacerations, abscesses, or ulcers.7 Common presentations of necrotizing fasciitis within 48 hours of infection include skin erythema and swelling at the affected site (97.6%), pyrexia (61.9%), hypotension (33.3%), altered consciousness (28.6%), bulbar lesions (26.2%), and crepitus (9.5%). Hypotension, altered consciousness, ventilator support, ALT > two-fold of normal, serum creatinine >177 µmol/L, thrombocytopenia (<100×109/L), and worsening symptoms within in 48 hours of admission have been associated with higher fatality rates.2 Lack of response to narrow-spectrum antibiotics, bullae formation, or a rapidly worsening clinical course, should heighten the suspicion for uncommon organisms like *S. marcescens*.

Reports of *S. marcescens* necrotizing fasciitis cases have increased in the literature. In this study, we identify characteristics, trends, and risk factors of those infections to better prepare the medical community and prevent poor outcomes.

**Materials and Methods**

PubMed was used to search for cases of necrotizing fasciitis caused by *Serratia marcescens* published in the English language literature between 1966 and 2013. Keywords that were used included: *Serratia marcescens*, necrotizing, and fasciitis. We identified eleven additional cases of necrotizing fasciitis due to *S. marcescens* in the literature since the latest review in 2001, which are included in Table 1.4,7-14,22 Cases were categorized as being healthcare-associated infections or community-acquired infections based on the Centers for Disease Control and Prevention (CDC) definition of healthcare-associated infections (HAIs) as infections that patients acquire during the course of receiving healthcare treatment for other conditions.22

**Case Report**

We recently identified a case involving a 51-year-old African-American woman who pre-
Table 1. *Serratia marcescens* necrotizing fasciitis cases, 1966 to present.

| Author                  | Year | Age | Sex | Risk factors                        | Precipitating factor                  | Site of infection | Type | *S. marcescens* cultures | Treatment                  | Outcome |
|-------------------------|------|-----|-----|-------------------------------------|---------------------------------------|-------------------|------|--------------------------|----------------------------|---------|
| Rimalho et al.          | 1987 | 74  | M   | Immuno-compromised                  | Dilofenac consumption                 | Leg               | CA   | Bacteremia               | None                                      | Died     |
| Bornstein et al.        | 1992 | 37  | F   | Renal failure on hemodialysis       | Pain during dialysis                  | Axilla and chest wall | HA   | Wound, bullae, blood    | Antibiotics and SD               | Recovered |
| Zipper et al.           | 1996 | 55  | F   | Diabetes                            | Left below-knee amputation             | Leg               | CA   | Wound                   | Antibiotics                | Recovered |
| Huang et al.            | 1999 | 73  | M   | Nephrotic syndrome                  | Steroid therapy                      | Lower leg          | HA   | Necrotic tissue, blood  | Antibiotics and SD               | Recovered |
| Huang et al.            | 1999 | 40  | M   | Uremia, peritoneal dialysis, SLE    | Pneumonia with cultures for *S. marcescens*, steroid and nabumetone | Left calf and thigh | CA   | Necrotic tissue, blood  | Antibiotics and SD               | Recovered |
| Liangpunsakul et al.    | 2001 | 66  | F   | Healthy                             | None                                  | Leg               | CA   | Blood                   | Antibiotics                | Died     |
| Newton et al.           | 2002 | 2   | F   | Healthy                             | Pharyngitis                           | Cervical spine     | CA   | Wound, blood            | Antibiotics and SD               | Died     |
| Bachmeyer et al.        | 2004 | 49  | M   | Small cell lung cancer, DM          | Chemotherapy and cellulitis           | Right leg          | HA   | Tissue, bullae, blood  | Antibiotics                | Recovered |
| Curtis et al.           | 2005 | 51  | M   | ESRD, T2DM, CHF                     | Scrapped knee on rock in river        | Left leg           | CA   | Wound                   | Antibiotics and SD               | Died     |
| Statham et al.          | 2006 | 6   | M   | Immunocompetent                     | Suspected pharyngitis                 | Oro-pharynx        | CA   | Wound                   | Antibiotics and SD               | Recovered |
| Motisiti et al.         | 2011 | 37  | M   | Healthy                             | Human bite                           | Forearm            | CA   | Wound                   | SD                                      | Died     |
| Naino-Galvan et al.     | 2012 | 57  | F   | CML, immuno-compromized             | Minor trauma                          | Right thigh        | HA   | Blister, blood          | Antibiotics                | Died     |
| Prelog et al.           | 2012 | 15  | F   | Acute lymphocytic leukemia          | Venous access port implantation       | Left axilla, venous HA access port site | Wound | Antibiotics and SD    | Recovered |
| Wen et al.              | 2012 | 40  | F   | Nephrotic syndrome, cyclosporine    | Chemotherapy 10 days prior            | Left leg           | CA   | Wound                   | Antibiotics                | Died     |
| Rehman et al.           | 2012 | 54  | F   | SLE, end-stage renal disease        | Central venous catheter, AV fistula ligation, steroid therapy | Chest wall         | HA   | Wound                   | Antibiotics and SD               | Died     |
| Present case            | 2012 | 51  | F   | DM, PVD                             | Bifemoral bypass and left distal femoral aneurysm repair | Lower abdomen      | CA   | Wound, blood            | Antibiotics and SD               | Recovered |
| Cope et al.             | 2013 | 97  | F   | Heat failure, CKD                   | Heart failure exacerbation             | Right leg          | HA   | Wound (post-mortem)    | Antibiotics                | Died     |

SD, Surgical debridement; CA, community-acquired infection; HA, healthcare-associated infection; SLE, systemic lupus erythematosus; ESRD, end-stage renal disease; T2DM, Type II, diabetes mellitus; PVD, peripheral vascular disease.
and 7 cases (41%) were healthcare-associated infections. An overwhelming number of cases (81.3%) had pre-existing open wounds. Seven cases (41%) were immunocompromized, 5 cases (29%) had kidney disease, and 4 cases (24%) had diabetes. A majority (59%) of the cases were among females. Three of the cases (18%) were children.

Nine out of seventeen cases (53%) died as a result of necrotizing fasciitis and its complications. Six of the cases (67%) that died had community-acquired infections. Individuals with positive versus negative blood cultures were more likely to die (88% vs. 75%). Patients who did not receive surgical debridement had inferior outcomes; they had a mortality of 71% compared to 40% among those who received surgical intervention as opposed to debridement. Additionally, all patients (18%, n=3) who received only antibiotics and the one patient who received only surgical debridement died. All previously healthy patients (18%, n=3) also died. A majority of those that recovered had a surgical procedure in the hospital prior to symptom onset such as below the knee amputation or venous access port implantation (75%, n=8).

Discussion and Conclusions

Necrotizing fasciitis is a deep infection of the subcutaneous tissue that results in progressive destruction of fascia and fat. The disease is classified as type I (polymicrobial infection), type II (monomicrobial) and type III gas gangrene, or clostridial myonecrosis. Type I infection involves anaerobic species in combination with one or more facultative anaerobic streptococci (other than group A) and members of the Enterobacteriaceae family. Type II infection is commonly caused by group A streptococci or other beta-hemolytic streptococci that are isolated alone or in combination with other species, most frequently S. aureus. These infections are also commonly referred as flesh eating infection. Among those with necrotizing fasciitis, the affected area is typically erythematous, swollen, warm, and exquisitely tender. The infection progresses rapidly over several days, with changes in skin color from red-purple to patches of blue-gray. Skin breakdown with bullae (containing thick pink or purple fluid) and frank cutaneous gangrene may be observed within three to five days.16 The development of anesthesia may precede the appearance of skin necrosis and provide a clue that the process is necrotizing fasciitis rather than cellulitis. In advanced infection, high fever and systemic toxicity are generally observed.

Individuals with positive blood cultures typically have poorer prognosis and higher mortality rates.125 Treatment of necrotizing infection consists of early and aggressive surgical exploration and debridement of necrotic tissue, together with broad spectrum empiric antibiotic therapy and hemodynamic support as necessary.126 Acceptable antibiotic regimens prior to identification of the causative organism(s) include administration of a carbapenem or beta-lactam/beta-lactamase inhibitor, together with clindamycin (600 to 900 milligrams intravenously every eight hours), as well as an agent with activity against MRSA. Definitive antibiotic treatment should be tailored according to blood and or tissue gram stain, culture, and sensitivity results when available.27 If S. marcescens is identified as the main pathogen, third-generation cephalosporins, fluoroquinolones, and imipenem/cilastatin are the antimicrobials of choice. Our patient’s fever on presentation was likely masked by the pain medications she was taking, though she did give a history of having fever at home. She most likely had a subacute course and thus lacked the classic features of rapid clinical progression and fulminant toxicity. An HIV test was not performed, as the patient did not have any risk factors or any previous sexually transmitted disease. Our review noted a mortality rate of 53% among those with S. marcescens-induced necrotizing fasciitis, which is higher than the mortality rate reported by Elliott et al.,26 in 1996 and nearly equivalent (53% vs. 50%) to the mortality rate documented by Nancy et al. in 2002.25 Patients who died were more likely to have positive blood cultures, present with a community-acquired illness, be previously healthy, experience minor trauma or sickness prior to symptom onset, and not receive both surgical debridement and antibiotic therapy. Similar observations of increased mortality among those with positive blood cultures or those not receiving appropriate antibiotic and surgical treatment were also found in other studies.126 Increased mortality among individuals with less severe injury, lack of traditional risk factors, and those who did not have a recent surgical procedure in the hospital may have been due to less vigilance about changes in health status and the fact that S. marcescens is common in the outdoor environment; these individuals were also more likely to have community-acquired infections (70% vs. 30%). Additionally, individuals who had been hospitalized recently were more likely to have received antibiotics in the near past and also more likely to be wary of any new symptoms. Healthy patients also tend to seek care later and receive less aggressive treatment; only one of the previously healthy patients received both antibiotics and surgical debridement.

Clinicians should be wary of necrotizing fasciitis among those with soft tissue infections presenting after environmental open skin exposure. Gram-negative bacilli, including S. marcescens, should especially be considered in cases of necrotizing fasciitis or cellulitis among immunocompromized persons with renal failure, steroid use, recent surgery or diabetes.25 Early diagnosis is key, but unfortunately is missed in 85 to 100% of cases since necrotizing fasciitis is often confused with cellulitis, myositis, or deep-seated abscess(es). A high index of suspicion is important in view of the paucity of specific cutaneous findings early in the course of the disease. Radiographic imaging studies may be useful in determining if muscle tissue is involved but should not delay surgical intervention. Our case also underscores the importance of preoperative and more so, postoperative wound care education, which could have potentially prevented this fatal complication.26 The most effective treatment regimen for suspected necrotizing fasciitis includes: early and aggressive surgical exploration and debridement, immediate broad spectrum empiric antibiotic therapy, and hemodynamic support as necessary.126

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