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学位論文内容の要旨

Abstract of the dissertation

博士の専攻分野の名称:博士(国際感染症学)氏名:ムワンガラ・ロナー・アカペルワ

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学位論文題名

The title of the doctoral dissertation

Molecular characterization of *Mycobacterium avium* clinical isolates from Japan and development of diagnostic tools

(日本の臨床分離トリ型結核菌株の遺伝子解析と診断法開発)

Mycobacterium avium (M.avium) is the leading cause of nontuberculous mycobacterial lung disease worldwide. Early detection and treatment are paramount for the timely control of M. avium infections. However, the lack of affordable rapid diagnostic tools and emergence of drug resistance among other factors has made the management of M. avium infections very challenging. And the emergence of macrolide resistance has resulted in the use of fluoroquinolones (FQ) to treat drug resistant M. avium infections. Recently, FQ resistance has been increasing and in Mycobacterium tuberculosis, resistance is largely attributed to mutations in FQ resistance associating genes, gyrA and gyrB. However, information on FQ resistance mechanisms in M. avium is limited.

In Chapter I, a rapid detection LAMP-based assay was developed targeting a widely used species-specific marker IS1245. The applicability of this assay was further assessed using human (n = 137) and pig (n = 91) *M. avium* isolates from Japan. The LAMP assay had a detection limit of 6 fg (equivalent to 1 genome copy) of *M. avium* DNA per reaction within 30 minutes. All 91 (100%) *M. avium* isolates from pigs were detected positive while all other tested bacterial species were negative. Interestingly, among the 137 clinical *M. avium* isolates, 41 (30%) were undetectable with this LAMP assay as they lacked the IS1245, the absence of which was revealed by PCR and whole-genome sequencing. These findings highlighted genotypic differences in *M. avium* strains from humans and pigs in Japan and how this diversity can influence the applicability of a detection tool across different geographic areas and hosts. Hence, it is critical to understand the local genetic make-up of a pathogen population before adopting internationally set standards. This fast and cost-friendly method will aid in the rapid diagnosis of *M. avium* especially in the Euro Americas which have reported high IS1245 carriage rates and result in timely treatment.

In Chapter II, the mechanisms of resistance to levofloxacin (LVX), a FQ, were investigated through minimum inhibitory concentration (MIC) determination and sequencing of *gyrA*, *gyrB*, *mfpA*, and *mfpB* in a total of 88 *M. avium* isolates from Japan. Among the isolates, 21.6 % (19/88)were susceptible to LVX, whereas 78.4 % (69/88), were resistant. Since only four of the resistant isolates

(4/88, 4.5 %) harbored resistance-conferring mutations (D94Y or D94G) within the QRDR of gyrA, the role of mfpA in LVX resistance was explored. Two mfpA genotypes were observed by whole-genome sequencing (WGS) and to simplify the analysis, a multiplex PCR was developed to rapidly detect these genotypes (C170deleted mfpA and Intact/Non-C170deleted mfpA). From the total 88 isolates analyzed in this study, 58 had the intact mfpA genotype while 30 had C170deleted mfpA type. Among the 58 isolates with intact mfpA, 53/55 (96.4%) were resistant to LVX.MIC determination revealed a significant association (p < 0.001) between isolates with intact-mfpA genotype and decreased LVX susceptibility in comparison with isolates bearing the C170deleted mfpA. This observation suggested that mfpA plays some role in FQ resistance in M.avium and therefore warrants more attention.

Overall, the findings of my studies are intended to contribute to the control of *M. avium* infections.