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*Full Length Research Paper*

## **DPPH radical scavenging activity of extracts from *Rhamnus prinoides***

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Hexane, chloroform, ethyl acetate and methanolic extracts from leaves and stem-bark of *Rhamnus prinoides* were evaluated for their antioxidant activity by DPPH radical scavenging assay. The leaves extracts showed scavenging activity ranging from  $03.33 \pm 0.89$  to  $55.03 \pm 3.40 \mu\text{g mL}^{-1}$  while the stem-bark extracts showed relatively strong scavenging activity ranging from  $03.65 \pm 1.02$  to  $59.55 \pm 2.27 \mu\text{g mL}^{-1}$ . The  $\text{IC}_{50}$  values of *R. prinoides* hexane leaves extract (RPHELs), *R. prinoides* chloroform leaves extract (RPCHLS), *R. prinoides* ethyl acetate leaves extract (RPEALS), *R. prinoides* methanolic leaves extract (RPMELS), *R. prinoides* hexane stem-bark extract (RPHEsB), *R. prinoides* chloroform stem-bark extract (RPCHsB), *R. prinoides* ethyl acetate stem-bark extract (RPEASB) and *R. prinoides* methanolic stem-bark extract (RPMESB) were found to be  $>3000$ ,  $>3000$ ,  $>3000$ ,  $950.42$ ,  $\sim 1500$ ,  $710.50$ ,  $\sim 1000$  and  $902.78 \mu\text{g mL}^{-1}$ , respectively. The positive control ascorbic acid showed an  $\text{IC}_{50}$  value of  $<200 \mu\text{g mL}^{-1}$ . From this study, we concluded that the extracts from *R. prinoides* showed promising antioxidant activity. *R. prinoides* finds therapeutic applications in the traditional medicine. Further research is required to commercialize products from this plant.

**Key words:** Antioxidant, ascorbic acid, *Rhamnus prinoides*, radical scavenging assay, methanolic extract, chloroform extract, hexane extract, ethyl acetate extract.

### **INTRODUCTION**

*Rhamnus prinoides* belongs to the Rhamnaceae family (Dale, 2000; Dlamini and Turner, 2002). *R. prinoides* is also known by other names such as African Dogwood, Glossy-leaf and miffi. *R. prinoides* is widely distributed in East and South African countries (Alemu et al., 2007; Abegaz et al., 1999) which include Ethiopia, Botswana, Eritrea, Lesotho, Namibia, South Africa, Swaziland,

Uganda and Kenya (Ashine, 2015). *R. prinoides* grows up to 4.5-m height and found in evergreen forests, in the wild and along water streams. Although, *R. prinoides* is a slow growing plant in low rainfall areas, it can grow 1 m per annum in wet areas (Ferede et al., 2018). The leaves begin with pale green and turn to dark and shiny on maturation. The roundish red berries attract bees and

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domestic fowl. *R. prinoides* casts a very deep shade such that it will not allow other plants to grow around it. *R. prinoides* flowers towards the end of the year and fruiting occurs at the beginning of the year. *R. prinoides* finds therapeutic applications in the traditional medicine. The decoction of roots has been used to treat pulmonary tuberculosis, pneumonia, bladder and kidney problems (Maliehe, 1997; Van Wyk and Moteetee, 2011). The bark has been used to induce vomiting. An extract of the root together with the bark of *Erythrina tomentosa* has been used to relief colic. The leaves have been applied as a liniment to simple sprains. *R. prinoides* has been used to provide a special aroma and flavor (Shale and Gashe, 1991; Abegaz et al., 1999). *R. prinoides* has also been used in the beer industry as a hopping agent. Geshoidin, a naphthalenic glycoside, present in the stem-bark is responsible for providing bitterness in alcoholic beverages (Nindi et al., 1999). The antioxidant activity of 97% ethanolic extracts from leaves of *R. prinoides* has been reported previously (Ashine, 2015). Methanolic and aqueous extracts from roots of *R. prinoides* have also evaluated for their DPPH radical scavenging activity (Kimondo et al., 2019). However, to the best of our knowledge, the scavenging activity of hexane, chloroform, ethyl acetate and methanolic extracts of the leaves and stem-bark of *R. prinoides* has not been reported previously, particularly the plant species gathered from the Kingdom of Lesotho. The aim of the present study was to evaluate the antioxidant activity of these extracts by DPPH radical scavenging assay and to determine their IC<sub>50</sub> values. The results obtained are communicated in this article.

## MATERIALS AND METHODS

### Plant materials

The leaves and stem-bark of *R. prinoides* were collected from the foothills of Popa and Popanyane Mountains at Mokhokhong village, Roma, Maseru district, the Kingdom of Lesotho, Southern Africa in January 2019. A voucher specimen viz. Santi/RPLS/2019 for leaves and Santi/RPSB/2019 for stem-bark were kept separately in the Organic Research Laboratory, Department of Chemistry and Chemical Technology, Faculty of Science and Technology, National University of Lesotho, Roma Campus, Maseru district, Kingdom of Lesotho, Southern Africa.

### Processing of materials

The leaves were allowed to air dry at room temperature for two weeks and then ground into powder using a commercial blender (Waring Blender, Blender 80119, Model HGB2WT93, 240V AC, 50-80 Hz, 3.6AMPS, Laboratory and Analytical Supplies). The chopped stem-barks were allowed to air dry at room temperature for two weeks and ground into powder using the blender.

### Preparation of plant extracts

The powdered leaves (300.043 g) of *R. prinoides* were extracted

with methanol under cold conditions for 3 days. The solution was filtered using a filter paper (Boeco, Germany). The solvent was removed by *vacuo* and the same procedure was repeated once again. Finally, the plant material was extracted with hot methanol. 40.1858 g of combined methanol extract was obtained after removal of solvent. The same extraction procedure was followed to get hexane (3.2274 g), chloroform (10.6285 g) and ethyl acetate (11.4763 g) extracts from 300.254, 300.131 and 299.921 g of powdered leaves, respectively. The powdered stem-bark (299.530 g) of *R. prinoides* was extracted first with methanol at room temperature for 3 days followed by a reflux condition for 6 h. 32.2047 g of combined crude methanol extract was obtained after removal of solvent. The same extraction procedures were followed to get hexane (2.5895 g), chloroform (5.4327 g) and ethyl acetate (8.1493 g) extracts from 300.014, 300.157, and 300.422 g of powdered stem-bark, respectively.

### Chemicals and solvents used

Ascorbic acid, DPPH, hexane (AR grade, 99.5%), chloroform (AR grade, 99.5%), ethyl acetate (AR grade, 99.5%) and methanol (AR grade, 99.5%) were all purchased from Sigma-Aldrich.

### Antioxidant activity

The antioxidant activity of the extracts was carried out using 1,1-diphenyl-2-picrylhydrazyl (DPPH) as described in literature (Kim et al., 2002). Briefly, stock solution of the methanolic extract was prepared at a concentration of 3.0 mg of extract in 1 mL of 50% methanol (v/v). Serial dilutions were made from this stock solution to obtain solutions with concentrations of 3000, 2000, 1500, 1000, 800, 500 and 200 µg mL<sup>-1</sup>. Solutions without extract concentration served as negative control. A solution of 3.94 mg of DPPH in 100 mL of methanol served as oxidant which was prepared just before use and stored in dark to minimize degradation. 0.1 mL sample of plant extract solution was mixed with 1.0 mL of 0.1 mM DPPH solution and 0.45 mL of 50 mM Tris-HCL buffer (pH 7.40). Similarly, stock solutions of hexane, chloroform and ethyl acetate extracts were prepared at a concentration of 3.0 mg of extract in 1 mL of 50% methanol (v/v). Further dilutions were made from these stock solutions to obtain solutions with concentrations of 3000, 2000, 1500, 1000, 800, 500, 200 and 0 µg mL<sup>-1</sup>. 0.1 mL each of extract was mixed separately with 1.0 mL of 0.1 mM DPPH solution and 0.45 mL of 50 mM Tris-HCL buffer (pH 7.40). A stock solution of ascorbic acid (0.3 g) in 50% methanol (v/v) was prepared and serial dilutions were made as previously and served as positive control. 0.1 mL was mixed with 1.0 mL of 0.1 mM DPPH solution and 0.45 mL of 50 mM Tris-HCL buffer (pH 7.40). The mixtures were incubated for 30 min and their optical density was measured at 517 nm. Percentage inhibition of DPPH free radical was calculated using the equation:

$$\text{DPPH Scavenged (\%)} = [(A_{\text{cont}} - A_{\text{test}})/A_{\text{cont}}] \times 100$$

where  $A_{\text{test}}$  = Absorbance in the presence of extract or positive control and  $A_{\text{cont}}$  = Absorbance of negative control.

The IC<sub>50</sub> value is defined as the concentration (in µg mL<sup>-1</sup>) of extract that inhibits the formation of DPPH radical by 50% (Moyo et al., 2013; Ndhala et al., 2013). A lower value of IC<sub>50</sub> represents higher antioxidant activity. The IC<sub>50</sub> values were calculated from graphs by plotting extract concentrations vs. percentage inhibition of DPPH radical using Microsoft Excel. Each experiment was carried out in triplicate and the averages of the three values were used to calculate IC<sub>50</sub> values. Standard deviation was calculated for each concentration from the three values of the experiment.

### Statistical analysis

Results were expressed as means of three determinations. One-way analysis of variance (ANOVA) was used to compare means at the significance level  $p < 0.05$ . All analysis was performed by Microsoft Excel software.

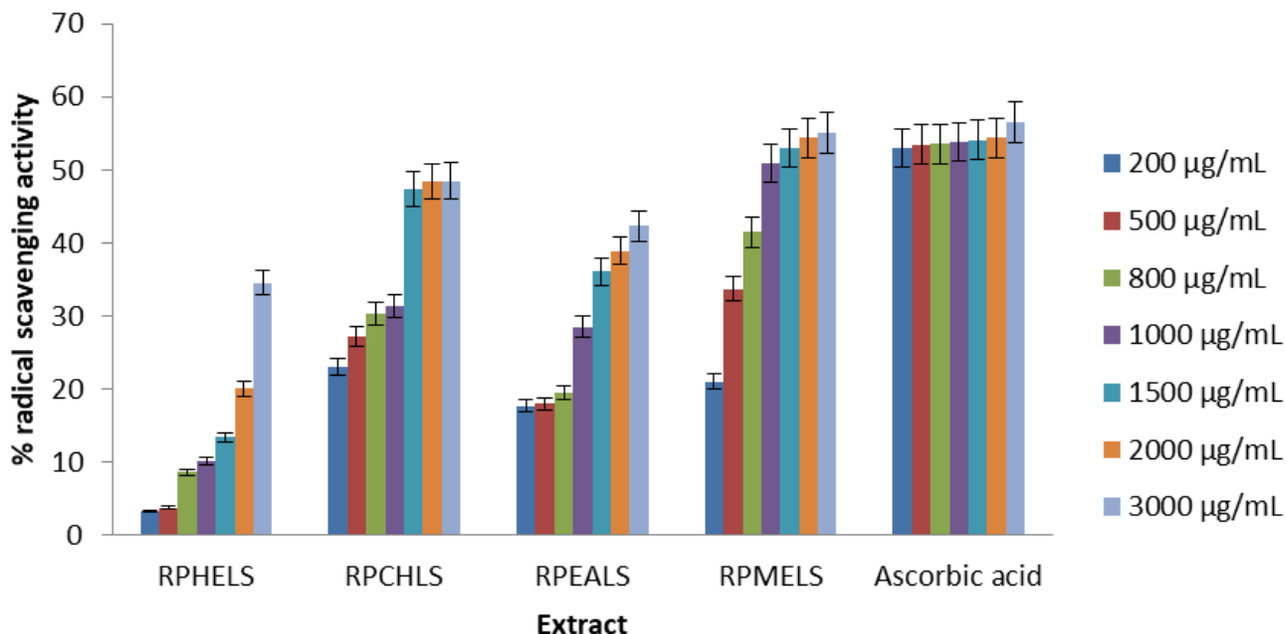
## RESULTS AND DISCUSSION

Table 1 summarizes the DPPH radical scavenging activity of hexane, chloroform, ethyl acetate and methanolic extracts of the leaves and stem-bark of *R. prinoides*. *R. prinoides* hexane leaves extract (RPHELS) showed  $3.33 \pm 0.89$ ,  $3.80 \pm 2.60$ ,  $8.61 \pm 1.39$ ,  $10.12 \pm 0.84$ ,  $13.39 \pm 2.94$ ,  $20.10 \pm 3.23$  and  $34.56 \pm 6.51\%$  of scavenging activity at concentrations 200, 500, 800, 1000, 1500, 2000 and  $3000 \mu\text{g mL}^{-1}$ , respectively. The positive control, ascorbic acid, showed  $53.01 \pm 3.98$ ,  $53.46 \pm 0.14$ ,  $53.51 \pm 0.77$ ,  $53.82 \pm 0.54$ ,  $54.12 \pm 1.64$ ,  $54.34 \pm 0.92$  and  $56.45 \pm 5.45\%$  of scavenging activity at concentrations 200, 500, 800, 1000, 1500, 2000 and  $3000 \mu\text{g mL}^{-1}$ , respectively. These results revealed that RPHELS showed very weak radical scavenging activity relative to positive control at all concentrations. *R. prinoides* chloroform leaves extracts (RPCHLS) showed  $23.01 \pm 3.44$ ,  $27.15 \pm 5.18$ ,  $30.35 \pm 1.02$ ,  $31.38 \pm 0.11$ ,  $47.37 \pm 4.14$ ,  $48.38 \pm 4.15$  and  $48.49 \pm 3.17\%$  of scavenging activity at concentrations 200, 500, 800, 1000, 1500, 2000 and  $3000 \mu\text{g mL}^{-1}$ , respectively. This result showed that RPCHLS has lower activity than positive control at all concentrations. *R. prinoides* ethyl acetate leaves extract (RPEALS) showed  $17.71 \pm 1.02$ ,  $17.96 \pm 0.82$ ,  $19.48 \pm 1.50$ ,  $28.51 \pm 0.62$ ,  $36.05 \pm 6.26$ ,  $38.94 \pm 5.75$  and  $42.33 \pm 5.88\%$  of scavenging activity at 200, 500, 800, 1000, 1500, 2000 and  $3000 \mu\text{g mL}^{-1}$ , respectively. This result revealed that RPEALS exhibited weak activity at low concentrations relative to positive control. However, at high concentration of  $3000 \mu\text{g/mL}$ , it showed higher scavenging activity of  $42.33 \pm 5.88\%$ . *R. prinoides* methanolic leaves extract (RPMELS) showed  $21.01 \pm 3.80$ ,  $33.74 \pm 1.42$ ,  $41.50 \pm 2.73$ ,  $50.93 \pm 5.27$ ,  $52.93 \pm 2.88$ ,  $54.34 \pm 2.70$  and  $55.03 \pm 3.40\%$  of scavenging activity at concentrations 200, 500, 800, 1000, 1500, 2000 and  $3000 \mu\text{g mL}^{-1}$ , respectively. This result revealed that RPMELS exhibited weak activity at low concentrations relative to positive control. However, RPMELS has comparable activity as that of positive control at high concentrations.

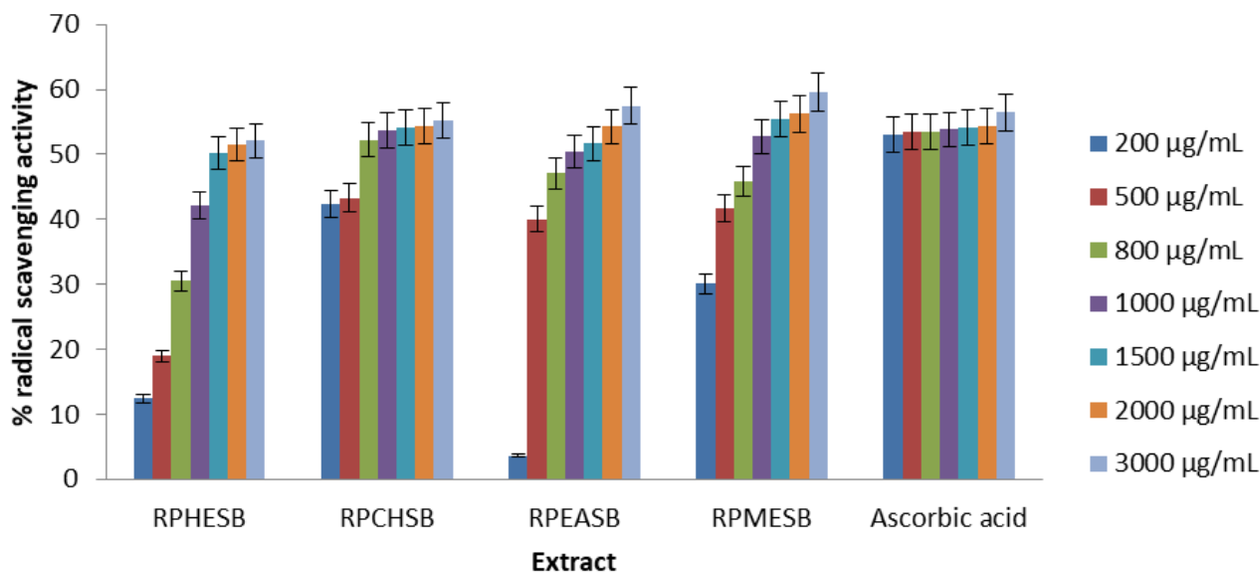
*R. prinoides* hexane stem-bark extract (RPHEBS) showed  $12.39 \pm 3.19$ ,  $18.93 \pm 1.04$ ,  $30.47 \pm 0.82$ ,  $42.15 \pm 4.06$ ,  $50.24 \pm 1.50$ ,  $51.52 \pm 2.47$  and  $52.15 \pm 1.06\%$  of scavenging activity at concentrations 200, 500, 800, 1000, 1500, 2000 and  $3000 \mu\text{g mL}^{-1}$ , respectively. This result revealed that RPHEBS exhibited weak activity at low concentrations relative to positive control. However, at higher concentrations such as 1500, 2000 and  $3000 \mu\text{g/mL}$ , RPHEBS exhibited comparable activity as that of positive control. *R. prinoides* chloroform stem-bark extract

(RPCHSB) showed  $42.37 \pm 5.65$ ,  $43.30 \pm 2.98$ ,  $52.23 \pm 2.46$ ,  $53.64 \pm 4.42$ ,  $54.06 \pm 1.41$ ,  $54.41 \pm 2.17$  and  $55.22 \pm 2.48\%$  of scavenging activity at concentrations 200, 500, 800, 1000, 1500, 2000 and  $3000 \mu\text{g mL}^{-1}$ , respectively. Thus, RPCHSB exhibited comparable activity as that of positive control at all concentrations except at concentrations of 200 and  $500 \mu\text{g mL}^{-1}$ . RPCHSB showed  $42.37 \pm 5.65$  and  $43.30 \pm 2.98\%$  of scavenging at concentrations 200 and  $500 \mu\text{g/mL}$ , respectively, while the positive control showed  $53.01 \pm 3.98$  and  $53.46 \pm 0.14\%$  of scavenging activity at concentrations 200 and  $500 \mu\text{g mL}^{-1}$ , respectively. *R. prinoides* ethyl acetate stem-bark extract (RPEASB) showed  $3.65 \pm 1.02$ ,  $40.04 \pm 1.50$ ,  $47.09 \pm 4.36$ ,  $50.31 \pm 7.80$ ,  $51.61 \pm 2.27$ ,  $54.23 \pm 1.83$  and  $57.43 \pm 3.28\%$  of scavenging activity at concentrations 200, 500, 800, 1000, 1500, 2000 and  $3000 \mu\text{g/mL}$ , respectively. Thus, RPEASB exhibited very weak activity relative to positive control at low concentration of  $200 \mu\text{g mL}^{-1}$ . *R. prinoides* methanolic stem-bark extract (RPMESB) showed  $30.09 \pm 5.26$ ,  $41.69 \pm 2.27$ ,  $45.80 \pm 2.54$ ,  $52.78 \pm 6.43$ ,  $55.37 \pm 3.90$ ,  $56.19 \pm 3.58$  and  $59.55 \pm 2.27\%$  of scavenging activity at concentrations 200, 500, 800, 1000, 1500, 2000 and  $3000 \mu\text{g mL}^{-1}$ , respectively. This result revealed that RPMESB exhibited weak activity at low concentrations relative to positive control. However, at high concentrations such as 2000 and  $3000 \mu\text{g/mL}$ , it showed higher scavenging activity of  $56.19 \pm 3.58$  and  $59.55 \pm 2.27\%$ , respectively. Among the extracts (RPHELS, RPCHLS, RPEALS, RPMELS, RPHEBS, RPCHSB, RPEASB and RPMESB) from *R. prinoides*, RPMESB showed the highest scavenging activity (Table 1). For comparison and clarity, the percentage of scavenging activity of these extracts at various concentrations are as shown in Figures 1 and 2. The  $\text{IC}_{50}$  values of hexane, chloroform, ethyl acetate and methanolic extracts of the leaves and stem-bark of *R. prinoides* are shown in Table 2. RPHELS, RPCHLS, RPEALS, RPMELS, RPHEBS, RPCHSB, RPEASB and RPMESB exhibited  $\text{IC}_{50}$  values of  $>3000$ ,  $>3000$ ,  $>3000$ ,  $950.42$ ,  $\sim 1500$ ,  $710.50$ – $1000$  and  $902.78 \mu\text{g mL}^{-1}$ , respectively. RPCHSB is the most potent with  $\text{IC}_{50}$  value of  $710.50 \mu\text{g mL}^{-1}$ . The positive control ascorbic acid showed an  $\text{IC}_{50}$  value of  $<200 \mu\text{g mL}^{-1}$ .

The DPPH radical scavenging activity of 97% ethanolic extract from leaves of *R. prinoides* has previously been reported and its maximum radical scavenging was found to be 81.148% at a concentration of  $24 \text{ mg mL}^{-1}$  (Ashine, 2015) and the  $\text{IC}_{50}$  value was determined to be  $5.2 \text{ mg mL}^{-1}$ . The positive control, ascorbic acid, showed 93.77% scavenging activity with an  $\text{IC}_{50}$  value of  $0.24 \text{ mg mL}^{-1}$  in the same assay (Ashine, 2015). Therefore, when compared with the present study on the hexane, chloroform, ethyl acetate and methanolic extracts from leaves of *R. prinoides*, this 97% ethanolic extract from leaves of *R. prinoides* showed higher radical scavenging activity and lower  $\text{IC}_{50}$  value. This 97% ethanol might have more extractive power of active constituents than



**Figure 1.** Percentage of radical scavenging of leaves extracts and ascorbic acid at various concentrations. RPHELS: *R. Prinoides* hexane leaves extract; RPCHLS: *R. prinoides* chloroform leaves extract; RPEALS: *R. prinoides* ethyl acetate leaves extract; RPMELS: *R. prinoides* methanolic leaves extract; Ascorbic acid in 50% methanol served as positive control.



**Figure 2.** Percentage of radical scavenging of stem-bark extracts and ascorbic acid at various concentrations. RPHEBS: *R. prinoides* hexane stem-bark extract; RPCHSB: *R. prinoides* chloroform stem-bark extract; RPEASB: *R. prinoides* ethyl acetate stem-bark extract; RPMESB: *R. prinoides* methanolic stem-bark extract; Ascorbic acid in 50% methanol served as positive control.

the solvents used in the present study. Additionally, the collection of plant materials at different geographic locations will also play a vital role in determining the

active constituents of extracts. Methanolic and aqueous roots extracts from *R. prinoides* have also been screened for their DPPH radical scavenging activity. Their  $IC_{50}$

**Table 1.** Percentage of radical scavenging activity of extracts from *R. prinoides* at various concentrations.

Extract	Concentrations ( $\mu\text{g mL}^{-1}$ )/ Percentage of inhibition						
	200	500	800	1000	1500	2000	3000
RPHELs	03.33 $\pm$ 0.89	03.80 $\pm$ 0.26	08.61 $\pm$ 1.39	10.12 $\pm$ 0.84	13.39 $\pm$ 2.94	20.10 $\pm$ 3.23	34.56 $\pm$ 6.51
RPCHLS	23.01 $\pm$ 3.44	27.15 $\pm$ 5.18	30.35 $\pm$ 1.02	31.38 $\pm$ 0.11	47.37 $\pm$ 4.14	48.38 $\pm$ 4.15	48.49 $\pm$ 3.17
RPEALS	17.71 $\pm$ 1.02	17.96 $\pm$ 0.82	19.48 $\pm$ 1.50	28.51 $\pm$ 0.62	36.05 $\pm$ 6.26	38.94 $\pm$ 5.75	42.33 $\pm$ 5.88
RPMELS	21.01 $\pm$ 3.80	33.74 $\pm$ 1.42	41.50 $\pm$ 2.73	50.93 $\pm$ 5.27	52.93 $\pm$ 2.88	54.34 $\pm$ 2.70	55.03 $\pm$ 3.40
RPHEsB	12.39 $\pm$ 3.19	18.93 $\pm$ 1.04	30.47 $\pm$ 0.82	42.15 $\pm$ 4.06	50.24 $\pm$ 1.50	51.52 $\pm$ 2.74	52.15 $\pm$ 1.06
RPCHsB	42.37 $\pm$ 5.65	43.30 $\pm$ 2.98	52.23 $\pm$ 2.46	53.64 $\pm$ 4.42	54.06 $\pm$ 1.41	54.41 $\pm$ 2.17	55.22 $\pm$ 2.48
RPEAsB	03.65 $\pm$ 1.02	40.04 $\pm$ 1.50	47.09 $\pm$ 4.36	50.31 $\pm$ 7.80	51.61 $\pm$ 2.27	54.23 $\pm$ 1.83	57.43 $\pm$ 3.28
RPMEsB	30.09 $\pm$ 5.26	41.69 $\pm$ 2.27	45.80 $\pm$ 2.54	52.78 $\pm$ 6.43	55.37 $\pm$ 3.90	56.19 $\pm$ 3.58	59.55 $\pm$ 2.27
Asc. acid	53.01 $\pm$ 3.98	53.46 $\pm$ 0.14	53.51 $\pm$ 0.77	53.82 $\pm$ 0.53	54.12 $\pm$ 1.64	54.34 $\pm$ 0.92	56.45 $\pm$ 5.45

RPHELs: *R. prinoides* hexane leaves extract; RPCHLS: *R. prinoides* chloroform leaves extract; RPEALS: *R. prinoides* ethyl acetate leaves extract; RPMELS: *R. prinoides* methanolic leaves extract; RPHEsB: *R. prinoides* hexane stem-bark extract; RPCHsB: *R. prinoides* chloroform stem-bark extract; RPEAsB: *R. prinoides* ethyl acetate stem-bark extract; RPMEsB: *R. prinoides* methanolic stem-bark extract; Asc. Acid = Ascorbic acid in 50 % methanol served as positive control. All experiments were conducted in triplicate (n=3) and reported as the mean of three values together with standard deviation,  $\pm$ SD.

**Table 2.** The IC<sub>50</sub> values of various extracts of *R. prinoides* by DPPH radical scavenging assay.

S/N	Extract	IC <sub>50</sub> ( $\mu\text{g mL}^{-1}$ )
1	RPHELs	>3000
2	RPCHLS	>3000
3	RPEALS	>3000
4	RPMELS	950.23
5	RPHEsB	~1500
6	RPCHsB	710.50
7	RPEAsB	~1000
8	RPMEsB	902.78
9	Asc. acid	<200

RPHELs: *R. prinoides* hexane leaves extract; RPCHLS: *R. prinoides* chloroform leaves extract; RPEALS: *R. prinoides* ethyl acetate leaves extract; RPMELS: *R. prinoides* methanolic leaves extract; RPHEsB: *R. prinoides* hexane stem-bark extract; RPCHsB: *R. prinoides* chloroform stem-bark extract; RPEAsB: *R. prinoides* ethyl acetate stem-bark extract; RPMEsB: *R. prinoides* methanolic stem-bark extract; Asc. Acid = Ascorbic acid in 50 % methanol served as positive control. All experiments were conducted in triplicate (n=3) and reported as the mean of three values together with standard deviation,  $\pm$ SD.

values were found to be 377.27 and  $\sim$ 250  $\mu\text{g mL}^{-1}$ , respectively (Kimondo et al., 2019). The positive control, ascorbic acid, showed an IC<sub>50</sub> value 50.32  $\mu\text{g mL}^{-1}$  in the same assay (Kimondo et al., 2019). Additionally, the kinetics of acetylcholinesterase (AChE) inhibitory activity of aqueous extract from *R. prinoides* has previously been reported (Catherine and Edward, 2009). The IC<sub>50</sub> value for *R. prinoides* was found to be 0.201 mg mL<sup>-1</sup>. The AChE inhibitory activity of *R. prinoides* was found to be higher than that of some Portuguese and Danish medicinal plants (Ferreira et al., 2006; Adersen et al., 2006). Biologically important secondary metabolites such as emodin, physcion, prinoidin, rhamnazin, geshodin and

many other emodin-derived compounds have been reported from *R. prinoides* (Van Staden and Drewes, 1994; Abegaz and Kebete, 1995). Alkaloids, flavonoids, terpenoids, anthraquinones, saponins, polyphenols, etc., classes of compounds have also been reported from various extracts of *R. prinoides* (Molla et al., 2016).

## Conclusion

DPPH radical scavenging activity of hexane, chloroform, ethyl acetate and methanolic extracts from leaves and stem-bark of *R. prinoides* collected from the Kingdom of

Lesotho have been evaluated. The leaves extracts showed scavenging activity ranging from  $03.33 \pm 0.89$  to  $55.03 \pm 3.40 \mu\text{g mL}^{-1}$  while the stem-bark extracts showed relatively strong scavenging activity ranging from  $03.65 \pm 1.02$  to  $59.55 \pm 2.27 \mu\text{g mL}^{-1}$ . The  $\text{IC}_{50}$  values of these extracts were also determined and were found to be in the range of 710.50 to  $>3000 \mu\text{g mL}^{-1}$ . *R. prinoides* finds therapeutic applications in the traditional medicine and showed promising antioxidant activity. Therefore, further studies will be useful to commercialize products from this plant.

## CONFLICT OF INTERESTS

The authors declared no conflict of interests.

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