1. Introduction

Cutaneous infections caused by *Mycobacterium marinum* have steadily increased, with over 850 additional cases reported in humans [1]. These cutaneous infections are generally attributed to aquarium or fish exposure after a break in the skin barrier [2]. In most reported cases, the upper limbs and fingers account for a majority of the infection sites [3, 4]. Treatment should be tailored to prevent progression to deeper infections and hasten recovery [4].

Optimal treatment usually involves appropriate drug therapy supplemented by surgical intervention for medical failure [4]. No prior case reports describe the aggressive nature of *M. marinum* resulting in a persistent necrotizing soft tissue infection of an affected extremity, ultimately requiring multiple aggressive wound debridements, followed by an amputation of the affected extremity, in order to hasten recovery.
Figure 1: Initial presentation of injury. Three weeks after a barb fish spine penetrated the volar aspect of the proximal portion of his ring finger on his right hand. The injury occurred while fishing along one of the Charleston charters in August. He complained of a gradual increase in pain and swelling of the finger. On physical examination, entry and exit points were noted on the affected finger. Marked swelling was present; however, no bruising, cellulitis, discharge, erythema, or induration was noted. He was able to extend and flex the finger.

to the adjacent fingers and lower hand, and an inability to completely flex and extend the affected digits. He denied any fevers, chills, or night sweats. On physical examination, vital signs were normal and he was afebrile. His right upper extremity was grossly normal; however, the volar aspect of the proximal phalanx of the right ring finger had two small scars with a slight firmness of the scar. There was extensive swelling, specifically along the base of the ring finger. There was no palpable foreign body, although there was exquisite tenderness over the A1 pulleys of the right fourth and fifth fingers. He was able to make a full fist without difficulty, and the affected fingertips were neurovascularly intact. No bruising, cellulitis, erythema, or induration of the area was noted, and there was no lymphadenopathy within the right upper extremity.

Surgical exploration of the flexor tendon sheath was performed, during which tenosynovitis was noted of the flexor digitorum superficialis (FDS) and flexor digitorum profundus (FDP) tendons (Figure 2). Clear nonpurulent fluid was noted around the tendons. A small puncture area was explored, and no foreign bodies were seen. Tissue samples were sent for pathology and routine acid fast bacilli (AFB) cultures. Due to the high concern for a potential mycobacterial infection, clarithromycin, moxifloxacin, and rifabutin were started empirically. Three days later, he discontinued these medications due to gastrointestinal side effects from rifabutin without informing his physicians. Exudative cultures were negative; however, AFB cultures were positive, consistent with Mycobacterium marinum.

Stain was positive, consistent with atypical mycobacterial involvement.

At one-month postoperative follow-up, he had worsening pain and swelling, discharge, and wound discoloration. Vital signs were normal and he was afebrile. The wound had opened spontaneously and was draining a clear to “dishwater” colored fluid. Physical examination noted significant edema of the right ring finger and the distal palm over the A1 and A2 pulleys. Erythema and induration surrounded the open wound, as well as nonviable wound edges that failed to bleed with manipulation, raising concern for a necrotizing soft tissue infection. He was immediately taken for surgical debridement and radical synovectomy of the right ring finger flexor tendon sheath (Figure 3), and tissue from the tendon sheath was sent for routine and mycobacterial cultures. Intravenous clarithromycin and moxifloxacin were initiated along with ceftriaxone to empirically cover potential superinfection. Twenty-four hours after admission, he was discharged to complete a six-month course of oral moxifloxacin and clarithromycin, in which he strictly adhered to this antimicrobial regimen, and close follow-up visits while on antibiotics were scheduled.

At the one-week after discharge follow-up visit, he had ongoing tenderness along the affected digit. Vital signs were normal and he was afebrile. Physical examination noted a closed incision with intact sutures. Erythema and induration were still present; however, no discharge was noted. There was moderate edema throughout his right ring finger, but the FDS and FDP tendons were intact with limited excursion. Approximately one month following the second surgery, he presented to clinic with persistent swelling, erythema, induration, and dark necrotic appearing tissue along the base of the affected finger, consistent with persistent necrotizing soft tissue infection. Of note, AFB cultures from the second surgery were again positive for persistent mycobacterial
In recent decades, most cases of aquatic sources. Minor traumas, such as Mycobacterium marinum. that failed initial attempts of both medical and surgical intervention.

In most reported cases, the upper limbs and fingers account for a majority of infection sites [3, 4]. Clinical manifestations of superficial M. marinum infection typically include painless clusters of superficial nodules or papules. In about one-third of cases, lesions spread proximally along the lymphatic system in a sporotrichoid fashion, without associated lymphadenopathy [4, 5]. Most infections follow an indolent course, although disseminated cases have been reported in immunocompromised patients [6–8]. Deeper infections, including osteomyelitis and tenosynovitis, can cause significant morbidity if untreated [5, 7, 9].

In general, the average inoculation period for M. marinum is around 2–4 weeks; however, as seen with our case, a longer inoculation period can be seen. Other case reports have also described a prolonged incubation period [3, 4]. Failure to obtain an adequate history of aquatic exposure often occurs, since the preceding trauma is generally minor and unrecalled by the patient [4, 5]. Histology and bacteriology are confirmatory tests of infection; however, they are often difficult to use [5].

Treatment should be tailored to prevent progression to deeper infections and hasten recovery [1–3]. Optimal treatment usually involves appropriate drug therapy supplemented by surgical intervention for medical failure. Currently, there is no consensus on definitive treatment regimens due to paucity of controlled trials [3]. However, most isolates are resistant to isoniazid and pyrazinamide and produce β-lactamase [3, 10].

Several studies have shown effective eradication with a single antibiotic for uncomplicated cases, although most
experts recommend 2-drug therapy [3, 10]. Clarithromycin alone, or in combination with ethambutol or rifampin, is preferred for superficial infections [2, 10]. Minocycline, doxycycline, or trimethoprim-sulfamethoxazole can be substituted for clarithromycin [10]. Duration of therapy ranges from 6 to 20 weeks, with some experts recommending continuation of treatment 1-2 months or longer, after resolution of skin lesions, as described in our case [3]. Deeper infections are best treated with combination of ethambutol and rifampin and surgical debridement [5].

While treatment of wound infections most commonly involves opening the incision, a necrotizing soft tissue infection, the most serious consequence of any wound infection, may require more aggressive surgical intervention [4, 11]. Wound infections resulting in a necrotizing soft tissue infection caused by M. marinum often present with a papule, nodule, or ulcer at the site of their trauma, along with a pertinent history of exposure to nonchlorinated water 2–4 weeks earlier [7]. With unrecognized or untreated cases, these infections often ascend up the finger or hand or spread to involve a local joint or tendon [7]. Over a short period of weeks to months, these localized cutaneous infections can spread to soft tissues, ultimately resulting in tissue necrosis [8]. Pertinent signs and symptoms typically include localized pain, edema, and induration surrounding an infected wound, with or without a nonpurulent exudate [11]. With the exception of immunocompromised hosts, fever, lymphadenopathy, and systemic infection are uncommon. Early intervention is prudent to prevent further extension of infection and, therefore, further tissue necrosis.

In our presenting case, prompt surgical wound debridement was performed, secondary to persistent erythema, swelling, a clear to dishwater colored fluid, and evidence of nonviable necrotic wound edges. Although closely followed up in our clinic with careful inspection and monitored antibiotic therapy, he did not improve. Continued debridement, with all efforts to salvage the affected finger, was undertaken; however, due to persistent Mycobacterium marinum involvement and evidence of nonviable necrotic wound edges, a ray amputation was performed to prevent further compromise of the entire right hand and adjacent fingers. Thus, aggressive surgical debridement in combination with judicious antimicrobial therapy is needed for necrotizing infections [12].

A high index of suspicion is needed in those with cutaneous lesions and a history of aquatic encounter. Since the organism can take several weeks to culture, patients should be preemptively treated with appropriate antibiotics to hasten the disease course. Misdiagnosis or delay in treatment may result in worsening of clinical symptoms and progression to deeper infection. Dual therapy is the mainstay of treatment, although selected cases may require surgical intervention.

Conflict of Interests

None of the authors declare any conflict of interests in the paper, including financial, consultant, institutional, and other relationships that might lead to bias or a conflict of interests. This paper is not under consideration by any other journal. All authors read and approved the final paper.

References


Consent

Written informed consent was obtained from the patient for publication of this case report and any accompanying images. A copy of the written consent is available for review by the editor of this journal.