Impact of Hypoxia on Drug Resistance and Growth Characteristics of *Mycobacterium tuberculosis* Clinical Isolates

Zhonghua Liu\(^1\)*, Yulu Gao\(^2\)*, Hua Yang\(^1\)*, Haiyang Bao\(^1\), Lianhua Qin\(^1\), Changtai Zhu\(^4\)*, Yawen Chen\(^3\)*, Zhongyi Hu\(^1\)*

\(^1\) Shanghai Key Laboratory of Tuberculosis, Shanghai Pulmonary Hospital, Tongji University School of Medicine, Shanghai, China, \(^2\) Department of Laboratory Medicine, Kunshan Hospital Affiliated to Nanjing University of Traditional Chinese Medicine, Kunshan, China, \(^3\) Department of Nursing, Shanghai Jiao Tong University Affiliated Sixth People’s Hospital, Shanghai, China, \(^4\) Department of Transfusion, Shanghai Jiao Tong University Affiliated Sixth People’s Hospital, Shanghai, China

\* These authors contributed equally to this work.

\* zct101@163.com (CTZ); yawench2211@163.com (YWC); shhuzy@163.com (ZYH)

Abstract

*Mycobacterium tuberculosis* (MTB) is a specific aerobic bacterium, but can survive under hypoxic conditions, such as those in lung cheese necrosis, granulomas, or macrophages. It is not clear whether the drug sensitivity and growth characteristics of MTB under hypoxic conditions are different from those under aerobic conditions. In this study, we examined the drug resistance and growth characteristics of MTB clinical isolates by a large sample of *in vitro* drug susceptibility tests, using an automatic growth instrument. Under hypoxic conditions, variance in drug resistance was observed in nearly one-third of the MTB strains and was defined as MTB strains with changed drug sensitivity (MTB-CDS). Among these strains, resistance in a considerable proportion of clinical strains was significantly increased, and some strains emerged as multi-drug resistant. Growth test results revealed a high growth rate and large survival number in macrophages under hypoxia in MTB-CDS. According to the results of fluorescence quantitative PCR, the expression of some genes, including *RegX3* (involving RIF resistance), *Rv0194* (efflux pump gene), four genes related to transcription regulation (*KstR*, *DosR*, *Rv0081* and *WhiB3*) and gene related to translation regulation (*DATIN*), were upregulated significantly under hypoxic conditions compared to that under aerobic conditions (\(p < 0.05\)). Thus, we concluded that some MTB clinical isolates can survive under hypoxic conditions and their resistance could change. As for poor clinical outcomes in patients, based on routine drug susceptibility testing, drug susceptibility tests for tuberculosis under hypoxic conditions should also be recommended. However, the detailed mechanisms of the effect of hypoxia on drug sensitivity and growth characteristics of MTB clinical isolates still requires further study.
Introduction

Tuberculosis (TB) is a serious health problem, and China is one of the countries with high burden of TB [1,2]. One of causes for the difficulty in controlling tuberculosis is TB drug resistance [3–6]. In recent years, there has been a rising trend in the number of multi-drug resistant *Mycobacterium tuberculosis* (MTB) clinical isolates [5–9].

Currently, to guide the clinically rational use of drugs, routine *in vitro* drug susceptibility tests for TB are commonly conducted in many laboratories [10–13]. However, despite therapy based on the test results, there are still some cases of treatment failure [14]. We speculate that one of the causes for this may be related to changes in MTB resistance under hypoxia.

The routine TB drug susceptibility tests in vitro are performed under aerobic conditions (21% of oxygen from the atmosphere). However, in vivo conditions such as granuloma, caseous necrosis tissue, or macrophages are hypoxic in nature [15–17]. If MTB resides in these tissues, the features of its biological metabolism might change [15–18], which may lead to variation in its resistance and growth characteristics. To identify whether the drug sensitivity and growth characteristics of MTB under hypoxic conditions are different from those under aerobic conditions, and to explore the related mechanisms, we conducted the following related research, using a large sample of MTB clinical isolates.

Methods and Materials

Ethics statement

This study was approved by the Shanghai Pulmonary Hospital Affiliated to Tongji University School of Medicine Ethics Committee. Subjects were treated in accordance with the Helsinki Declaration on the participation of human subjects in medical research. Written informed consent was obtained from each participant.

Collection of strains and drug susceptibility testing

Two hundred and forty-five MTB clinical isolates were obtained from the Department of Clinical Laboratory Medicine at Shanghai Pulmonary Hospital between 2013 and 2015 (S1 Fig). Routine drug susceptibility tests were performed using a commercial microplate kit (Yibaishi Biotech, Inc., Shen-zhen, China) on the BACTECT 960 System (Becton Dickinson, Franklin Lakes, NJ) under conventional aerobic conditions, and hypoxic drug susceptibility testing was conducted under hypoxic conditions by covering 50 μl of paraffin oil. In this study, susceptibility to first-line drugs like isoniazid (INH), ethambutol (EMB), rifampicin (RMP), and streptomycin (SM) was analyzed. The minimum inhibitory concentrations (MICs) were recorded. The standard laboratory strain H37Rv was purchased from the Chinese National Institute for Food and Drug Control.

Establishment of the hypoxia model and analysis of growth characteristics of MTB

Aerobic cultures (NRP-2) by Wayne’s method were used to establish the hypoxia model described in this report [15,19]. Population growth curves were determined by a Bioscreen Growth Curve Instrument (Bioscreen C, Helsinki, Finland), using a honeycomb plate with 100 wells (Bioscreen C, Helsinki, Finland). Briefly, 300 μl of 7H9 medium was added to each well, and cultured with shaking at 37˚C. The optical density was measured as absorbance at 600 nm after every 2 hours. Hypoxia conditions were established by covering 50 μl of paraffin oil. If both growth curves under the aerobic and hypoxic conditions were completely in accord with
the Webster’s model, the model of hypoxia was considered successful. The growth characteristics analysis was performed under aerobic as well as hypoxic conditions.

**Enumerating MTB clinical strains in macrophages**

Clinical isolates of MTB were cultured under both aerobic and hypoxic conditions. MTB isolates grown to the logarithmic phase were used to infect THP1 macrophages. At different time points after infection, the infected cells were washed twice with PBS and lysed for 10 minutes using Triton X-100. Subsequently, the lysed mixture was inoculated onto 7H9 solid plates and cultured at 37°C for 4 weeks. Finally, MTB clones were counted.

**Effect of hypoxia on gene expression in MTB clinical isolates**

After reviewing previous studies [11,15–43], the relevant regulatory genes for efflux pumps and transcription factors likely to affect drug resistance in MTB (Table 1) were selected and their expression was quantified by RT-PCR. Briefly, 25 ml of culture was used to extract RNA upon cell lysis via the TRIzol bead beater method and phenol extraction [35]. RNA concentrations were determined using a Nanodrop 2000 (NanoDrop Technologies). RNA was treated with DNase as previously described [35] and reverse-transcribed using the High Capacity RNA-to-cDNA kit (Applied Biosystems) following the manufacturers’ instructions. RT PCR was performed using Power SYBR Green PCR Master Mix (Applied Biosystems), as previously described [35]. PCR primers for the above genes are specified in S1 Table. The tests in this study were performed on the 7500 Fast Dx Real-Time PCR System (Applied Biosystems).

| Category                         | Gene name | Rv number | Description                                                                 |
|----------------------------------|-----------|-----------|-----------------------------------------------------------------------------|
| INH regulation gene              | FbpC      | Rv0129c   | Diacylglycerol acyltransferase/mycolyltransferase Ag85C                      |
| INH regulation gene              | EfpA      | Rv2846c   | MFS-type transporter EfpA                                                   |
| INH regulation gene              | FadE23    | Rv3140    | Acyl-CoA dehydrogenase FadE23                                               |
| Rif regulation gene              | RegX3     | Rv0491    | Two component sensory transduction protein RegX                             |
| Rif regulation gene              | SigF      | Rv3286c   | RNA polymerase sigma factor SigF                                            |
| Rif regulation gene              | MrpA      | Rv0981    | Two-component response regulator MrpA                                       |
| Efflux pump regulation gene      | Rv0194    | Rv0194    | Multidrug ABC transporter ATPase/permease                                   |
| Efflux pump regulation gene      | WhiB7     | Rv3197A   | Transcriptional regulator WhiB7                                              |
| Efflux pump regulation gene      | Rv2136c   | Rv2136c   | Undecaprenyl-diphosphatase                                                   |
| Efflux pump regulation gene      | Rv1410c   | Rv1410c   | Aminoglycosides/tetracycline-transport integral membrane protein            |
| Transcript regulation gene       | Rv0081    | Rv0081    | HTH-type transcriptional regulator                                           |
| Transcript regulation gene       | Rv0324    | Rv0324    | Transcriptional regulator                                                   |
| Transcript regulation gene       | LipY      | Rv3097c   | Triacylglycerol lipase Lip                                                   |
| Transcript regulation gene       | LipX      | Rv1169c   | Lipase LipX                                                                 |
| Transcript regulation gene       | Pks3      | Rv1180    | Polyketide beta-ketoacyl synthase                                            |
| Transcript regulation gene       | MutA      | Rv1492    | Methylmalonyl-CoA mutase small subunit                                       |
| Transcript regulation gene       | AccA3     | Rv3285    | Bifunctional protein acetyl-/propionyl-CoA carboxylase subunit alpha AccA    |
| Transcript regulation gene       | WhiB3     | Rv3416    | Redox-responsive transcriptional regulator WhiB3                            |
| Transcript regulation gene       | KstR      | Rv3574    | HTH-type transcriptional regulator KstR                                     |
| Transcript regulation gene       | Lsr2      | Rv3597c   | Iron-regulated H-NS-like protein                                             |
| Transcript regulation gene       | DevR      | Rv3133c   | Two component transcriptional regulator DevR                                 |
| Transcript regulation gene       | IciA      | Rv1985c   | HTH-type transcriptional regulator                                           |
| Transcript regulation gene       | DATIN     | Rv0079    | Dormancy Associated Translation Inhibitor                                    |

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reaction conditions were as follows: 95˚C for 60 s, 40 cycles of 95˚C for 5 s, 62˚C for 8 s, and 72˚C for 20 s, followed by a melt curve analysis.

Statistical analysis
SPSS software (SPSS Inc., Chicago, IL, USA) was used for the statistical analysis of results. Chi-square test, non-parametric test, and one-way analysis of variance (ANOVA) were performed to analyze the different data in this study. A p-value less than 0.05 was considered statistically significant.

Results
Drug resistance of clinical strains under hypoxic conditions
In this study, the MICs of 245 MTB clinical isolates determined by the microplate method were found to be identical with those obtained by the BACTECT 960 System (S2 Table).

Addition of liquid paraffin to the microplate allowed the formation of a curve similar to that in the Wayne’s aerobic model, as determined by OD (S2 Fig). Therefore, construction of a hypoxic growth model in the microplate was successful.

Upon analyzing the MICs of 245 clinical isolates, 68.2% (167/245) were determined as MTB strains of unchanged drug sensitivity (MTB-UDS), whereas others (31.8%, 78/245) belonged to MTB strains of changed drug sensitivity (MTB-CDS; Fig 1).

The stratified analysis of the drug susceptibility test result showed that the overall sensitivity in MTB-CDS was significantly lower than that in MTB-UDS (12.8% vs. 48.5%, p < 0.05). Resistance to single drugs in MTB-CDS was also significantly higher than that in MTB-UDS (35.9% vs. 10.2%, p < 0.05); multi-drug resistance in MTB-CDS was also significantly higher than that in MTB-UDS (23.1% vs. 12%, p < 0.05; Fig 2).

Overall, under hypoxic conditions, the rate of MTB clinical isolates for which the MICs of first line drugs were increased was 38.5% (30/78), whereas that for which the MICs were decreased was 23% (18/78; S3 Fig).

Growth characteristics of MTB clinical isolates under hypoxic conditions
Based on the growth curve analysis, the growth of MTB-CDS was slower than that of MTB-UDS under aerobic conditions; whereas under hypoxic conditions, the growth curves of MTB-CDS nearly coincided with those of MTB-UDS, (Fig 3) indicating that MTB-CDS can suitably grow under hypoxia.

Analysis of survival ability of MTB clinical strains in macrophages under hypoxic conditions
Examination of the MTB survival ability in macrophages at different times under aerobic conditions showed that the difference in survival between the MTB-CDS and MTB-UDS with increasing time was not statistically significant (p > 0.05; Fig 4); whereas under hypoxic conditions, the survival of the MTB-CDS was significantly higher than that of MTB-UDS (p < 0.05; Fig 5).

Gene expression potentially related to resistance in MTB
According to the results of fluorescence quantitative PCR, the expression of some genes, including RegX3 (involving RIF resistance), Rv0194 (efflux pump gene), four genes related to transcription regulation (KstR, DosR, Rv0081 and WhiB3) and gene related to translation
regulation (DATIN), was significantly upregulated under hypoxic conditions compared to that under aerobic conditions \( (p < 0.05; \text{S4, S5, S6 and S7 Figs}). \)

**Discussion**

Previous studies have shown that MTB can survive under hypoxic conditions \([23, 44, 45]\). In this study, MICs of 245 MTB clinical isolates, determined by the microplate method, were found to be identical with those obtained from the BACTECT 960 System under aerobic conditions. The Wayne model is currently recognized as a reliable *in vitro* model for simulating *in vivo* environments such as MTB dormancy or hypoxic environments. In this study, we...
observed almost coinciding curves upon comparing the growth curve of the MTB culture systems with H37Rv, which was cultured under both aerobic and anaerobic environments. Therefore, the growth model of hypoxia was successfully established. By analyzing the MICs of 245 clinical isolates, we found that about one-third of the MTB strains were MTB-CDS, whereas the others belonged to MTB-UDS.

Further stratified analysis of the results of drug sensitivity testing showed that the overall sensitivity of MTB-CDS was significantly lower than that of MTB-UDS, and the resistance to single drugs and multiple drugs in MTB-CDS were both significantly higher than those in MTB-UDS. Overall, under hypoxic conditions, the proportion of MTB clinical isolates for which the MICs of first line drugs were increased was 38.5%.

Therefore, we can conclude that there was a significant change in drug resistance of some MTB strains under hypoxic conditions. Considering that routine drug susceptibility testing is unable to completely reflect the actual resistance in MTB, we suggest that drug susceptibility tests should also be conducted under hypoxic conditions to exclude hypoxia resistance in MTB, especially for cases showing initial therapeutic failure.

Based on the growth curve analysis, the growth of MTB-CDS was slower than that of MTB-UDS under aerobic conditions, whereas under hypoxic conditions, the growth curves of MTB-CDS were nearly similar to those of MTB-UDS, indicating that MTB-CDS are more suitable for survival under hypoxia.

By determining the survival of MTB in macrophages at different times, we found that the difference in survival under aerobic conditions was not statistically significant between the MTB-UDS and MTB-CDS with increasing time; however, under hypoxic conditions, the
survival of MTB-UDS was significantly higher than that of MTB-CDS. This further proves that MTB-CDS can survive under hypoxic conditions.

A previous study has shown that the dormancy survival regulon (DosR regulon) is chiefly responsible for encoding the dormancy related functions of MTB, and appears to be involved in translation regulation through the interaction of its product with bacterial ribosomal subunits [46]. One of the DosR regulon proteins, DATIN, encoded by the gene Rv0079, can stimulate macrophages and peripheral blood mononuclear cells to secrete important cytokines, which may be significant in granuloma formation and maintenance [47]. Additionally, the expression of DATIN in \textit{M. bovis} BCG was found to be upregulated under pH stress and micro-aerobic conditions [47]. This indicates that MTB survival under hypoxia is closely associated with gene regulation, and some relevant regulatory genes related to drug resistance, such as efflux pumps and transcription factors, are likely involved in MTB drug resistance under hypoxia. According to the results of fluorescence quantitative PCR, the expression of some genes, including RegX3 (involved in RIF resistance), Rv0194 (efflux pump gene), four genes related to transcription regulation (KstR, DosR, Rv0081 and WhiB3) and gene related to translation regulation (DATIN), was significantly upregulated under hypoxic conditions compared to that under aerobic conditions. As for other genes such as IciA, we didn’t find difference in gene expression. However, the previous study showed that, IciA has a bearing on the functional role of the important regulator of MTB chromosomal replication in the context of latency [48]. According to the report, synergy between the N-terminal and C-terminal domains of HupB is essential for its DNA-binding ability, and to modulate the topological

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**Fig 3. Growth curves of MTB clinical isolates under aerobic and hypoxic conditions.** MTB-UDS: changed MTB strains of drug sensitivity; MTB-CDS: unchanged MTB strains of drug sensitivity.

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Fig 4. Survivability of changed MTB strains of drug sensitivity and unchanged MTB strains of drug sensitivity in macrophage cell under aerobic conditions. MTB-UDS: changed MTB strains of drug sensitivity; MTB-CDS: unchanged MTB strains of drug sensitivity.
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Fig 5. Survivability of changed MTB strains of drug sensitivity and unchanged MTB strains of drug sensitivity in macrophage cell under hypoxic conditions.
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features of DNA, which has implications for processes such as DNA compaction, gene regulation, homologous recombination, and DNA repair [49]. Thus, we believe that the mechanisms of hypoxia resistance in MTB clinical isolates are associated with the expression of some genes. Moreover, we speculate that the difference in the expression of some genes could result in better survival of MTB-CDS under hypoxia compared to that of MTB-CDS.

Efflux pumps are thought to have emerged for the expulsion of noxious substances from the bacterial cell, thus allowing its survival; increased expression of efflux pumps is associated with resistance to alien substances, including drugs [50]. These pumps are often called multi-drug efflux pumps [50]. In addition, the expression of drug resistance in MTB is linked with some transcription regulation genes. However, the exact mechanisms involved in hypoxia drug resistance of MTB remain unknown.

In summary, in our study, some MTB clinical isolates showed very good adaptability under hypoxic conditions, better survival in macrophages, and resistance to some anti-tuberculosis drugs. This resulted in incorrect results from routine MTB drug susceptibility testing; hence, we recommend that drug susceptibility tests should be simultaneously conducted under hypoxic conditions, especially for cases with initial therapeutic failure. The mechanisms underlying the effect of hypoxia on drug sensitivity and growth characteristics of MTB still require further study.

Supporting Information
S1 Fig. Flow chart of the M. tuberculosis clinical isolates included in this study. (TIF)
S2 Fig. Growth curves of MTB clinical isolates under aerobic or hypoxic conditions. (TIF)
S3 Fig. Drug resistance characteristics of MTB clinical isolates under aerobic or hypoxic conditions. (TIF)
S4 Fig. Expression of Rv0194 and RegX3 genes under aerobic or hypoxic conditions. (TIF)
S5 Fig. Expression of KstR and WhiB3 genes under aerobic or hypoxic conditions. (TIF)
S6 Fig. Expression of DosR and Rv0081 genes under aerobic or hypoxic conditions. (TIF)
S7 Fig. Expression of DATIN genes under aerobic or hypoxic conditions. (TIF)
S1 Table. PCR primers and conditions used in this study. (XLS)
S2 Table. Overall drug resistance characteristics of the MTB clinical isolates. (XLS)

Author Contributions
Conceptualization: ZHL CTZ YLG ZYH.
Data curation: ZHL.
Formal analysis: YWC CTZ.
Funding acquisition: ZYH ZHL CTZ.
Investigation: ZHL HYB HY LHQ.
Methodology: ZHL HYB HY LHQ.
Project administration: ZYH ZHL CTZ.
Resources: YLG YWC ZYH.
Software: CTZ ZHL.
Supervision: ZHL CTZ YWC.
Validation: ZHL HYB HY LHQ.
Visualization: CTZ YWC YLG ZHL.
Writing – original draft: CTZ ZHL ZYH.
Writing – review & editing: ZHL CTZ ZYH YWC YLG.

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