# Antiviral Activity of Three Plant Species, Rhus coriaria, Chenopodium quinoa, and Ailanthus altissima against Tobacco Mosaic Virus

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## ABSTRACT

Research on natural compounds provides new alternatives for effective and sustainable control of plant viral pathogens. Herein, we prepared and investigated the *in vitro* antiviral activity of 60 plant species from 22 families. The hydroethanolic extracts of *Rhus coriaria*, *Chenopodium quinoa* and *Ailanthus altissima* have strong inhibitions on *Tobacco Mosaic Virus* (TMV) infection. Hydroethanolic extract of *C. quinoa* with half-maximal Effective Concentration ( $EC_{50}$ ) value of 1.64 mg mL<sup>-1</sup> exhibited the highest inhibitory effect against TMV. The extracts of *R. coriaria* and *A. altissima* with  $EC_{50}$  values of 2.82 and 4.42 mg mL<sup>-1</sup>, being compared with *C. quinoa*, showed an anti-TMV activity at higher concentrations, respectively. The systemic assay indicated that all of the three extracts reduced the symptoms and negative effects of TMV on tobacco plants. The chemical analysis of *C. quinoa* extract demonstrated a rich profile of saponins and anthocyanins, while *A. altissima* and *R. coriaria*, and *A. altissima* extracts had significant antiviral activity, and could be used as suitable sources for discovering new antiviral agents.

Keywords: Antiviral agents, Inhibitory effects, Plant viruses, Screening plant extracts.

## **INTRODUCTION**

Plant viruses cause diverse diseases in a wide range of crop plant species and account for a large portion of crop disease epidemics. It is well documented that these pathogens cause significant damages to quantity and/or quality of products in a wide range of crops worldwide. Despite the ambiguity existing about the clear data on the economic impact of plant virus diseases in agriculture, the annual worldwide yield reduction assignable to plant viruses is roughly worth \$30 billion (Sastry and Zitter, 2014). Therefore, development of effective disease management strategies against plant viruses remains a major concern for growers. Common control strategies, including resistant hosts and chemical control of vectors, have limited success in controlling plant viruses (Jones and Naidu, 2019; Ritzenthaler, 2005). Researches on bioactive

natural products provide new alternatives for developing effective and sustainable control of plant virus diseases. Noteworthy, some natural plant products, in addition to the ability of direct interference with viral replication, can also lead to the induction of immune response against viruses. (Ma *et al.*, 2020; Guo *et al.*, 2020). During the last decades, accumulated data have shown several plant products with potent antiviral activity, which implies the potentiality of plant products for the development of effective and sustainable control measures for plant virus diseases (Zhao *et al.*, 2017).

Tobacco Mosaic Virus (TMV), a member of Tobamovirus genus, is one of the most common viral pathogens in plants. Tobamoviruses collectively have a very wide host range and cause significant yield losses in many crops such as solanaceous, brassicas and cucurbits (Jewehan *et al.*,

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2021; Zhang *et al.*, 2016; Alishiri *et al.*, 2013; Choi *et al.*, 2009); Furthermore, various characteristics of TMV, including infection of the host plant by mechanical inoculation and showing local lesion phenomenon in some host plants, have made it particularly amenable as a preferred standard model for screening and evaluating the antiviral activity of numerous compounds (Scholthof, 2004).

Iran has a diverse climate in different geographical areas and provides a unique environment for growing a comprehensive source of plant species. Some of these plant species have been reported to possess antiviral effects, and several of them have been practically used for treating humans and animals infected with several viral diseases (Asadi-Samani et al., 2016; Salehi et al., 2018); Nevertheless, despite the rich biodiversity of unique natural plant species in different Iranian geographical regions, only a few studies have screened and isolated plant-based antiviral compounds (Sanjarian et al., 2021; Biniaz et al., 2021; Gholizadeh, 2019).

Despite the importance of identifying biogenic antiviral compounds with the advantage of high activity and low environmental side-effects, the antiviral effects of plant extracts on plant viruses have been studied in a few cases. In the current study, using ethnomedical approach and literature-based follow-up of the existing studies (Pan *et al.*, 2013), we selected 60 plant species, and evaluated hydroethanolic extracts of their antiviral activity on TMV and characterized extracts with the most antiviral effect.

## MATERIALS AND METHODS

## Virus and Plant Materials

*Tobacco Mosaic Virus* (TMV) Shiraz, Iran, was propagated in *Nicotiana tabacum* L. var. Turkish and was purified as described by Gooding and Hebert (1967). The concentration of the virus was measured using a Nanodrop

(ND-1000 Spectrophotometer) [virus concentration = (A260) /

 $E_{1cm}^{0.1\%,260 \text{ nm}}$ ]. The purified virus was stored at -20°C until use.

The plant materials (used for evaluation of antiviral activity) were collected from the Eram Botanical Garden (Fars Province, Iran), Bamu National Park (Fars Province, Iran), and Dena National Park (Kohgiluyeh and Boyer-Ahmad Province, Iran) in spring and summer 2018, and were identified by the Department of Biology, Faculty of Sciences, Shiraz University, Shiraz, Iran.

## **Preparation of Plant Extracts**

The air-dried plant samples were ground to a fine powder using a grinder. Then, about 10 grams of the powdered leaves were extracted using 80% ethanol solvent at room temperature (24 hours). The extracts were then filtered through gauze and were clarified by centrifugation at 12,000×g for 15 minutes. The supernatant was collected and concentrated using a rotary evaporator (Heidolph. Germany) under reduced pressure. Then, the dried plant extracts were weighed, dissolved in a small volume of Dimethyl Sulfoxide (DMSO), and diluted to 20 mg mL<sup>-1</sup> with distilled water. The DMSO concentration did not exceed 2% (v/v) for any of the solutions (Jing et al., 2012).

#### Local Lesion Assay

Tobacco (*Nicotiana glutinosa*) plants at the 5-6 leaves stage were used to evaluate the plant extracts' antiviral properties against TMV. Each sample (20 mg mL<sup>-1</sup>) was mixed with an equal volume of TMV solution (10  $\mu$ g mL<sup>-1</sup>) and was immediately inoculated on the half leaves of tobacco, whereas the opposite half leaves were mock (distilled water with 2% DMSO) inoculated, using 500-mesh carborundum as abrasive. The antiviral activity of the plant extracts was estimated according to the inhibition rates of local lesions on the leaves of *N. glutinosa*. The local lesions were numbered 3-4 days after inoculation. The MAP30 protein of *Momordica charantia*, a type I ribosome-inactivating protein and an antiviral agent (Moghadam *et al.*, 2016), was used as the positive control. The inhibition rate of viral infection was calculated using the following equation 1 (Verma *et al.*, 1996):

Inhibition rate (%) =  $(1 - T/C) \times 100$  (1)

Where, T is the average mean lesion number of treated half-leaves, and C is the average lesion number of the control halves. Seven replicates of each treatment were randomized on the leaves of test plants. This experiment was performed at least three times.

#### Leaf-Disc Assay

The extracts that showed more than 50% inhibition of viral infection by the local lesion assay on N. glutinosa were further tested with the leaf-disc method. For this method, the growing tobacco (N. tabacum L. var. Turkish) leaves were mechanically inoculated with TMV (10 µg mL<sup>-1</sup>). After 72 h, 12-mm diameter leaf discs that were smooth and thin, and had no major veins, were cut from the leaf surface. The leaf discs were hovered on the solution of each sample  $(10 \text{ mg mL}^{-1})$  in a petri dish and were then incubated for 48 hours in growth chamber with light intensity of 3,000 lx ( $25\pm2^{\circ}C$ ). The discs that were floated on mock were used as the control. After 48 hours, leaf discs were ground in coating buffer, and using antiserum developed in Plant Virology Research Center, Iran, their viral accumulation were assessed by Enzyme-Linked Immunosorbent Assay (ELISA) (Clark and Adams, 1977) using antiserum developed in Plant Virology Research Center, Iran. Measurements were conducted by absorbance Plate Reader (model ELx808LBS, Biotek, USA), and the mean absorbance value (OD 405 nm) of three replicates for each experimental condition was taken. The inhibition rate was calculated

according to Equation (2) (French and Towers, 1992):

Inhibition rate (%) =  $(1 - C/C_0) \times 100$  (2)

Where, C is the virus accumulation in the treated leaf discs, and  $C_0$  is the viral accumulation in the control, using the A405 value of TMV at concentrations of 10, 5, 2.5, 1.25, and 0.625  $\mu$ g mL<sup>-1</sup>. TMV accumulation level was estimated from a standard curve generated.

#### Systemic Assay and Phytotoxicity

Healthy tobacco plants (N. tabacum L. var. 'Turkish') were sprayed with 5 mL (10 mg mL<sup>-1</sup>) of plant extracts, Ailanthus altissima (AAE), Chenopodium quinoa (CQE) and Rhus coriaria (RCE) at the 5-6 leaves stage. After 24 hours, the treated leaves were uniformly inoculated with TMV inoculum  $(10 \ \mu g \ mL^{-1})$  and were kept under greenhouse conditions (25±2°C, 12-hour photoperiods). The experiment involved five CQE+TMV, treatments: AAE+TMV, RCE+TMV, mock (Distilled water with 2% DMSO)+TMV and mock+water (healthy plants). Three plants were used for measuring growth parameters, i.e., height and the fresh weight of aerial parts at 40days post-inoculation (dpi). The experiment was performed three times.

In a phytotoxicity bioassay, ethanol extracts of the three plant species were used at four concentrations (10, 20, 30, and 40 mg mL<sup>-1</sup>) in an experimental greenhouse under 12-hr photoperiod, at  $25\pm2^{\circ}$ C conditions. One week after treatment, the leaves were visually examined for recording chlorotic and necrotic injury levels (Frackowiak *et al.*, 2019).

## Determination of Half Maximal Effective Concentration (EC<sub>50</sub>)

Dose-dependent inhibition analyses were further carried out to examine the antiviral potential of the selected extracts. The hydroethanolic extract of each species was diluted to 20, 15, 10, 5, 4, 2, 1, and 0.5 mg mL<sup>-1</sup> with distilled water containing 2% DMSO: it was mixed with an equal volume of TMV inoculum (10  $\mu$ g mL<sup>-1</sup>) and immediately inoculated on the half leaf of tobacco (*N. glutinosa*). A mixture of DMSO solvent and the inoculum was rubbed on the opposite half leaf as a negative control. The number of developed local lesions was recorded at 3-4 days post-inoculation. The inhibition rate of viral infection was recorded and calculated by equation (1).

### **LC-ESI-MS** Analysis

For the Liquid Chromatography (LC)Electrospray Ionization (ESI) mass Spectrometry (MS) analysis, a Perkin-Elmer API 165 (Norwalk, CT, USA) single quadrupole MS instrument with Turbo-Ion spray interface was applied in the negative and positive ion modes scan spectra (150 to 1000 amu). ESI settings were as follows: temperature 350°C ~ ion spray 4500V, curtain gas 8 psi, and CEM detector 2300V in negative mode. The MS detector was coupled to Hewlett Packard (Palo Alto, CA, USA) 1100 HPLC system consisting of a highpressure mixing pump, autosampler, column oven, and DAD. Online UV spectra were determined at 220-500 nm. Firstly, solutions  $(0.12 \ \mu g \ \mu L^{-1})$  were extracted in formic acid:  $H_2O$ : acetonitrile (1:5:94, v/v), and centrifuged at 13,000×g for 3 minutes. Then, five microliters were injected into LC-MS. Linear gradient elution was carried out using H<sub>2</sub>O with 0.1% formic acid as eluent A and acetonitrile 0.1% formic acid as B with an elution rate of 500  $\mu$ L min<sup>-1</sup>.

#### **Statistical Analysis**

Using the paired Student's t-test at P-value< 0.05, the significant difference between the mean values of treatments and controls were analyzed statistically. One-way ANOVA analysis was used for the virus

effect assessment on growth parameters data. Group means values were compared using the Tukey method (P < 0.05). Statistical analysis was performed using SAS 9.4 statistical software.

## RESULTS

#### **Local Lesion Assay**

The concentration of plant extracts used in this research exhibited different inhibition rates at 10 µg mL<sup>-1</sup> against TMV in the local lesion assay (Table 1). Out of 60 extracts tested, 14 crude extracts had no positive effect on preventing the virus infection; though, the other evaluated hydroethanolic extracts had relative antiviral activity. Among these all, the leaves and flowers of *Ailanthus altissima* (80.1%), seeds of *Chenopodium quinoa* (90.3%), and leaves of *Rhus coriaria* (89.6%) showed higher inhibition rate against TMV infection; whereas MAP30 protein at 100 µg/mL resulted in 73.4% inhibition rate.

### Leaf-Disc Assay

Based on the results of the local-lesion assay, 8 crude extracts were assessed using the leaf-disc assay. Except for Ferula aucheri (-3.55%) and Rydingia persica (Burm.f.) Scheen and V. A. Albert (-.51%), all the other crude extracts had inhibition effects against TMV replication with the inhibition rate ranging from 21.63% to 42.93%, repectively. The extract of Ziziphus spina-christi leaves had the lowest inhibition rate (21.63%), while Ailanthus altissima (AAE, 36.44%). Chenopodium quinoa (CQE, 42.02%), and Rhus coriaria (RCE, 42.93%) extracts had the highest inhibition rate (Table 2).

#### Systemic Assay

Our results showed that three extracts could reduce the negative impact (including,

Family	Plant name	Location GPS	Part of plant used	Inhibition rate (%)
Amaranthaceae	Amaranthus blitoides S.Watson	293650 N, 524215 E	AP	37.4
Anacardiaceae	Rhus coriaria L.	293813 N, 523127 E	L	89.6
Apiaceae	Echinophora platyloba DC.	293604 N, 524505 E	AP	8.5
Apiaceae	Foeniculum vulgare Mill.	293813 N, 523127 E	S	42.4
Apiaceae	Ferulago angulata	305214 N, 513021 E	ĀP	NI
Apiaceae	Ferula aucheri	310810 N, 505516 E	L	72.2
Apiaceae	Prangos ferulacea (L.) Lindl.	304133 N, 513827 E	AP	19.0
Apiaceae	Trachyspermum ammi (L.) Sprague	293813 N, 523127 E	S	NI
Apocynaceae	Vinca major L.	293813 N, 523127 E	L	7.9
Araceae	Arum rupicola Boiss.	304138 N, 513828 E	L	42.2
Asteraceae	Lactuca serriola L.	,	L L	42.2 NI
	Calendula officinalis	294342 N, 523520 E	AP	
Asteraceae		293813 N, 523127 E		32.9
Asteraceae	Cichorium intybus	293813 N, 523127 E	L	28.0
Asteraceae	Achillea millefolium L	293813 N, 523127 E	AP	NI
Asteraceae	Artemisia absinthium L.	293813 N, 523127 E	AP	31.1
Asteraceae	Tanacetum polycephalum Sch.Bip.	304140 N, 513827 E	AP	NI
Asteraceae	Achillea santolinoides Lag.	304138 N, 513827 E	AP	32.5
Asteraceae	Artemisia sieberi Besser	293637 N, 525321 E	AP	37.4
Asteraceae	Dahlia pinnata Cav.	293813 N, 523127 E	AP	24.0
Boraginaceae	Onosma rostellatum Lehm.	293637 N, 525321 E	AP	NI
Boraginaceae	Heliotropium europaeum L.	294134 N, 525331 E	AP	21.5
Chenopodiaceae	Chenopodium quinoa Willd.	294346 N, 523515 E	S	90. 3
Chenopodiaceae	Atriplex leucoclada Boiss.	293650 N, 524215 E	AP	48.5
Chenopodiaceae	Salsola imbricata Forssk.	293650 N, 524215 E	WP	44.1
Euphorbiaceae	Euphorbia erubescens Boiss.	304139 N, 513828 E	AP	42.0
Hypericaceae	Hypericum perforatum L.	294354 N, 523535 E	AP	NI
Lamiaceae	Thymus vulgaris L.	294353 N, 523536 E	AP	19.8
Lamiaceae	Thymus daenensis Celak	293813 N, 523127 E	AP	26.1
Lamiaceae	Stachys aucheri Benth.	304141 N, 513829 E	AP	49.4
Lamiaceae	Satureja bachtiarica Bunge	293637 N, 525321 E	AP	26.8
Lamiaceae	Ajuga austroiranica Rech.f.	293637 N, 525321 E	WP	23.6
Lamiaceae	Hyssopus officinalis L.	294353 N, 523536 E	AP	36.3
Lamiaceae	Nepeta persica	293813 N, 523127 E	AP	9.3
Lamiaceae	Melissa officinalis	294353 N, 523536 E	L	40.0
Lamiaceae	Zataria multiflora	294353 N, 523536 E	AP	30.2
Lamiaceae	Lamium album L.	304140 N, 513829 E	L	NI
Lamiaceae	Salvia officinalis	293813 N, 523127 E	AP	51.6
Lamiaceae	Stachys benthamiana Boiss.	293637 N, 525321 E	AP	59.6
Lamiaceae	Marrubium vulgare L.	293925 N, 525312 E	AP	29.7
Lamiaceae	<i>Rydingia persica</i> (Burm.f.) Scheen &	293637 N, 525321 E	R	29.7 56.7
Lannaceae	V.A.Albert		К	50.7
Lamiaceae	Micromelia persicae	293637 N, 525321 E	AP	NI
Lamiaceae	Hyoscyamus bornmuelleri Khat.	293637 N, 525321 E	AP	37.1
Lamiaceae	Lophanthus depauperatus (Benth.) Levin	293925 N, 525312 E	AP	NI
Lamiaceae	Ballota aucheri Boiss.	293925 N, 525312 E	AP	42.6

Table 1. The inhibition rate of plant extracts against Tobacco Mosaic Virus (TMV) infection	on on Nicotiana glutinosa
by local lesion assay. <sup><i>a</i></sup>	

<sup>*a*</sup> NI: No Inhibition; L: Leaves; R: Roots; S: Seed; WP: Whole Plant; B: Bulb; AP: Aerial Part, F: Flower. Inhibition rate (%) was calculated as outlined in the text. <sup>*b*</sup> The MAP30 protein was used as the positive control.

Table 1 continued...



Family	Plant name	Location GPS	Part of	Inhibition
			plant used	rate (%)
Lamiaceae	Hyssopus officinalis L.	294353 N, 523536 E	AP	33.0
Liliaceae	Fritillaria imperialis L.	304138 N, 513830 E	В	16.5
Malvaceae	Malva sylvestris L.	304138 N, 513827 E	AP	NI
Malvaceae	Malva neglecta Wallr.	294342 N, 523520 E	AP	43.6
Malvaceae	Alcea kurdica Alef.	303633 N, 513642 E	F	38.3
Meliaceae	Melia azedarach L.	293734 N, 523141 E	L	19.3
Nitrariaceae	Peganum harmala L.	293637 N, 525321 E	AP	28.0
Plantaginaceae	Veronica orientalis Mill.	304138 N, 513827 E	AP	41.9
Plantaginaceae	Plantago major L.	293813 N, 523127 E	S	NI
Polygonaceae	Rumex crispus L.	293814 N, 523127 E	L	12.4
Pteridaceae	Adiantum capillus-veneris	293813 N, 523127 E	WP	31.1
Urticaceae	Parietaria judaica L.	293925 N, 525312 E	AP	NI
Rhamnaceae	Ziziphus spina-christi (L.) Desf.	293813 N, 523127 E	L	54.9
Rosaceae	Crataegus ambigua C.A.Mey. ex A.K.Becke	293813 N, 523127 E	L	NI
Simaroubaceae	Ailanthus altissima (Mill.) Swingle	293816 N, 523127 E	L, F	80.1
Zygophyllaceae	Tribulus terrestris	294353 N, 523536 E	AP	41.3
· · · ·	MAP30 <sup>b</sup>			73.4

**Continued of Table 1.** The inhibition rate of plant extracts against *Tobacco Mosaic Virus* (TMV) infection on *Nicotiana glutinosa* by local lesion assay.<sup>*a*</sup>

<sup>*a*</sup> NI: No Inhibition; L: Leaves; R: Roots; S: Seed; WP: Whole Plant; B: Bulb; AP: Aerial Part, F: Flower. Inhibition rate (%) was calculated as outlined in the text. <sup>*b*</sup> The MAP30 protein was used as the positive control.

Name	$OD_{405}^{a}$	Viral concentration (µg mL <sup>-1</sup> )	Inhibition rate (%)	
Ailanthus altissima	0.706 <sup>a</sup>	3.52	36.44	
Chenopodium quinoa	0.614 <sup>a</sup>	3.21	42.02	
Ferula aucheri	1.305 <sup>b</sup>	5.53	-3.55	
Rhus coriaria	0.599 °	3.16	42.93	
Rydingia persica	1.281 <sup>b</sup>	5.45	-0.51	
Salvia officinalis	$0.804^{\rm d}$	3.85	30.49	
Stachys benthamiana	0.906 <sup>d</sup>	4.19	24.30	
Ziziphus spina-christi	0.950 <sup>d</sup>	4.34	21.63	
Control	1.307 <sup>b</sup>	5.54		

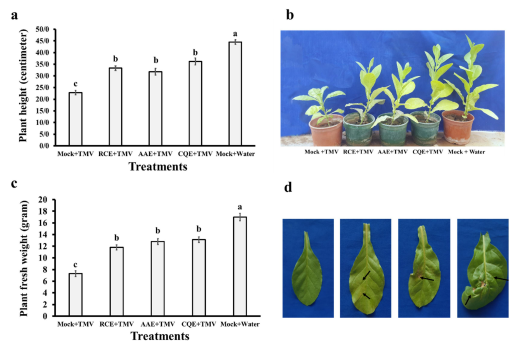
Table 2. The antiviral activity of 8 crude extracts on the viral accumulation in the leaf-disc assay.

<sup>*a*</sup> Data are expressed as the mean of three biological replicates. Different letters indicate statistically different values (P < 0.05, Tucky test).

symptoms and reduction in plant growth) of the virus in infected plants (Figure 1). As a result of applying different extracts treatments, significant differences were observed between samples. As compared with the infected controls, these significant differences were in plant height and fresh weight. Forty days after inoculation, the significant differences of plant height (36.9%) occurred in infected plants pre-treated with CQE in comparison with the control (mock+TMV), followed by those pre-treated with RCE and AAE (31.6 and

28.3%, respectively) (Figures 1-a and -b). The plants treated with CQE were heavier (44.4%) than those treated with RCE and AAE (38.1 and 43%, respectively). As shown in Figure 1, there were no significant differences in height and fresh weight between these three groups. The growth parameters of infected plants pre-treated with extracts increased variably compared to the control (mock+TMV).

Phytotoxicity assay of each extract separately showed various symptoms on tobacco leaves including leaf spotting, leaf



**Figure 1.** The effects of three plant extracts, *Rhus coriaria* (RCE), *Ailanthus altissima* (AAE), and *Chenopodium quinoa* (CQE) on growth parameters of TMV-infected tobacco plants at 40 days postinoculation. (a) Plant height; (b) Phenotypical modification of tobacco plants in response to five different treatments;(c) Plant fresh weight; (d) Visual symptoms of phytotoxicity (Arrow) in leaves of tobacco sprayed with mock (distilid water+2% DMSO) and extracts of *Chenopodium quinoa, Rhus coriaria* and *Ailanthus altissima*, respectively, from left to right, 7 days after exposure to treatments. Data are expressed as the mean±SE (error bars) of three independent experiments. Different letters indicate statistically different values (P<0.05, Tucky test).

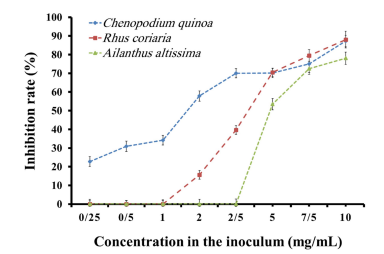
deformation, and scorch or dead tissue at concentrations of 20, 30, and 40 mg mL<sup>-1</sup> (Figure 1-d). In addition, phytotoxicity was observed in the leaves treated with all three extracts; though phytotoxicity intensity was higher in the leaves treated with AAE extract than in the other two extracts.

## EC<sub>50</sub>

Among the crude extracts with the highest antiviral activity, the  $EC_{50}$  values of CQE, RCE, and AAE were evaluated by local lesion assay. CQE exhibited the most inhibitory activity ( $EC_{50}$ , 1.64 mg mL<sup>-1</sup>) on TMV, followed by RCE and AAE showed  $EC_{50}$  values 2.82 and 4.42 mg mL<sup>-1</sup>, respectively (Figure 2).

#### Analysis of Secondary Metabolites

Mass spectrometry analysis was performed on the hydroethanolic (ethanol 80%) extracts of CQE, RCE, and AAE (Table 3) and showed higher antiviral activity than the other plant extracts. In particular, CQE showed a rich profile of saponins and anthocyanins, with a high amount of oleanolic acid. The main anthocyanins content of CQE was petunidin 3-*O*-glucoside (11.09 mg g<sup>-1</sup>), cyanidin pentoside (9.12 mg g<sup>-1</sup>), and 7-*O*-methylated anthocyanins (8.56 mg g<sup>-1</sup>). In AAE, with the highest amount of the phenolic compounds, high concentrations were found of epicatechin, ferulic acid, and galloylglucose. Compared to other analyzed extracts, RCE demonstrated a



**Figure 2.**  $EC_{50}$  values of selected extracts against tobacco mosaic virus using local lesion bioassay in *N. glutinosa*. Error bars show the standard error of the average of three independent experiments with three replicates.

rich profile of phenolic compounds, with a significant concentration of ferulic acid, *p*-coumaric acid, sinapic acid, and isorhamnetin (Table 3).

## DISCUSSION

Identifying a large number of valuable biogenic substance with insecticidal, antimicrobial, fungicidal, nematocidal, and antiviral activity in different plant species suggests an excellent opportunity for designing a novel biorational pesticides (Gonçalves et al., 2021; Kaur and Chandi, 2021; Pushpa et al., 2013). Therefore, owing to the limitation of the existing management of plant viruses, there is a global tendency to identify antiviral agents and search for biogenic product alternatives for the purpose of controlling viruses (Jing et al., 2012; Sharma et al., 2021).

In the present study, 60 plant extracts were screened for their antiviral activity against TMV as an experimental model, and their anti-TMV inhibitory effects were compared with that of MAP30 protein *in vivo* (Table 1). Extracts from six species, *C. quinoa, A. altissima, R. coriaria, Salvia officinalis,* 

Stachys benthamiana, and Ziziphus spinachristi, had significant inhibition against TMV replication. Among these six plants, R. coriaria (Ashoori et al., 2020), S. officinalis (Ghorbani and Esmaeilizadeh, 2017), C. quinoa (Pereira et al., 2020), and Z. spinachristi (Owayss et al., 2020) have been reported to have antimicrobial activity and have already been used for the prevention and treatment of some human and animal microbial infections. However, few studies have explored the antiviral activity of plant extracts, except that on A. altissima demonstrated potent antiviral activity (Ni et al., 2019). The results of local lesion assays support the possibility of using A. altissima extract.

This study is the first report of potential antiviral activity of crude extracts of *S. benthamiana, S. officinalis, R. coriaria, C. quinoa,* and *Z. spina-christi* against plant viruses. Notably, *R. coriaria, C. quinoa,* and *A. altissima* extracts showed the highest inhibitory activity. Besides, as compared to the controls, the investigated growth parameters in systemic infection of TMV showed the potentiality of *R. coriaria, C. quinoa,* and *A. altissima* extracts to decrease TMV infection. In addition, results indicated

**Table 3.** Mass spectrometry analysis of the secondary metabolites from the hydroethanolic extracts of three plants, *Rhus coriaria* (RCE), *Ailanthus altissima* (AAE), and *Chenopodium quinoa* (CQE).<sup>*a*</sup>

Type of	Compound	MW	Formula	RT (min)	CQE	AAE	RCE
phytochemicals	1				$(mg g^{-1})$	$(mg g^{-1})$	$(mg g^{-1})$
Phenolic compounds	Ferulic acid	194	$C_{10}H_{10}O_4$	3.15	0.25	5.83	2.19
	Gallic acid	170	$C_7H_6O_5$	3.36	0.41	1.64	ND
	Chlorogenic acid	354	$C_{16}H_{18}O_9$	11.14	ND	1.84	1.16
	Galloylglucose	332	$C_{13}H_{16}O_{10}$	17.26	0.32	4.92	0.77
	Ellagic acid	302	$C_{14}H_6O_8$	22.16	0.11	4.11	0.89
	Vanillic acid	168	$C_8H_8O_4$	25.25	0.32	2.76	1.16
	p-Coumaric acid	164	$C_9H_8O_3$	27.40	0.06	1.22	1.55
	Syringic acid	198	$C_9H_{10}O_5$	42.16	ND	ND	ND
	Sinapic acid	224	$C_{11}H_{12}O_5$	44.29	0.44	ND	1.84
	Quercetin	302	$C_{15}H_{10}O_7$	23.48	0.24	0.04	ND
	Rutin	610	C27H30O16	24.43	ND	1.19	1.12
	Catechin	290	$C_{15}H_{14}O_{6}$	27.36	0.02	ND	ND
Flavonoids and	Apigenin	270	$C_{15}H_{10}O_5$	30.30	0.03	0.96	0.15
their	Epicatechin	442	$C_{22}H_{18}O_{10}$	21.71	ND	24.03	1.36
glycosides	Luteolin glucoside	448	$C_{21}H_{20}O_{11}$	29.74	ND	1.12	ND
	Quercetin 3-O-galactoside	626	$C_{27}H_{30}O_{17}$	30.02	0.53	3.50	1.66
	Isorhamnetin	316	$C_{16}H_{12}O_7$	34.17	ND	0.51	1.14
	kaempferol	286	$C_{15}H_{10}O_{6}$	41.31	0.02	ND	ND
	Petunidin 3-O-glucoside	479	$C_{22}H_{23}O_{12}$	6.51	11.09	0.74	0.62
	Malvidin O-glucoside	493	$C_{23}H_{25}ClO_{12}$	8.25	5.15	1.48	1.36
Anthocyanins	Cyanidin 3-O-β- dgalactopyranoside	449	$C_{21}H_{21}O_{11}^{+}$	9.31	6.84	0.19	0.77
	Cyanidin 3-O-galactoside	449	$C_{2l}H_{2l}O_{1l}^{+}$	11.36	7.39	0.63	0.32
	Ginsenoside	801	$C_{42}H_{72}O_{14}$	3.22	ND	0.21	0.02
	Notoginsenoside R4	1241	$C_{59}H_{100}O_{27}$	3.61	ND	0.36	ND
	Floranotoginsenoside	1095	C53H90O23	3.95	1.05	ND	0.95
Saponins and their aglycones	Oleanolic acid	456	$C_{30}H_{48}O_3$	4.15	9.12	ND	ND
	Notoginsenoside R3	933	$C_{48}H_{82}O_{19}$	4.29	ND	0.17	0.25
	Hederagenin	472	$C_{30}H_{48}O_4$	4.49	5.3	ND	ND
	Gypenoside XIII	755	$C_{41}H_{70}O_{12}$	4.66	ND	1.32	0.38
	Malonyl-vinaginsenoside	1033	$C_{51}H_{84}O_{21}$	4.83	ND	ND	ND
	Yesanchinoside	1093	$C_{53}H_{88}O_{23}$	5.11	ND	0.71	0.37
	Serjanic acid	500	$C_{31}H_{48}O_5$	5.26	2.07	ND	0.61
	Quinquenoside	819	$C_{42}H_{74}O_{15}$	5.34	0.11	ND	ND
	Ginsenoside Mc	755	$C_{41}H_{70}O_{12}$	5.47	0.02	1.65	0.05
	Phytolaccagenic acid	516	$C_{31}H_{48}O_6$	5.53	3.51	ND	0.4

<sup>*a*</sup> ND: Not Detected); MW: Molecular Weight, RT: Retention Time. Retention time was calculated as the time from injection to detection.

that treatment with extracts reduced the symptoms caused by TMV compared with the control group.

Assay of  $EC_{50}$  value highlighted the strong antiviral activity of *C. quinoa* extract compared to other extracts. The experimental results indicate that these plant species might potentially be sources of natural antivirus or might be used to develop new antiviral agents. Differences in the effects of crude extracts could be attributable to the diversity of plant samples, structural diversity of biogenic substances and the relative concentrations of bioactive components in the extracts (Shen *et al.*, 2007). Exploring various lead compounds, there has been a strong interest in natural product research, which may be used as models for developing new biorational compounds in the pharmaceutical and agrochemical industries. Due to restrictions on antiviral activity screening techniques, there is an immediate need for developing new approaches and methodologies to reduce the screening costs and time. In particular, integrating standard methods with computer-based techniques would be very useful. With

the reduced number of possible bioactive compounds by in-silico methods, *in-vitro* and *in vivo* assays would further evaluate these components efficacy (Siqueira *et al.*, 2020).

Besides, we investigated chemical components present within the hydrohalic extract by LC-ESI-MS analysis (Table 3). The major components of R. coriaria, C. quinoa, and A. altissima extracts were ferulic acid  $(2.19 \text{ mg mL}^{-1})$ , petunidin 3-O-glucoside  $(11.09 \text{ mg mL}^{-1})$ , and epicatechin (24.03 mg  $mL^{-1}$ ), respectively. The extracts of C. quinoa demonstrated a rich profile of saponins and anthocyanins. In the extracts of A. altissima and R. coriaria, a significant concentration of phenolic compounds including ferulic acid, galloylglucose, epicatechin, p-coumaric acid, sinapic acid, and isorhamnetin were observed (Table 3). Flavonoid glycosides from Clematis lasiandra demonstrated obvious antiviral activities against TMV with multiple modes of action (Li et al., 2021). Three triterpene saponins were isolated from a MeOH extract of the leaves of Ilex oblonga, and showed potent antiviral activities against TMV (Wu et al., 2007). Although several reports show that compounds of epicatechin, ferulic acid, galloylglucose, and oleanolic acid have antiviral properties (Behrendt et al., 2017; Wang et al., 2017; Su and D'Souza, 2013), crude when working with extracts, constituents responsible for bioactivity are often unknown. Therefore, to improve the efficacy of bioactive substances, they should be comprehensively characterized, and the identities of components contributing to the biological activity must be investigated. (Caesar and Cech, 2019). Therefore, to compare the profile of these metabolites among the evaluated plants, chemical analysis would be essential. This chemical profiling could provide clues to link the detected bioactivity with the natural abundance of some metabolites.

## CONCLUSIONS

In this study, we have demonstrated the antiviral activity of 60 medicinal plants

against TMV. Six of these plant samples showed potent antiviral activity against TMV. R. coriaria, C. quinoa, and A. altissima extracts demonstrated the most potent antiviral effect amongst the selected plant. To isolate and characterize the bioactive constituents responsible for the antiviral activity of these plant extracts, some further research is required. To the best of our knowledge, this is the first report of the potential antiviral activity of Stachys benthamiana, Salvia officinalis, Rhus coriaria, Chenopodium quinoa, and Ziziphus spina-christi extracts against the TMV. Therefore, these plant extracts can be considered good candidates in developing natural virucides.

#### ACKNOWLEDGEMENTS

The authors acknowledge help from Prof. A.R. Khosravi, plant taxonomist, Shiraz University, in collecting and identifying plant species. Also, the authors are grateful to Dr. A. Moghadam, Institute of Biotechnology, Shiraz University, for providing MAP30 protein.

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فعالیت ضد ویروسی سه گونه گیاهی *Chenopodium quinoa ، Rhus coriaria و* Ailanthus altissima بر روی ویروس موزاییک تنباکو

ی. بی نیاز، ف. احمدی، ع. نیازی، و ع. افشاری فر

## چکیدہ

تحقیقات بر روی ترکیبات طبیعی، فرصتهای جدیدی برای کنترل موثر و پایدار بیمارگرهای ویروسی گیاهان فراهم نموده است. در این پژوهش، ما عصاره هیدرواتانولی ۶۰ گونه گیاهی از ۲۲ خانواده مختلف را آماده و فعالیت ضد ویروسی آنها را بررسی نمودیم. از میان عصارههای گیاهی تهیه شده، عصارههای Rhus آماده و فعالیت ضد ویروسی آنها را بررسی نمودیم. از میان عصارههای گیاهی تهیه شده، عصارههای Rhus تمودند. عصاره Quinoa و Chenopodium و Ailanthus altissima ویروس موزاییک توتون را به خوبی مهار نمودند. عصاره Ailanthus altissima و Chenopodium quinoa. ویروس موزاییک توتون را به خوبی مهار آلودگی ویروس موزائیک توتون داشت. عصاره ی گرم بر میلی گرم بر میلی لیتر بالاترین تاثیر را در کاهش آلودگی ویروس موزائیک توتون داشت. عصاره ی گیاهان C. quinoa و antizism مه به ترتیب با مقادیر موزائیک توتون را مهار نمودند. نتاج بررسی فعالیت ضد ویروسی عصارهها بر روی ویروس موزائیک توتون در موزائیک توتون را مهار نمودند. نتاج بررسی فعالیت ضد ویروسی عصارهها بر روی ویروس موزائیک توتون در میزبانان سیستمیک نشان داد که هر سه عصاره به طور معنی داری در کاهش علائم ویروس در گیاهان توتون میزبانان سیستمیک نشان داد که هر سه عصاره به طور معنی داری در کاهش علائم ویروس در گیاهان توتون میزبانان سیستمیک نشان داد که هر سه عصاره به طور معنی داری در کاهش علائم ویروس در گیاهان توتون میزبانان سیستمیک نشان داد که هر سه عصاره به طور معنی داری در کاهش علائم ویروس در گیاهان توتون میزبانان میستمیای ویروس در گیاهان توتون میزبانان میستمیای معصاره می می داری می داری ترکیبات ساپونین و آنتوسیانین را نشان داد در حالی که عصاره های R. coriaria می می داری ترکیبات ساپونین و آنوسیانین مانشان