Quality control and quantitation of catechin and epicatechin from leaves of *Maytenus rigida* Mart

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**Abstract:** *Maytenus rigida* Mart belongs to the family Celastraceae, formed by 106 genera and 1,300 species, that are widely distributed in tropical and subtropical regions. The species is distributed in Caatinga region of northeastern Brazil in the states of Sergipe, Paraíba and Pernambuco. *Maytenus rigida* is popularly known as bom nome or chapéu de couro. It is used in folk medicine for gastritis and gastric ulcer. The main secondary metabolites reported for this species are pentacyclic triterpenes, flavonoids and alkaloids. This work aimed to evaluate the quality control of raw material and the quantification of catechin and epicatechin by High Performance Liquid Chromatography (HPLC) in *Maytenus rigida* crude ethanolic extract (CEE) obtained from leaves. The plant material was collected in the municipal district of Boa Vista - PB in June of 2018. Physicochemical assays were developed with leaves: Determination of loss on drying, total ashes, limit assay for heavy metals, bulk density, following the recommendations of the 5th Brazilian Pharmacopoeia. The content of total phenolic compounds was determined using the Folin-Ciocalteu method and total flavonoids was measured using the colorimetric method with AlCl\(_3\) with CEE. The quantitation of (+)-Catechin and (-)-Epicatechin was performed with a validated method previously developed. *M. rigida* leaves presented 5.9% ± 0.143 of water content, 5.10% ± 0.136 of total ashes, < 10 ppm of heavy metals and 0.40 g/mL of bulk density. The total phenolic content was 187.52 ± 2.01 µg EAG/mg and the flavonoid content was 88.80 ± 1.47 µg EQ/mg. The evaluation of CEE showed the presence of 7.79 µg of (+)-Catechin/mg and 6.04 µg of (-)-Epicatechin/mg. The quality control analyzes performed with *M. rigida* established a better profile for the characterization of the drug. This study confirms the applicability of method for the quantification of phenolic compounds in extracts of *Maytenus rigida*.

**Keywords:** Maytenus rigida; quality control; catechins

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1. Introduction

*Maytenus rigida* Mart., Celastraceae, is an evergreen tree that is endemic to the Caatinga region from Northeastern Brazil and is found in the states of Sergipe, Paraiba and Pernambuco. This species is popularly known as “bom-nome” and “chapéu-de-couro” and is used in folk medicine as analgesic, anti-inflammatory, healing and for the treatment of gastritis and gastric ulcer (Santos et al., 2011).

There are many reports about the phytochemical composition of *M. rigida* leaves and barks, especially catechin, epicatechin, terpenoids and proanthocyanidins (Almeida et al., 2005; Estevam, 2006) and pharmacological activities, such as antimicrobial, antinociceptive, antioxidant, antiulcerogenic and antidiarrheal (Estevam, 2006; Santos, 2008, Santos et al., 2011).

One important aspect for the efficiency of the treatment is the quality control of herbal medicine products and the determination of its chemical markers (Marques et al., 2013).

The aim of this work was to evaluate the quality control of raw material and the quantification of chemical markers by HPLC from *M. rigida* crude ethanolic extract (CEE) obtained from its leaves.

2. Results and Discussion

The determination of loss on drying and water content are important to evaluate the quality of vegetal raw materials, as moisture content favors the development of microorganisms, as well as enzymatic activities. The values found for the plant drug *M. rigida* (5.9% ± 0.143) are in accordance with the specified limits in Brazilian Pharmacopoeia (Brazil, 2010).

The second parameter evaluated was the total ashes content, this evaluation aims to determine the content of inorganic impurities and constituents contained in organic substances. The raw material of *M. rigida* presented (5.10% ± 0.136) of total ashes, content below the specified in the compendium (8 - 14%).

Bulk density consists in an important parameter to the extraction process control of vegetable raw materials, the results obtained for dry material in study was 0.40 g/mL. While there are no reference parameters for this species, the values found might be considered normal.

The proximity of highways as well as water quality by pollutants that are conducted by rainwater, can affect plant (Freire, 2005). The limit assay for heavy metals provided satisfactory results (below 10 ppm).

The total phenolic content was 187.52 ± 2.01 µg EAG/mg and the flavonoid content was 88.80 ± 1.47 µg EQ/mg. These results confirm the abundance of the polyphenolic content found in the species and demonstrate its pharmacological potential, since these metabolites are mainly related to antimicrobial, antiulcerogenic and antioxidant activities (Santos, 2008; Santos et al., 2011). The evaluation of CEE showed the presence of 7.79 µg of (+)-Catechin/mg and 6.04 µg of (-)-Epicatechin/mg, these are important results considering that flavan-3-ols may be involved in the anti-inflammatory and gastroprotective process (Santos, 2008).

3. Materials and Methods

The leaves of *Maytenus rigida* were collected in the city of Boa Vista, Paraiba, Brazil, in June of 2018. The plant materials were dried at temperature of 40 °C, for 96 hours, and then powdered in a mill.

**Physicochemical characterization of the plant material**

The physicochemical assays were developed: determination of loss on drying, total ashes, limit assay for heavy metals and bulk density in accordance with the 5th Brazilian Pharmacopoeia (Brazil, 2010).

**Obtaining plant extracts**

The leaves of *Maytenus rigida* were submitted to an extraction process by maceration method, with ethanol 95%. Three extractions were developed, replacing the solvent every 72 hours. The extraction solution obtained was submitted to rotary evaporator at an average temperature of 40 °C.

**Determination of total phenolic compounds and total flavonoids**

The content of total phenolic compounds (TPC) was measured based on the method described by Singleton et al (1999). The total
flavonoids (TF) were determined using the colorimetric method by metallic complexation described by Schmidt and Ortega (1983). All samples were performed in triplicates.

Quantitation of (+)-Catechin and (-)-Epicatechin

The markers were quantified in CEE of *M. rigida* by HPLC coupled with diode array detection (HPLC-DAD). Analytical separation was performed on a Prominence Shimadzu LC-20AT quaternary pump, a photodiode array detector SPD-M20 A and a Shim-pack® column C18 (150 x 4.6 mm x 5 µm), with pre-column Shim-pack G® ODS (4 x 3.0 mm x 5 µm). HPLC data acquisition was performed by LC Solution software. The optimized analytical separations of (+)-Catechin and (-)-Epicatechin were carried out using a mobile phase of 0.1% phosphoric acid in water (A) and acetonitrile (B) with the following method: 0–30 min: 5 to 25% of B; 30–40 min: 25 to 100% of B; 40-50 min: 100% of B; 50-55 min: 100 to 5% of B; 55–60 min: 5% of B. A flow rate of 1.0 mL/min at 40 °C and an injection volume of 20 µL were employed. The UV spectra were recorded at 280 nm. (+)-Catechin and (-)-Epicatechin standards (> 99%) were purchased by Phytolab®. The markers were quantified by the equation obtained by linear regression. (+)-Catequin y = 7426.2x + 607.0 (r = 0.9999) and (-)-Epicatechin y = 17877x – 10588 (r = 0.9997).

4. Conclusions

The quality control analyzes performed with *M. rigida* established a better profile for the characterization of the plant material. The results demonstrated that the evaluated parameters were in accordance with the 5th Brazilian Pharmacopoeia specifications. The TPC and TF contents supported its applicability in folk medicine. The previously validated HPLC method for quantitation of (+)-Catechin and (-)-Epicatechin is suitable to measure the markers and can be used on another species of *Maytenus* genera.

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Conflicts of Interest

The authors declare no conflict of interest.

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