Revealing antibiotic-tolerance of the Mycobacterium smegmatis Xanthine/Uracil Permease Mutant using Microfluidics and Single-Cell Analysis

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Abstract: To reveal rare phenotypes in bacterial populations conventional microbiology tools should be advanced to generate rapid, quantitative, accurate and high-throughput data. The main drawbacks of widely used traditional methods for antibiotic studies include low sampling rate and averaging data for population measurements. To overcome these limitations microfluidic-microscopy systems have great promise to produce quantitative single-cell data with high sampling rates. Using Mycobacterium smegmatis cells we applied both conventional assays and a microfluidic-microscopy method to reveal antibiotic-tolerance mechanisms of wild type and the msm2570::Tn mutant cells. Our results revealed that the enhanced antibiotic tolerance mechanism of the msm2570::Tn mutant was due to the low number of lysed cells during the antibiotic exposure compared with wild-type cells. This is the first study that characterized the antibiotic-tolerance phenotype of the msm2570::Tn mutant that has a transposon insertion in the msm2570 gene encoding a putative xanthine/uracil permease, which enrolls in uptake of nitrogen compound during nitrogen limitation. The experimental results indicate that the msm2570::Tn mutant can be further interrogated to reveal antibiotic killing mechanisms, in particularly, antibiotics those targets cell wall integrity.

Keywords: antibiotics; conventional; microbiology; microfluidics; microscopy; mycobacterium smegmatis; population; single cell

1. Introduction

Antibiotic tolerant bacteria, persisters, are among the key players in several refractory infectious diseases. Antibiotic tolerance might contribute to treatment failures, recurrences of infectious or extremely long duration of antimicrobial therapy that might often lead evolution of antibiotic resistance [1-4]. Bacterial persisters are low-frequency, metastable phenotypic variants in the clonal populations [4,5]. Several laboratories have used different models to investigate persisters and various hypothesis have been examined to understand whether persister cells are dormant, slowly killed by antibiotics, slowly divide or non-divide in the presence of antibiotics, tolerate single or multiple antibiotic stresses, regrow upon antibiotic treatment, or live in the higher antibiotic concentrations than the minimum inhibitory concentrations (MIC) on the class of antibiotics tested [5-12]. Still, very little is known about the single-cell kinetics and the underlying molecular mechanisms of these phenomena. Persistence has been mostly interrogated according to model systems more than naturally occurred persister cells using conventional methods. One of the biggest shortcomings of the conventional tools is their inadequate detection limits to identify persisters or other rare cells in the heterogenous cell populations [9,10].
In the past decade, microfluidics and microscopy methods have arisen to overcome the limitations of traditional tools [11-23]. The advantages of microfluidic assays over macro-scale, traditional batch culture methods include increased limit of detection, higher sampling rate, higher specificity, and compatibility with several imaging techniques [11-23]. Among these pioneering studies, Goormaghtigh and Van Melderen observed Escherichia coli (E. coli) cells that generated ofloxacin persisters at single-cell level using a commercially available CellASIC ONIX B04A-03 Microfluidic Plate (Millipore, USA) and the CellASIC ONIX Microfluidic System [14]. They showed that persister cells did not constitute a slowly growing subpopulation as previously reported [15-17]. In their experimental model, persisters originated from metabolically active cells and formed long polymucleoid filaments and exhibited maximum SOS induction upon removal of ofloxacin. This study nailed down that persister subpopulation was still heterogeneous. To investigate persistence, case-by-case analysis was required at the single-cell level. On the contrary, Pu et al. reported that dormancy was a passive defense mechanism of E. coli persisters that should be activated prior to enhanced efflux activity. They used single-cell time-lapse microscopy to measure fluorescence intensities of the cells in the microchannels [17]. Widels et al. first filtered persister cells, which exhibited β-lactam-induced filamentation and then monitor awakening of E. coli cephalaxin persisters using a microfluidic mother machine device [18]. Along the same line, Manuse et al. also sorted E. coli cells with low levels of energy-generating enzymes and then observed ATP levels using time-lapse images of the isolated cells in the mother machine device. They showed that cells with low level of ATP survived in the presence of ampicillin [19].

On the other hand, McKinney group mostly focused on mycobacterial persistence and widely investigated behavior of Mycobacterium smegmatis (M. smegmatis), model organism of Mycobacterium tuberculosis (M. tuberculosis, Mtb) in the presence of first-line antituberculosis drugs [11,13,20,21]. Their ground-breaking studies showed that persistence was a dynamic behavior of antibiotic-stressed mycobacteria based on single-cell analysis using a microfluidic-microscopy system [13]. As a recent innovative approach, Baron and co-workers developed a microfluidic acoustic-Raman platform to monitor in real-time lipid concentrations of M. smegmatis under the influence of isoniazid (INH). Their findings presented that INH increased the lipids in the mycobacteria which might render isoniazid tolerance of the cells [22]. Moreover, Bielecka et al. used bioelectrospray technology to generate three-dimensional (3D) microsphere system that incorporated primary human cells, M. tuberculosis cells, and type I collagen, where they could both mimic cellular aggregation and upregulate mycobacterial stress genes. Next, they combined microspheres with a microfluidic platform to study real-time pharmacokinetic modeling [23]. The more efficient experiment systems have been developed than standard in vitro culture and animal models. The better understanding of antibiotic tolerance or resistance mechanisms would be obtained, the probability of discovering new antibiotics would be increased.

In this study, we used a microfluidic chip integrated with dialysis membrane, published in [13,20,21]. We elucidated an increased antibiotic-tolerance mechanism for M. smegmatis. We performed both batch and microfluidic culturing for wild type and the msm2570::Tn mutant of M. smegmatis cells. Our approach produces detailed quantitative data at higher sampling rates, which is more accurate and precise comparing to conventional batch culture methods. To our knowledge, no one has previously characterized the phenotype of the msm2570::Tn transposon mutant that has an insertion in the msm2570 gene encoding a putative xanthine/uracil permease. Besides, the persistency phenotypes of wild-type and the msm2570::Tn mutant cells have been shown to be in good agreement with those by traditional methods. This approach can be improved for automatization and the generated data can be extended to high-throughput for deep-learning approaches.

2. Results

The msm2570::Tn transposon mutant was characterized using conventional assays including growth curves, drug-mediated killing, and stress-response assays. Upon
understanding whether or not the phenotype of the msm2570::Tn transposon mutant was antibiotic-specific, microfluidic cell culture experiments were conducted using fluorescent time-lapse microscopy. We quantified the behavior of the msm2570::Tn transposon mutant at single-cell resolution using the acquired time-lapse videos.

2.1. Conventional Assays

The first approach was to determine the growth rate of the msm2570::Tn transposon mutant measuring culture turbidity (OD\textsubscript{600nm}) at the indicated time points, Figure 1. Growth of the msm2570::Tn transposon mutant in 7H9 liquid medium was indistinguishable from wild-type bacteria, Figure 1. Therefore, transposon insertion did not alter the proliferation rate of the cells.

Figure 1. Growth of the msm2570::Tn mutant in 7H9 medium. Wild type (○), the msm2570::Tn mutant (●). Results are representative of at least two experiments.

To determine the impact of msm2570 disruption on drug-mediated killing, we measured the killing rate of the msm2570::Tn mutant against the first-line and second-line antituberculosis drugs including isoniazid (INH), ethionamide (ETH), ethambutol (EMB), rifampicin (RIF). \textit{M. smegmatis} wild-type and msm2570::Tn transposon mutant strains were exposed to (a) INH 50 µg ml\textsuperscript{-1}, (b) ETH 200 µg ml\textsuperscript{-1}, (c) EMB 5 µg ml\textsuperscript{-1}, or (d) RIF 200 µg ml\textsuperscript{-1}, Figure 2. Serial dilutions of the treated cultures were plated at the indicated time points to determine the percent survival (CFU, colony formation assay). Results are means ± standard errors from three independent cultures. Figure 2 illustrates that the msm2570::Tn transposon mutant cells have higher tolerance to INH and EMB compared to wild-type cells.

Figure 2. Response of the msm2570::Tn transposon mutant to antibiotics in batch cultures. \textit{M. smegmatis} wild-type (○) and msm2570::Tn mutant (●) strains were treated with (a) INH 50 µg ml\textsuperscript{-1}, (b) ETH 200 µg ml\textsuperscript{-1}, (c) EMB 5 µg ml\textsuperscript{-1}, or (d) RIF 200 µg ml\textsuperscript{-1}. Serial dilutions of the treated cultures were plated at the indicated time points to determine the percent survival (CFU, colony formation assay). Results are means ± standard errors from three independent cultures.
ETH 200 µg ml\(^{-1}\), (c) EMB 5 µg ml\(^{-1}\), (d) RIF 200 µg ml\(^{-1}\). Results are the means ± standard errors from three independent cultures.

To evaluate INH sensitivity of the \textit{msm2570::Tn} mutant, we determined the minimum inhibitory concentration (MIC) of INH using both agar and microdilution assays. The MIC values for INH were 3.125 µg ml\(^{-1}\) and 6.250 µg ml\(^{-1}\) for wild type and the \textit{msm2570::Tn} mutant, respectively. This result confirmed that transposon insertions of the \textit{msm2570::Tn} mutant did not make the mutants more sensitive or resistant to INH. Besides, the drug concentrations used in killing rate experiments were 10-fold higher than the MIC values, Figure 2, Figure 3.

\textbf{Figure 3.} Complementation of the \textit{msm2570::Tn} mutant. (a) Chromosomal locus encoding the \textit{msm2570} gene in \textit{M. smegmatis}. (b) INH killing assay for complemented \textit{msm2570::Tn} mutant in batch culture. \textit{M. smegmatis} wild-type (○), the \textit{msm2570::Tn} mutant (*), complemented strain (Φ) strains were treated with INH 50 µg ml\(^{-1}\). Results are the means ± standard errors from three independent cultures.

When the wild-type locus encoding the \textit{msm2570} gene inserted back in the mutant, it perfectly complemented the mutant. It confirmed that the transposon insertion was responsible for the INH-tolerance phenotype, and another mutation did not occur. Upon restoring wild-type phenotype by complementation and assessing antibiotic-specific killing rate, we tested the sensitivity of the \textit{msm2570::Tn} transposon mutant to environmental stresses including 45 °C heat shock, pH 4.5 acidic condition, %1 sodium dodecyl sulfate (SDS) detergent, phosphate buffered saline (PBS) nutrient starvation, Figure 4.
Figure 4. Sensitivity of the msm2570::Tn mutant and wild-type cells to thermal, SDS, pH and PBS. Percent survival of WT and the msm2570 mutant at 45 °C (a), after incubation in 7H9 in the presence of 0.1 % SDS (b), pH 4.5 (c), PBS (d).

The msm2570::Tn transposon mutant did not show increased sensitivity to any of these stresses (i.e., the mutant was equally sensitive as wild-type cells), Figure 4.

2.1. Microfluidic-Microscopy Assay

We investigated underlying dynamics of the enhanced persistency of the msm2570::Tn mutant by analyzing the acquired single-cell resolution, time-lapse fluorescence videos. We used the microfluidic platform which was explained in detailed in [13] and shown in Figure 5.

In this system, the polydimethylsiloxane (PDMS) microfluidic device allowed stable nutrient flow and rapid change in growth conditions of bacteria, which was sandwiched in between coverslip and dialysis membrane Figure 5a, Figure 5b. The device was assembled using two parallel polymethylmethacrylate (PMMA) plates and six screws, next mounted on an inverted, automated, time-lapse fluorescent microscope (Olympus IX75) equipped with a Hamamatsu camera (ORCA-AG CCD). Medium flow (25-30 µl/min) was controlled by a syringe pump. Figure 5c illustrates the acquired images using 100x oil-immersion objective for 0-24, 24-48, 48-64, 64-72 hours. Since wild-type M. smegmatis cells were expressing green fluorescent protein (GFP), the images were obtained using both phase and green-fluorescent channels (150 µs).

Using the microfluidic-microscopy assay we acquired image stacks of premixed (1:1), wild-type (GFP-expressing) and the msm2570::Tn mutant (red fluorescent protein (RFP), RFP-expressing) cells. We manually counted cytolysis and division events of cells for 4-hours sampling time using ImageJ cell counter plugin. Figure 6 shows the total cell number, the numbers of divided and being killed cells during the antibiotic treatment using the microfluidic-microscopy system. GraphPad Prism 4 was used for visualization of the data.
3. Discussion

Microfluidic devices and microscopy techniques have been widely used to reveal antibiotic tolerance mechanisms of bacteria, but only a few studies have reported their observations or findings with underlying genetic mechanisms [11-23]. We investigated the antibiotic tolerances of M. smegmatis wild-type and the msm2570::Tn mutant cells using microfluidic bacteria culture platform which is capable of providing stable flow and rapid medium change [13]. Using microfluidic-microscopy assay we elucidated that the msm2570::Tn mutant displayed higher tolerance to isoniazid in comparison to wild-type cells. The mutation in the msm2570 gene is responsible for xanthine and uracil permease and involved in uptake of nitrogen compound during nitrogen limitation [24,25]. In this context, there was only Petridis and co-workers’ study that reported the msm2570 gene in the list of differentially expressed genes that were involved in uptake of nitrogen compounds during nitrogen limitation in M. smegmatis. Interestingly, they also revealed that the msm2570 gene differentially expressed under nitrogen stress conditions using a continuous culture system. However, when Williams et al. investigated the response of M. smegmatis cells to nitrogen stress using batch culture tools, the msm2570 gene was not explicitly reported [26]. Both studies performed growth measurement and gene expression profiling, none has demonstrated the behavior of the msm2570::Tn mutant cells in the presence of antibiotics. Still correlated results generated using our microfluidic culture and Petridis’ continuous culture chemostat exhibited the limitations of batch culture methods [24].

Our study for the first time reports the importance of the msm2570 gene in the antibiotic stewardship. The msm2570::Tn mutant displayed the most specific hyper persistence to INH and then enhanced survival to EMB, Figure 2, Figure 6. INH and EMB are one of the first lines of treatment for tuberculosis [26,27]. Both INH and EMB are interfering with the biosynthesis of cell membrane in M. smegmatis. It can be speculated that killing mechanisms of INH and EMB might be affected due to altered cell membrane composition of the msm2570::Tn mutant [26]. Thus, further research is needed to reveal the role of the msm2570 gene against the action mechanisms of antibiotics those target cell wall integrity in M. smegmatis. It is important to emphasize that the msm2570::Tn mutant and wild-type cells showed similar proliferation rate in the normal growth medium, Figure 1 and comparable killing profile against ETB and RIF, Figure 2. Moreover, the msm2570::Tn transposon mutant survived slightly better than wild-type under detergent (SDS) stress. This result might be correlated with the antibiotic responses of the msm2570::Tn mutant, which only showed higher antibiotic tolerance to drugs those target cell membrane.
However, for the other stresses including, heat shock, acidic pH, nutrient starvations, the mutant responded similarly to wild-type cells.

When the antibiotic responses of wild type and the msm2570::Tn mutant cells were interrogated using a microfluidic-microscopy assays, the obtained data was in good agreement with those by conventional methods. However, microfluidic-microscopy system allowed long-term, real-time observation of single cells before and during the antibiotic exposure. Therefore, it overcomes the limitations of batch culture assays. It provides more reliable growth measurements in comparison to optical density measurements, which relies on turbidity of cell population without eliminating contribution of death cells and cell debris. Notably, optical density measurements are lacking single-cell level data, they provide the mean value for the growth of the population assuming that a population always has a normal distribution. On the other hand, this method can generate rapid, quantitative, real-time data for automation and high-throughput readouts.

 Colony-Forming Unit (CFU) assays remain the gold standard for evaluating effectiveness of antibiotics. To reveal the underlying drug killing mechanisms, accurate, real-time, and high-rate sampling of cell division and lysis data are required, which makes CFU assay not feasible for automation and digitalization. Our single-cell data obtained using microfluidic-microscopy system was correlated with CFU assay, Figure 2, Figure 6. CFU assay is limited to decipher the viable cells those cannot form colonies [28,29]. Real-time monitoring of cells might address this problem and provides more accurate data with higher sampling frequencies. Besides, CFU assay provided the number of cells which can form colonies at 0, 24, 48, and 72 hours, Figure 2. In our study, we evaluated the behavior of single cells in each image sequence with 4 hours of sampling rate. Figure 6 shows the number of divided and lysed cells (Figure 6c, Figure 6d) with the total cell numbers (Figure 6a, Figure 6b). The cumulative behavior of wild-type M. smegmatis (Figure 6a) and the msm2570::Tn mutant (Figure 6b) cells followed the similar killing profile as they displayed in batch culture (Figure 2a, Figure 2c). Notably, single-cell analysis made it possible to understand behavior of the cells when the cell numbers are at the steady state between two consecutive time intervals in CFU assays. By monitoring the bacterial behavior in the presence of antibiotics at the single-cell level, the difference in their survival strategies can be elucidated as shown for wild-type cells (Figure 6c) and the msm2570::Tn mutant cells (Figure 6d). First, the cumulative cell number was increased both for wild-type and the msm2570::Tn mutant cells due to higher division than lysis events. Next, the number of lysed cells were higher than the number of divided cells for both cell populations. However, compared with wild-type, the msm2570::Tn mutant exhibited less cell lysis. Consequently, the total cell number was higher for the msm2570::Tn mutant, Figure 6. Interestingly, single-cell analysis explained that the steady state survival of wild-type cells between the time points of 20-48 hours in CFU assay (Figure 2) was due to the equal number of division and lysis events in the population (Figure 6c). Although this system can be widely applied for administration of different antibiotics to different cell types, manually counting the number of divided cells and lysed cells was cumbersome and can still lead to inaccurate enumeration in single-cell analysis. Therefore, accurate and precise segmentation of mycobacterium cells is still the great challenge to overcome to make microfluidic-microscopy systems more effective.

4. Materials and Methods

4.1 Growth Rate Measurements

The growth rate of the M. smegmatis cells in Middlebrook 7H9 medium (BD/Difco) at 37°C, 3 g was determined by measuring time-dependent changes in optical density at 600 nm (OD600) (Thermo Scientific biomate 5). Middlebrook 7H9 medium contains 0.5% albumin, 0.2% glucose, 0.085% NaCl, 0.5% glycerol, and 0.05% Tween-80.

4.2 Minimum Inhibitory Concentration (MIC)
The MIC values of INH for the msm2570::Tn mutant was determined using both agar proportion and microdilution assays. For both methods cells were grown to mid-log phase (OD$_{600}$ of 0.5-1). For the agar proportion method, the OD$_{600}$ of cells were diluted to 0.05 in fresh 7H9 medium. Then 10-fold serial dilutions were plated on Luria-Bertani (LB) agar solid plates containing various concentrations of INH (2-fold serial dilutions: 100 – 0.024). Plates were incubated at 37 °C and the lowest concentration of drug required to detect minimum bacterial CFU was determined. For the micro dilution method, cells were diluted to OD$_{600}$ of 0.005 in fresh 7H9 medium. Then, 2-fold serial dilutions of INH ranging from 100 µg ml$^{-1}$ to 0.024 µg ml$^{-1}$ was prepared with 3 ml of cultures those were aliquoted into 15 ml test tubes. As a control, one tube of culture was left drug-free, and one tube was fresh 7H9 medium as a blank. Tubes were incubated at 37 °C for 3 days. Turbidity of the cultures was read using OD$_{600}$ absorbance on a TECAN plate reader. For the plate reader, 200 µl of cells were aliquoted from each tube into Costar 96-well transparent flat-bottom plates. When the measurements were done, the absorbance of the blank was subtracted from readouts from each of the wells. MIC was defined as the lowest concentration of drug required to detect minimum bacterial growth.

4.3 Killing Rate Measurements

The killing rate of the msm2570::Tn mutant was investigated using a conventional CFU assay. The wild-type control strain and the msm2570::Tn mutant cells were grown in 7H9 medium overnight at 37 °C ~3 g. When the cultures were in log phase (OD$_{600}$ ≈ 0.5), the wild-type and mutant strains were diluted to OD$_{600}$ 0.05 with standard 7H9 medium. At time zero, antibiotics were added to the diluted cell cultures, at 6-, 24-, 48-, and 72-hours survival was assessed by withdrawing aliquots of the cultures, plating 10-fold serial dilutions on LB agar, and scoring colonies after 3-4 days at 37°C. The killing rate measurements were performed for isoniazid (INH) 50 µg ml$^{-1}$, rifampicin (RIF) 200 µg ml$^{-1}$, ethionamide (ETH) 200 µg ml$^{-1}$, and ethambutol (EMB) 5 µg ml$^{-1}$. The drug concentrations used in our experiments were 10-fold higher than the MIC values.

4.4 Complementation Assay

The c2570F and c2570R primer pair was designed with the following sequences, respectively, GCT AGC ATG ACC ATT CCT TCT GCC and GAT ATC TCA CGA AAG TCG GTC GGA GAC. The msm2570 gene was PCR-amplified using these primers with NheI and EcoRV restriction sites. The amplified gene was sub-cloned into PCR2.1 TOPO, checked by digestion, and sequenced to check for errors. The pND200_Strep integrative plasmid was ligated with the msm2570 gene [32,33]. The streptomycin resistance encoding pND200_strep_msm2570 plasmid was transformed into DH5α E. coli cells.

4.5 Sensitivity Assay

Sensitivity of the msm2570::Tn mutant to other stresses was assessed. First, thermal stress was tested at 45 °C for 6 hours. Next, cells were exposed to 0.1% detergent, sodium dodecyl sulfate (SDS) for 24 hours. Then, nutrient starvation of the msm2570::Tn mutant was determined by incubation of cells in phosphate-buffered saline (PBS) for 24 hours. Afterwards, the msm2570::Tn mutant was cultured in the medium with acidic conditions, pH 4.5 for 24 hours. The 0, 6, 24 hours survival of the msm2570::Tn mutant was assessed by withdrawing aliquots of the cultures, plating 10-fold serial dilutions on JALB (Luria-Bertani) agar, and scoring colonies after 3-4 days at 37°C.

4.6 Microfluidic-Microscopy Assay

The RFP-expressing msm2570::Tn mutant cells and GFP-expressing wild-type M. smegmatis cells were mixed at 1:1 ratio. The mixture of the cells was sandwiched in between the coverslip (#1) and the semipermeable membrane (Spectrum Lab, MWCO:8) which is under the PDMS (Sylgard 184) microfluidic channels (50 µm x 50 µm) [10]. The
microfluidic chip provided growth of the cells via continuous medium flow (25 - 35 µl min⁻¹, silicone tubing ID/OD: 0.076/0.165 cm, HelixMark, syringe pump: World Precision Instruments) and possibility of medium switching for on-chip drug exposures. Design and operation of the microfluidic device were detailedly explained in [13]. The microfluidic device loaded with cells was mounted on the microscope inside a temperature-controlled chamber (37°C). The microfluidic assay included three steps, 24-hour growth of cells in 7H9 medium, 72-h drug exposure using 50 µg/ml INH (as applied in the CFU assay), and 24-hour regrowth of the cells in the absence of INH.

4.7 Live-Cell Imaging and Single-Cell Analysis

Live-cell imaging was performed using an inverted, automated, time-lapse fluorescent microscope (Olympus IX75) equipped with a Hamamatsu ORCA-AG CCD camera. Using 100x oil-immersion objective (UPLFLN) images were acquired on phase and fluorescent channels (TRIS-red, GFP-green, exposure: 150 µs). Then, images were converted into time-lapse fluorescence movies for image analysis. We used ImageJ cell counter plugin to count the number of divided and lysed cells for every 4-hour time frame. Visualization of the data was performed in GraphPad Prism 4 software.

4.8 Statistical Analysis

Student’s unpaired t-test (two-tailed) was used to assess statistical significance of pairwise comparisons. P values < 0.05 were considered significant. P values were calculated using GraphPad Prism 4 software.

5. Conclusions

In this study, we implemented both a microfluidic-microscopy method and batch culture assays to better understand antibiotic-tolerance behavior of wild type and the msm2570::Tn mutant of M. smegmatis cells. Our findings were confirmed by conventional methods including optical density measurements and CFU assays. Moreover, microfluidic-microscopy techniques revealed that the enhanced antibiotic tolerance mechanism of the msm2570::Tn mutant was due to the low number of lysed cells during the antibiotic exposure compared with wild-type cells. This is the first study that characterized the phenotype of the msm2570::Tn transposon mutant that has a transposon insertion in the msm2570 gene encoding a putative xanthine/uracil permease. Our technique produced quantitative data at higher sampling rates in comparison to traditional methods. Therefore, this technique is convenient to be improved for automation and digitalization process. The first step of this process requires development of rapid, precise, and accurate image segmentation tools for M. smegmatis cells. Segmentation of label-free mycobacteria cells by naked eye is cumbersome and still lead to inaccurate and imprecise results. Therefore, there is an urgent need for development of rapid, robust, and effective deep learning approaches for segmentation of label-free mycobacteria.

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