**Abstract**

*Caralluma fimbriata* (Wall.) (Asclepiadaceae), is mentioned as vegetable in Indian Materia Medica and an affluent resource of saponins. It is reported in conventional medicine method of India as well as Arabia that *C. fimbriata* was extensively used for cancer treatment. Current study was planned to assess anticancer potential of saponin rich fraction from *C. fimbriata* using *in silico* and *in vitro* assays. Caratubersides A-G, a pregnane glycosides found in *C. fimbriata* were taken for *in silico* examination and processed through PASS Online software for the prediction of structure dependent pharmacological actions. Docking was carried out using Autodock Tool and Autodock Vina, revealed antineoplastic action of caratubersides along with apoptogenic potential. MTT assay was performed on MCF-7 cell line. Shell less chicken embryo culture assay was done for anti-angiogenic properties at different concentrations (1.5µg/ml, 3µg/ml, and 6µg/ml). Chromosomal aberrations assay was carried out in cultured human blood. And apoptogenic potential was estimated on MCF-7 cells using cleaved caspases 3 and caspase 8 cell based ELISA assay kit. Results of study showed that IC50 of saponin rich fraction of *C. fimbriata* was at 3µg/mL. Considerable (p <0.05) decreases were observed in angiogenic properties. Insignificant chromosomal aberrations were found in normal cells. Treatment of saponin rich part improved levels of caspases 3 as well as caspase 8 (ODs 1.35 and 1.68 respectively). From the study, saponin rich portion obtained from *C. fimbriata* displayed antiproliferative, anti-angiogenic actions along with apoptogenic prospective and no significant chromosomal aberrations were found in normal human cells.

**Key Words:** *Caralluma fimbriata*, *in silico*, MTT assay, Angiogenesis, Chromosomal Aberrations assay, Apoptosis.

**Introduction**

Upsurge in numbers of deaths related to cancer along with unfavourable and deadly side effects of cancer treatments like radiation and chemotherapy, the interest in searching for novel cancer treating agents, especially of plants origin, is on rise (1). *C. fimbriata* has synonym *Caralluma ascendens* is a widespread, juicy cactus as well as feral remedial plant belongs to Asclepiadaceae family (2). In Western it is also called Shindula makadi, Makad shenguli, Kullimudayan, and Ranshabar (3). The plant is widely found in Africa, India, the Canary Islands, Southern Europe and Arabia (4).

*Caralluma fimbriata* is marked as a hunger suppressant, vegetable and thirst quencher in Indian Materia Medica and grow as an arid herb in waterless places of India (5). *C. fimbriata* is also used to make boundaries around gardens. Plant is also used to make chutneys and pickles in numerous centuries (6). *C. fimbriata* is reported as an antiobesogenic agent and metabolic regulator (7). It is considered to be “famine food” which repress hunger and satisfy thirst. It is a “portable food” which prevents starvation in period of frantic requirement of food while travelling (8). *C. fimbriata* is safe and utilized conventionally for treating paralysis, rheumatic disorders, inflammation and diabetes along with leprosy. It showed antimalarial, antiproliferative, antiulcer, antinoceptive, antitypanosomal, antioxidant activities, antihyperglycaemic, hypolipidaemic and hepatoprotective activity (9, 10). Devi and her friends in 2016 reported that the plant is extensively useful in the Indian traditional medication system for anticancer activity (11). Qayyum in 2018 reported that traditionally *Caralluma* species used for cancer treatment (12). Zari and his colleagues 2018 reported that in the Indian and Arabic traditional medicine *Caralluma* species are used in the cancer therapy (13). Earlier Shenai et al., observed that ethanolic extract of *C. fimbriata* leaves having cytotoxicity in COLO 320 cells (1). Also Al-Faifi investigated in vitro antiproliferative action of *Caralluma spp.* in HEPG2 and MCF-7 cell lines (14).

*Caralluma fimbriata* is a rich source of pregnane glycosides, megastigmagine glycosides, steroidal glycosides, saponins, flavonoids and flavone glycosides (15), pregnane steroids, hexadecanoic acid, oleic acid, aromatic and nonaromatic bitter principles, alkaloids, unsaturated and saturated hydrocarbons, volatile compounds along with β-sitosterol. There are also few...
active elements present in *C. fimbriata* like Caratubersides (16), Boucerosides and Tomenkogenin (17).

Saponins are obtained as secondary metabolites from plants and they are glycosides in nature which exerts various pharmacological effects. Previous studies demonstrated anti-proliferative, anti-angiogenic, and antimetastatic activity of saponins through induction of apoptosis and cell differentiation. Saponins also showed reversal of multidrug resistance in some cancers (18). Moreover, steroidal class of saponins, Pregnan glycosides, was reported to have potent anticancer and pro-apoptotic activity. Caratubersides available in *C. fimbriata* are pregnane glycosides which were ill reported for their cytotoxic or apoptogenic potential. This promoted us to explore anticancer effects of saponin rich extract of *C. fimbriata* using in vitro models (19).

Here, structural dependent biological activities of caratubersides (A to G) were predicted using PASS Online and Swiss Target Prediction Software and structural docking was done via Autodock Tools and Autodock Vina. Additional saponin rich fraction was isolated from *C. fimbriata* leaves using water and n-butanol fractionation and studied further for anticancer actions. Cell viability and cell toxicity (MTT) assay was performed to evaluate antiproliferative activity on MCF-7 cells. Shell less chicken embryo culture assay was performed to check anti-angiogenic property. Effects on physical reliability of genetic materials were studied using cultured human blood in Chromosomal aberrations assay. Apoptogenic prospective was estimated on MCF-7 cells through cleaved caspase 3 as well as caspase 8 cell based ELISA assay kit.

**Materials and Methods**

**Materials**

Breast cancer MCF-7 cell line was attained from NCCS, Pune, Maharashtra.

Dried hydroalcoholic *C. fimbriata* leaves extract was procured from Navchetna Kendra, New Delhi, India.

Prolific eggs obtained through egg seller. And ELISA assay kit was obtained from Cell Signaling Technology.

**Isolation of Saponin from *Caralluma fimbriata***

In hydroalcoholic extract of *C. fimbriata* leaves (5gm), 50 ml distilled water was added and shifted the mixture in separating funnel. The same amount of n-butanol (10:10 v/v) was placed in separating funnel and permitted them for overnight separation of n-butanol fraction at room temperature. At 45°C collected n-butanol part was evaporated to get saponin rich extract (20).

**Fingerprinting of *Caralluma fimbriata***

Fingerprinting study of *C. fimbriata* was performed by the use of CAMAG HPTLC equipment prepared by Linomat V applicator, TLC scanner 3, Reprostar 3 as well as 12 bit CCD camera for documents of photos and restricted with software WinCATS-4. The samples (30µL) were placed in band form which containing 6 mm width by the use of Camag microlitre plunger on 200 µm thick silica gel coated aluminum plate (5×10 cm). Leaves extract as well as β-sitosterol standard were solubilized in small quantity of methanol. Samples were prepared in concentration of 10µg/10µl. The plate was developed till 85mm distance in a glass chamber 30min formerly saturated with toluene: ethyl acetate: methanol: glacial acetic acid (8:1:0:5:0.3v/v/s) solvent system. Air dried the plat for removal of mobile phase and then derivatized the plat by dripping in anisaldehyde-H2SO4 reagent till 2 seconds. After that dried the plate in air and heated it till 10 minutes at 110°C. Then scanning was executed at λ= 584nm with Camag TLC scanner (21).

**In silico studies**

**Prediction of structure base biological activities in PASS Online**

For caratubersides A to G, structures of them were draw in ChemDraw Ultra 10.0 and then save them in .cdx file formats which were then opened in Chem3D Ultra 10.0 to converted them in .mol file extension. These MDL Mol file were then opened in PASS online software for prediction of activities. For caspase 3 as well as for caspase 8 stimulation resultant values Pa and Pi (probability to be active and probability to be inactive respectively) were search and adopted from different predictable pharmacological actions (22, 23, 24).

**Molecular docking**

For docking studies .mol files of caratubersides (A-G) were changed to .pdb format (PDB) by OpenBabel 2.4.1software. Casepase 3 (3DEI) as well as caspase 8 (3KJQ) configurations was acquired in .pdb file format through RCSB data bank. After that they were opened through AutoDock Tools 4.2.6 and processed for obtaining .pdbqt files. MetaPocket 2.0 was used to recognize different binding sites for ligands on caspase 3 and caspase 8 and to choose parameters for grid box. Values of grid box were utilized for docking of related proteins and caratubersides in Autodock Vina (25, 26).

**MTT assay**

MTT assay was carried out for evaluation of antiproliferative activity of saponin rich part from *C.fimbriata* in MCF-7 cells line with different (0.001µg/ml, 0.01µg/ml, 0.1µg/ml, 1µg/ml and 10µg/ml) concentrations. Saponin rich fraction of *C.fimbriata* 1mg/ml was solubilized in to DMSO, then consecutively diluted by growth media for the different. Concentration of DMSO was reserved less than 0.1%. MCF-7 cell line was maintained within DMEM having 10% (v/v) bovine serum from fatal calves, were placed into 96 well plates. Treatments of different concentrations of saponin wealthy portion was given to cells and place the plate in incubator at 37°C and 5% CO2 till 4 days. After 96 hours cell were treated with MTT dye and again incubated till 4 hours. Cells were formed purple colour formazan products and then
DMSO was added in wells. DMSO dissolved the formazan products which were measured spectrophotometrically on 550nm to obtain IC50 value (27).

Shell less chicken embryo culture assay
Fertilized eggs cooled till 25 minutes and cleaned through 70% alcohol for reduction of outside effluence after 72 hrs of incubation. Albumin of non-fertile eggs was placed in sterilized Petridish, to provide pad for egg contains. After that incubated eggs were break from top by using spatula and filling of eggs were smoothly discharge above albumin bed in Petridish. Effectively placed embryos were separated in the control and treatment groups. In treatment groups treatment with various concentrations (1.5µg/ml, 3µg/ml, and 6µg/ml) of saponin loaded part of C. fimbriata was given in a dosage of 10µl on blood vessels developed area. All Petridish were enclosed and again incubated at 37 ± 0.5°C and 80% humidity. Later than 3hours and 6 hours, differences in quantity of extremities, junctions, nodes, branches, total branches length were examined via angiogenesis analyzer ImageJ 1.50 b software (28).

Chromosomal aberrations (CA) assay
Fresh blood samples were collected in four vials from healthy, non-smoking and erratically chosen female and male (2:1) between 20-25 years of age. One vial was marked as untreated control and remaining three were marked as treated vial. Treatment by three different concentrations (1.5µg/ml, 3µg/ml and 6µg/ml) was given in a dosage of 50µl. In to each vials 5ml PBMAX TM karyotyping growth medium, 50µl Heparin and 0.6 ml fresh blood was added and allowed to incubate them till 72 hours. Later at 24 hours, treatment with various concentrations of saponin wealthy portion was given. On 69th hour cultures were treated with100µl colchicine. On the completion of 72 hours, the vials were centrifuged till 8 minutes on 1200 rpm and discarded the supernatants. After that cells were treated with 5ml of 0.56% KCl and centrifuge tubes were permitted to place in incubator for 25 min at 37°C for the swelling of cells. The tubes were centrifuged again on 1200 rpm till 8 min. Supernatants were removed, cells were fixed by addition of 6 ml of carnoy’s fixative and placed them into refrigerator till 1 hour. The samples were centrifuged again for 8 min at 1200 rpm, supernatants were discarded and using 3ml of fixative, the washing step was continued for obtaining white pellet. After that slides were prepared through placing 5-7 drops of white pallet from suitable altitude on priorly cooled clean slides. Then coded them and used for scoring after staining with 2% giemsa stain. 100 metaphases were scored in each group using compound light microscope at 100 x magnifications for chromosomal aberrations (29).

Caspase assay
Stimulation of caspase 3 as well as caspase 8 was estimated by using ELISA assay kit in breast cancer MCF-7 cell line. The cells were placed into 96 well plates, treated with 10µg/ml concentration of saponin rich fraction and allowed them to incubate till 6 hours at 37°C and 5% CO2. At the end of 6 hours development media was discarded through wells and washed the cells by PBS. Washed cells were again rinsed and treated by cleaved caspase-3 (Asp175) as well as cleaved caspase- 8 (Asp 391) till couple of hours. Rinsed cells 3 times using PBS and treated with corresponding HRP conjugated secondary antibodies till half an hour. After 30 minute cells were rinsed three times, identify through TMB substrate and calculated optical densities using wavelength 450 nm by Fluostar (BMG Germany) (30).

Statistical analysis
Results were uttered as mean ± S.E.M. for each investigational group. Arithmetical examination was carrying out by IBM SPSS Statistics 22 statistical software. Arithmetic assessment among various treatments and control were executed using ANOVA pursued via turkey’s post hoc test. Minimum significant level was recognized on p<0.05.

Results
Isolation of Saponins
C. fimbriata is a rich source of saponins. We obtained the extraction yield of 35% of saponins from hydroalcoholic extract of C. fimbriata leaves.

Fingerprinting of Caralluma fimbrita
Chromatograms attained by β-sitosterol and C. fimbriata extract using solvent system toluene: ethyl acetate: methanol: glacial acetic acid (8:1:0.5:0.3 v/v) were shown in figure 1 and 2. The Rf value at 584 nm was found to be 0.55 (Fig. 1a and 2A, 2B).

Figure 1. 3D HPTLC chromatogram for β-sitosterol and C. fimbriata extract

Figure 2. HPTLC chromatograms of
(A) β-sitosterol
(B) C. fimbriata extract
In Silico studies
Prediction of biological activities using in silico methods
On the basis of structures biological activities of caratubersides (A- G) were predicted using PASS Online and SwissTargetPrediction Softwares. PASS Online predictions with Pa values greater than 0.7 were enlisted. The anticancer effect related predictions which were common for all caratubersides were found to include antineoplastic (breast cancer), antiangiogenic activity, apoptosis agonist and caspase 3 as well as caspase 8 stimulation. Also by SwissTargetPrediction apoptotic regulation activities were expected for all caratubersides. For caspase 3 and caspase 8 stimulation values of Pa along with Pi were stated separately. (Table 1).

Table 1. Biological activity prediction of caratubersides using in silico methods.

<table>
<thead>
<tr>
<th>Sr. No.</th>
<th>Triterpenoids / Structures</th>
<th>PASS Online Prediction</th>
<th>Binding energy (Kcal/mol)</th>
<th>Amino acid residue</th>
<th>Hydrogen Bond</th>
<th>PASS Online Prediction</th>
<th>Binding energy (Kcal/mol)</th>
<th>Amino acid residue</th>
<th>Hydrogen Bond</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Pa</td>
<td>Pi</td>
<td></td>
<td></td>
<td></td>
<td>Pa</td>
<td>Pi</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>Caratuberside A</td>
<td>0.857</td>
<td>0.004</td>
<td>-7.9</td>
<td>Glu123, Gly122, Rxb300, Gly165, His121, Tyr204</td>
<td>1</td>
<td>0.718</td>
<td>0.003</td>
<td>-7.6</td>
</tr>
<tr>
<td>2</td>
<td>Caratuberside B</td>
<td>0.773</td>
<td>0.007</td>
<td>-7.9</td>
<td>Glu123, Gly122, Rxb300, Gly165, His121, Tyr204</td>
<td>1</td>
<td>0.691</td>
<td>0.004</td>
<td>-6.3</td>
</tr>
<tr>
<td>3</td>
<td>Caratuberside C</td>
<td>0.545</td>
<td>0.022</td>
<td>-9.3</td>
<td>Glu123, Gly122, Rxb300, Gly165, His121, Tyr204</td>
<td>1</td>
<td>0.445</td>
<td>0.032</td>
<td>-8.0</td>
</tr>
<tr>
<td>4</td>
<td>Caratuberside D</td>
<td>0.495</td>
<td>0.028</td>
<td>-9.4</td>
<td>Glu123, Gly122, Rxb300, Gly165, His121, Tyr204</td>
<td>1</td>
<td>0.354</td>
<td>0.084</td>
<td>-7.5</td>
</tr>
<tr>
<td>5</td>
<td>Caratuberside E</td>
<td>0.398</td>
<td>0.052</td>
<td>-9.0</td>
<td>Glu123, Gly122, Rxb300, Gly165, His121, Tyr204</td>
<td>1</td>
<td>0.354</td>
<td>0.084</td>
<td>-7.9</td>
</tr>
</tbody>
</table>
MTT assay:
Antiproliferative MTT assay for saponin rich part from *C.fimbriata* was performed on MCF-7 cells line displayed IC50 at 3µg/ml (Table 2).

<table>
<thead>
<tr>
<th>Caratuberside F</th>
<th>6</th>
<th>0.470</th>
<th>0.032</th>
<th>-9.1</th>
<th>Glu123, Gly122, Rxb300, Gly165, His121, Tyr204</th>
<th>1</th>
<th>0.435</th>
<th>0.036</th>
<th>-6.8</th>
<th>Thr149, Pro415, Tyr412, Ser316, His317</th>
<th>11</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caratuberside G</td>
<td>7</td>
<td>0.689</td>
<td>0.012</td>
<td>-8.5</td>
<td>Glu123, Gly122, Rxb300, Gly165, His121, Tyr204</td>
<td>1</td>
<td>0.510</td>
<td>0.016</td>
<td>-7.4</td>
<td>Thr149, Pro415, Tyr412, Ser316, His317</td>
<td>11</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Concentration</th>
<th><em>C. fimbriata</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>68.75</td>
</tr>
<tr>
<td>1</td>
<td>32.41</td>
</tr>
<tr>
<td>0.1</td>
<td>22.18</td>
</tr>
<tr>
<td>0.01</td>
<td>2.36</td>
</tr>
<tr>
<td>0.001</td>
<td>1.28</td>
</tr>
<tr>
<td>IC50 value µg/ml</td>
<td>3</td>
</tr>
</tbody>
</table>

Shell less chicken embryo culture assay
There is a considerable (p<0.05) decreases in quantity of angiogenic properties after 0-3hour and 0-6 hour of exposure to different concentration of saponin loaded portion (Table 3 and Fig. 3).
Table 3. Effect of saponin rich fraction of *C. fimbriata* on Angiogenic parameters in Shell less chicken embryo culture.

<table>
<thead>
<tr>
<th>Concentrations</th>
<th>0 - 3 hours</th>
<th>0 - 6 hours</th>
<th>0 - 3 hours</th>
<th>0 - 6 hours</th>
<th>0 - 3 hours</th>
<th>0 - 6 hours</th>
<th>0 - 3 hours</th>
<th>0 - 6 hours</th>
<th>0 - 3 hours</th>
<th>0 - 6 hours</th>
</tr>
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<tr>
<td>Control</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>1.5 µg/ml</td>
<td>26.85 ± 0.99*</td>
<td>41.96 ± 0.93*</td>
<td>28.75 ± 0.83*</td>
<td>48.20 ± 1.07*</td>
<td>34.04 ± 1.32*</td>
<td>52.62 ± 1.87*</td>
<td>37.74 ± 1.47*</td>
<td>55.73 ± 0.95*</td>
<td>28.81 ± 0.82*</td>
<td>44.61 ± 1.23*</td>
</tr>
<tr>
<td>3 µg/ml</td>
<td>36.57 ± 1.08*</td>
<td>60.37 ± 1.32*</td>
<td>35.97 ± 0.95*</td>
<td>60.68 ± 1.52*</td>
<td>40.19 ± 0.74*</td>
<td>59.02 ± 0.84*</td>
<td>45.83 ± 0.83*</td>
<td>61.00 ± 1.34*</td>
<td>37.08 ± 0.72*</td>
<td>59.88 ± 1.37*</td>
</tr>
<tr>
<td>6 µg/ml</td>
<td>42.93 ± 1.04*</td>
<td>67.12 ± 1.01*</td>
<td>46.57 ± 1.01*</td>
<td>67.38 ± 1.64*</td>
<td>50.25 ± 1.37*</td>
<td>68.04 ± 2.03*</td>
<td>51.12 ± 1.25*</td>
<td>69.00 ± 0.75*</td>
<td>51.54 ± 1.00*</td>
<td>68.09 ± 2.45*</td>
</tr>
</tbody>
</table>

Each values stated as mean ± S.E.M. (n=6) per group. Statistical analysis was performed using one-way ANOVA pursued via Turkey’s post hoc test. *P < 0.05 contrasts to control.

Chromosomal aberrations (CA) assay:
50 µl of various concentrations (1.5µg/ml, 3µg/ml, 6µg/ml) of saponin wealthy part was placed in cultured vials showed insignificant change in amount of entire chromosomal aberrations contrasts to control group (Table 4 and Fig 4).

Table 4. Chromosomal aberrations after treatment

<table>
<thead>
<tr>
<th>Aberration</th>
<th>Normal (Control)</th>
<th>Saponin rich fraction of <em>C. fimbriata</em> (1.5 µg/ml)</th>
<th>Saponin rich fraction of <em>C. fimbriata</em> (3 µg/ml)</th>
<th>Saponin rich fraction of <em>C. fimbriata</em> (6 µg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chromatid Break</td>
<td>5.00 ± 0.57</td>
<td>4.66 ± 1.76</td>
<td>6.33 ± 0.88</td>
<td>5.66 ± 0.88</td>
</tr>
<tr>
<td>Chromosome Break</td>
<td>3.66 ± 0.66</td>
<td>3.33 ± 0.88</td>
<td>5.00 ± 0.57</td>
<td>6.00 ± 1.15</td>
</tr>
<tr>
<td>Chromosome Gap</td>
<td>5.66 ± 0.88</td>
<td>6.33 ± 0.88</td>
<td>7.00 ± 0.57</td>
<td>4.33 ± 0.33</td>
</tr>
<tr>
<td>Dicentric Fragments</td>
<td>1.00 ± 0.57</td>
<td>2.33 ± 0.33</td>
<td>3.33 ± 1.45</td>
<td>1.00 ± 0.57</td>
</tr>
<tr>
<td>Acentric Fragment</td>
<td>3.33 ± 0.88</td>
<td>2.33 ± 0.88</td>
<td>4.33 ± 1.22</td>
<td>4.66 ± 1.20</td>
</tr>
<tr>
<td>Ring</td>
<td>0.33 ± 0.33</td>
<td>0.00 ± 0.00</td>
<td>0.00 ± 0.00</td>
<td>0.33 ± 0.33</td>
</tr>
<tr>
<td>Premature Separation</td>
<td>1.00 ± 1.00</td>
<td>2.00 ± 1.15</td>
<td>1.66 ± 0.88</td>
<td>4.00 ± 2.08</td>
</tr>
<tr>
<td>Endoreduplication</td>
<td>0.66 ± 0.66</td>
<td>0.33 ± 0.33</td>
<td>0.00 ± 0.00</td>
<td>0.00 ± 0.00</td>
</tr>
<tr>
<td>Hypodiploid</td>
<td>8.66 ± 0.88</td>
<td>9.00 ± 1.52</td>
<td>11.66 ± 0.66</td>
<td>13.33 ± 0.33</td>
</tr>
<tr>
<td>Hyperdiploid</td>
<td>2.00 ± 0.57</td>
<td>2.33 ± 0.88</td>
<td>1.33 ± 0.33</td>
<td>0.66 ± 0.66</td>
</tr>
</tbody>
</table>

Each values stated as mean ± S.E.M. (n=6) per group. Statistical analysis was performed using one-way ANOVA pursued via Turkey’s post hoc test. *P < 0.05 contrasts to control.

Figure 4. Representative pictures of Chromosomal Aberrations

A: Normal metaphase: Number of chromosomes is from 44 to 48; B: Chromatid break: Break in one sister-chromatid at any one locus; C: Chromosomal break: Breaks in both sister-chromatids at any one locus; D: Chromosomal gap: Gap between the chromosomes from centromere; E: Dicentric fragment: Abnormal chromosome with two centromeres. It is formed through the fusion of two chromosome segments; F: Acentric fragment: A segment of a chromosome that lacks a centromere; G: Ring: A portion of a chromosome has broken off and formed a circle or ring; H: Premature separation: Premature loss of cohesion between centromeres results in their independent segregation at meiosis I; I: Endoreduplication: Replication of the nuclear genome in the absence of mitosis, which leads to elevated nuclear gene content and polyploidy; J: Hypodiploidy: Number of chromosomes are <44; K: Hyperdiploidy: Number of chromosomes are >48.
Caspase assay

Caspases 3 as well as 8 activations was observed in MCF-7 cancer cell line after treatment. Table 5 and Figure 5 represent the data of stimulation of caspases 3 and 8 (Table 5 and Fig 5).

Table 5. Apoptosis assay of saponin rich fraction of C. fimbriata for caspases 3 and 8

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Concentrations</th>
<th>OD450 (Caspase 3)</th>
<th>OD450 (Caspase 8)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saponin rich part of C. fimbriata</td>
<td>10 µg/ml</td>
<td>1.35</td>
<td>1.68</td>
</tr>
<tr>
<td>Staurosporine</td>
<td>1 µg/ml</td>
<td>2.47</td>
<td>2.6</td>
</tr>
</tbody>
</table>

Figure 5. The ODs at 450nm in consequence of cleaved Caspase 3 (Asp 175) and Caspase 8 (Asp 391).

Discussions

Innovation in the dealing and avoidance of cancer requires the continuous expansion of new cancer protective and curative agents (31). Numbers of constituents isolated from herbs are being used as curing agents against various cancers including breast cancer (32). The worldwide demand of herbal products is increased in past few decades for the management of chronic as well as incurable ailments (33).

Saponins are secondary metabolites usually exist in plants with composite arrangement play a role of defense mechanism and accounted to be active in various dreadful ailments (34). C. fimbriata is a rich source of Saponin and used in this study.

Bioinformatics techniques are gaining importance in the field of new drug development (35). It empowers the researchers to predict pharmacological activities of compounds and allows them to practically monitor a database of millions of different compounds to identify lead molecules. PASSOnline and SwissTargetPredictions are such powerful softwares which could forecast the probabilities of test component to be active against vast number of targets in the body. PASSOnline assigns two probability scores, one for ‘Active’ and other for ‘Inactive’. A compound assigned bigger Pa score and smaller Pi score toward a specific pharmacological activity will be considered for further analysis. SwissTargetPrediction will directly depict the probability to be active score. Structures of Caratubersides A to G were processed through PASSOnline and SwissTargetPrediction softwares. The predictions related to anticancer activity, which commonly found for all caratubersides, such as antineoplastic for breast cancer, antiangiogenic activity, apoptosis agonist along with caspases 3 and 8 stimulations. Caratuberside A showed the probability of 85.7% and 71.8% for stimulating the Caspase 3 and Caspase 8 respectively. While Caratuberside B showed 77.3% and 69.1% probability as a stimulator of Caspase 3 and Caspase 8 activity respectively. Rest of the compounds showed probability in decreasing order like Caratuberside G > Caratuberside C > Caratuberside D > Caratuberside F > Caratuberside E.

Further, we processed these structures for molecular docking by AutoDock Tool and AutoDock Vina. These are freeware which forecasts the affinity of ligands towards the target protein using Lamarckian Genetic algorithm. In molecular docking, ligands are docked with target proteins and the affinities are reported in the form binding free energy. Caratubersides were docked with the identified targets from pharmacological activity predictions i.e. Caspase 3 and Caspase 8. Interestingly lowest binding free energy i.e. -9.4 was found for Caratuberside D with Caspase 3 and -7.9 for Caratuberside E with Caspase 8. Caratubersides A and B, which showed maximum probability to be active in pharmacological activity prediction, showed binding free energy of -7.9 and -7.9 respectively for Caspase 3 while -7.6 and -6.3 respectively for Caspase 8 (Table 1).

MTT assay is the most preferred and relatively simple method used for cytotoxicity studies. MTT is yellow colored water-soluble dye which is metabolized by the alive cells in purple colored formazan products via NAD(P)H-dependent cellular oxidoreductase enzymes (36). We treated MCF-7 cells by different concentrations of saponin rich fraction of C. fimbriata till 96 hours and inhibitions in percentage were measured. Graph of percentage inhibition against concentration was designed and IC50 was found to be 3 µg/ml. Earlier Shenai et al., observed the cytotoxicity of ethanolic extract of C. fimbriata leaves in COLO 320 cells, but no reports were found regarding cytotoxicity of saponins from C. fimbriata leaves (1).

Recently, much attention was gain by angiogenesis in cancer biology (37). Oxygen supply and nutrients required for growth and survival of cancer was maintained through sprouting of new blood vessels from the established vasculature and creating a new vascular system within the tumor (38). Moreover, this neovascularization also plays significant role in tumor growth, invasion and metastasis (39). Cancer is considered to be angiogenic dependent disease and most of the studies successfully targeted various regulators of angiogenesis. We exposed shell less chicken embryo cultures at various concentrations (1.5, 3 and 6 µg/ml) of saponin rich fraction and observed considerable (p<0.05) decreases into quantity of angiogenic properties among 0-3 hours as well as 0-6 hours of contact to drug. The test compounds not only prevented the neovascularization but also decreased the number along with length of existing blood vessels.
Evaluating the occurrence of chromosomal aberrations in peripheral blood is a susceptible cytogenetic test to discover potency of test compounds against mutagenesis and carcinogenesis (40). This kind of assays is generally used in the genotoxicity assessment in human subjects (29). Aberrations in chromosomes were observed, with and without treatment of saponin rich fraction in cultured blood samples. End result revealed non-significant correlation between treated and control groups in chromosomal aberration assay.

Within organisms, pathophysiological conditions can be overcome by various cell death modalities (41). Apoptosis is one such mechanism where genetically synchronized death of the cell acting a vital task in exclusion of infected, injured as well as other unwanted cells from the body (42). Biochemically, the key feature of apoptosis includes activation of cascade of caspase and DNA fragmentation. In human, apoptosis can be instigated by (1) the extrinsic pathway, which can cause caspase 8 stimulation; (2) the intrinsic pathway, started through cellular tension pursued via stimulation of caspase 9; or (3) the granzyme B pathway (43). All such paths meet to a general finishing stage of apoptosis with the aim of proteolytic stimulation of caspases 3. Here saponin rich fraction was assessed for its apoptogenic potential in MCF-7 cells using ELISA. We observed significant stimulation of caspases 3 (OD of 1.35 on 450nm) as well as caspase 8 (OD of 1.68 on 450nm) within test compound treated groups. Caspase 8 is synthesized as inactive zymogen procaspase and gets activated either in extrinsic pathway due to activation of death receptors or it is recruited and activated at the outer membrane of mitochondria in mitochondria mediated apoptosis. It is not clear by which mechanism (intrinsic or extrinsic pathway) saponin rich factor of \textit{C. fimbriata} increases cleaved Caspase 8, but once getting activated Caspase 8 further increases levels of Caspase 3.

**Conclusion**

Results of presented study displayed antiproliferative, cytotoxic, antiangiogenic as well as apoptotic effects of saponin rich fraction of \textit{C. fimbriata}. CAs in normal human blood was non-significant, which suggests absence of genotoxicity in normal cells. In silico studies predicted the apoptotic prospective of caratubersides by stimulating caspase 3 and caspase 8. The results obtained for saponin rich fraction of \textit{C. fimbriata} is in accordance with this prediction and showed significantly increased levels of caspases 3 along with caspase 8 in breast cancer MCF-7 cell line.

**Conflict of interest**

The authors declared no conflict of interest.

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