

EVALUATION OF *INVITRO* AND *INSILICO* ANTILITHIATIC AND ANTIOXIDANT POTENTIAL WITH THE ACETONIC EXTRACT OF *PUNICA GRANATUM* WHOLE FRUIT

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ABSTRACT

Stones that develop anywhere in the urinary sections, are referred to as urolithiasis. The Punicaceae plant family comprises *Punica granatum*. Pomegranate fruit is utilised in ailment of diarrhoea, dysentery, and intestinal parasites and has anti-inflammatory and antibacterial qualities. The objectives of our study is to evaluate the antilithiatic activity by *invitro* and *insilico* molecular analysis of whole fruit acetonic extract of *Punica granatum* Cystone used as a standard drug in the present study. Plant screening resulted the presence of tannins, terpenoids, alkaloids, glycosides, carbohydrates, saponins and steroids. The acetonic extract of *Punica granatum* was proved to possess prominent antioxidant activity by reducing power assay. By reducing the concentration of reactive oxygen species i.e., measured by the transformation of Fe^{3+} to Fe^{2+} , increased absorption of reaction mixture indicates increased reducing power. The nucleation and aggregation processes of CaC_2O crystallisation were both hindered by the administration of PGAE and cystone coupled with calcium chloride dihydrate; the decrease in solution turbidity in both phenomena indicated percentage inhibition. *In vitro* antilithiatic activity by Homogenous precipitation method, PGAE and Cystone dissolved calcium oxalate precipitate. The compounds present in *Punica granatum* whole fruit acetonic extract they are docked with protein 5G3N. Ellagic acid, Alpha tocopherol, Brevifolin, Kaempferol, Maslinic acid shown good docking score Overall results explain us that PGAE has proven *in-vitro* antilithiatic activity and antioxidant activity.

Key words: *Punica granatum*, Homogenous precipitation method, docking.

I INTRODUCTION

The term "urolithiasis," which is drawn out from Greek terms "ouros" (for urine) plus "lithos" (for stone), describes the buildup of tough, solid, & non-metallic particles with in urinary system. Aphorisms of Hippocrates describe the clinical development and expression of this chronic and widespread ailment (1). Defined also as a kidney calculus, a renal stone is a hard crystalline that develops in the kidney via minerals within urine. The preponderance of stone formation are made up of calcium salt crystals (2).

Five years after the first episode with symptomatic renal lithiasis, recurrence chances have been found to be approximately to 50% (3). The nucleation & aggregation assay and the homogenous precipitation method are *in vitro* models for antilithiatic action. Cystone is the standard medication employed in the trial. Allopathic drugs could eliminate healthy microorganisms and have unfavourable side effects that make other therapies less successful (4). Traditional medical systems are still used extensively on many fronts. For broad range of ailments in human were treated with herbal materials as a source that has received more attention as a result of factors including population growth, insufficient drug supply, restrictive cost of therapies, adverse reactions of many pharmaceutical chemicals, and resistance emergence to presently utilised medicine for infectious ailments.

Punicaceae is the family of plant which comprises the pomegranate, scientific name is *Punica granatum* L. Its therapeutic and nutritional qualities make it a significant fruit. The pomegranate plant's various elements have medicinal benefits, including the fruit's anti-inflammatory and antibacterial properties, the seed oil's inhibitory effect on skin and breast cancers, and it shows presence of phytoestrogenic compounds. The fruit is also abundant in phenols, which have potent antioxidant properties. It is employed against dysentery, intestinal parasites, diarrhoea, heart and throat tonic. It is used to treat haemorrhoids and halt nose and gum bleeding (5).

A crucial technique in molecular biology and computer-aided drug design is *insilico* study of molecular docking. Predicting the dominant linkage mode(s) of a receptor with a protein with a recognized three-dimensional configuration is objective ligand-protein docking.

The goal of a study to perform *invitro* analysis of antilithiatic activity and antioxidant activity by *Punica granatum* whole fruit extract.

II MATERIALS AND METHODS

The process of establishing a technique entails a number of processes carried out in a methodical manner in order to accomplish the desired outcomes in accordance with the established standards and rules.

A Phytochemical screening of plant

The herb serves as a laboratory of biosynthetic producing a variety of chemicals, including glycosides, alkaloids, volatile oils, tannins, and others that have physiological & medicinal effects in addition to chemical molecules like carbohydrates, protein, and lipids.

B Evaluation of antioxidant assay by *in vitro* method

Punica granatum whole fruit's acetonetic extract was used to test the antioxidant activity *in vitro* using a reducing power assay methodology.

i. Reducing power assay

Enhanced absorbance is a sign that antioxidant activity has elevated. The methodology is based on the idea that reaction combinations absorb additional light. Potassium ferricyanide (Fe^{3+}) combines with substances that have the ability to reduce it to make potassium ferrocyanide (Fe^{2+}), that eventually interacts with FeCl_3 to create ferric complex, which contains absorption peak at 700 nm. Phosphate buffer pH 6.6, Potassium dihydrogen phosphate (0.2 M) solution, Sodium hydroxide solution (0.2M) solution, Potassium ferric cyanide (1% w/v) solution and Ferric chloride solution (0.1% w/v) are the reagents required for the assay.

1. Add potassium ferricyanide 2.5 mL (1 percent by weight) and phosphate buffer of 205 mL pH 6.6 to 1 mL of the standard and test substances, and then incubate at 500 C for about 30 minutes.
2. To stated supernatant liquid of 2.5 mL, distilled water 205 mL, and 0.5 mL of a 0.1 percent w/v FeCl_3 solution were added.
3. Employing an Ultra violet - visible analyser, the absorbance of complex (ferric ferrous) was determined utilising phosphate buffer at pH 6.6 as a reference control at 700 nm and the increase in absorbance was calculated (6).

The present increase in reducing power was calculates using the following equation,

Where 'Abs_{test}' is absorbance of test solution: 'Abs_{blank}' is absorbance of blank.

C Evaluation of antilithiatic activity by *invitro* method

Punica granatum whole fruit's acetonetic extract was used to test the antilithiatic activity *in vitro* using the nucleation and aggregation assay and homogenous precipitation technique.

i. Nucleation and aggregation assay

A freshly made combination of 200 mM NaCl and 10 mM sodium acetate trihydrate, together with 10 mM CaCl_2 . dihydrate and 1.0 mM sodium oxalate, was brought to pH 5.7. A flowing water bath was used to conduct each experiment at a temperature of 37 °C. Sodium oxalate of 25 mL solution was allowed into beaker and maintained at 37 C by placing in a hot plate magnetic stirrer (Model 2MLH,REMI) upon stirring continuously at about 800 rpm, for crystallisation studies. Before adding 25 mL of calcium chloride solution, an additional 1 mL of distilled water, the standard (Cystone), and the extract were added. Following the addition of a calcium-containing solution, the (OD) optical density was examined at 620 nm using a spectrophotometer, first every 15 s over 5 minutes followed by every 1 minute over 10 minutes. The studies were all carried out in triplicate. To evaluate the intensity of constituted crystals in the finished solutions, they were observed under a light microscope (Olympus,USA).

With the use of following formula, the percentage of inhibition by the influence of Cystone or PGAE was compared to the control. The formula used to determine the percentage inhibition was

$$1 - \frac{T_{si}}{T_{sc}} * 100$$

Where T_{sc} , the turbidity slope of the control; and T_{si} , the turbidity slope in the presence of the inhibitor (7)

ii. Homogenous method of precipitation

Step 1: Generation of experimental calcium oxalate kidney stones using homogeneous precipitation

Equimolar solutions of sodium oxalate (AR) in 10 mL of 2 N H_2SO_4 and CaCl_2 . dihydrate in distilled water were combined and permitted to respond in a beaker with enough distilled water. Calcium oxalate was the precipitate that was obtained. Ammonia solution cleans precipitate of sulphuric acid residue. It was rinsed in distilled water and dried for four hours at 60 degrees Celsius.

Step-2: Semi-permeable membrane preparation employing farm eggs

A semi-permeable layer over an egg sits between its calcified outside shell and it's inside yolk and albumin. By soaking the eggs using 2M Hydrochloric acid for quite a night that resulted in full decalcification, the shell being chemically removed from the eggs.

was gently poked thoroughly

$$\text{Percentage increase in reducing power (\%)} = \frac{\text{Abs}_{\text{test}} - \text{Abs}_{\text{blank}}}{\text{Abs}_{\text{blank}}} * 100$$

Additionally, the egg with a pointed tip, washed using

distilled water, then had the contents strained out apart out from decalcified egg. After properly washing the layer of egg membrane with water, it was immersed inside an ammonia solvent and kept moist for a time before being completely rinsed. Preserved in refrigerator with a pH of 7.3–7.4.

Step-3: Titrimetry determination of calcium oxalate

1 mg of calcium salt of oxalic acid and 10 mg of an extract, chemical, or standard were precisely weighed, suturing them together in a membrane obtained from eggs. A titration flask holding 100 mL of 0.1M TRIS buffer was used to suspend this. One group served as the negative control (contained calcium oxalate of 1 mg). All conical flasks in an incubator were warmed for 2 hours, or for roughly 7-8 hours, to 37 C. Each group's semi-permeable membrane's contents were taken out and placed in a test tube. The end point was reached by adding 2 millilitres of 1N sulphuric acid and titrating with 0.9494 N KMnO_4 .

$$1 \text{ mL of } 0.9494\text{N } \text{KMnO}_4 = 0.1898 \text{ mg of } 4 \text{ Calcium}$$

To determine how much calcium oxalate really dissolved the test chemical, the quantity of undissolved calcium oxalate was deducted from the total quantity utilised in the experiment at the beginning (s) (8).

D *Insilico* analysis: Molecular Docking Studies

A specific type of bioinformatics simulation called molecular docking includes the combination of more than one molecule to produce a stable compound. It makes predictions about the three-dimensional configuration of every complex based on the binding characteristics of ligand as well as target. Utilizing the score feature of the mCULE programme, distinct potential compound structures obtained using molecular docking are graded and categorised. Biochemical docking needs a dataset to search for targets with the correct PDB format and a mechanism to design compounds in the mCULE. In a ligand-receptor complex, the intermolecular reactions are

crucial and challenging modelling exercises. Typically, the ligand molecules are permitted to vary while the receptor is kept stiff or somewhat rigid (9). Through the discovery studio visualizer, the resulting docking orientations were seen. The glide score technique was used to choose the best docked structures. The binding is more favourable, greater negative is the score.

III RESULTS

Using appropriate *in-vitro* models and *insilico* studies, extract of *Punica granatum* whole fruit by acetone was investigated for its antioxidant and anti-lithiatic activity. All the data achieved in the study were presented below.

A Phytochemical screening of plant

The acetonic extract contained tannins, terpenoids, alkaloids, glycosides, carbohydrates, saponins, and steroids, according to phytochemical analysis.

B Evaluation of antioxidant assay by *in vitro* method

i. Reducing power assay

This assay was used to measure the antioxidant activity of *Punica granatum* acetonic extract.

Table 1: Antioxidant activity of *Punica granatum* Acetone extract whole fruit by using reducing power assay

S. No	Compounds	Concentration ($\mu\text{g} / \text{mL}$)	% Inhibition (Mean \pm SEM)	IC ₅₀ value ($\mu\text{g} / \text{mL}$)
1	PGAE	10	18.33 \pm 0.98	34
		20	23.33 \pm 0.72	
		30	36.66 \pm 0.54	
		40	58.6 \pm 0.72	
		50	68.3 \pm 0.72	
2	Ascorbic acid	10	21 \pm 0.47	29
		20	34.3 \pm 0.98	
		30	56.6 \pm 0.98	
		40	69 \pm 0.47	
		50	73.3 \pm 0.98	

With increasing doses, PGAE has demonstrated a greater percentage of free radical inhibition, and its IC₅₀ value was discovered to be 40 $\mu\text{g}/\text{mL}$. The extract's potential was still on track with that of the reference ascorbic acid, and its IC₅₀ value was determined to be 29 g/mL .

C Evaluation of antilithiatic activity by *invitro* method

i. Nucleation and aggregation assay:

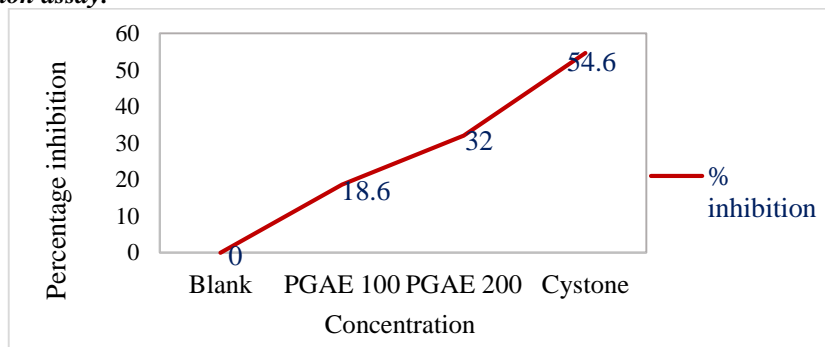


Figure 1: Effect of PGAE and cysteine in *in-vitro* nucleation and aggregation assay method.

The antilithiatic activity of the acetonic extract of *Punica granatum* was carried out by employing *in-vitro* nucleation and aggregation assay. Blank group showed high turbidity so the percent inhibition was found to be 0 %. PGAE has shown increase in percent inhibition, decrease in turbidity with increase in dose, PGAE 100 – 18.6 % and PGAE 200 – 32 %. The potential of the extract was comparable with standard Cysteine and percent inhibition value was found to be 54.6.

ii. Homogenous method of precipitation

Step 1: Generation of experimental calcium oxalate kidney stones using homogeneous precipitation



A - Preparation of equimolar solutions of sodium oxalate & CaCl_2 ,

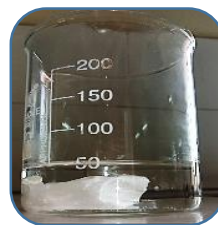


B - Mixer of Equimolar solutions in distilled water



C – CaC_2O_4 ppt filtered,

Step 2: Semi-permeable membrane preparation employing farm eggs



D –Eggs covered with HCl overnight, **E** – Squeezing of contents from egg, **F** – Ammonia solution occupied with membrane.

Step 3: Titrimetry determination of calcium oxalate



G – CaC_2O_4 + 10 mg control/PGAE/Cystone sutured in membrane and immersed in conical flask with tris buffer and

H - Estimation of percent dissolution of calcium oxalate by titrimetric.

Figure 2: *In vitro* antilithiatic activity by Homogenous method of precipitation

Table 2: Outcome of PGAE on percent dissolution CaC_2O_4 by homogenous method of precipitation.

Groups	KMnO ₄ (mL)	Dissolved Calcium (mg)	Undissolved calcium (mg)	Percent dissolution calcium oxalate (%)
Blank	-	-	1	0
PGAE	2.5	0.45	0.55	45
Cystone	3.2	0.57	0.42	57.6

The homogeneous precipitation technique was used to test *Punica granatum's* acetonic extract's *in-vitro* antilithiatic activity. The solubility of calcium oxalate in the blank group was determined to be zero percent. Calcium oxalate dissolution has increased to 45%, according to PGAE. The PGAE potential was comparable with regular Cystone, and 57.6 percent of calcium oxalate was observed to dissolve.

D *Insilico* analysis: Molecular docking studies

Initially the protein was downloaded from PDB was prepared by removing extra chains. Attributes of spheres are prepared and noted. Molecules identified from GC-MS were selected. Later molecules drawn in mCULE and ligprep was created. Protein is uploaded with sphere attributes and the structures were docked against 5G3N protein. Docking indicated that some of our compounds have good binding ability with phospholipase A2 inhibitor protein (PDB ID: 5G3N). Following are the ligand interactions of compounds present in *Punica granatum* whole fruit with 5G3N protein.

Table 3: Mcule docking scores

Compounds	Score
Ellagic acid	-8.8
Gallic acid, Punicalin	-5.5
Oleic acid	-5.6
Linoleic acid, Vanilic acid	-5.9
Caffeic acid	-6.2
Ferulic acid	-6.4
Alpha tocopherol	-7.4
Beta carotene	-5.5
Kaempferol	-7.1
Brevifolin	-8.1
Maslinic acid	-7.2
Cystone A	-7.8

The greater negative the score the more favourable the binding.

in turbidity with increase in dose. In homogenous precipitation method PGAE has shown enhanced percent dissolution calcium oxalate lower growth of crystal. The compounds' ability to inhibit PLA2 in the cytosol was validated by molecular docking experiments since they occupied the binding site and displayed interactions. The experimental PLA2 inhibitory action was represented in the docking score order for all compounds.

In future, to understand active component responsible for anti-lithiatic action and to investigate the mechanism, *in-vivo* activity and the isolation of active ingredients are required.

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