Peganum harmala seed extract can prevent HSV-1 replication in vitro

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Abstract: Peganum harmala L. (Zygophylaceae) has been used in a variety of practical applications in medical science. Our objective in the current study was to determine the effects of the seed extract on Herpes simplex virus type 1 (HSV-1) replication in Vero cells. Different concentrations (100, 500, 571, 667, 800 and 1000 µg/ml) of the extract were examined. Experiments were carried out using Vero cells. P.harmala seed extract was found to be non-toxic to Vero cells up to a concentration of 667 µg/ml. The antiviral activity of non-toxic concentration against HSV-1 was tested. The replication of HSV-1 was inhibited, indicating that the P.harmala L. extract contains an anti-HSV-1 substance.

Keywords: Peganum harmala L.  Vero cells  TCID50  Cytotoxicity

Introduction

Herpes simplex viruses (HSV) are ubiquitous agents which cause a variety of diseases ranging in severity from mild to severe, and in certain cases, these may even become life threatenings, especially in immunocompromised patients. After primary infection, HSV persists in the host for the lifetime. HSV infection is thus considered lifelong infection. Nucleoside analogues such as acyclovir (ACV), pencyclovir, are the only approved drugs for the treatment of HSV infections. However, the widespread use of nucleoside based drugs has led to the emergence of resistance in HSV especially among immunocompromised patients (1). Herbal remedies used in the traditional folk medicine provide an interesting and still largely unexplored source for the creation a development of potentially new drugs for chemo-therapy which might help to overcome the growing problem of resistance and also the toxicity of the currently available commercial antibiotics. The traditional medicinal methods, especially the use of medicinal plants, still play a vital role to cover the basic health needs in the developing countries. Therefore, it is of great interest to carry out a screening of these plants in order to validate their use in folk medicine and to reveal the active principle by isolation and characterization of their constituents (2). A great number of medicinal plants are being used to treat microbial infections, particularly in the rural areas of Iran where the traditional folk medicine remains a major source to cure minor ailments.

The Peganum harmala L. (Syrian rue) is a wild-growing flowering plant belonging to the Zygophyaleae family and is found abundantly in Middle East and North Africa (3). From ancient times, it has been claimed to be an important medicinal plant. Its seeds are known to possess hypothermic and hallucinogenic properties (4, 5). It has been used traditionally as an emmenagogue and an abortifacient agent in the Middle East and North Africa (6). There are several reports in the literature indicating a great variety of pharmacological activities for Peganum harmala L such as anti-bacterial, antifungal and monoamine oxidase (MOA)-inhibition (7). It has also been known to interact with α2-Adrenoceptor subtypes (8) and have hallucination potency and be effective in the treatment of dermatosis (9), hypothermic (10) and cancer (11). Thus the aim of this study was to elucidate the possible mechanism underlying the action of alkaloid extract of peganum harmala seeds on herpes simplex virus type 1 replication in cell culture.

Material & Methods

Plant material

The plant material was collected from Tabriz, Iran on July 2007. Five grams of dried and powdered seeds of peganum harmala fruits was macerated
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four times with 50 ml methanol at 5°C in water bath for 1h. The extracts were combined and evaporated to dryness. The residue was dissolved in 1N HCl at concentration of 2% then filtered. The filtrate was extracted twice with 20 ml petroleum ether. The aqueous acid layer was basified (pH 10) with 1N NaOH and extracted four times with 50 ml chloroform. The chloroform layer was combined and evaporated to dryness, and then the residue was dissolved in 25 ml methanol. The solution of alkaloid extract was passed through 0.45 µm filter and stored at 4°C refrigerator for further use.

**Cell culture & Virus**

Vero cells (African green monkey kidney cells) were cultured as monolayers in DMEM supplemented with 10% heat-inactivated fetal bovine serum (FBS), 100 IU/ml penicillin, and 100 µg/ml streptomycin. The cells were maintained at 37°C in a humidified atmosphere with 5% CO₂ and were subcultured twice a week. HSV-1 was isolated from patients, and identified by monoclonal anti-HSV-1 antibodies. Viruses were propagated in Vero cells and quantified in tissue culture plates in mentioned concentration for 2 and 5h at 37°C and 5% CO₂. After removal of the extract, the cells were washed with phosphate-buffered saline (PBS) and then infected with HSV-1 at a multiplicity of infection (MOI) of 0.1. After 1h incubation the unadsorbed virus was removed, the cell monolayer was washed with PBS and further incubated in DMEM with 2% FBS. Controls consisted of Vero cells untreated alone and Vero cells infected with HSV-1.

**Cytotoxicity assay**

In order to test the effects of the extract on Vero cells, 5x10⁴ cells (in 1ml DMEM, supplemented with 10% FBS) were seeded into each well of microplates, cultured for 6 h at 37°C, and cells were allowed to grow for additional 48 h in the presence of increasing amounts of extract (100, 500, 571, 667, 800 and 1000 µg/ml). The cytotoxicity of extracts was determined on a conventional hemocytometer using the trypan blue exclusion method (13). The highest noncytoidal (on Vero cells) concentration of plant extract was determined to be 667 µg/ml. Therefore, the 667 µg/ml concentration was used for the determination of antiviral activities.

**Virus yield inhibition assay**

Nontoxic concentration of test drug was checked for antiviral property by virus yield reduction assay (14) against virus challenge dose of 100 TCID₅₀/ml. In virus yield assay, reduction in the yield of virus when cells were treated with the plant extract was determined.

**Pre-incubation of cell monolayer with extract before virus infection**

The extract was dissolved in serum free DMEM and incubated with semi-confluent cell in 96-well tissue culture plates in mentioned concentration for 2 and 5h at 37°C and 5% CO₂. After removal of the extract, the cells were washed with PBS before virus infection. The assay was performed as described above, with the exception that extract was added together with the virus. Virus stock was mixed with mentioned concentration of extract in equal volume, and used for infecting cells. After an incubation time of 1 h, the solutions containing both extract and virus were removed, the cell monolayer was washed with PBS and further incubated in DMEM with 2% FBS.

**Incubation of virus with extract before virus infection**

The assay was performed as described above, with the exception that extract was added together with the virus. Virus stock was mixed with mentioned concentration of extract in equal volume, and used for infecting cells. After an incubation time of 1 h, the solutions containing both extract and virus were removed, the cell monolayer was washed with PBS and further incubated in DMEM with 2% FBS.

![Fig. 1: Effect of the extract pre-, during and post-infection on virus yield in Vero cells.](image)

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Incubation of cell monolayer with extract after virus infection
The cell monolayer was infected with the virus. After 1 h incubation the unadsorbed virus was removed, the cell monolayer was washed with PBS and then incubated with mentioned concentration of the extract in DMEM with 2% FBS after 1, 4, 8, and 12 h post infections. After 48h incubation at 37 °C in presence of 5% CO₂, supernatant was removed and semi-confluent cell monolayer, grown in 96-well plate, was inoculated with 10-fold dilutions of the supernatant for 1 h at 37 °C. After removal of the inocula, monolayer was washed once with PBS and then incubated in DMEM containing 2% FBS for 48 h. Virus titer was determined by the endpoint dilution method and expressed as TCID50 according to the Reed–Muench formula. The results were expressed as reduction ratio of virus titers by comparison with virus control.

Immunofluorescence
Confluent Vero cells grown on 13mm coverslips were infected with HSV-1 at MOI of 1 TCID50 for 16h in the presence of 667 µg/ml of the extract compared to untreated infected cells. The cover slips were removed, washed in PBS and fixed in acetone at 4 °C; then they were stained indirectly with fluoresceine conjugated anti-human IgG using specific viral antiserum. The cover slips were mounted in glycerol buffer and examined in a UV equipped microscope.

Polymerase Chain Reaction
To determine presence of viral DNA in extract-treated infected cells, DNA was extracted from cells and PCR was performed using HSV-1 specific primers. Preparation of the reaction mix was based on the protocol of HSV-1 diagnostic PCR kit (Dynamic BioScience). The DNA was amplified in a thermal cycler for 40 cycles. Before the first cycle, an early denaturation step of 3 min at 94 °C was performed to complete the denaturation. Each cycle consisted of a 30 sec denaturation step at 94 °C, a 30 sec annealing step at 50.5 °C, and a 30 sec elongation step at 72 °C. After the last cycle, a final elongation step of 10 min at 72 °C was performed to complete the elongation. The samples were then kept at 4 °C until analysis. Amplified products were visualized under ultraviolet light after electrophoresis for 1 h at 100 V through a 1.5% (w/v) agarose gel, containing ethidium bromide, and photographed. A molecular size marker included in the gel was GelPilot 100 bp plus ladder (QIAGEN).

Results
Cytotoxicity
Cytotoxicity of Pharmala seed extract was examined by means of trypan blue exclusion method. Microscopic observations showed that no change occurred in cell growth and morphology in the presence of extract up to a concentration of 667 µg/ml. Trypan blue exclusion method showed that the total viable cell numbers were approximatley 98% as compare with the control cells (data not shown).

Virus yield reduction assay
The inhibition of virus yield by the extract was evaluated by TCID50 assay in Vero cells. We
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examined the antiviral activity of the extract when incubated with cells prior to infection with the virus; the virus titer of the supernatant did not show any significant change in this case. To examine the direct effect of the extract to virus, HSV was mixed with nontoxic concentration of the extract and the mixture was used for infecting cells; the result of virus yield inhibition showed no distinct drop in virus titer. In order to study the antiviral activity after viral adsorption, the extract was incubated with the infected cell monolayer 1, 4, 8, and 12h after infection. Results are shown in Fig.1. The most inhibitory effect of the extract was shown 1h after infection, when the virus titer of the supernatant dropped from $10^{6.5}$ TCID50/ml to $10^{3.5}$ TCID50/ml. Inhibitory effect of the extract was not significant in other cases, except for 4h that reduced virus titer from $10^{6.5}$ TCID50/ml to $10^{4.5}$ TCID50/ml. As it is shown in fig.1, treating infected cells 1h after infection was not completely inhibited virus growth, but after third passage in presence of the extract, virus production was completely inhibited.

**Immunofluorescence**

To confirm the inhibitory effect of P. harmala seed extract on HSV-1 replication in cell culture, coverlip-grown Vero cells infected with HSV-1 in the presence and absence of the extract, were fixed and processed for indirect Immunofluorescent assay. The expression of HSV-1 proteins was evidenced after 16h of infection in untreated group and its characteristic cytopathic effects were apparent. Repeating the same experiment in the presence of the extract, expression of HSV-1 proteins became very weak and did not show significant fluorescence in 3rd passage.

**Viral DNA synthesis**

To determine presence of viral DNA in extract of treated infected cells, DNA was extracted from cells and PCR test was performed using HSV-1 specific primers. Although viral DNA was detected at first passage, there was no detectable DNA after third passage.

**Discussion**

Human herpes viruses are found worldwide and are among the most frequent causes of viral infections in immunocompromised patients (15). HSV infection is one of these infectious diseases, which occurs in far-ranging region of body. HSV-I mainly causes infection on the craniofacial skin and mucosa, and HSV-II results in the infection of genitals and anus. HSV induces a variety of diseases, for example, herpesencephalitis, herpetic keratitis, herpetic gingivitis, herpes genitalis, infection of newborn and fetal malformation, some of which are lethal (16). Currently, the primary drugs used for HSV infection are nucleoside analogues. Among these, acyclovir is the preferred medicine in clinical treatment, but the drug resistance develops after longterm administration (17). Meanwhile, the extracts of some medicinal herbs were investigated in term of searching hopeful candidates for treating HSV infection (18, 19).

Fig. 3: Effect of the extract on viral DNA synthesis; Lane 1: HSV-1 positive control; Lane 2: MW Marker; Lane 3: first passage; Lane 4: second passage; Lane 5: third passage; Lane 6: negative control; Lane 7: MW Marker.
Peganum harmala is a plant known since the first century A.D. and still, currently used for therapeutic purposes. Harmaline, the active principle of the plant seeds, and its derivatives, cause visual troubles, loss of coordination, agitation and delirium, and, at high doses, it can produce paralysis (20). The present study was initiated to evaluate inhibitory effect of the extract of P. harmala on HSV-1 replication in vitro.

In this study, the effect of different concentrations of the extract on Vero cells was determined after 72h incubation by Trypan blue exclusion method. Pharmala seed extract had no cytotoxicity on Vero cells up to concentration of 667µg/mL. The inhibition of virus yield showed that treating the cells with the extract 1h after infection can significantly reduce virus titer in the first passage and inhibit completely virus production in the third passage. Regarding the HSV replication cycle, soon after infection, approximately 2 to 4 hours postinfection (hpi), the α or immediate early (IE) genes are expressed. Although transcription of α genes requires no prior viral protein synthesis, an HSV protein brought in with the virion tegument, VP16, stimulates transcription of α genes (21). Therefore, inhibitory effect of the extract 1h after infection can be caused by preventing α genes protein synthesis or possibly repressing function VP16 tegument protein.

Evaluation of HSV-1 protein expression in treated infected cells by Immunofluorescence assay showed significant reduction in protein synthesis at first passage and complete repression in third passage which means that the extract can prevent viral gene expression in transcription or translation level.

Finally, for investigating the effect of the extract on viral DNA synthesis, PCR test was performed on the supernatant of three passages. Although there were detectable DNA bands in 1st and 2nd passages, we could not detect any DNA band in 3rd passage, indicating that viral DNA synthesis was completely inhibited in third passage. Regarding to this results, we can conclude that inhibitory effect of Pharmala seed extract is not clearly understood. Further study is required to determine the mechanism of viral growth inhibition.

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