Research Article

Activity of Medicinal Plant Extracts on Multiplication of \textit{Mycobacterium tuberculosis} under Reduced Oxygen Conditions Using Intracellular and Axenic Assays

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Aim. Test the activity of selected medicinal plant extracts on multiplication of \textit{Mycobacterium tuberculosis} under reduced oxygen concentration which represents nonreplicating conditions.

Material and Methods. Acetone, ethanol and aqueous extracts of the plants \textit{Acorus calamus} L. (rhizome), \textit{Ocimum sanctum} L. (leaf), \textit{Piper nigrum} L. (seed), and \textit{Pueraria tuberosa} DC. (tuber) were tested on \textit{Mycobacterium tuberculosis} H37Rv intracellularly using an epithelial cell (A549) infection model. The extracts found to be active intracellularly were further studied axenically under reducing oxygen concentrations.

Results and Conclusions. Intracellular multiplication was inhibited ≥60% by five of the twelve extracts. Amongst these 5 extracts, in axenic culture, \textit{P. nigrum} (acetone) was active under aerobic, microaerophilic, and anaerobic conditions indicating presence of multiple components acting at different levels and \textit{P. tuberosa} (aqueous) showed bactericidal activity under microaerophilic and anaerobic conditions implying the influence of anaerobiosis on its efficacy.

\textit{P. nigrum} (aqueous) and \textit{A. calamus} (aqueous and ethanol) extracts were not active under axenic conditions but only inhibited intracellular growth of \textit{Mycobacterium tuberculosis}, suggesting activation of host defense mechanisms to mediate bacterial killing rather than direct bactericidal activity.

1. Introduction

Globally, 8 million cases of tuberculosis (TB) are reported annually [1]. TB is characterized by actively dividing \textit{Mycobacterium tuberculosis} (M.tb) in the alveolar macrophages and their dissemination through aerosols generated via cough/droplet nuclei.

Whilst axenic multiplication of M.tb requires aerobic conditions, a number of publications have reported that the bacterium can switch from aerobic/microaerophilic to anaerobic respiratory pathways [2] and may exist in three different respiratory states depending on the microenvironment encountered [3]. It has been postulated that M.tb under reduced oxygen conditions assumes dormancy in which it can remain viable for prolonged periods [4]. This dormancy is not a consequence of constant reactivation and destruction of the bacilli but rather of slower metabolism adopted to enter the stationary phase [5]. Despite quantitative metabolic shutdown dormant bacilli retain environmental responsiveness, indicating continuing metabolic integrity [6].

These nonreplicating, dormant bacilli represent a major reservoir for developing active TB. The current chemotherapeutic regimen mainly targets rapidly multiplying M.tb with the exception of pyrazinamide which is known to act on nonreplicating M.tb [7, 8].

Rapid acquisition of drug resistance in M.tb [9] demands alternative therapeutic approaches. Use of medicinal plants in TB treatment has been explored to strengthen chemotherapeutic regimens [10]. Most of these studies are targeted towards actively dividing bacteria and do not address the pool of “persisters.”

The current study aims to explore the efficacy of selected medicinal plants on slowly dividing M.tb. An assay system using alveolar epithelial cell line (A549) and an axenic...
anaerobic setup based on the modified Wayne method [11] was established to assess the activity of the plant extracts on H37Rv. Airway epithelial cells (AECs) harbor significantly higher numbers of intracellular bacteria and act as a preliminary defense against pathogen invasion by acting as a structural barrier. They also initiate and augment host airway defense through secretion of numerous antimicrobial agents, reactive oxygen species, nitric oxide, and proinflammatory chemokines and cytokines. The cytokines and chemokines secreted by AECs play an important role in attracting other cell types to the site of infection and thus assist in granuloma formation. Additionally in alveolar type II cells it has been reported that bacterial replication is higher compared to human or mouse macrophages in vitro [12,13]. Thus we chose to explore an intracellular model based on type II AECs, represented by A549 cells.

2. Materials and Methods

2.1. Plant Material. Acorus calamus, Ocimum sanctum, Piper nigrum, and Pueraria tuberosa were selected on the basis of their broad antibacterial and immune-modulatory properties (Table 1). Plants were collected and authenticated by Dr. P. Tetali, Naoroji Godrej Centre for Plant Research (NGCPR). Voucher specimens were deposited at the Botanical Survey of India (BSI), Western Center, Pune, India, and NGCPR.

2.2. Extract Preparation. Coarsely powdered plant material was sequentially extracted [14] with acetone, ethanol, and distilled water using the Soxhlet apparatus. Respective solvent (300 mL) was continuously refluxed with 25 g of plant material for a period of 24–30 hours for efficient extraction of the phytoconstituents. After ethanol extraction and evaporation of the solvent, the aqueous extract was prepared by boiling the plant material until the volume of water was reduced to 25%. The aqueous extract was lyophilized (Thermo Fisher Scientific, USA) and the acetone and ethanol extracts were allowed to air-dry. The extracts were reconstituted at 20 mg/mL in dimethyl sulfoxide (DMSO), filtered through 0.2 µm, 25 mm DMSO resistant Acrodisc syringe filters ( Pall Corporation, USA), and stored at −20°C.

2.3. Bacterial Culture. The reference M.tb laboratory strain H37Rv was used. H37Rv was susceptible to the first-line drugs, isoniazid, ethambutol, rifampicin, and pyrazinamide.

2.4. Cytotoxicity Testing Using A549 Cell Line. The human lung carcinoma epithelial cell line, A549 (National Center for Cell Sciences, Pune, India), was used. Cells were grown in DMEM supplemented with 10% fetal calf serum (FCS) and 4 µg/mL of gentamycin. The cytotoxicity testing was carried out by the neutral red uptake assay [15]. Briefly, 100 µg/mL of the plant extract prepared in DMEM + 10% FCS was incubated overnight on a 24 h culture of A549 cells and following incubation subjected to the neutral red assay. The assay was carried out in triplicate and repeated twice.

2.5. Intracellular Assay. A549 cells were grown in DMEM supplemented with 10% fetal calf serum (FCS) and 4 µg/mL of gentamycin. 10^5 cells/well were seeded in a 96-well plate and were infected with M.tb at a multiplicity of infection (MOI) of 1:1 for 6 h. After washing off the extracellular H37Rv, the cells were treated with amikacin (50 µg/mL) for 2 h to kill the remaining extracellular bacteria. After amikacin treatment, the M.tb infected cells were incubated overnight with the plant extracts at a concentration of 25 µg/mL, as no toxicity was observed on A549 cells with this concentration. On the following day the plant extract was washed off. On 0, 3rd, 5th, 7th, and 10th day after infection the cells were lysed with 0.1% sodium dodecyl sulphate to release the intracellular bacteria. The lysate was threefold serially diluted with PBS and 10 µL of the highest two dilutions was spotted onto MiddleBrook 7H11 (MB7H11) agar plates supplemented with OADC (Becton Dickinson, USA) and 0.5% glycerol, the plates were incubated at 37°C for 3 weeks, and the CFUs (Colony Forming Units) were enumerated.

2.6. Axenic Assay under Differential Oxygen Concentration. MiddleBrook 7H9 broth (10 mL/tube), supplemented with ADC and 0.5% glycerol, was aliquoted and the bacterial suspension containing 10^6 CFUs/mL was inoculated into the tubes. Since our earlier study had shown that these plant extracts were inactive at lower concentrations under aerobic axenic conditions [16] 100 µg/mL concentration of the plant extracts was used. Positive (viable M.tb, VMTB) and medium (MB7H9) controls along with a rifampicin (1 µg/mL) control were maintained. The above setup in triplicate was subjected to differentially reducing oxygen concentration, namely, aerobic, microaerophilic, and anaerobic conditions. The microaerophilic conditions were obtained using the candle jar method [17]. The anaerobic conditions were achieved in an anaerobic jar with a gas pack (HiMedia, India) and confirmed using the indicator tablets provided by the manufacturer. The sets were incubated for a period of 10 days. After incubation all of the tubes were vortex mixed and serially diluted 10-fold and 10 µL of this dilution was spotted on MB7H11 agar plate supplemented with OADC and 0.5% glycerol. The plates were incubated at 37°C and under aerobic conditions to ensure growth of bacteria.

3. Results

3.1. Effect of Plant Extracts on Intracellular Growth of H37Rv. The plant extracts were assessed at 100 µg/mL for their toxicity in an in vitro assay system using A549 cells and were found to be nontoxic (Table 2). The growth of H37Rv in an intracellular model using A549 is demonstrated in Figure 1. Twelve plant extracts (25 µg/mL) were tested for their ability to inhibit H37Rv intracellularly. Greater than 90% inhibition of bacterial growth was observed only with P. nigrum aqueous and acetone extracts. Other plant extracts showed inhibition of bacterial growth ranging from 16% to 86% (Figure 2).

Five plant extracts which showed more than 60% inhibition of intracellular bacterial growth (Figure 2) were considered active. The active extracts were tested further for
Table 1: Selection criteria for plants and their percentage yield.

| Sr. number | Botanical name (Herbarium number) | Acronym | Common name | Area of collection | Parts used | Selection criteria* | Percent yield† | Acetone | Ethanol | Aqueous |
|------------|-----------------------------------|---------|-------------|-------------------|-----------|---------------------|----------------|---------|---------|---------|
| 1          | *Acorus calamus* Linn. (BSI-131716) | AC      | Sweet flag  | Shindewadi, Satara district, Maharashtra, India | Rhizome   | Antibacterial, immune-modulatory | 10             | 7.04    | 10.8    |
| 2          | *Ocimum sanctum* Linn. (BSI-131712) | OS      | Indian tulsi | Shindewadi, Satara district, Maharashtra, India | Leaves    | Immune-modulatory   | 6.0            | 3.62    | 24.0    |
| 3          | *Piper nigrum* Linn. (BSI-131714) | PN      | Black pepper | Kottayam, Kerala, India | Seed      | Antibacterial       | 3.16           | 6.64    | 22.64   |
| 4          | *Pueraria tuberosa* DC. (NGCPR-670) | PT      | Indian kudzu | Mulshi, Pune district, Maharashtra, India | Rhizome   | Immune-modulatory   | 6.72           | 5.92    | 22.83   |

*The plants were selected for the present study based on their biological activities mentioned under the column “selection criteria.”

†Percentage yield w/w after sequential extraction using acetone, ethanol, and aqueous extracts.
Table 2: Cytotoxicity of the four plants used in intracellular assay using A549 cells.

| Plant      | Extract (100 μg/mL) | % viability of A549 cells* |
|------------|---------------------|---------------------------|
| A. calamus | Acetone             | 96 ± 3                    |
|            | Ethanol             | 113 ± 7                   |
|            | Aqueous             | 102 ± 32                  |
| O. sanctum | Acetone             | 97 ± 5                    |
|            | Ethanol             | 73 ± 14                   |
|            | Aqueous             | 114 ± 19                  |
| P. nigrum  | Acetone             | 90 ± 7                    |
|            | Ethanol             | 98 ± 19                   |
|            | Aqueous             | 118 ± 11                  |
| P. tuberosa| Acetone             | 118 ± 20                  |
|            | Ethanol             | 97 ± 19                   |
|            | Aqueous             | 82 ± 14                   |

*Values represent mean ± SD of % viable A549 cells from two independent assays.

Figure 1: Intracellular growth of H37Rv over a period of 10 days in A549 based epithelial cell model. The results are represented as mean ± SD of three experiments. An increase of ~1.5 log is observed in growth.

Figure 2: Intracellular inhibition of growth of H37Rv. Extracts of 4 plants were tested on intracellular growth of H37Rv in A549 cells at 25 μg/mL with MOI of 1:1. The extracts showing greater than 60% inhibition of bacterial growth 10 days after infection were selected for testing under anaerobic conditions in an axenic setup.

Figure 3: Activity of plant extracts under reducing oxygen concentration against H37Rv. The 5 intracellularly active plant extracts were tested on H37Rv under reducing oxygen concentration. Viable M.tb acted as the positive control whereas rifampicin was used as the experimental control. P nigrum acetone extracts showed inhibition of bacterial growth under all 3 conditions. The results are represented as mean ± SD of three experiments.

4. Discussion

Due to increasing drug resistance, alternative approaches for treatment of TB and MDR TB are being sought. There is ample clinical and animal experimental evidence that M.tb can persist in tissues for months to decades without replicating and resumes growth to cause active TB [18].
One-third of the population in TB endemic regions acts as reservoirs of these dormant, nonreplicating bacilli [19]. While previous studies [10, 20] focus on activity against actively dividing bacteria, very few report efficacy against the non-replicating bacteria. Hence it is important to study the efficacy of plant extracts on the latter.

It has been observed that when pure isolated compounds are tested individually at a concentration at which they occur in a plant, they are often less active than the plant itself implying that crude plant extracts may be exerting their activity through synergistic/additive action [21–26]. Thus the present study preferred crude extracts over individual active components.

A. calamus (aqueous, ethanol) and P. nigrum (aqueous) were unable to inhibit the growth of H37Rv under axenic condition. However, in the presence of these extracts, intracellular growth was abrogated. The most likely reason would be that the extracts activated the host cell to kill the bacteria. Nevertheless, it cannot be ruled out that the extracts could be modified/altered by the host cell to exhibit bactericidal action.

On the other hand efficacy of P. tuberosa (aqueous) under intracellular, microaerophilic, and anaerobic conditions but not under axenic aerobic conditions may be indicative of its activity only under reduced oxygen concentration. It has been suggested that low oxygen levels can enhance the activity of essential oils present in plant extracts. Diminished oxygen supply leads to fewer oxidative changes in the essential oils and/or the fact that cells obtaining energy via anaerobic metabolism are more sensitive to the toxic action of essential oils [27].

P. nigrum (acetone) was active both intracellularly and in axenic cultures under varying oxygen conditions (aerobic, microaerophilic, and anaerobic) suggesting either a single mode of action that affects multiple bacterial physiological states or the presence of multiple components acting at different levels.

The active extracts identified through this preliminary study with the standard laboratory strain H37Rv need to be screened on clinical isolates. If confirmed, the potential of adjunct therapy of TB with such extracts is viable.

**Ethical Approval**

Clearance for this study was obtained from the Foundation for Medical Research (FMR) Institutional Ethics Committee (20.07.2001/01).

**Disclosure**

The funders had no role in the study design; in the collection, analysis, and interpretation of data; in the writing of the paper; and in the decision to submit the paper for publication.

**Conflict of Interests**

The authors declare no financial/personal relationship with other people or organizations that could inappropriately influence (bias) this work.

**Authors’ Contribution**

Purva D. Bhattar has been involved in designing the experiments, conducting them, and analyzing them and writing the paper; Pooja D. Gupta has been involved in designing the experiments, conducting them, and analyzing them and writing the paper; Tannaz J. Birdi has been involved in designing the assays and editing the paper. The manuscript has been read and approved by all the authors.

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