Treatment of Methicillin-Resistant *Staphylococcus aureus* (MRSA) Pneumonia with Ceftaroline Fosamil in a Patient with Inhalational Thermal Injury

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ABSTRACT

A 48-year-old female, who was found unresponsive and suffered inhalation injury secondary to a house fire, was transferred to our burn center for definitive treatment. Post tracheostomy, the patient became febrile and tachycardic. On hospital day (HD) 5, the patient expressed thick yellow secretions during suctioning and diffuse rhonchi was noted on physical exam. Blood cultures and a culture from the broncheo-alvelolar lavage grew Gram-positive cocci in clusters and the patient was started on empiric vancomycin. Despite aggressive vancomycin dosing (1750 mg intravenously every 6 h), the patient’s status continued to deteriorate. The organism was identified as methicillin-resistant *Staphylococcus aureus* (MRSA) with a vancomycin minimum inhibitory concentration (MIC) of 2 mg/L. Based on the potential for drug–drug interactions with linezolid, the patient was started on ceftaroline fosamil (MIC = 0.5 mg/L) 600 mg intravenously every 8 h with a prolonged 2-h infusion to anticipate suboptimal concentrations secondary to thermal burn injury. Post change in antibiotic therapy, a rapid clinical improvement was observed with the patient becoming afebrile at 48 h after initiation of ceftaroline. The patient completed a total of 14 days of ceftaroline therapy and was subsequently weaned from the ventilator on HD 22 and decannulated 2 days later. To our knowledge, this is the first report of the use of ceftaroline for the treatment of MRSA pneumonia in a patient with thermal injury.
Keywords: Burn; Ceftaroline; Inhalational; Pharmacokinetics; Pneumonia; Methicillin-resistant Staphylococcus aureus (MRSA); Staphylococci

INTRODUCTION

Infections caused by methicillin-resistant Staphylococcus aureus (MRSA) are a frequent occurrence in most medical centers, especially in patients admitted to the intensive care unit (ICU) for prolonged periods of time. These infections can be secondary to implanted catheters, skin and soft tissue infections, and pneumonia. Patients with extensive thermal injuries are at high risk of developing infections from MRSA and other multi-drug-resistant pathogens due to their need for prolonged ICU care, frequent receipt of antimicrobial therapy, and the immunological and physiological disturbances encountered as a result of their injuries. Historically vancomycin, a glycopeptide antimicrobial, has been the treatment of choice for MRSA infections. However, the clinical utility of vancomycin has recently been questioned when employed in the treatment of MRSA infections when the mean inhibitory concentration (MIC) of vancomycin is greater than 1 mg/L, with published literature reporting increased treatment failure and mortality in patients due to inability to attain currently established pharmacokinetic (PK) and pharmacodynamic (PD) targets [1–3].

Ceftaroline fosamil (Teflaro®; Actavis Pharmaceuticals), an advanced cephalosporin with activity against MRSA, was approved by the Food and Drug Administration (FDA) for the treatment of acute bacterial skin and skin structure infections and community-acquired bacterial pneumonia (CABP). We describe a case of MRSA pneumonia in a patient with inhalational thermal injuries who did not respond to initial vancomycin therapy (vancomycin MIC = 2 mg/L) but responded clinically to treatment with ceftaroline. Currently, there are limited data available to support the utilization of ceftaroline for the treatment of pneumonia caused by MRSA. There are no data that we are aware of with respect to treatment of patients with burn-related infections. This article does not contain any new studies with human or animal subjects performed by any of the authors.

CASE REPORT

A 48-year-old Caucasian female was found unresponsive at the scene of a house fire with burns to the face [<1% total body surface area (BSA)] and inhalation injury. She was intubated via endotracheal tube at an outside facility and subsequently transferred to our burn center for definitive care. The initial bronchoscopy showed diffuse carbonaceous sputum with pale friable mucosa extending throughout the airways distally. The patient was intubated for airway protection, which required adequate sedation and neuromuscular blockade. Due to the inability of the patient to provide a detailed medical history, the patient’s sister and psychiatrist supplied the relevant information, which included a diagnosis of severe depression with psychotic features. On admission, the patient’s list of home medications included paliperidone palmitate 234 mg once monthly as an intramuscular injection, paliperidone 6 mg by mouth daily, citalopram 40 mg by mouth daily, trazodone 150 mg by mouth at bedtime, risperidone 2 mg by mouth twice daily, and clonazepam 1 mg by mouth twice
The patient had no known drug allergies or other relevant medical history. She was a chronic smoker and substance abuser, but on admission the urine drug screen was negative.

Upon admission to the burn ICU, the patient had a mild leukocytosis (white blood cell count 15.8 k/mm³) but was afebrile, normotensive, and demonstrated adequate urine output (>0.5 mL/kg/hr). She had evidence of acute respiratory distress syndrome (ARDS), with a ratio of partial pressure arterial oxygen and fraction of inspired oxygen of less than 200, and was started on pressure control ventilation. Due to the severity of the inhalation injury and the need for prolonged mechanical ventilation with aggressive pulmonary hygiene, a tracheostomy was performed on the second hospital day (HD). Post-operatively, the patient became febrile with an associated tachycardia. By HD 5, the patient expressed thick yellow secretions during suctioning and on physical examination diffuse rhonchi were appreciated, worse in the right lung than the left, corresponding with a worsening right-sided, patchy infiltrate seen on chest X-ray. A culture from the broncho-alveolar lavage (BAL) on HD 6 revealed gram-positive cocci in clusters (Table 1). Blood cultures obtained on HD 4 also grew Gram-positive cocci in clusters, and the patient was started on empiric vancomycin and cefepime on HD 6 for presumed

| Date | Site            | Organism     |Susceptibility (mg/L) |
|------|-----------------|--------------|----------------------|
| HD 4 | Blood           | MRSA         | Daptomycin: MIC < 0.5|
|      |                 |              | Gentamicin: MIC < 4  |
|      |                 |              | TMP/SMX: MIC < 0.5/9.5|
|      |                 |              | Vancomycin: MIC = 1  |
| HD 4 | Respiratory     | MRSA         | Gentamicin: MIC < 0.5|
|      |                 |              | Linezolid: MIC < 4   |
|      |                 |              | TMP/SMX: MIC < 0.5/9.5|
|      |                 |              | Vancomycin: MIC = 1  |
| HD 6 | Quantitative BAL| >180,000 CFU/mL MRSA   | Gentamicin: MIC < 4  |
|      |                 |              | Linezolid: MIC < 4   |
|      |                 |              | TMP/SMX: MIC < 0.5/9.5|
|      |                 |              | Vancomycin: MIC = 2  |
|      |                 |              | Ceftaroline: MIC = 0.5|
| HD 10| Quantitative BAL| >110,000 CFU/mL MRSA   | Gentamicin: MIC < 4  |
|      |                 |              | Linezolid: MIC < 4   |
|      |                 |              | TMP/SMX: MIC < 0.5/9.5|
|      |                 |              | Vancomycin: MIC = 2  |
| HD 10| Blood           | No growth   |                      |

*BAL* broncho-alveolar lavage, *CFU* colony forming units, *HD* hospital day, *MIC* mean inhibitory concentration, *MRSA* methicillin-resistant *Staphylococcus aureus*, *TMP/SMX* trimethoprim/sulfamethoxazole
hospital-acquired pneumonia with associated bacteremia. However, despite an aggressive vancomycin dosing schedule [1750 mg intravenous (IV) every 6 h] and adequate, supratherapeutic, vancomycin trough concentrations of 26 mg/L, the patient’s clinical status did not improve. She continued to be febrile and tachycardic, requiring increased ventilatory support and oxygenation requirements. Repeat urine and blood cultures obtained on HD 5 were negative for the Gram-positive pathogen, which was at this time identified as MRSA. The repeat bronchoscopy on HD 10 demonstrated mild improvement in secretions, but the Gram-stain from the BAL subsequently yielded growth in culture of >110,000 colony forming units/mL gram-positive cocci in clusters, later identified as MRSA with a vancomycin MIC of 2 mg/L by automated susceptibility testing. Due to the patient’s deteriorating clinical status on HD 10 while on high-dose vancomycin therapy, the decision was made to modify the antibiotic regimen. At the Detroit Medical Center, in accordance with suggested treatment pathways, MRSA bacteremia and/or pneumonia and a vancomycin MIC of ≥2 mg/L constitute a reason to change therapy to a suitable alternative. However, with consideration of the patient’s medication history, and current medication regimen of antipsychotics and a selective serotonin reuptake inhibitor (SSRI), linezolid was not employed due to the possibility of drug–drug interactions with the psychiatric medications. After consultation with the infectious diseases (ID) pharmacist and the ID medical consult service, a decision was made to initiate ceftaroline fosamil (Teflaro) for the treatment of this patient’s MRSA pneumonia.

Ceftaroline fosamil (Teflaro) is a new advanced generation cephalosporin approved by the FDA for use in acute bacterial skin and skin structure infection (ABSSSI) including MRSA and CABP caused by susceptible bacteria. Ceftaroline binds to the penicillin-binding proteins including PBP 2a associated with methicillin resistance in S. aureus and prevents the cross-linkage of peptidoglycan in bacterial cell walls [4–6]. Ceftaroline fosamil, a prodrug, is converted to the active form of ceftaroline via serum phosphatases and undergoes minimal cytochrome P450 oxidation. Approximately 88% of a dose of ceftaroline is excreted renally, with only 6% of the drug recovered in the feces [4]. The prescribing information indicates the approved dose of ceftaroline fosamil administered intravenously is 600 mg every 12 h for ABSSSI and CABP [7]. However, due to the increased volume of distribution ($V_d$), renal clearance, and urine output in burn injury patients, the patient’s regimen was empirically modified to 600 mg IV every 8 h, and administered as a 2-h infusion to ensure that adequate drug concentrations be maintained throughout the dosing interval. The clinical MRSA isolate tested had a ceftaroline MIC of 0.5 mg/L by Etest® (bioMérieux SA). This antimicrobial regimen was continued for a total of 14 days. The patient became afebrile 48 h after initiation of ceftaroline and remained afebrile for the length of the hospital stay. Rapid clinical improvement was seen after initiation of ceftaroline; the patient was subsequently weaned from the ventilator on HD 22 and decannulated 2 days later. However, because of her psychiatric illness and auditory/visual hallucinations, hospital discharge was delayed to allow for psychiatric placement.

Due to the relative lack of clinical data supporting the use of ceftaroline in burn patients, serum concentrations of ceftaroline were obtained for PK characterization and
potential dosage adjustment. Serum drug levels were obtained on day five of ceftaroline therapy. Concentrations were obtained 30 min post-infusion, 2 h later, and a final concentration drawn 30 min prior to the next dose (Table 2).

Serum Drug Assay and PK Characterization

The ceftaroline bioassay was performed according to previously published methods [8]. Briefly, quarter-inch disks were placed on agar plates (antibiotic medium number 11) pre-swabbed with Bacillus subtilis ATCC 6633. The disks were spotted with 10 µL of ceftaroline standards (2.5, 10, and 40 mg/L) or plasma samples. Each standard and sample was tested in duplicate. Plates were incubated for 18–24 h at 37 °C, after which the zone sizes were measured using a protocol reader (Protocol; Microbiology International, Frederick, MD, USA). Ceftaroline half-life, $C_{\text{max}}$ (peak), and $C_{\text{min}}$ (trough) mg/L were determined from concentration-versus-time plots assuming a one-compartment model (Table 2). Patient-specific concentrations and time above MIC ($T > \text{MIC}$) were calculated utilizing first-order elimination concepts (Table 3). The elimination half-life, peak and trough concentrations, and area under the curve (AUC) were calculated using PK Analyst Software (version 1.10; MicroMath Scientific Software, Salt Lake City, UT, USA).

DISCUSSION

MRSA is a frequently encountered pathogen in burn units and poses a serious health risk. As aforementioned, this risk is not limited to burn site infections but can be associated with bacteremia and pneumonia. Due to the severity of burn injuries, patients may require multiple surgical procedures, numerous invasive central catheters, and prolonged periods of mechanical ventilation. These risk factors, in addition to the decreased immunological response, can dramatically increase the mortality rates in burn patients. Treatment of these infections in this patient population can present a PK challenge for clinicians, and patients frequently require high

Table 2 Patient-specific ceftaroline levels

| Time       | Ceftaroline level (mg/L) |
|------------|--------------------------|
| 0615–0815  | Infusion 600 mg IV ceftaroline |
| 0845       | 21.9*                    |
| 1045       | 7.3                      |
| 1330       | 4.2                      |

$IV$ intravenous
* 30 min post-infusion

Table 3 Patient-specific ceftaroline PK parameters versus package insert

|                  | $C_{\text{max}}$ (mg/L) | $C_{\text{min}}$ (mg/L) | $T_{1/2}$ (h) | $V_d$ (L/kg) | AUC$_{0-\infty}$ (µg h/mL) | $T > \text{MIC}$ (h) | Clearance (L/h) |
|------------------|--------------------------|--------------------------|---------------|--------------|-----------------------------|-------------------|-----------------|
| Burn patient     | 27.5                     | 1.69                     | 1.5           | 0.42         | 87.6                        | 8                 | 10              |
| Package insert   | 21.3                     | 2.3–2.9                  | 0.31–0.45     | 56.3         |                             |                   |                 |

It should be noted that the AUC$_{0-\infty}$ for the burn patient is for 0–8 and 0–12 h per the package insert (i.e., the dosing interval)

$AUC$ area under the curve, $C_{\text{max}}$ maximum serum concentration, $C_{\text{min}}$ minimum serum concentration, MIC mean inhibitory concentration, $T_{1/2}$ half-life, $V_d$ volume of distribution
doses of antimicrobials to maintain adequate serum concentrations [9–14].

Patients that suffer from a thermal injury have numerous pathophysiological changes, which can dramatically alter the PK/PD of prescribed medications. The physiological changes vary based on the time elapsed from the initial burn injury. The acute phase occurs within the first 72 h of the burn injury and results in decreased cardiac and urine output, and increased systemic vascular resistance, which can affect the distribution and elimination of medications [15]. These alterations in cardiac output, renal clearance, and protein binding can significantly alter the PK properties of most medications including absorption, distribution, and elimination [15, 16]. The hypermetabolic phase occurs around 72 h after the burn injury. During this second phase of burn injury, patients will have increased levels of catecholamines, prostaglandins, glucagon, and cortisol which result in increased cardiac output and glomerular filtration rates [15]. Decreased albumin and increased acute-phase proteins resulting from the acute injury can also potentially alter the effectiveness of highly protein bound drug molecules since only free drug is available to elicit an effect [15, 16]. Additionally, the Vd in significantly burned patients (>20% total BSA), or patients with an inhalation injury, can be altered due to extensive fluid loss from the burned tissue [16]. Finally, hepatic metabolism of medications varies after a burn injury [12, 13, 15, 16]. In the hypermetabolic phase, a decrease in phase 1 metabolism can increase the half-life of medications that are hepatically metabolized and cleared, which ultimately increases the risk of toxicity to the patient. However, phase 2 metabolism, or glucuronidation, does not appear to be affected during the hypermetabolic phase of burn injury [17].

Alterations in the physiological response to a burn make dosing medications, especially antibiotics, challenging in burn patients. Although not typically discussed in burn literature, an inhalation injury elicits the same physiological response as any other thermal injury [18]. Lovering and colleagues [19] evaluated the dosing regimens of linezolid in patients with >20% total BSA. In their study, the burn patients had increased non-renal clearance of the drug and elimination rate constants. However, the Cmax concentrations and the Vd of linezolid were similar between the burn patients and healthy volunteers. A case report of a 27-year-old male with a 52% total BSA burn demonstrated that the typical dose of linezolid 600 mg IV every 12 h and meropenem 1 g IV every 8 h produced subtherapeutic serum concentrations in the patient, necessitating an increase in the frequency of dosing to obtain adequate concentrations of the antibiotics in serum, including T>MIC [20]. Daptomycin Cmax concentrations and AUC values have also been reported to be less (44% and 47%, respectively) in patients with thermal injury, with these patients experiencing a 77% increase in mean clearance values compared to healthy volunteers [12]. This study found that daptomycin doses would need to be increased to 10–12 mg/kg/dose to approximate drug exposures in healthy volunteers (dosed at 6 mg/kg/dose). Overall, the alterations in PK parameters in these antibiotics are variable based on burn size and renal function [16, 21]. In one study evaluating cefepime, the renal clearance and Vd were reportedly 10–30% higher in burn patients than non-burn patients [9]. As a result of available data in burn patients, the practice of increasing the
dose, as well as the frequency of administration, may be reasonable to maintain therapeutic serum concentrations of antibiotics [9, 20, 21]. In this case, alternative antimicrobial therapy was sought due to the patient’s lack of clinical response to vancomycin therapy coupled with an elevated vancomycin MIC (2 mg/L). As previously mentioned, although literature supports linezolid in the treatment of nosocomial pneumonia, it was not chosen due to the potential for subtherapeutic concentrations resulting from pathophysiological changes. In addition, due to the patient’s medication history, there was a potential for drug–drug interactions with the patient’s prescribed antipsychotic medication, which included an SSRI and linezolid. Case reports have cautioned against the use of this combination as near-fatal serotonin syndrome has been reported with linezolid’s inhibition of MAO-A and B [22]. Therefore, ceftaroline was initiated using the 600 mg dose at an 8-h dosing schedule, administered as a 2-h infusion, to maximize the PK/PD of the cephalosporin class (i.e., time-dependent killing) and allowing for infusion of other IV medications simultaneously.

This dosing regimen has been studied in vitro PK/PD models and shown to be highly effective against MRSA, while the 12-h regimen has been studied in complicated skin and skin structure infections and for treatment of CAbbP [23–27]. Bhavnani et al. [28] used Monte Carlo Simulation (MCS) to simulate 2000 patient exposures with the 600 mg IV every 12 h dose, evaluating the ability of this dosing regimen to attain a free drug time above the MIC (fT>MIC) target up to 51% of the dosing interval against S. aureus isolates, utilizing a wide range of MICs (0.125–16 mg/L). Interestingly, against simulated isolates with MICs typically seen at this institution based upon internal data (MIC<sub>90</sub> 1 mg/L), it was reported that 600 mg IV every 12 h would only achieve this PD endpoint in 76.2% of the simulated patients with normal renal function [28]. Recently, a published report where 600 mg IV every 8 h was initiated in 5 out of 6 patients with S. aureus bacteremia (3 endocarditis, 1 urinary tract infection, 1 ethmoid osteomyelitis with uveitis, and 1 prostatitis with septic thrombophlebitis) demonstrated resolution of infection in 5 out of the 6 patients, including one patient with a vancomycin-intermediate (vancomycin MIC = 4 mg/L) isolate [29]. Recently, Casapao and colleagues [30] described 527 patients who received ceftaroline therapy, including 241 patients infected with MRSA. In this observational study of patients who received ceftaroline for various indications, 76 (14.4%) received an off-label dose. Side effects were noted to be slightly higher in these patients (17.1%) but all side effects reported in this study were noted to be ≤7.8% [30]. Although most patients in these reports received multiple agents active against MRSA, the results support further research into the efficacy and safety of ceftaroline at this increased dose. Extended infusion beta-lactam therapy has shown promise in the treatment of ICU infections, notably with piperacillin/tazobactam (given as a 4-h infusion) [31]. Infusion times longer than 2 h were not feasible in this patient due to a combination of drug stability (manufacturer labeling indicates 6-h stability at room temperature) and shortage of infusion sites from demand for other non-antimicrobial parenteral infusions.

Serum concentrations (obtained to characterize the pathophysiological changes associated with inhalational thermal injuries) demonstrated that serum drug concentrations remained above the MIC (0.5 mg/L by Etest) for the full duration of the dosing interval. The 8-h dosing regimen was continued due to the potential for increased drug clearance noted in
this patient, as the high inoculum infection, which could impact the efficacy of ceftaroline. This “inoculum effect” has been noted with other beta-lactams, including cephalosporins as well as glycopeptide antimicrobials [32–34]. Our patient demonstrated similar drug clearance and \( V_d \) compared to healthy adults. However, this patient also demonstrated a much shorter half-life (1.5 versus 2.66 h) and increased AUC (87.6 versus 56.3 \( \mu g \) h/mL), when compared with data reported in the ceftaroline prescribing information. This finding was dissimilar to previously published reports characterizing the increased clearance of beta-lactam antimicrobials in patients with thermal injury [9, 11, 14]. This could potentially be due to the limited amount of burn injury (<1% total BSA) and time period from the burn injury itself, as the ceftaroline therapy was initiated in this patient many days into their admission. Curiously, the concentration obtained 30-min post-infusion was higher than reported in the prescribing information and cannot be completely explained other than this patient exhibited altered ceftaroline PK distribution. The patient completed a total of 14 days of ceftaroline therapy and responded appropriately with resolution of signs and symptoms of infection, and more importantly with no reported toxicity from antimicrobial therapy. Ultimately, the patient was liberated from mechanical ventilation on HD 22 and discharged to a psychiatric care facility on HD 48.

CONCLUSION

In conclusion, we describe a case of an inhalational thermal burn injury patient successfully treated with ceftaroline 600 mg IV every 8 h for documented MRSA pneumonia with a vancomycin MIC of 2 mg/L. We also characterized the PK of ceftaroline in a patient with inhalational thermal injuries utilizing a higher than normal dose of medication with the aim of optimizing serum concentrations due to their disease pathophysiology. Our patient demonstrated higher calculated AUC values resulting from the higher dose and increased frequency of dosing, as well as shorter serum half-life. Larger studies are needed to further evaluate this increased dose for safety and efficacy. Clinicians should recognize these pathophysiological changes in burn injury patients and, if possible, adjust treatment accordingly, thus allowing for improved patient outcomes.

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Disclosures. Janie Faris declares she has no conflict of interest. Ryan Mynatt has served on the advisory board for Theravance Pharmaceuticals. Ashley Snyder is currently employed by Merck & Co, Inc. but was a fellow at the time the work was conducted. Michael J. Rybak has received grants, speaking honoraria or consulting fees for Actavis, Cempra, Cubist (a wholly owned subsidiary of Merck & Co, Inc.), The Medicines Company, Theravance, and partial funding support from NIH. It is important to note that the treatment described in this case report was prescribed at the treating physician’s sole discretion without
the knowledge of Actavis (formerly Forest Laboratories, Inc.), the marketer of ceftaroline fosamil, or its affiliates. The bioassay results were generated without Actavis’s knowledge or authorization. Upon learning of the proposed publication, Actavis was unable to confirm these bioassay results.

**Compliance with ethics guidelines.** This article does not contain any new studies with human or animal subjects performed by any of the authors. The institutional review board (IRB) of Wayne State University does not require IRB approval for publication of a single case report. This was a retrospective case report, so no consent was required.

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