In vitro Anti Tubercular Activity and Physicochemical Standardization of Selected Medicinal Plant Extracts

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Globally tuberculosis is a major health problem. The main question with the current tuberculosis treatment is adverse effects and drug resistance. The present study aimed to identify safe and effective alternative herbal drugs for the treatment of tuberculosis. This study involves the standardization of selected medicinal plant extracts and in vitro antitubercular activity. Based on the literature survey, four medicinal plant extracts were selected i.e. Glycyrrhiza glabra (Yastimadhu), Piper longum (Pippali), Curcuma longa (Haridra) and Adhatoda vasica (Vasaka). To evaluate the authenticity and quality control aspect, chromatographic fingerprinting and physicochemical analysis were carried out. Quantification of phytomarkers was analyzed by High performance liquid chromatography and in vitro anti-mycobacterial activity was performed by agar well diffusion method on Mycobacterium smegmatis. All necessary physicochemical parameters were performed and found to comply with specified limits. Extracts were authenticated by qualitative High-performance thin layer chromatography fingerprinting in comparison with suitable raw herbs and phytomarkers. Quantification by High performance liquid chromatography showed that Glycyrrhiza glabra contains 10 % Glycyrrhizin, Piper longum contains 7 % Piperine, Curcuma longa contains 82.41 % Curcumin and Adhatoda vasica contains 2.3 % Vasicine. In vitro activity of Glycyrrhiza glabra, Piper longum and Curcuma longa showed 20 mm, 16 mm and 14 mm of maximum Zone of inhibition respectively. Adhatoda vasica did not show any activity against Mycobacterium smegmatis.

The present study demonstrates that the selected plant extracts have the potential to inhibit mycobacterial growth. Thus, these standardized herbal extracts could be used in the design and development of a polyherbal formulation for the treatment of primary stage tuberculosis.

Key words: Tuberculosis, anti-tubercular activity, standardization, high performance liquid chromatography, medicinal plants

Tuberculosis (TB) is an infectious bacterial disease and is a threat in many regions of the world, especially in developing countries. According to WHO, Tuberculosis is a virulent bacterial disease caused by Mycobacterium tuberculosis, which is most commonly affects the lungs. It is transferred from person to person through droplets. The development of paleopathology and paleoepidemiology in infectious disease has proven the very ancient origin of this disease[1]. Every year around 10 million people developed TB disease and 1.3 million deaths from TB. Globally 90 % were adults (aged ≥15 y), two-thirds were in eight countries: India (27 %), China (9 %), Indonesia (8 %), the Philippines (6 %), Pakistan (5 %), Nigeria (4 %), Bangladesh (4 %) and South Africa (3 %) are suffering from TB[2]. Approximately over 50 new Mycobacterium species are reported to cause Mycobacterium infections[3]. Currently available treatments have various adverse effects like the time duration of therapy makes patient compliance difficult and such patients become a potent source of drug-resistant strains, another problem of current therapy is that majority of the TB drugs available today are ineffective against pertinacious bacilli, excluding for Rifampicin and Pyrazinamide. Rifampicin is effective against both actively growing and slow metabolizing non-growing bacilli, whereas

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Pyrazinamide is active against semi-dormant non-growing bacilli. However, there are still pertinacious bacteria that are not killed by currently available TB drugs[4]. Three countries India (24 %), China (13 %), Russian Federation (10 %) accounted for Multi drug resistance (MDR). Extensively drug-resistant (XDR) TB strains are resistant to first-line and second-line anti-tubercular antibiotics. XDR-TB is now present in over 100 countries. Immunosuppressed people are more prone to XDR-TB, especially HIV infected. Thus, there is a need to discover new anti-tubercular drugs that are effective in MDR and XDR tuberculosis.

Approximately 60 % of the world’s population still believes in medicinal plants for their primary healthcare. In the Ayurvedic system of medicine, TB is known as ‘Rajayakshma’. Several plant species have been reported in Ayurveda for the treatment of TB and its related disorders. But the major concern in Ayurvedic formulations like tablet, capsules are made with plant powders which are not standardized and may have geographical variation which can also affect the potency and stability of finished goods. Since many years, Ayurvedic and herbal medicines has had to compete with modern medicines, which are proven to be quick acting, strong and effective and a recent trend in ayurvedic and herbal manufacturing are reduction in bulk dosage form and enhancement of potency by using standardized plant extracts, patient compliance and acceptability by improving palatability. Hence an effective and alternative standardized anti-tubercular herbal drug has to be developed with the help of standardized herbal extracts which is essential to maintain safety and consistent efficacy of the product. For this study, we have selected four medicinal plants extract i.e. Glycyrrhiza glabra (Yastimadhu), Piper longum (Pippali), Curcuma longa (Haridra), Adhatoda vasica (Vasaka). Glycyrrhiza glabra (Yastimadhu) dry extract, Piper longum (Pippali) dry extract, Curcuma longa (Haridra) dry Extract and Adhatoda vasica (Vasaka) dry extract were obtained from Vasu Healthcare Pvt. Ltd. Vadodara, Gujrat, India. The Mycobacterial Strain i.e. Mycobacterium smegmatis-NCIM 5138/MTCC 6 was procured from the National Collection of Industrial Microorganisms (NCIM) Pune and other AR grade chemicals were procured from Merck, Germany.

In the present study, the above-described medicinal plant extracts were individually identified; physio chemically standardized and quantified their active constituents by High performance liquid chromatography (HPLC). Further, each standardized plant extracts were used at different concentrations for in vitro anti-mycobacterial activity by using surrogate Mycobacterium smegmatis.

**MATERIALS AND METHODS**

**Plant extracts and chemicals:**

Glycyrhiza glabra (Yastimadhu) dry extract, Piper longum (Pippali) dry extract, Curcuma longa (Haridra) dry Extract and Adhatoda vasica (Vasaka) dry extract were obtained from Vasu Healthcare Pvt. Ltd. Vadodara, Gujarat, India. The Mycobacterial Strain i.e. Mycobacterium smegmatis-NCIM 5138/MTCC 6 was procured from the National Collection of Industrial Microorganisms (NCIM) Pune and other AR grade chemicals were procured from Merck, Germany. Lowenstein Jensen Medium slant (LJ Medium) ready to use (Hi-media) and Mueller Hinton Agar (MHA) was used as a media. The medium base was Modified Middle Brooke 7H9 broth with indicator ready to use (Hi-media) and Modified Selective Enrichment (Lyophilized).

**HPTLC fingerprinting for identification:**

High-Performance Thin Layer Chromatography (HPTLC) was carried out by CAMAG Linomat 5-Applicator, silica gel 60 F 254 pre-coated aluminum foil plate with layer thickness 0.2 mm (Merck, Germany). The mobile phase used for Glycyrrhiza glabra (Yastimadhu) was n-butanol: Water: Glacial acetic acid (7:2:1). Mobile phase for Piper longum (Pippali) was Toluene:Diethyl ether: Dioxane (6.25:2.15:1.6), Mobile phase of Curcuma longa (Haridra) was Chloroform:Methanol:Glacial acetic acid (9.4:0.5:0.1) and Mobile phase of Adhatoda vasica...
(Vasaka) DE\textsuperscript{23} was Ethyl acetate:Methanol:Ammonia (8:2:1). Individually applied each sample concerning their powders and standards into the chromate plate and developed the chromate plates in the above mobile phase about 8 cm from point of application. After removal, dry the plate, then evaluated.

**Physicochemical tests:**

Loss on drying (LOD), pH Value, Water-soluble extractive, Alcohol soluble extractive, Total ash and Acid insoluble ash were performed as per Ayurvedic Pharmacopoeia of India\textsuperscript{24}.

**Quantification of phytomarkers by HPLC:** The instrument used for quantification of phytomarkers was Shimadzu make prominence HPLC with Shim Pack Solar column C18 (250 mm×4.6 mm id, 5μ particle) and photo diode array (PDA) detector. ‘Lab solution’ software was used for data collection.

**Quantification of glycyr rhiz in *Glycyrrhiza glabra* (Yastimadhu)\textsuperscript{20}:** For sample preparation, 500 mg of *Glycyrrhiza glabra* (Yastimadhu) dry extract was taken in 100 ml of volumetric flask and 100 ml of HPLC grade methanol was added in it, further sonicated it for 15 min & filtered with 0.22 μ syringe filtration unit. For reference solution, 100 mg of glycyr rhiz was taken in 100 ml of volumetric flask and the volume made up with HPLC grade methanol (1000 ppm), after that sonicated and filtered with a 0.22 μ syringe filtration unit. Sample and reference solution was run in Glacial Acetic acid: Methanol: 0.2 M Ammonium acetate (1:67:33) mobile phase and results were detected at UV/PDA (280 nm).

**Quantification of piperine in *Piper longum* (Pippali)\textsuperscript{21}:** For sample preparation, 2 gm of *Piper longum* (Pippali) dry extract was taken in 100 ml of a volumetric flask, 50 ml of HPLC grade methanol was added. After that sonicate it for 3 min and heated on a boiling water bath for 25 min. Started cooling and diluted it with 100 ml of HPLC grade methanol, further filtration had done with a 0.22 μ syringe filtration unit. For reference solution, 100 mg of piperine had incorporated in 100 ml of volumetric flask and the volume made up with HPLC grade methanol, further took 1 ml of this solution and diluted it with 10 ml HPLC grade methanol (100 ppm), after that sonicate it and filtered with 0.22 μ syringe filtration unit. Sample and reference solution was run in mobile phase solution i.e. A: 0.136 g of potassium dihydrogen orthophosphate in 900 ml of Demineralised Water (DM water), adjusted to the pH 2.5 with orthophosphoric acid and dilute to 1000 ml with water; B: Acetonitrile and results were detected at UV/PDA (345 nm).

**Quantification of curcumin in *Curcuma longa* (Haridra)\textsuperscript{24}:** For sample preparation, dissolved 50 mg *Curcuma longa* (Haridra) dry extract in 50 ml of HPLC grade Acetonitrile, sonicate it for 10 min. and the volume made up to 100 ml by HPLC grade water. Then filter it through a 0.22 μ syringe filtration unit. For reference solution, 10 mg of curcumin was taken into a 100 ml volumetric flask, then approx. 50 ml HPLC grade Acetonitrile was added sonicate it for 10 min and the volume made up by HPLC grade water, after that filter it through a 0.22 μ syringe filtration unit. Sample and reference solution was run in the mobile phase solution that is A: Acetonitrile (40) B: 2 % Acetic acid (60) and results were detected at UV/PDA (425 nm).

**Quantification of vasicine in *Adhatoda vasica* (Vasaka)\textsuperscript{20}:** For sample preparation, 2 gm of *Adhatoda vasica* (Vasaka) dry extract was dissolved in 100 ml of HPLC grade methanol and refluxed it for 30 min then filtered it through 0.22 μ syringe filtration unit, further 5 ml of this solution was taken and diluted with 10 ml of HPLC grade methanol. For reference solution, 100 mg of vasicine was taken in 100 ml of volumetric flask and the volume made up with HPLC grade methanol, further 1 ml of this solution was taken and diluted to 10 ml of HPLC grade methanol (100 ppm). 0.1 % Orthophosphoric acid: 100 % Acetonitrile (95:5) was taken as a mobile phase and results were detected at UV/PDA (280 nm).

**Invitro anti-tubercular activity of selected plant extracts:**

**Culture preparation:** Reconstituted the Modified selective enrichment with 1 ml of sterile distilled water. Transferred reconstituted Modified selective enrichment aseptically to the Middle brook 7H9 Broth base. Sample ready to use was taken as Middle brook 7H9 Broth with Indicator medium. Then incorporated lyophilized culture into the above medium in aseptic condition, mixed well the sample and incubated at 37° for 48 h. After 2 d, checking the purity of culture by striking a loopful culture from broth on Lowenstein Jensen (LJ) slant. The slant was incubated at 37° for 48 h. After incubation, observed the light-yellow color colony on LJ medium slant and confirmed it by
Ziehl-Neelsen stain (ZN) Acid-fast staining under the microscope.

**Preparation of bacterial inoculum:** Picked one loopful colony of *M. smegmatis* and inoculate into Middle brook 7H9 broth with indicator medium and used as a bacterial inoculum.

**Sample preparation:** All plant samples at different doses were taken individually in a 250 ml Iodometric flask (Table 1). After that methanol and Dimethyl sulfoxide (DMSO) in a ratio of 7:3 had added. Further sonicated all the individual samples for proper mixing then refluxed it on a water bath for 30 min at 80°. Samples were cooled and centrifuged at maximum rpm for three times. Filtered the supernatants and used it as test samples. Each experiment was performed in duplicate. Sterile Mueller Hinton Agar (MHA) media was cooled up to 55°. After cooling, 15 ml of it was poured by a sterile measuring cylinder into sterile petri plates and then the plate was allowed to solidify on a smooth surface. In the rest of the media, 10 μl cultures of bacteria were added and mixed slowly. Then the media was poured on above MHA containing plate. The plate was solidified and then wells were made in MHA plate at a proper distance by sterile borer and labeled. All individual samples at different concentrations were poured in a well.

Solvent (Methanol & DMSO) was used as a negative control. An experiment was performed in duplicate. When samples were diffused completely in well, all MHA plates were incubated into a Bacteriological incubator at 35°-37° for 48 h and observed the zone of inhibition.

**RESULTS AND DISCUSSION**

Chromatographic fingerprinting of herbal dry extracts was used for the authenticity and quality control of herbal medicines. Retention factor (Rf) Value of Glycyrrhizin Standard, *Glycyrrhiza glabra* dry extracts and *Glycyrrhiza glabra* powder were found to be 0.28 (fig.1). Physico-chemical parameters of *Glycyrrhiza glabra* dry extract are detailed in Table 2. *Glycyrrhiza glabra* dry extract contains 10 % of Glycyrrhizin. Retention time (RT) of *Glycyrrhiza glabra* dry extract was 10.150 and Glycyrrhizin standard was 10.281 which are shown in fig. 2 and fig. 3. Rf Value of Piperine Standard, *Piper longum* dry extract & *Piper longum* powder were found to be 0.65 (fig. 4). Hence confirm the presence of Piperine. Physico-chemical parameters of *Piper longum* dry extract are detailed in Table 3. *Piper longum* dry extract contains 7 % of Piperine active content. Retention time (RT) of *Piper longum* dry extract is 25.879 and Piperine standard is 25.867 which are presented in fig. 5 and fig. 6. Rf Value of Curcumin Std., *Curcuma longa* dry extract and Curcumin powder were found to be 0.73 (fig.7). Physico-chemical parameters of *Curcuma longa* dry extract are detailed in Table 4. *Curcuma longa* dry extract contains 82.41 % of Curcumin. Retention time (RT) of *Curcuma longa* dry extract was 38.549 and Curcumin std. was
Vasicine standard, *Adhatoda vasica* DE and *Adhatoda vasica* powder were found to be 0.65 (fig.10). Physicochemical parameters of *Adhatoda vasica* dry extract contain 2.3% of Vasicine. Retention time (RT) of *Adhatoda vasica* dry extract was 5.877 and Vasicine standard was 5.807 which are shown in fig.11 and fig.12. Anti-tubercular activity of *Glycyrrhiza glabra* DE (Yastimadhu) at 80 mg (fig.13), *Piper longum* DE (Pippali) at 50 mg (fig.14) and *Curcuma longa* at

| Parameters                      | Results          |
|--------------------------------|------------------|
| Description                    | Yellowish-brown color powder with a characteristic taste |
| LOD                            | 5.89%            |
| pH (1% solution)               | 3.89             |
| Water Soluble extractive       | 76.36%           |
| Alcohol Soluble extractive     | 11.51%           |
| Total Ash                      | 7.51%            |
| Acid insoluble ash             | 6.86%            |
| % Active of Piperine by HPLC   | 7.0%             |

**TABLE 3: PHYSICOCHEMICAL EVALUATION OF Piper longum DE**

DE=Dry extract; LOD=Loss on drying; HPLC=High performance liquid chromatography

80 mg (fig.13), *Piper longum* DE (Pippali) at 50 mg (fig.14) and *Curcuma longa* at 38.644 which are shown in fig. 8 and fig. 9. Rf Value of Vasicine standard, *Adhatoda vasica* DE and *Adhatoda vasica* powder were found to be 0.65 (fig.10). Physicochemical parameters of *Adhatoda vasica* dry extract are detailed in Table 5.
10 mg (fig.15) shows the maximum zone of inhibition. *Adhatoda vasica* DE did not show any activity (fig.16).

Above results demonstrated that selected standardized plant extracts have the potential to treat tuberculosis. However, *Adhatoda vasica* DE did not give any activity against *Mycobacterium smegmatis* but Ayurvedic literature review and latest scientific publications revealed that *Adhatoda vasica* is one of the major and common ingredient in Classical formulations which is commonly used for the treatment of “Kshyaroga” (TB) and has a potential role as an anti-tuberculosis drug. A Combination of *Adhatoda vasica* with other standardized plant extracts may be helpful in symptomatic treatment. The major problem with long-term use of existing allopathic anti-TB drugs are drug resistance and adverse effects. Drug-resistant tuberculosis is widespread. Research studies

| Parameters | Results |
|------------|---------|
| Description | Yellow-orange color fine powder with a characteristic odor |
| LOD | 1.45 % |
| pH (1% solution) | 5.10 |
| Water Soluble extractive | 15.05 % |
| Alcohol Soluble extractive | 73.00 % |
| Total Ash | 0.52 % |
| Acid insoluble ash | 0.06 % |
| % Assay of Curcumin by HPLC | 82.41 % |

**TABLE 4: PHYSICOCHEMICAL EVALUATION OF Curcuma longa DE**

DE=Dry extract; LOD=Loss on drying; HPLC=High performance liquid chromatography

| Parameters | Results |
|------------|---------|
| Description | Brown color powder with characteristic odor and bitter taste |
| LOD | 3.52 % |
| pH (1% solution) | 4.82 |
| Water Soluble extractive | 95.96 % |
| Alcohol Soluble extractive | 38.99 % |
| Total Ash | 6.40 % |
| % Active of Vasicine by HPLC | 2.3 % |

**TABLE 5: PHYSICOCHEMICAL EVALUATION OF Adhatoda vasica DE**

DE=Dry extract; LOD=Loss on drying; HPLC=High performance liquid chromatography
have been reported that the main advantage of natural antimicrobial agents is that they do not promote the development of resistance. Various research articles and Ayurvedic literature showed that natural products derived from medicinal plants may play a significant role in the discovery of new anti-TB drugs and can be used as an alternative therapy. The quality control point of view, Identification, Standardization, and Quantification of selected plant extracts are mandatory for the formulation of any dosage form like tablet, capsule, syrup, powder etc. In vitro anti-tuberculosis activity was performed on *Mycobacterium smegmatis*.
by Agar well diffusion method, the advantage of using this bacterial strain is naturally resistant to isoniazid and rifampin and has the same profile to MDR strains of Mtb. Formulation development is an important criterion for patient compliance and consistent efficacy. So, this study has been further executed in different combinations of plant extracts for formulation of novel dosage form and efficacy and quality parameters will be evaluated for developed polyherbal formulation and present in upcoming publication.

**Conflict of interests:**

The authors declare no conflict of interest.

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