Anti-mycobacterial Activity of Selected Indian Botanicals Using Surrogate Mycobacterium Strains

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Introduction

Tuberculosis continues to be one of the leading infectious causes of death in the world especially in a developing and densely populated country like India. The main cause of this disease is Mycobacterium tuberculosis (M. tuberculosis). Each year 8 million people are infected by M. tuberculosis, and 2-3 million patients die from TB. It has been estimated that by 2020, nearly 1 billion people will become infected, 200 million will develop disease and 35 million will die from TB, although recent figures suggest that the global epidemic is on the threshold of decline [1]. In India the prevalence of all forms of TB is significantly limited by the usage and efficacy of presently prescribed anti-TB drugs such as isoniazid and rifampicin. Owing to the evolving emergence of drug resistant TB which ultimately leads to poor compliance and complications in the management of TB [4]. The emergence of Extensive Drug Resistant TB (XDR-TB) has been considered as a more serious threat owing to the resistance of XDR-TB strains towards first- and second-line drugs. The limitations and failure of XDR-TB treatment are more frequently observed in immunocompromised people, especially living with HIV infections [5]. Therefore the emerging MDR and XDR crisis in the mainstream of management of tuberculosis has given a new impulse for searching novel, effective and safe antituberculoid agents.

Exploring natural resources especially medicinal plants has always remained a significant hope in the ‘target rich lead poor’ scenario of the present-day drug discovery research. Several natural compounds with anti-mycobacterium activity have been described from plants, fungi and marine organisms [6,7].

The first hand information from the traditional healers from the Kinwat and Mahur forest range of Nanded district (MS), India and the circumstantial literature describing the traditional usage of the medicinal plants in the management of infectious diseases was used as a rational for selecting the botanicals for the present study. For example, the folk practitioners traditionally use the Amorphophyllus sylvaticus as a remedy for healing piles, abscesses are the result of an acute infection in the internal glands of the anus. Andrographis paniculata commonly called as ‘King of Bitter’ has been found to be an ingredient of at least 26 Ayurvedic...
formulations against variety of human ailments. Besides suggested remedy against series of human diseases, traditionally it has been also recommended for the treatment of infections associated with diarrhea, dysentery, cholera, pneumonia, swollen lymph nodes, leprosy, bronchitis, sore throats and against variety of chronic and infectious diseases [9]. Besides the traditional applications Corallocarpus epigaeus in the treatment of chronic rheumatism and snake bite, it has been also recommended for the treatment of infectious diseases like dysentery and syphilitic disorders. Currently the Bhil, Meena, Garasia, Damor and Kathodi community, Rajasthan, India uses tubers of Corallocarpus epigaeus for the treatment of typhoid [10].

The tubers of different species of genus Dioscorea such as D. oppositifolia, D. pentaphylla, D. bulbifera, D. hispida etc. are basically eaten as vegetables by forest inhabiting tribal communities in India [10], however they are also used as traditional medicine to cure different types of diseases including infections associated with piles, syphilis, asthma and ulcers [11]. In Bangladesh, a neighbor country of India, D. bulbifera has been traditionally used for the treatment of leprosy: caused by Mycobacterium leprae [11]. Nevertheless, a report from Dschang, Cameroon (Central Africa) demonstrated the efficacy of bulbils of D. bulbifera against the M. smegmatis and M. tuberculosis ATCC and MTCS2 strains. In Cameroon, this plant has traditional reputation for treating microbial infections [12]. The tribal communities such as Hill-Kharia, Mankirdia, Santhal, Ho, Kolha, Munda and Bhumi of Odisha state, India traditionally use the parts of D. pentaphylla as a remedy for treating infectious diseases [13]. Perhaps the circumstantial literature and traditional usage inspired us to select different species of Dioscorea for testing against surrogate Mycobacterial strains.

According to Ayurvedic literature the fresh leaves of Enicostema axillare has been traditionally used as bitter tonic and is a remedy for typhoid and arthritis. The leaves extract of E. axillare has been reported to possess broad spectrum antibacterial activity [14]. The tribal communities like menthonni or Karal vatti and kalihari in north India traditionally use the tuber paste of Gloriosa superba for the treatment of leprosy, parasitical infections and remedy for other human diseases. The starch obtained from the tuberous root is given internally in gonorrhoea [15]. In the current literature Morinda citrifolia commonly known as “Noni” and also called as “Indian Mulberry” has established its reputation as health promoting plant. Tribes of Andaman and Nicobar Islands in India, have used the ripe and unripe fruit as food and medicine. Traditionally it is also suggested as remedy for treating dysentery and skin infections, moreover the antibacterial activity of various parts of the plant are well documented [16].

In the present study, above described botanicals were sequentially extracted in water, ethanol and hexane along with their alkaloid fractions. The extracts of the individual plants were further evaluated as possible growth inhibitors of surrogate Mycobacterium strains such as M. smegmatis and M. phlei. The surrogate Mycobacterium strains are commonly used as alternatives to the virulent species, M. tuberculosis, to address the anti-mycobacterial potential of plant extracts and other natural products [17]. We have designed the present studies with a hope of discovering new potential antimycobacterial leads from the selected medicinal plants having reputation as traditional medicines.

Materials and Methods

Materials

The Mycobacterium strains such as Mycobacterium smegmatis (M. smegmatis, NCIM 5138 (Figure 1), suggested 'neotype strain' and grows on medium 25a or 48a, 37°C) and Mycobacterium phlei (M. phlei, NCIM 2240 (Figure 2), strain "Crotin" and grows on medium 33, 37°C) were obtained from National Collection of Industrial Microorganisms (NCIM), National Chemical Laboratory Pune (MS), India and were subcultured and maintained onto Lowenstein Jensen media as described previously [18]. Other chemicals like L-asparagine (s.d. fine chemical Ltd. Mumbai), Rifampicin (HiMedia Laboratories Pvt. Ltd. Mumbai), solvents, reagents used were of AR grade and were obtained from commercial sources.
Collection, identification and preparation of plant material

The selected medicinal plants such as tubers of *Amorphophallus sylvaricus* Roxb Kunth (Araceae) (A9), leaves of *Andrographis paniculata* Burm Wall (Acanthaceae) (A10), tubers of *Corallocarpus epigaeus* Rottl and Wld Hook (Cucurbitaceae) (A11), tubers of *Dioscorea oppositifolia* Linn (Dioscoreaceae) (A12), tubers of *Discroa pentaphylla* Linn (Dioscoreaceae) (A13), tubers of *Dioscorea bulbifera* Linn (Dioscoreaceae) (A14), tubers of *Dioscorea hispida* Dennst (Dioscoreaceae) (A15), whole plant of *Enicostema axilläre* Lam Raynal (Gentianaceae) (A16), tubers of *Gloriosa superba* Linn (Liliaceae) (A17) and unripe fruits of *Morinda citrifolia* Linn (Rubiaceae) (A18), were collected from the different localities of Marathwada region (MS), India, in the month of September 2015. The collected plants were identified and authenticated by RNG, Head Department of Botany of the host institute using ‘Flora of Marathwada region’ [19]. The voucher specimens of the collected plants (A9-A18) were deposited in the Herbarium centre of the host institute. Shade dried, powdered plant specimens of the collected plants (A9-A18) were deposited in the Herbarium centre of the host institute. Shade dried, powdered plant samples (10 g) were sequentially extracted in water, ethanol, chloroform and hexane using Soxhlet’s apparatus up to 8 hours. The solvents were evaporated under reduced pressure at approximately 40°C and the extracts were stored in refrigerator until needed for further experiments.

Quantitative determination of the alkaloids and flavonoids

**Estimation of alkaloids content:** Quantitative determination of total alkaloids was performed by the method of Harborne [20]. Briefly, 5 g of the individual plant sample was mixed with 200 ml of 10 % acetic acid (in ethanol) and allowed to stand for 4 h. The cocktail was filtered and the extract was concentrated on a water bath to one-quarter of the original volume. Concentrated ammonium hydroxide was added drop wise to the extract until the precipitation was achieved. The precipitate obtained was washed with dilute ammonium hydroxide and then filtered. The residue of alkaloid was dried and weighed.

**Estimation of flavonoids content**

The total flavonoids content was estimated by referring the method of Bohm and Kocipai-Abyazan [16,21]. Briefly, 10 g of the individual plant sample was extracted repeatedly with 100 ml of 80 % aqueous methanol at room temperature. The solution was filtered through Whatman filter paper No. 42 (125 mm). The filtrate was later transferred into a crucible and evaporated up to dryness over a water bath at 60°C and weighed.

**Determination of antimycobacterial activity of selected plant samples**

Stock solutions of the individual plant extract along with their flavonoid and alkaloid fractions were prepared in 0.5 % dimethyl sulfoxide (DMSO) and diluted to the final concentrations of 30, 40 and 50 mg/ml in sterilized distilled water. As a part of experimental standardization, initially 1 mg/ml concentration of plant extract was used for antimycobacterial analysis and it was further extended up to 10 or 20 mg/ml; however no clear zone of inhibition was observed under experimental conditions. Considering the respective results of the optimization experiments, the concentrations of plant extract were kept on higher side (30-50 mg/ml) in order to have clear demonstrable antimycobacterial effects in terms of zone of inhibition [22].

The sensitivity of test *Mycobacterium* strains to the various fractions of the selected botanicals was demonstrated by agar diffusion method [17]. In short, with help of sterile cork borer (7 mm) the holes were made into the inoculum seeded solidified nutrient agar medium. A 50 µl volume of individual (30, 40 and 50 mg/ml) plant extracts was loaded into the labeled well in the prepared media plate using sterile pipette. Three parallel sets were carried out simultaneously. The plates thus prepared were kept in a refrigerator for proper prediffusion of the sample and incubated at 37°C for 48 hours. Growth of *M. smegmatis* and *M. phlei* was observed after the incubation of 48 hours and the diameter of inhibition zone was measured subtracting the well size. Rifampicin (600 µg/ml) was used as a standard antibiotic for antimicrobial assay. The activity of rifampicin was not demonstrated by agar diffusion because of whole plate inhibition at 30-50 mg/ml concentrations.

**Determination of the minimum inhibitory concentration (MIC) of the plant extracts**

The MIC of the various plants extract including alkaloids and flavonoids fraction were performed using the broth microdilution assay against the test *Mycobacterium* strains [23]. Tests were performed in sterile 96-well microplates by adding a total volume of 200 µl, comprising 100 µl of standardized suspension of either *Mycobacterium* strains (1 × 10^6 cells/ml) and 100 µl of different concentrations of plant extracts/alkaloids/flavonoids and incubated up to 48 h at 37°C. The plates were observed for microbial growth by measuring the absorbance at 620 nm using Thermo make Automatic Ex-Microplate Reader (M 5118170). The MIC was defined as the lowest concentration of the sample that inhibited the growth of test microorganisms.

**HPTLC profiling**

The finger print of flavonoids and alkaloids of the selected plants were developed using a CAMAG (Germany) make instrumental high performance thin layer chromatography (HPTLC) unit available at Anchrome Pvt. Ltd. Mumbai (MS), India. The flavonoid and alkaloid fractions of the individual sample, 10 µl (2 mg/ml) were spotted separately onto the activated TLC plates using a ‘Linomat 5’ application system. Rutin hydrate (1, 5 and 10 µg/ml) was used as a marker flavonoid compound. The flavonoids were separated using a mobile phase of ethyl acetate:formic acid:glacial acetic acid:water (100:11:1:27), while alkaloids separation was carried out using a mobile phase of toluene:ethyl acetate:diethyl amine (70:20:10). After running the TLC plates, the Anisaldehyde sulphuric acid reagent and Dragendorff reagent were used as derivatizing agents for flavonoids and alkaloids respectively. After derivatization the spots were visualized under CAMAG UV cabinet (254 and 366 nm) and were digitized using CAMAG photo documentation unit [24].

**Results**

The amount of total alkaloids present in *M. citrifolia*, *A. paniculata* and *E. axilläre* were estimated to be 13, 26, 18 mg/gm respectively, while the quantity of total flavonoids content was estimated to be and 73.3, 54.5 and 152.3 mg/gm respectively. The antimycobacterial activity of selected botanicals was demonstrated using agar diffusion assay and the efficacy was calculated in terms of MICs. Amongst the ten selected medicinal plants, only the extracts of *M. citrifolia*, *A. paniculata*, and *E. axilläre* showed antimycobacterial activity against the selected
Mycobacterium strains (Figures 1-3). No antimycobacterial activity was observed with the water soluble fraction of either of the plant samples. Amongst the plants showing promising anti-mycobacterial activity, the chloroform extract of A. paniculata was graded as the most effective antimycobacterial agent against the M. smegmatis and M. phlei showing MIC values of 10 mg/ml for both Mycobacterium strains. It is interesting to note that the similar results were observed with the flavonoid fraction of the A. paniculata. The alkaloid fractions of M. citrifolia, A. paniculata, and E. axillare also exhibited significant antimycobacterial activity towards both the strains of Mycobacterium, showing MIC in a range of 10-20 mg/ml. With exception of certain fractions, M. citrifolia and E. axillare also demonstrated considerable anti-mycobacterium activity with a MIC range of 10-30 mg/ml (Figure 3).

The HPTLC fingerprints of flavonoid (Figure 4) and alkaloid fractions (Figure 5) of individual plant samples were carried out along with calculation of the R_f values (Table 1). The HPTLC analysis of flavonoid fractions of the individual plant samples showed the diversity of flavonoids (Figure 4), but the presence of rutin (R_f 0.39, marker compound) was only observed in E. axillare. However, some of the very close R_f values such as 0.16, 0.18, 0.17; 0.23, 0.28, 0.28; 0.42, 0.43, 0.39; 0.60, 0.55, 0.56 and 0.93, 0.92, 0.81 were observed in the flavonoids fraction of the M. citrifolia, A. paniculata and E. axillare respectively. The compounds associated with concerned R_f values shared by these plants might be responsible for showing antimycobacterial activity. The observed R_f values shared by the flavonoids fraction of the M. citrifolia, A. paniculata and E. axillare can be correlated with their MIC values of 20, 10, 15 mg/ml for M. smegmatis and 15, 10, 10 mg/ml for M. phlei respectively (Table 1).

**Table 1:** The ratio of distance travelled by individual fraction to the distance travelled by solvent was calculated as R_f value. The underlined R_f value (0.73) indicates the presence of a common compound in the alkaloid fraction of Morinda citrifolia, Andrographis paniculata and

<table>
<thead>
<tr>
<th>S.No</th>
<th>Name of Plants</th>
<th>Name of the fraction</th>
<th>Band Number (As per the direction of solvent flow)</th>
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<tr>
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<td>R_f values of respective band</td>
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<tr>
<td></td>
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<td>1</td>
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<tr>
<td>1</td>
<td>Morinda citrifolia</td>
<td>Alkaloid</td>
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<td></td>
<td></td>
<td>Flavonoid</td>
<td>0.16</td>
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<tr>
<td>2</td>
<td>Andrographis Paniculata</td>
<td>Alkaloid</td>
<td>0.10</td>
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<td></td>
<td></td>
<td>Flavonoid</td>
<td>0.15</td>
</tr>
<tr>
<td>3</td>
<td>Enicostema axillare</td>
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<td></td>
<td></td>
<td>Flavonoid</td>
<td>0.17</td>
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<td>4</td>
<td>Marker compound</td>
<td>Rutin</td>
<td>0.39</td>
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</table>
Enicostema axillare and the Rf 0.39 of Enicostema axillare indicates the presence of rutin in its flavonoid fraction. The values marked as * indicates the presence of similar type of flavonoids shared by M. citrifolia, A. paniculata and E. axillare.

The HPTLC fingerprints of alkaloids (Figure 5) of M. citrifolia, A. paniculata and E. axillare possessing promising antimycobacterial activity have a common unique band of alkaloid (Rf 0.73), which possibly might be an active ingredient responsible for imparting antimycobacterial activity. The Rf 0.73 shared by the alkaloid fractions of the M. citrifolia, A. paniculata and E. axillare can be correlated with their MIC values of 20, 15, 20 mg/ml for M. smegmatis and 20, 10, 15 mg/ml for M. phlei respectively. Moreover, the HPTLC fingerprints of alkaloid fraction of A. paniculata apparently shows more concentration of Rf 0.73 compound, which possibly might be a reason for showing lower MIC values than M. citrifolia and E. axillare. The above relations clearly demonstrate the possibility of Rf 0.73 compound as potential antimycobacterial agent. The pharmacological investigations of alkaloids from plant origin are also reported to possess antimycobacterial activities [25,26].

The possible mechanism of anti-mycobacterium activities exhibited by these plants could be traced to the possession of diverse phytochemicals such as alkaloids, flavonoids, saponins, tannins, and other nature phenolic compounds or free hydroxyl groups which are classified as active antimicrobial agents [24]. The HPTLC fingerprint of alkaloids of M. citrifolia, A. paniculata and E. axillare possessing promising anti-mycobacterium activities have a common unique band of alkaloid, which possibly might be the active ingredient responsible for imparting anti-mycobacterium activity. Moreover the pharmacological investigations of alkaloids from plant origin are also reported to possess antimicrobial activities [25,26]. Perhaps, the isolation and characterization of the active principles from these plants might help in translating these medicinal herbs as therapeutic modality.

Conclusion

In conclusion, the results of the present study clearly demonstrates the efficacy of A. paniculata, E. axillare and M. citrifolia as growth inhibitors of surrogate Mycobacterium strains such as M. smegmatis and M. phlei. These plants can be explored for further studies which may result in identification of a novel lead/s arresting the growth of Mycobacterium tuberculosis. The results may find useful in converting these botanicals into therapeutic modalities. Further studies on isolation & characterization of active principles from the A. paniculata, M. citrifolia and E. axillare and testing them in vitro and in vivo (animal model) against the virulent Mycobacterium tuberculosis are necessary for ascertaining the practical utility of these plants as anti-tuberculous agents. Nevertheless the results of the present studies hold importance in the midst of MDR-XDR crisis in the management of TB.

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References


