Development and validation of a RP-HPLC method for the simultaneous determination of Embelin, Rottlerin and Ellagic acid in Vidangadi churna

Rakesh K. Patel, Vishal R. Patel, Madhavi G. Patel

Department of Pharmacognosy, Baroda College of Pharmacy, Parul Group of Institutes, Limda, Vadodara, Gujarat 391760, India
Parul Institute of Pharmacy, Parul Group of Institutes, Limda, Vadodara, India

Received 7 September 2011; accepted 6 March 2012
Available online 14 March 2012

Abstract Vidangadi churna is a popular Ayurvedic formulation described in the chapter Krimikisita of the Ayurvedic literature Cakradatta for the treatment of Krimiroga. The preparation is a composite mixture of the fine powder of fruits of Vidang (Embelia ribis), glandular trichomes of the fruits of Kamala (Mallotus philippensis), mature fruits of Harde (Terminalia chebula), Saindhava and Yavakshara. The use of reversed phase C18 column eluted with gradient mobile phase of acetonitrile and water enabled the efficient separation of the chemical markers in 22 min. Validation of the method was performed in order to demonstrate its selectivity, accuracy, precision, repeatability and recovery. All calibration curves showed good linear correlation coefficients ($r^2 > 0.995$) within the tested ranges. Three markers in Vidangadi churna were quantified with respect to Embelin (0.647%, w/w), Rottlerin (4.419%, w/w), and Ellagic acid (0.459%, w/w). Intra- and inter-day RSDs of retention times and peak areas were less than 3.12%. The recoveries were between 99.66% and 102.33%. In conclusion, a method has been developed for the simultaneous quantification of three markers in Vidangadi churna. The RP-HPLC method was simple, precise and accurate and can be used for the quality control of the raw materials as well as formulations.

1. Introduction

Standardization and analysis of the chemical markers of the ayurvedic and other poly herbal formulations is always difficult. Quantitative determination of chemical markers of each ingredient in the poly herbal preparation required optimal separation techniques by which these markers are separated with the...
highest resolution and the least interferences from each other. For botanicals and herbal preparations, there is a requirement for scientific evidence and clinical validation with chemical standardization, biological assays, animal models and clinical trials [1]. Herbal medicine has been enjoying renaissance among the customers throughout the world. However, one of the impediments in the acceptance of the ayurvedic medicines is the lack of standard quality control profiles. The quality of herbal medicine, i.e. the profile of the constituents in the final product, has implication in efficacy and safety. Due to the complex nature and inherent variability of the chemical constituents of plant-based drugs, it is difficult to establish quality control parameters. Modern analytical techniques are increasing to overcome these problems [2,3]. Separation, identification and determination of chemical components are very difficult for such polyherbal formulations [4]. The advances in chromatographic separation techniques made it possible to quantify the chemical constituents in a mixture with comparatively little clean-up [5]. Particularly, methods using gradient elution high performance liquid chromatography (HPLC) with reversed phase columns are most commonly applied for the analysis of multiple constituents present in medicinal plants and herbal preparations [6].

Vidangadi churna is a well known Ayurvedic preparation described in the chapter krimicikitsa of the ayurvedic literature cakradatta for the treatment of krimiroga [7]. It consists of the mixture of the fine powder of the fruits of Vidang (*Embelia ribes* Burm., F. Myrsinaceae), glandular trichomes of the fruits of Kamala (*Mallotus philippensis* Muell., F. Euphorbiaceae), mature fruits of Harde (*Terminalia chebula* Retz. F. Combretaceae), Saindhava (Rock salt) and Yavakshara (water soluble ash of the grains of *Hordium vulgare* Linn., F. Gramineae).

In the present investigation, we have developed a simple, optimized and validated HPLC method for the standardization of Vidangadi churna. Three chemical markers were selected for quantification, and one was from each medicinal herb used as raw materials, Emelin for Vidang, Rotterlin for Kamala and Ellagic acid for Harde. These markers are responsible for the physiological action of the respective plants [8]. The method was validated on the basis of its selectivity, linearity, precision, accuracy, limit of detection (LOD) and limit of quantification (LOQ) according to the International Conference on Harmonization (ICH) requirements. Profiles of the individual ingredients of Vidangadi churna were also recorded and the markers present in them were quantified.

2. Experimental

2.1. Materials

Vidangadi churna and its individual components were procured from a local market in Ahmedabad city, Gujarat and authenticated by comparison with herbarium specimens [9–11]. Emelin was isolated according to the method of Indian Herbal Pharmacopoeia [12] while Rotterlin was isolated according to the method described by Khorana and Motiwala [13]. The isolated markers were identified by FTIR, MASS and $^1$H NMR spectroscopy. Ellagic acid was obtained from Sigma-Aldrich (St. Louis, Mo, USA). HPLC grade acetonitrile, water and methanol were obtained from Merck (Darmstadt, Germany).

2.2. Instrumentation

Analysis was performed on a Shimadzu LC-20AD HPLC system equipped with an online degaser DGU-20 As, an Rhenodyne 7725 injection valve furnished with 20 $\mu$L loop, an SPD-M20A photodiode array detector and a Class-VP software. Separation was carried out using a Phenomenex column (250 mm $\times$ 4.6 mm i.d., 5 $\mu$m pore size). The column was maintained at 27 °C throughout analysis and the UV detector was set at 298 nm.

Figure 1  HPLC chromatogram of the mixture of Emelin, Rotterlin and Ellagic acid.
2.3. Sample preparation

500 mg powder of Vidangadi churna and 100 mg powder of its three ingredients, Vidang, Kamala and Harde, were extracted three times with 100 mL methanol. The extracts were combined and concentrated at reduced temperature (50 °C) on a rotary evaporator (Equitron rotevar, Medica Instrument Mfg. Co.) to 100 mL. Prior to use, all samples were filtered through a 0.45 μm nylon membrane filter.

2.4. Calibration

The contents of the markers were determined using a calibration curve established with six dilutions of each standard, at concentrations ranging from 10 to 60 μg/mL. Each concentration was measured in triplicate. The corresponding peak areas were plotted against the concentrations of the markers injected. Peak identification was achieved by comparison of both the retention time (Rt) and UV absorption spectrum with those obtained for standards. The reference substances employed to construct the calibration curves were Embelin, Rottlerin and Ellagic acid.

2.5. Validation parameter

The method was validated according to ICH guideline for linearity, precision, accuracy, selectivity, LOD and LOQ [14]. Selectivity was checked using an extract of Vidangadi churna and a mixture of standards in order to optimize separation and detection. Linearity of the method was performed by analyzing a standard solution of markers by the method in the concentration range of 10–60 μg/mL. The accuracy of the

![HPLC chromatogram of Vidangadi churna](image)

**Figure 2** HPLC chromatogram of Vidangadi churna.

![HPLC chromatogram of Vidang](image)

**Figure 3** HPLC chromatogram of Vidang.
3. Results and discussion

3.1. Optimization of HPLC chromatographic conditions

Optimum chromatographic conditions were obtained after running different mobile phases with a reversed phase C18 column. Water was preferred over methanol as a mobile phase because its use resulted in improved separation. Many different gradient systems of mobile phases were tried to achieve the best separation of peaks. Selecting 298 nm as the detection wavelength resulted in an acceptable response and enable the detection of all three compounds used in this study. The temperature of column was maintained at 27 °C throughout analysis. An HPLC fingerprint for Vidangadi churna was developed. Elution was carried out at a flow rate of 0.5 mL/min with acetonitrile as solvent A and water as solvent B using
gradient elution in 0–2 min with 50% A, 2–4 min with 50–60% A, 4–6 min with 60% A, 6–8 min with 60–65% A, 8–10 min with 65–40% A, 10–13 min with 40–35% A, 13–17 min with 35–30% A and 17–22 min with 30–10% A. Each run was followed by a 10 min wash with 10% acetonitrile and an equilibration period of 15 min.

3.2. Quantification of markers present in Vidangadi churna

The three markers were found in Vidangadi churna and they were quantified with respect to Embelin (0.647%, w/w), Rottlerin (4.419%, w/w), and Ellagic acid (0.459%, w/w). The chromatograms of a mixture of Embelin, Rottlerin and Ellagic acid (Fig. 1) and Vidangadi churna (Fig. 2) showed complete separation of the three markers. The chromatograms of the individual components of Vidangadi churna, Vidang (Fig. 3), Kamala (Fig. 4) and Harde (Fig. 5), were obtained. The ingredients were also quantified with respect to the following standards. Vidang contained Embelin (3.126%, w/w), Kamala contained Rottlerin (21.215%, w/w) while Harde contained Ellagic acid (2.218%, w/w). The results obtained are shown in Table 1.

3.3. Method validation for HPLC fingerprinting method

The HPLC method was validated by defining the selectivity, linearity, accuracy, precision, LOD and LOQ. For quantitative purposes, the method was evaluated by taking into account the precision in the retention time and selectivity of marker compounds eluted. A high repeatability in the retention time was obtained for standards and extracts even at high concentrations. For quantitative purpose, linearity, accuracy, precision, LOD and LOQ were evaluated. LOD and LOQ values were 0.12 μg/mL and 0.5 μg/mL for Embelin, 0.45 μg/mL and 1.9 μg/mL for Rottlerin and 0.66 μg/mL and 2.4 μg/mL for Ellagic acid, respectively. Linear correlation was obtained between peak area and concentration of three markers in the range of 10–60 μg/mL. Values of the regression

| Table 1 | Quantification of Embelin, Rottlerin and Ellagic acid in Vidangadi churna and its ingredients by HPLC method. |
| Sample | Amount of Embelin (%, w/w) | Amount of Rottlerin (%, w/w) | Amount of Ellagic acid (%, w/w) |
|--------|--------------------------|-----------------------------|-------------------------------|
| Vidang powder | 3.13±0.01 | – | – |
| Kamala powder | – | 21.25±0.11 | – |
| Harde powder | – | – | 2.22±0.05 |
| Vidangadi churna | 4.15±0.08 | 0.46±0.04 | 0.65±0.01 |

*Mean±SD (n=6).

| Table 2 | Regression parameter, linearity, limit of detection (LOD) and limit of quantification (LOQ) of the proposed HPLC method. |
| Compound | Concentration range (μg/mL) | Rt (min) | Regression equation | R² | LOD (μg/mL) | LOQ (μg/mL) |
|----------|--------------------------|--------|-------------------|---|-------------|-------------|
| Embelin  | 10–60 | 14.10±0.14 | y=42150x−22788 | 0.995 | 0.12 | 0.5 |
| Rottlerin | 10–60 | 17.88±0.03 | y=39764x+91252 | 0.996 | 0.45 | 1.9 |
| Ellagic acid | 10–60 | 7.01±0.06 | y=21019x+33669 | 0.998 | 0.66 | 2.4 |

*Mean±SD (n=6).

| Table 3 | Repeatability and recovery study for the three markers in Vidangadi churna. |
| Compound | Contents (mg/g) | Added amount (mg) | Recorded amount (mg) | Recovery rate (%) | RSD (%) |
|----------|----------------|-----------------|---------------------|------------------|--------|
| Embelin  | 6.47±0.01 | 3 | 9.34±0.11 | 95.66±1.08 | 1.12 |
| Rottlerin | 41.49±0.08 | 20 | 61.69±1.07 | 100.33±0.73 | 0.73 |
| Ellagic acid | 4.59±0.04 | 2 | 6.67±0.17 | 102.33±2.30 | 2.24 |

*Mean±SD (n=3).
coefficients ($r^2$) of the markers were higher than 0.99, thus confirming the linearity of the methods (Table 2). The high recovery values (95.66–102.33%) indicated a satisfactory accuracy. Relative standard deviation of all the parameters was less than 3.5% for the degree of repeatability, indicating the high repeatability of the developed method (Table 3). The low coefficient of variation values of intra-day and inter-day precision revealed that the method is precise (Table 4). Therefore, this HPLC method can be regarded as selective, accurate and precise.

4. Conclusion

The results indicate that Vidangadi churna contains a number of markers that may be responsible for its therapeutic activity. The developed HPLC method will assist in the standardization of Vidangadi churna using biologically active chemical markers. The developed HPLC method for simultaneous determination of Embelin, Rottlerin and Ellagic acid from Vidangadi churna is accurate, precise, reproducible and repeatable. Vidangadi churna also contains a number of other constituents, which are currently the subject of further investigation, apart from those standards studied. In addition, profiles of the individual components in Vidangadi churna have been recorded as a standardization tool. With the growing demand for herbal drugs and increased belief in the usage of herbal medicine, the development of a standardization tool will help in maintaining the quality of this important Ayurvedic preparation.

Acknowledgments

The authors thank to S.K. Patel College of Pharmaceutical Education & Research (SKPCPER), Ganpat University, Mehsana and for providing facility to carry out this research.

References

[1] E.S. Ong, Extraction methods and chemical standardization of botanicals and herbal preparations, J. Chromatogr. B 812 (2004) 23–33.
[2] L.V. Asokar, K.K. Kakkar, O.J. Chakra, Glossary of Indian medicinal plants with active principles, Publication and Information Directorate, New Delhi, 1992, p. 122.
[3] M.N. Ravishankara, N. Shrivastava, H. Padh, et al., HPTLC method for the estimation of alkaloids of Cinchona officinalis stems bark and its marketed formulations, Planta Med. 67 (2001) 294–296.
[4] K. Dhalwal, Y.S. Biradar, M. Rajani, TLC densitometric method for simultaneous quantification of phyllanthin, hypophyllanthin, gallic acid and ellagic acid in Phyllanthus amarus using HPTLC, J. AOAC Inter. 89 (2006) 619–623.
[5] Quality Standards of Indian Medicinal Plants, Vol. I, Indian Council of Medical Research, New Delhi, 2003, pp. 10–50.
[6] S.J. Sheu, K.L. Li, Liquid chromatographic determination of the constituents in Shao-yao-tang and related Chinese herbal preparations, J. High Resol. Chromatogr. 21 (1998) 569–573.
[7] The Ayurvedic Formulary of India. Part II. first ed., Government of India, Ministry of Health and Family Welfare, New Delhi, 2000, p. 128.
[8] The Ayurvedic Pharmacopoeia of India, Vol. I, first ed., Government of India, Ministry of Health and Family Welfare, New Delhi, 2001, pp. 47–48, 55, 123–124.
[9] Quality Standard of Indian Medicinal Plants, Vol. IV, Indian Council of Medicinal Research, New Delhi, 2006, pp. 130–136.
[10] P.K. Mukherjee, Quality Control of Herbal Drugs, Business Horizons, New Delhi, 2005, pp. 741–743.
[11] Quality Standard of Indian Medicinal Plants, Vol. I, Indian Council of Medicinal Research, New Delhi, 2003, pp. 205–211.
[12] Indian Herbal Pharmacopoeia, Indian Drug Manufacturer’s association, Mumbai, 2002, pp. 206–213.
[13] M.L. Khorana, D.K. Motiwala, Anthelmentic activity of Kamala and its constituents, Indian J. Pharm. 11 (1949) 37–43.
[14] Guideline on Validation of Analytical Procedure-Methodology. International Conference on Harmonization, Geneva, Switzerland, 1996.