Background Buruli ulcer is one of the neglected tropical diseases. It is a chronic, debilitating, necrotising disease of the skin and soft tissue caused by Mycobacterium ulcerans. Most times, the pattern of presentation is neglected by the infected because it is regarded as a disease of the poor who have little or no access to healthcare. Living in rural often inaccessible areas and suffering from a triad of ignorance, stigma and poverty, this poor population fails to present early to a hospital.

Methods A retrospective review of patients who accessed care at the infectious disease clinic of Nnewi Diocesan Hospital, Nnewi, Southeast Nigeria, between 1 January to 31 December 2017. To achieve a complete inference, the results of laboratory wound swab culture of all patients were collated and matched with the clinical presentation. All cultures were done by a trained scientist of the German Leprosy and TB Relief Association (GLRA).

Results Review of data showed a total of 10 120 patients of which 6402 were outpatients and 3718 were inpatients; they were between 1 and 86 years of age. There were 60 cases of limb ulcers of which wound swab culture was done. Fifty-four (54) were diabetic foot ulcers while five (5) were venous ulcers. Acid-fast bacilli were detected with Ziehl-Neelsen staining in one specimen and confirmed by the reference center.

Conclusion Most of the Buruli ulcer patients are found incidentally following late presentation at hospitals with a questionable ulcer/wound with a high index of suspicion on clinical examination. If Buruli ulcer is to be eradicated, an intensive rural epidemiological identification programme must be implemented to isolate the infected. The vicious cycle of awareness and education campaigns.

Abstracts

CULTURE-FREE APPROACHES FOR THE DIAGNOSIS AND MANAGEMENT OF PATIENTS WITH RIFAMPICIN RESISTANT TUBERCULOSIS: THE DIAMA PROJECT

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The TB Supranational Reference Laboratory of Benin leads a consortium of 11 partners involved in multiple-drug resistant TB care in Africa. The DIAMA project will explore the feasibility and accuracy of: i) diagnosing TB resistance to first and second line drugs through novel molecular multiplex assays developed by the company Genoscreen; ii) setting-up alternative culture-free approaches for the monitoring of patients’ response to rifampicin-resistant treatment; iii) piloting whether the implementation of software by Data2Care Technologies for real-time monitoring of molecular test results can reduce delays between diagnosis and treatment of RR-TB patients. This project is funded by EDCTP for a period of five years.

Conclusion Together, these advances could dramatically improve the currently dismal prognosis of multiple-drug-resistant TB in health systems in resource-poor settings. Through this presentation, we will share the background information, the design of this project and its progress.

REGULATING CLINICAL TRIALS DURING AN EBOLA EMERGENCY: THE LIBERIAN EXPERIENCE

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Background Effective clinical trials oversight is a major function of a fully functional national medical products regulatory system. However, exercising clinical trial oversight in a resource-limited environment is challenging, in particular during an Ebola outbreak or health emergency. Until the devastating Ebola virus disease (EVD) outbreak in 2014, the Liberia Medicines and Health Products Regulatory Authority (LMHRA) had no capacity for effective clinical trial regulation. This presentation describes the main challenges encountered by LMHRA in regulating clinical trials in Liberia during the largest EVD outbreak that affected West Africa in 2014 and 2015.

Methods By carefully documenting activities during the EVD outbreak, interviewing key stakeholders, and discussions among the LMHRA clinical trial committee, key challenges observed during the outbreak were identified and documented.

Results Limited financial resources, lack of expertise in clinical trials, inaccurate and insufficient information about the functions of the LMHRA, poor coordination among key stakeholders, and the lack of a well-developed regulatory framework, adversely influenced the LMHRA clinical trial oversight performance during the EVD outbreak.

Conclusion It is true that several challenges need to be addressed when regulating a clinical trial in a limited-resource environment during any disease outbreak or international medical emergency. However, the importance of building local expertise in clinical trials through mentorship and training cannot be overemphasised. By taking advantage of grants from developmental partners, national medicines regulatory authorities in resource-limited environments can develop capacity for clinical research oversight.