Tunga penetrans causing a rapidly progressing foot ulcer in a patient with uncontrolled type 2 diabetes mellitus

Ashabilan A. Ebrahim1, Elisia P. Mpango2, Joseph A. Temba1, Zulfiqarali G. Abbas3 and Fredirick L. Mashili4,1

1Department of Physiology, Muhimbili University of Health and Allied Sciences (MUHAS), Dar es Salaam, Tanzania
2Department of Surgery, Amana Regional Hospital, Dar es Salaam, Tanzania
3Department of Medicine, Abbas Medical Centre (AMC), Dar es Salaam, Tanzania
4Correspondence address. Department of Physiology, Muhimbili University of Health and Allied Sciences (MUHAS), PO Box 65001, Dar es Salaam 00255, Tanzania. Tel: +255-752255949; E-mail: fredirick@gmail.com

Abstract
Tungiasis is a parasitic disease resulting from infestation by a female flea Tunga penetrans. The parasites are endemic in the tropics and can infect patients with diabetes mellitus (DM). Augmented by uncontrolled hyperglycemia and pre-existing neuropathy, the parasite may trigger a locally spreading inflammation, which may aggravate the trauma introduced during its extraction, leading into a rapidly progressing foot ulcer. To the best of our knowledge, no such cases in patients with type 2 diabetes have ever been published from Tanzania and likely none worldwide. This case report shows that, in diabetic patients, the wound resulting from the extraction of T. penetrans may get infected and aggravated by the ongoing inflammatory reaction, rapidly evolve into limb-threatening condition and mortality. Preventive measures are necessary and should be emphasized in patients with DM. Studies are needed to increase our understanding of the pathophysiology, proper management and sequelae of ulcers of this nature.

INTRODUCTION
Tungiasis refers to a cutaneous ectoparasitic infection of the skin with the female flea, Tunga penetrans. The disease is a fairly common infection in sub-Saharan Africa [1]. Infection often occurs in the lower limbs, and the infected site presents with severe inflammation and ulceration with secondary bacterial infections that may lead to sepsis, osteomyelitis and eventually gangrene [2].

Diabetic patients are at risk of developing foot ulcers from a combination of hyperglycemia, trauma, underlying diabetic peripheral neuropathy (DPN) and peripheral arterial disease (PAD) [3]. In diabetes, T. penetrans infection and accompanied trauma following its extraction may predispose to rapidly progressing ulcers [3]. We report such a case in a 50-year-old diabetic woman of African descent who presented with an ulcer on the plantar surface of the first metatarsal-phalangeal joint.

CASE REPORT
A 50-year-old woman presented to us with a 2-week history of a progressing ulcer on her right foot following a cutaneous infestation by a sandflea, T. penetrans. The patient had traveled to a known tungiasis endemic area in Tanzania [4]. She was a small-scale businesswoman, known type 2 diabetic for 5 years and not on regular medication.

She reported to have noticed a whitish ‘nodular’ lesion at the tip of the second toe, with no accompanying pain or itching. Removal of the flea using nonsterile sharps was done, leaving an initial small wound. She later observed a similar lesion across the first right metatarsal-phalangeal joint, which was also punctured to remove another flea. Following the removal, an abscess developed around the wounds, extended to involve the medial and lateral aspects of the second and big toes, respectively. At this point, there was accompanying pain and swelling of the foot. The abscess ruptured spontaneously turning into an ulcer that occupied the medial and lateral aspects of the second and big toes, respectively.

Within 2 weeks, the wound had progressed into a foul-smelling ulcer. It extended to the planter aspect of the big toe across the metatarsal phalangeal joint and eventually involved the antero-medial aspect of the right foot. In parallel, she experienced fever, generalized malaise and episodes of fatigue.

On examination, there was a deep-seated foul-smelling ulcer on the plantar surface between the first and second metatarsal-phalangeal joint extending between the second and big toes (5 × 2 cm) (Fig. 1). Also, there was a narrow extension across the first metatarsal phalangeal joint with an expansion on the medial side (4 × 3 cm) (Fig. 2).
Figure 1. Deep-seated ulcer on the plantar surface between the first and second metatarsal-phalangeal joints (5 x 3 cm) extending to the medial side of the right foot.

Figure 2. A narrow extension across the first metatarsal phalangeal joint with an expansion on the medial side (4 x 3 cm).

Examination of the peripheral arterial function using an automated device (Smartdop XT, Kodymedics, India) revealed bilateral impairments in peripheral arterial function, and these were letter confirmed by Doppler studies. The patient had moderate DPN as measured by vibration perception threshold (25v). Her random blood glucose and glycated hemoglobin (HbA1C) were 18 mmol/dl and 8.9%, respectively. She was diagnosed to have a neuro-ischemic diabetic foot ulcer secondary to T. penetrans and was admitted for further management.

Patient’s blood glucose was controlled using insulin, and infection was treated empirically with initial parenteral (iv Ceftriaxone and Metronidazole), then oral antibiotics (Amoxicillin and Metronidazole) and dressing. Symptoms and signs of infection subsided 3 days later, and her blood glucose was between 6 and 8 mmol/l in three consecutive days. Nevertheless, despite successive infection and blood glucose control, the ulcer progressively increased in size. Due to worsening peripheral arterial status (infected toes turning gangrenous), the patient was scheduled for disarticulation of the two affected toes.

Following the decision, counseling was done. However, after repeated attempts, she firmly refused to have the procedure and voluntarily requested to be discharged from the hospital. She continued with regular dressing and oral antibiotics (amoxicillin) while at home. One week later, the patient was brought back and readmitted at the Emergency Department. She presented with fever and loss of consciousness and, having been stabilized, she was rushed to a tertiary hospital for intensive care. The patient was managed at the intensive care and apparently died 24 hours later.

**DISCUSSION**

Tungiasis is a parasitic disease common and endemic in specific localities in the tropics [1, 2]. Being common in the tropics, it may coexist with diabetes mellitus (DM). Despite logical speculations from the pathophysiology of the two diseases pointing to their synergy in causing diabetic foot ulcers [1, 5–7], evidence for this concurrence is still lacking. To the best of our knowledge, we report for the first time, a case of rapidly progressing foot ulcer following infestation by T. penetrans in a patient with poorly controlled DM.

This case, with poor glucose control due to neglected diabetes treatment, presented with severe DPN and PAD. Together, these co-morbidities might have synergistically played a role and facilitated the rapid evolvement and advancement of the ulcer [1, 6, 7]. The wound was caused by trauma that resulted from the extraction of the parasite, while its progression was aggravated by the widespread inflammation and infection possibly caused by the parasite (Table 1).

The reported patient noticed an already enlarged nodule with a grown flea, with no previous itching and pain. In tungiasis, growth of the flea usually triggers acute inflammatory reaction and cause itching and pain [2]. In this patient, however, the pre-existing DPN likely obscured these symptoms and was the reason for the delay in noticing the existence of the parasite. Evidence from previous reports show that T. penetrans infestation can introduce both aerobic and anaerobic bacteria [8]. Additionally, the use of non-sterile sharps to extract the parasite may also introduce bacteria [8]. Therefore, it is likely that the flea, unsterile sharps, or both, introduced bacteria into the wound and caused infection.

Moreover, this patient had PAD that caused ischemia and was advancing to cause gangrene. PAD is fairly common in DM [9]. Nevertheless, T. penetrans is implicated in causing ischemia and gangrene [2] and could have contributed to its progression. Due to advancing ischemia, evident by early signs of gangrene, the two affected toes were scheduled for disarticulation.

This case shows that tungiasis can trigger a rapid development and progression of a wound in a patient with uncontrolled DM. Effective foot care should be emphasized in those with DM and residing in tungiasis-endemic areas. Furthermore, in the tropics, research
Table 1. Full blood count results showing high total white blood cells (WBCs) signifying infection. Neutrophil predominance points toward bacterial infection. Additionally, there is microcytic hypochromic anemia.

| Indices | Absolute count | Reference range | % | Interpretation |
|---------|----------------|-----------------|---|---------------|
| WBC     | $16.3 \times 10^3/\mu l$ | 4.5–11          |   | H             |
| NEU     | 13.6           | 2–9             | 83.5% | H             |
| LYM     | 1.11           | 1–3.3           | 6.82% | N             |
| MONO    | 1.43           | 0–1             | 8.79% | H             |
| EOS     | 0.031          | 0–0.7           | 0.192% | H             |
| BASO    | 0.122          | 0–0.15          | 0.746% | N             |

to uncover the effective and appropriate management of tungiasis-related foot ulcers in patient with DM is mandated.

ACKNOWLEDGEMENTS

We would like to thank the nurses and other hospital staff who contributed to the care and treatment of the presented patient at the Amana regional and Muhimbili national hospitals. We also thank the patient and her relatives who gave consent for this case to be published. Special thanks to Drs Rhona Scott, Maria Davy and Juliet Addo from the GlaxoSmithKline R&D (Africa Non-Communicable Disease Open lab) for reviewing the manuscript and for their constructive comments.

AUTHORS’ CONTRIBUTIONS

All authors have been involved in all stages of the preparation of this case report and they have all read and approved the final version of this report.

CONFLICT OF INTEREST STATEMENT

None declared.

CONSENT

Written informed consent was obtained from the patient for her participation into the study. An additional consent was obtained for publication of this case report. Copies of the written consents are available for review from the editor-in-chief of this journal.

ETHICAL APPROVAL

Since this case was recruited in a research set-up, ethical approval for conducting research and publication was obtained from Muhimbili University of Health Ethical Review Board. A copy of the approval is available for review from the editor-in-chief of this journal.

FUNDING

This case was first noticed from an ongoing recruitment of patients into a longitudinal study on diabetic foot ulcers [10]. The study is funded by a project grant from the GSK Africa Non-Communicable Disease Open Lab (project number: 8806) and the GlaxoSmithKline R&D (Africa Non-Communicable Disease Open lab grant). The funder did not participate in data collection or any activity that is directly related to the execution of the research. Authors retained control of the final content of the publication.

GUARANTOR

Dr Zulfiqarali G. Abbas.

REFERENCES

1. Feldmeier H, Heukelbach J, Ugboroiko US, Sentongo E, Mbabazi P, von Samson-Himmelstjerna G, et al. Tungiasis—a neglected disease with many challenges for global public health. PLoS Negl Trop Dis 2014;8. https://doi.org/10.1371/journal.pntd.0003133.
2. Brothers WS, Heckmann RA. Tungiasis in North America. Cutis 1980;25:636–8.
3. Bandyk DF. The diabetic foot: pathophysiology, evaluation, and treatment. Semin Vasc Surg 2018;31:43–48. https://doi.org/10.1053/j.semvascsurg.2019.02.001.
4. Dassoni F, Polloni I, Margwe SB, Veraldi S. Tungiasis in northern Tanzania: a clinical report from Qameyu village, Babati District, Manyara Region. J Infect Dev Ctries 2014;8:1456–60. https://doi.org/10.3855/jidc.4324.
5. Veraldi S, Valsecchi M. Imported tungiasis: a report of 19 cases and review of the literature. Int J Dermatol 2007;46:1061–6. https://doi.org/10.1111/j.1365-4632.2007.03280.x.
6. Shah BR, Hux JE. Quantifying the risk of infectious diseases for people with diabetes. Diabetes Care 2003;26:510–3. https://doi.org/10.2337/diacare.26.2.510.
7. Muller LMAJ et al. Increased risk of common infections in patients with type 1 and type 2 diabetes mellitus. Clin Infect Dis 2005;26:510–3. https://doi.org/10.1086/431587.
8. Feldmeier H, Heukelbach J, Queiroz Sousa A, Barbosa LMM, Carvalho CBM. Bacterial superinfection in human tungiasis. Trop Med Int Heal 2002;7:559–64. https://doi.org/10.1046/j.1365-3156.2002.00904.x.
9. Barnes JA, Eid MA, Creager MA, Goodney PP. Epidemiology and risk of amputation in patients with diabetes mellitus and peripheral artery disease. Arterioscler Thromb Vasc Biol 2020;7:559–64. https://doi.org/10.1161/ATVBAHA.120.314595.
10. Mashili F, Joachim A, Aboud S, Mchembe M, Chiwanga F, Addo J, et al. Prospective exploration of the effect of adiposity and associated microbial factors on healing and progression of diabetic foot ulcers in Tanzania: study protocol of a longitudinal cohort study. 2019;9:2019–031896. https://doi.org/10.1136/bmjopen-2019-031896.