Original article

Chemical composition and pharmacological bio-efficacy of Parrotiopsis jacquemontiana (Decne) Rehder for anticancer activity

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Abstract

Consistent STAT3 (Single transducer and activator of transcription 3) activation is observed in many tumors and promotes malignant cell transformation. In the present investigation, we evaluated the anticancer effects of Parrotiopsis jacquemontiana methanol fraction (PJM) on STAT3 inhibition in HCCLM3 and MDA-MB 231 cells. PJM suppressed the activation of upstream kinases i.e. JAK-1/2 (Janus kinase-1/2), and c-Src (Proto-oncogene tyrosine-protein kinase c-Src), and upregulated the expression levels of PIAS-1/3 (Protein Inhibitor of Activated STATs-1/3), SHP-1/2 (Src-homology region 2 domain-containing phosphatase-1/2), and PTP-1b (Protein tyrosine phosphatase 1b) which negatively regulate STAT3 signaling pathway. PJM also decreased the levels of protein products conferring to various oncogenes, which in turn repressed the proliferation, migration, invasion, and induced apoptosis in cancer cell lines. The growth inhibitory effects of PJM on cell-cycle and metastasis were correlated with decreased expression levels of CyclinD1, CyclinE, MMP-2 (Matrix metalloproteinases-2), and MMP-9 (Matrix metalloproteinases-9). Induction of apoptosis was indicated by the cleavage and subsequent activation of Caspases (Cysteine-dependent Aspartate-directed Proteases) i.e. caspase-3, 7, 8, 9, and PARP (Poly (ADP-ribose) polymerase) as well as through the down-regulation of anti-apoptotic proteins. These apoptotic effects of PJM were preceded by inhibition of STAT3 cell-signaling pathway. STAT3 was needed for PJM-induced apoptosis, and inhibition of STAT3 via pharmacological inhibitor (Stattic; SC-203282) abolished the apoptotic effects. Conclusively, our results demonstrate the capability of PJM to inhibit cancer cell-proliferation and induce apoptosis by suppressing STAT3 via upregulation of STAT3 inhibitors and pro-apoptotic proteins whereas the down-regulation of upstream kinases and anti-apoptotic protein expression. In future, one-step advance studies of PHM regarding its role in metastatic inhibition, immune response modulation for reducing tumor, and inducing apoptosis in suitable animal models would be an interesting and promising research area.

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Abbreviations: STAT3, Single transducer and activator of transcription 3; PARP, Poly (ADP-ribose) polymerase; XIAP, X-linked inhibitor of apoptosis protein; JAK-1/2, Janus kinase-1/2; SHP-1/2, Src-homology region 2 domain-containing phosphatase-1/2; PIAS-1/3, Protein Inhibitor of Activated STATs-1/3; BAK, Bcl-2 homologous antagonist/killer; BAX, Bcl-2-Associate X protein; BCL-XL, B-cell lymphoma-extra large; MTT, 3-(4, 5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazoliumbromide; MMP-2/9, Matrix metalloproteinases-2/9.

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1. Introduction:

Natural products deliver invaluable opportunities for discovery of new drugs conferring to the availability of unmatched chemical diversity. The bulk of world’s population (>80%) rely on folk medicines to meet their primary healthcare requirements (Zahra et al., 2021a, Zahra et al., 2021b). Traditional medicine systems i.e. Kampo, Chinese, Unani, and Ayurveda have been using medicinal plants for treating various ailments and this knowledge is further explored in modern medicine by novel drug discoveries. Medicinal plants serve as useful sources of novel anticancer drugs (Kashyap et al., 2019; Shannugam et al., 2017). It is to be noted that many herbal preparations have been used in cancer treatment for thousands of years owing to their traditional acceptability, less toxicity, biocompatibility, cost-effectiveness, consistent supply and relatively fewer side-effects (Batool et al., 2019; Batool et al., 2020, Trang et al., 2020). So far, different medicinal plants have been investigated for different phytochemicals (alkaloids, flavonoids, tannins, saponins etc) and their bioactivities and have shown promising biological potentials against different kinds of ailments (Iqbal et al., 2017, Iqbal et al., 2018a,b, Abbasi et al., 2019).

Cancer is declared as a disease with high prevalence and mortality rate worldwide (Iqbal et al., 2018a). According to the new global cancer data, one in every five male while one in every six female are likely to develop cancer hence leading to one in every eight male whereas one in every ten female deaths, respectively (Costa et al., 2020). Among the women, breast cancer is considered as the most frightful disease globally after lung cancer due to its easy metastasis and poor prognosis, which makes it even more deadly (Iqbal et al., 2018b, Abbasi et al., 2018). During the migration and invasion process, the tightly-packed epithelial cells of breast cancer become loose and produce pseudopodium transforming into mesenchymal cells. This process is known as epithelial-mesenchymal transformation (EMT) and follows alterations in a sequence of pathways, detected at molecular level (Cheng et al., 2019; Lee et al., 2019). Once cancer cells are transformed into mesenchymal cells, penetration to the vascular endothelium becomes easier and they can move from primary sites to distal secondary sites, thereby increasing the risk of recrudescence (Xing et al., 2019). Likewise, hepatocellular carcinoma (HCC) ranks as the fifth solid tumor found commonly around the globe and third leading cause of mortality with an incidence rate of more than two-thirds of currently available drugs are either obtained directly from natural origin or are developed based on the knowledge of their natural ingredients (Byahatti et al., 2018; Mohan et al., 2020). Several reports correlate the anticancer activity exhibited by medicinal plants to the presence of diverse metabolites. The study of discovering new natural products having anticancer potential is therefore important in synthesizing new chemicalderivatives established on the bioactivity and mechanism of action (Gómez et al., 2016, Iqbal et al., 2019, Abbasi et al., 2020). One such naturally occurring plant holding medicinal value is Parrotiopsis jacquemontiana (Decne) Rehder, (Hamamelidaceae) reported for alleviating several health disorders viz. liver and skin ailments, oxidative stress, inflammation, pain (Malik et al., 2011; Ali et al., 2017), wounds (Mir, 2014), dermatitis (Kumar et al., 2015), and microbial infections (Ali et al., 2018). In this study, six different extracts of P. jacquemontiana were screened for cytotoxic potential against different cancer cell lines and the best fraction to show maximum cytotoxic potential on the most susceptible cancer cell lines were chosen for further examining the mode of cell-death, cell-cycle arrest, migration, and invasion capabilities. Western blot analysis using specific antibodies for analyzing apoptotic, anti-apoptotic, cell-cycle arrest, and metastatic mechanisms was also conducted along with specific inhibition through STAT3 pharmacological inhibitor (Stattic).

2. Materials and methods

2.1. Plant sampling and processing

The leaves of P. jacquemontiana (Hamamelidaceae) were collected in the month of May and June 2018 from the forests located in upper Dir (altitude: 4,660 ft), KPK, Pakistan. The plant of interest was recognized, collected, and confirmed by the help of Dr. Syed Afzal Shah, a well-known plant taxonomist and botanist in the department of plant sciences, Quaid-i-Azam University (QAU), Isb, Pakistan. A voucher specimen #63214 was allocated at Pakistan herbarium, QAU, Isb, Pakistan. The plant material (P. jacquemontiana leaves) was shade-dried at room temperature and converted to fine- textured powder through Wiley mill (80 mesh-size). The obtained plant powder was three times (1:8; w/v)
extracted with 95% methanol for complete extraction of bioactive constituents. The filtrate obtained after evaporation of methanol using rotary evaporator (+40 °C) was termed as PJM. PJM extract (50 g) was suspended in water and the subsequent fractions were attained in order of escalating polarity by solvent-olvent extraction including n-Hexane (PHJ), Chloroform (PIC), Ethyl-acetate (PJE), Butanol (PJB), and Aqueous (PJA) fractions. Finally, the respective solvents were completely evaporated from fractions via rotary evaporator and the resulting plant extracts were stored at +4 °C.

2.2. Chemical reagents used

Tris-glycine, Sodium dodecyl sulphate (SDS), Ammonium persulphate (APS), EDTA, β-mercaptoethanol, Triton X-100, Crystal violet dye, Dulbecco’s modified eagle’s medium (DMEM), Trypsin EDTA, MTG reagent, Ribonuclease A (RNase A), Propidium iodide (PI), and Annexin-V FITC were purchased from Sigma-aldrich (St. Louis, MO: United States). Western blot membrane, TEMED, Laemmli sample buffer, and Acrylamide (40%) were obtained from BIO-Ras (Herculu, CA: United States). Likewise, Fetal bovine serum (FBS) was attained from Hyclone (Loughborough: United Kingdom). PBS from Vivantis technologies (Selangor: Malaysia), chemiluminescent substrate for western blot from Advansta (Menlo park, CA: United states), and Tween-20 from Merck & Co., Inc. Different antibodies including STAT3, p-STAT3 (Y705), p-STAT3 (Ser727), JAK1, p-JAK1 (Y1022/1023), JAK2, p-JAK2 (Yr1007/1008), Src, p-Src (Y416), cleaved Caspase-3 (Asp175), cleaved-PARP (Asp214), cleaved Caspase-8 (Asp391), cleaved Caspase-7 (Asp198), Cyclin D1, XIAP, Survivin, and β-actin belonged to Cell-Signaling Technology (Massachusetts: U.S.A) whereas PTP1b, Caspase-9, Caspase-8, Caspase-3, BCL-2, BAX, MPP-9, BAK, Cyclin E, MPP-2, C-myc, goat anti-mouse antibody-HRP, and goat anti-rabbit antibody-HRP were obtained from Santa-Cruz Biotechnology (Texas: U.S.A).

2.3. Cell lines and culture techniques used

Human hepatocellular carcinoma (HCCLM3) cell line was a gift from Professor. Zhao-You Tang, working at Liver-Cancer Institute of Zhongshan Hospital (Shanghai, China). Human mammary cancer (MDA-MB 231), normal mammary epithelial (MCF-10a), and normal liver cell line (LO2) was acquired from ATCC (American type culture collection), LO2, MDA-MB 231, and HCCLM3 were sub cultured in DMEM (Biowest, Nuaille, France), while MCF-10a cells were sub cultured in MEGM (mammary epithelial cell growth medium; Lonza Corporation). All the culture media’s were supplemented with FBS (10%; Biowest, Nuaille, France) and penicillin/streptomycin (1% of 100X; Thermofisher scientific, U.S.A). Respective cell lines were placed and maintained in humidified incubator provided with 5% CO2 at +37 °C.

2.4. Preparation of drug for cell culture

All fractions of P. jacquemontiana (80 mg) were suspended in 1 ml DMSO (dimethyl sulfoxide; sigma-aldrich, U.S.A) to make a stock solution and stored at + 20 °C. Upon use, the stock solution was diluted in culture media according to its requirement for different anticancer assays, but retaining the final concentration of DMSO to be administered as > 0.1% during the study course. Similarly, a stock solution (50 mM) of SC-203282 (Santa-Cruz Biotechnology, Dallas: USA) was prepared in DMSO and stored at + 4 °C. The serial dilutions of SC-203282, a pharmacological STAT3 inhibitor were prepared in cell culture medium prior to use in different anticancer assays following the same procedure as used for plant drugs.

2.5. MTT cell proliferative assay

MTT assay was used to determine the cell viability after treating with respective plant formulations against cancer cell lines. The cancer cells were seeded (10 × 10³ cells per well) in triplicates in 96-well plates overnight and then treated with 25, 50, 100, 200, and 400 μg/ml of P. jacquemontiana extracts for a time interval of 24, 48, and 72 hrs (hrs). After indicated time points, each well was provided with 20 μl MTT (5 mg/ml) and incubated for 3 hrs. The medium of wells was replaced by 100 μl DMSO to dissolve formazan crystals, after which optical density of each well was recorded at 570 nm using microplate reader (SPARK™ 10 M, TECAN: Switzerland). Cell viability was deliberated using the equation;

\[
\text{Cell viability} \% = \frac{\text{Abs. (sample)}}{\text{Abs. (control)}} \times 100
\]

2.6. GC/MS (gas chromatography/mass spectroscopy) analysis

The methanol extract of P. jacquemontiana (PJM) was analyzed via GC/MS for the manifestation of active components employing “Thermo GC-Trace Ultra Version: 5.0” gas chromatograph fixed with, “Thermo MS DSQ-II” for determination of mass. The components in test sample were separated by “Agilent DB-5, MS capillary Non-polar Column” with 60 m length and 0.25 μm film thickness. The temperature during the experiment was raised from 70 to 260 °C at 06 °C/min frequency. The injection volume of test sample was 1 μl and carrier gas was helium (1.5 ml/min flowrate). The mass spectra and respective retention times were matched with accurate samples acquired from Wiley Libraries spectra/NBS/NIST and the chemical constituents were identified (Cha et al., 2005).

2.7. AV-FITC/PI staining assay

HCCLM3 and MDA-MB 231 cells (0.5–0.6 × 10⁶) were seeded in culture petri-dishes (60 mm) overnight after which they were treated with PJM (100, 150, and 200 μg/ml) at 24, 48, and 72 hrs time interval. The drug treated/untreated cells were harvested through trypsinization procedure, gently washed with 1X-PBS (Phosphate Buffered Saline) and finally stained with Annexin-V-FITC Kit (Milenyi Biotech, Bergisch-Gladbach: Germany) for 15–20 min (min) according to manufacturers’ protocol. The samples to be tested were filtered and apoptosis cell percentage was analyzed using LSRFortessa™ flow cytometer (BD-Biosciences, San Diego: U.S. A). All the test samples were analyzed within 1 hr of Annexin-V/PI staining where each trial detected 10,000 events for each sample.

2.8. Cell-cycle arrest analysis assay

HCCLM3 and MDA-MB 231 cells were seeded (0.5–0.6 × 10⁶) overnight in culture cell dishes (60 mm) followed by specified drug (PJM) treatment at 24, 48, and 72 hrs time interval. The treated/untreated cells were harvested, fixed via ethanol (70%) for half an hrs at + 4 °C and stored at – 20 °C until required further. The ethanol-fixed cells were then washed following centrifugation and resuspended in 0.5 ml 1X PBS containing 1 μl/ml RNase A (Thermo-Fisher Scientific, Waltham: U.S.A) and 7 μl/sample PI (Santa-Cruz Biotechnology, Dallas: U.S.A). After placing the test samples in dark for 15 to 20 min, they were filtered and the cell distribution was investigated in various phases of cell-cycle through LSRFortessa™ flow cytometer (BD-Biosciences, San Diego: U.S.A).
U.S.A) and Summit-4.3 software (Beckman-Coulter, Inc.). All samples to be tested were analyzed within 1 hr of PI staining.

2.9. Wound scratch assay

HCCLM3 and MDA-MB 231 cells were seeded (7 × 10^4 cells/well) in micro dish-culture inserts (Ibidi-GmbH: Germany) overnight and incubated at +37 °C till the culture flask was confluent. The next day, micro dish-culture inserts were removed straight upward followed by careful washing with 1X-PBS (1 ml) to create cell-free gap area in the center. The respective cancer cell lines were treated with PJM (100, 150, and 200 μg/ml) and the pictures were captured using magnification lens (10X) at zero, 04, 08, and 12 hrs time interval employing Bright-field microscopy (Olympus: DP-70: Japan). The captured images were analyzed through Fiji Software (Schindelin et al., 2012), and normalized wound healing was deliberated using the equation:

Normalized wound healing (%) = \(\frac{\text{Distance of closed gap (treated \ μ dishes)}}{\text{Distance of closed gap (control \ μ dishes)}} \times 100\)

2.10. Matrigel invasion analysis assay

The Bio-coated Matrigel invasion chambers (Discovery, Labware Bedford U.S.A) were first hydrated and then equilibrated according to manufacturers’ guidelines. HCCLM3 and MDA-MB 231 cells were seeded at a density of 7 × 10^4 cells/well and resuspended in DMEM serum-free media after treating with indicated PJM concentrations (100, and 150 μg/ml) and the pictures were captured using magnification lens (10X) at zero, 04, 08, and 12 hrs time interval employing Bright-field microscopy (Olympus: DP-70: Japan). The captured images were analyzed through Fiji Software (Schindelin et al., 2012), and normalized wound healing was deliberated using the equation:

Normalized wound healing (%) = \(\frac{\text{Distance of closed gap (treated \ μ dishes)}}{\text{Distance of closed gap (control \ μ dishes)}} \times 100\)

2.11. Western blot analysis

HCCLM3 and MDA-MB 231 cells were seeded (1.3–1.7 × 10^6) overnight in 60 mm culture cell dishes followed by treatment with specified drug (PJM) concentration at indicated time intervals. The treated/untreated cells were harvested and lysed in M-PER (Thermo-Fisher Scientific, Waltham: U.S.A) coupled with addition of 1% 100X Halt-Protease and Phosphatase Inhibitor cocktail (Thermo-Fisher Scientific, Waltham: U.S.A) for 25 to 30 min placed on ice. The lysates were centrifuged at 13,300 rpm for 15 min to remove any insoluble material if present and the resultant protein was quantified through Bradford protein assay (Sigma-Aldrich, St. Louis: U.S.A). The respective lysates were diluted by adding 4X-Laemmli sample buffer (Bio-Rad Laboratories, California: U.S.A) followed by heating at 95 °C for 5 min for protein denaturation. The samples processed were stored at −20 °C till further required. For gel loading, equal quantity of protein (30–60 μg) for each sample was resolved onto SDS-PAGE gels (8–10 %) and transferred electrophoretically onto PVDF/nitrocellulose membrane (Bio-Rad Laboratories, California: U.S.A). After gel transferring, the membrane was blocked with Blocking One (Nasalai-Tesque, Kyoto: Japan) and finally probed with desired primary antibodies at + 4 °C overnight. The next day, respective blots were washed three times, incubated in HRP-conjugated (anti-rabbit/anti-mouse) secondary antibody for 1 hr and subsequently detected via Western-Bright Sirius Chemiluminescent-Detection Kit (Advanta, California: U.S.A). Chemi-Doc touch image system (Bio Rad Laboratories, California: U.S.A) was used to take chemiluminescent images. The blots were stripped in Restore™ stripping buffer (Thermo-Fisher Scientific, Waltham: U.S.A) followed by blocking in Blocking buffer, and reprobed with relevant primary antibodies where necessary. Fiji Software was used for densitometry analysis.

Fig. 1. Anti-proliferative effect of *P. jacquemontiana* fractions against (A) HCCLM3, and (B) MDA-MB 231 cell line at 24, 48, and 72 hrs. The results are communicated as Mean ± S.D (n = 3) of three independent experiments.
**3. Results**

### 3.1. PJM inhibits proliferation of cancer cell lines

MTT assay screening of *P. Jacquemontiana* fractions alongside cancer cell lines revealed significant anti-proliferative effects based upon dose and exposure time. The least IC50 value indicated more potent anti-proliferative activity while higher IC50 represented weak anti-proliferative activity. Among different *P. Jacquemontiana* extracts, PJM and PJC exhibited least IC50 values alongside HCCLM3 (corresponding to 188.2 ± 0.62, 118.7 ± 0.7, 170.9 ± 1.04 and 281.4 ± 5.58, 117.5 ± 5.33, 206.9 ± 4.15 μg/ml at 24, 48, and 72 hrs respectively). The least IC50 value indicated more potent anti-proliferative activity while higher IC50 represented weak anti-proliferative activity. Among different *P. Jacquemontiana* extracts, PJM and PJC exhibited least IC50 values.
However, the best anti-proliferative activity against MDA-MB 231 was displayed by PJM and PJE followed by PJC, PJH, PJB, and PJA (Fig. 1B). Likewise PJM dominated in all derived *P. jacquemontiana* fractions against HEPG2 and MCF-7 revealing an IC50 of > 1000, 804.7 ± 3.29, 650.8 ± 5.96 and 258.4 ± 2.83, 219.2 ± 2.77, 172.9 ± 4.33 μg/ml at 24, 48, and 72 hrs respectively. None of the extracts proved cytotoxic to normal liver and mammary cell line (LO2 and MCF-10A) (Supplementary Table 1, 2, and 3).

### 3.2. GC/MS analysis of PJM

A total of 51 compounds were identified via GC/MS analysis in the crude methanol fraction of *P. jacquemontiana* (PJM) corresponding to 93.9% of total detected compounds (Table 1). The chromatogram (Supplementary Fig. 1) displayed noticeable peaks in the specified retention time ranging from 11.0 to 41.3 min. The detected compounds in PJM belonged to 19 major classes including 5 carboxylic acids (35.7%), 7 fatty acids (11.9%), 4 phenols (11.2%), 8 esters (7.1%), 4 terpenoids (5.2%), 4 alcohols (5.0%), 4 amines (2.9%), 1 acridine (2.6%), 4 benzenoids (1.9%), 2 tocopherols (1.4%), 1 hydrazide (1.4%), 1 Ureas (1.2%), 1 benzo-furan (1.2%), 1 alkene (0.3%), 1 alkyn (0.3%), 1 azole (0.2%), 2 ketones (0.2%), 1 pyridine (0.06%), and 1 alkane (0.04%) respectively.

### 3.3. PJM induces apoptosis in HCCLM3 and MDA-MB 231 cells

The ability of PJM to initiate apoptosis was studied via flow-cytometry after treating HCCLM3 and MDA-MB231 with PI and Annexin V-FITC (Fig. 2). The cell population was distributed into four quadrants (Q1 to Q4) of which Q1 corresponded to necrotic cells, Q2 to late apoptotic cells, Q3 to live cells, and Q4 to early apoptotic cells. PJM induced apoptosis in cancer cell lines following a dose and time-dependent pattern. The dosage increase (100, 150, and 200 μg/ml) raised the percentage of early apoptotic cells as well as total apoptotic percentage with an increase in exposure time (24, 48, and 72 hrs). The maximum apoptotic percentage was noted for PJM (200 μg/ml) alongside MDA-MB231 conferring to 33.6 ± 2.0, 49.3 ± 4.0, and 73.6 ± 3.85 at 24, 48, and 72 hrs whereas 13.13 ± 1.95, 34.13 ± 3.95, and 72.36 ± 3.30 values were recorded besides HCCLM3 at similar time interval.

Fig. 2. PJM induces apoptosis in (A,B) HCCLM3 and MDA-MB 231 cells, (C,D) Graphical representation of PJM-induced apoptosis in HCCLM3 and MDA-MB 231 cells at 24, 48, and 72 hrs. The results are communicated as Mean ± S.D (n = 3) and the data was processed by one-way ANOVA followed by Dunnett’s multiple-comparison test. *** (significance level at p < 0.001) was noted for PJM concentrations compared to control.
3.4. PJM induced G0/G1 arrest in HCCLM3 and MDA-MB 231 cells

To verify whether the anti-proliferative effects of PJM alongside HCCLM3 and MDA-MB 231 cells were due to arrest of cells in cell-cycle, the flow-cytometer analysis was performed at 24, 48, and 72 hrs post treatment (Fig. 3). The treatment with PJM revealed a dose and time-dependent cell-cycle (G0/G1) growth arrest and apoptosis in respective cell lines. Though all doses of PJM (100, 150, and 200 \(\mu\)g/ml) induced highly significant alterations in cell cycle relative to untreated however, PJM (200 \(\mu\)g/ml) exhibited the maximum cell-growth arrest in subG1 phase conferring to 20, 34.5, and 71.9% alongside HCCLM3 and 16.5, 41.2, and 62.4% alongside MDA-MB 231 at specified time intervals, respectively. Treatment with PJM induced apoptosis at all concentrations, which was apparent from a reduced cell distribution in G1 and S phases of cell-cycle whereas an increased cell-accumulation in subG1 at 48 and 72 hrs interval.

3.5. PJM suppresses the migration of HCCLM3 and MDA-MB 231 cells

In vitro wound scratch assay was performed to determine anti-metastatic potential of PJM alongside HCCLM3 and MDA-MB 231 cell lines. Fig. 4 shows the treated cells with PJM (100, 150, and 200 \(\mu\)g/ml) to dramatically decrease (p > 0.001) wound-closure rates dose-dependently compared to control. All PJM concentrations were effective against HCCLM3, particularly PJM (200 \(\mu\)g/ml) showed 0.36, 2.15, and 2.87% wound closure rate compared to untreated exhibiting 13.1, 26.3, and 53.1% wound closure rates at 04, 08, and 12 hrs, respectively. Likewise, PJM (200 \(\mu\)g/ml) revealed minimum wound closure rates corresponding to 1.11, 2.23, and 5.57% relative to untreated showing 22.5, 53.9, and 84.8% against MDA-MB231 at similar time interval.

3.6. PJM suppresses the invasion of HCCLM3 and MDA-MB 231 cells

The anti-metastatic aptitude of PJM was evaluated by an in vitro metastasis model including Matrigel invasion chamber. Stromal cell derived factor-1 (SDF-1) was used as invasion stimulant for HCCLM3 and MDA-MB 231 cell lines. As presented in Fig. 5, the untreated cells spontaneously invaded through basement membrane in the absence of SDF-1 however the invasion through matrigel membrane was considerably increased (p > 0.001) in the presence of SDF-1. Co-treatment of PJM (100 and 150 \(\mu\)g/ml) with and without SDF-1 significantly reduced the invasive ability compared to untreated cells. HCCLM3 invaded cells after PJM (150 \(\mu\)g/ml) treatment were quantified as 17.4% and 24.9% to invade the artificial membrane provided without and with SDF-1 relative to untreated, taken as 100%. Likewise, PJM (150 \(\mu\)g/ml) revealed minimum wound closure rates corresponding to 15.8% and 32.8% of MDA-MB 231 cells to invade the matrigel membrane relative to untreated (100%).

3.7. PJM suppresses JAK/STAT3 signaling pathway in HCCLM3 and MDA-MB 231 cells

Western blot analysis of whole cell-lysates was done to identify the potential effect of PJM on JAK/STAT3 signaling pathway as shown in Figs. 6A and 7A. The protein levels of both the phosphorylated forms of STAT3 viz. p-STAT3 (Y705) and p-STAT3 (S727) were found to decline as early as 02 hrs post PJM treatment in HCCLM3 and MDA-MB231 cell lines. The levels of p-JAK1 and p-JAK2, which are the upstream proteins to phosphorylate STAT3, were also observed to decrease within 02 hrs of PJM treatment in both cell lines. Besides this, the p-Src level was reduced whereas STAT3 inhibitor levels (PTP1b, SHP1, SHP2, PIA51, and PIA53) were found to increase in both the cell lines, respectively. The total...
STAT3, Src, JAK1, and JAK2 levels remained constant in both cell lines even after 08 hrs of treatment suggesting PJM to downregulate JAK/STAT3 pathway.

3.8. JAK/STAT3 inhibitor studies of PJM

Owing to the PJM-induced inhibition of p-STAT3 within 2 hrs post treatment in western blot analysis assay and keeping in mind the expression of STAT3 in uncontrollable proliferation and apoptosis evasion, MTT proliferation assay along with G0/G1 flow-cytometry analysis was executed again but with Static (SC-203282; a pharmacological inhibitor of STAT3). This was performed to confirm whether the cytotoxic and apoptotic effects of PJM were mediated by STAT3 suppression. The effectiveness of Static (SC-203282) was established as an initial step through western blot analysis in HCCLM3 and MDA-MB231 cells. As presented in Figs. 6C and 7C, the p-STAT3 level of PJM in combination with Static is significantly higher than PJM only treated cells but similar with untreated cells. The results put forward the effectiveness of 10 μM of Static (SC-203282) in activating PJM-induced suppression of STAT3 in HCCLM3 and MDA-MB231 cell lines.

In MTT assay, treatment of PJM (200 μg/ml) in combination with Static (SC-203282) significantly reduced the anti-proliferative action against HCCLM3 cells at 24 and 48 hrs whereas PJM (100 and 150 μg/ml) did not exhibit any difference in cytotoxic effects compared to PJM alone (Fig. 6D). In case of MDA-MB231 cell line, PJM (200 μg/ml) in combination with Static (SC-203282) markedly reduced the anti-proliferative effects at 24, 48, and 72 hrs whereas PJM (100 and 150 μg/ml) only demonstrated noticeable cytotoxicity at 24th hr relative to PJM alone treatment as evident in Fig. 7D. Similar results were displayed by flow cytometry.
analysis which revealed a significant decline in subG1 population of HCCLM3 cells after collective treatment of PJM (150 and 200 µg/ml) and Stattic (SC-203282) compared to PJM alone treatment at 48th hr. While all the selected PJM (100, 150, and 200 µg/ml) concentrations in combination with Stattic (SC-203282) clearly decreased the distribution of MDA-MB231 cells in subG1 compared to PJM alone treatment at same time interval (Figs. 6E and 7E). The results obtained suggest the reduced anti-proliferative and apoptotic effects with PJM-induced suppression of STAT3 which provide some evidence that anti-cancer effect of PJM is partially mediated by STAT3 inhibition.

3.9. PJM regulates anti-metastatic and cell-cycle proteins

The level of those proteins heavily connected with tumor metastasis and invasion i.e. MMP9 and MMP2 proteins was quantified alongside HCCLM3 and MDA-MB231 cell lines after treating with PJM (100, 150, and 200 µg/ml) as shown in Figs. 8A and 9A. Both MMP9 and MMP2 expression levels were distinctly reduced in a dose-dependent style, suggesting PJM drug to have an anti-migration and anti-invasive role in both cancer cell lines. Besides this, the expression levels of cyclinD1 and cyclinE were also quantified based upon the key role they play in regulating the transition of cells in cell-cycle from G1 to S phase. PJM clearly decreased the protein expression of cyclinD1 and cyclinE dose-dependently (100, 150, and 200 µg/ml) against selected cancer cell lines (Figs. 8A and 9A) suggesting its potential inhibitory effects on cell-cycle progression.

3.10. PJM regulates pro-apoptotic and anti-apoptotic proteins

The pro-apoptotic pathway of PJM was further illustrated in HCCLM3 and MDA-MB 231 cell lines through western blotting analysis of those proteins associated with apoptosis in addition to flow cytometric analysis assays. Accordingly, the expression levels of main pro-apoptotic proteins including BAK, BAX,
cleaved-PARP, cleaved caspase 9, and cleaved caspase 3 were monitored dose-dependently (100, 150, and 200 μg/ml of PJM) as presented in Figs. 8B and 9B. The main anti-apoptotic proteins i.e. XIAP, survivin, and BCL-XL, known to play fundamental roles in cell survival and cell proliferation was also detected. In addition to this, cleaved caspase-7 and 8, well-recognized to play significant roles in apoptotic pathway was also investigated. The attained results revealed the expression levels of pro-apoptotic proteins to increase substantially whereas the anti-apoptotic protein levels to decrease noticeably in a concentration-dependent pattern of PJM treatment. Expression levels of cleaved caspase-7 and 8 were also markedly elevated in a dose-dependent style. Given altogether, these results coincide with the former observed pro-apoptotic effects of PJM analyzed via flow cytometry, respectively.

4. Discussion

Since ancient times, medicinal aromatic plants have been used traditionally in every part of the world and nowadays an upsurge of interest is found by using these plant and their derived products.
virtually in all sectors and industries (Gezici et al., 2017; Tewari et al., 2015). In addition to using medicinal plants for traditional medical purposes, the plants are viewed by researchers as major sources of interest for the isolation and identification of bio-active compounds leading to the development/production of novel drugs with efficacy and safety (Tavakoli et al., 2017; Lee et al., 2014). The secondary metabolites found in medicinal plants in the form of crude extracts are well-recognized for their biological activities including chemo-preventive agents against cancer. Many researches have shown the historical medicinal background of Asian countries to utilize medicinal plants and their products in cancer treatment (Sajid et al., 2018).

The present study reports the molecular mechanistic studies of the anticancer aptitude of P. jacquemontiana most effective extract via MTT, Annexin-V/PI, G0/G1 cell-cycle arrest, anti-metastatic, and western blot analysis assays alongside selected cancer cell lines (HCCLM3 and MDA-MB 231). Here the term “most effective extract” means the fraction having maximum toxicity against cancer cell lines and the one providing least IC50 value contributing to 50% of cell death. MTT cell-viability assay depicted the methanol extract of P. jacquemontiana (PJM) to exhibit greater cytotoxicity relative to other plant extracts, conferring to least IC50 value shown against all tested cancer cell lines (HCCLM3, MDA-MB 231, MCF-7, and HEPG2) at 24, 48, and 72 hrs. At the same time, the toxic effect of PJM against normal cell lines (LO2 and MCF-10A) appeared less due to a great difference between the IC50 values calculated for cancer and normal cells. The co-treatment of PJM along with conventional chemo-therapeutic drugs may be useful in lessening the potential dosage of the chemo-therapeutic drugs and eliminating the adverse side-effects; however further studies regarding this perspective would be required.

The chemical composition analysis of PJM (P. jacquemontiana methanol fraction) was determined via GC/MS. A total of 53 compounds were detected in PJM of which Octadecanoic-acid and
Octadecanoic acid methyl-ester are reported as antifungal, antibacterial, and anti-tumor agents (Abou-Elela et al., 2009; Hsouna et al., 2011). α-linolenic acid methyl-ester possesses antimicrobial, antioxidant, hypercholesterolemic, and anticancer activity (Kumar et al., 2010). It is also reported previously to inhibit proliferation of ER-negative and ER-positive breast cancer cells and is a well-documented potent antiangiogenic agent in colorectal cancer and HUVEC cells (Oyugi et al., 2011). However, the most abundant compound identified in PJM was Gallic acid (29.6%), which has numerous industrial applications especially its role in pharmaceutical industry as a standard for the determination of phenolic constituents in analytes. Gallic acid and its derivatives have combatting potential against oxidative stress, microbial infestations, and cancer manifestations. Moreover, gallic acid is reported as a well-known antioxidant, hepatoprotective, antiviral, antimicrobial, anti-inflammatory, and anticancer agent (Nayeem et al., 2016).

An emerging strategy for cancer treatment nowadays is the capability of inducing cellular apoptosis in cancer cells and involves various signaling pathways (Lou et al., 2019). The cells undergoing apoptosis shows the externalization of phosphatidylserine (PS) towards the outer side from inner layer and is considered the bio-chemical feature of cellular apoptosis. For determination of apoptosis, Annexin-V (a lipoprotein specifically binding to
PS) was tagged with FITC (Komonrit and Banjerdpongchai, 2018). In our study, PJM-treated HCCLM3 and MDA-MB 231 cells for 24, 48, and 72 hrs were stained with Annexin-V-FITC/PI and analyzed via flow cytometry. Dot-blot analysis of PJM-treated HCCLM3 and MDA-MB 231 cells showed regulated cell death by an increase in both late (Q2) and early apoptotic (Q4) stained cells following a dose-dependent pattern but without a significant effect on cell-necrosis quadrant (Q1). This hypothesizes that cell death was most probably regulated via cellular apoptosis mechanism and not by necrosis.

The increase in apoptotic cells was further confirmed by G0/G1 cell-cycle arrest analysis assay, which indicated a considerable accumulation of SubG1 arrested cells post PJM treatment at 24, 48, and 72 hrs whereas a consequent decrease in G1 phase cells. Cell-cycle progression is a main biological event, which has controlled regulation in normal cells but becomes almost universally deregulated or aberrant in neoplastic and transformed cells (Birjandian et al., 2018). Regarding this, the potential prognostic roles played by cell-cycle regulators and natural products in cancer therapy are under focus. Cell-cycle arrest caused by PJM treatment
in HCCLM3 and MDA--MB 231 cancer cells was evident from decreased protein expression levels of CyclinD1 and CyclinE. Both of these proteins are required essentially by cell-cycle for pushing the cells from G1 to S phase (Kannaiyan et al., 2011), whereby their inhibition blocks the entry of cells to further progress into other phases of cell-cycle and hence causes the greater accumulation of cells in SubG1 phase. This regulation of cell-cycle is an important method for controlling propagation of tumors (Xavier et al., 2009).

Besides this, in the cells undergoing apoptotic cell death, caspases are stimulated leading to activation of further reactions. Among caspases, caspase-3 functions to proteolyze the structural cytoskeleton proteins (laminin, actin, and fodrin) and leads to apoptosis accompanied with typical morphology (Komorrit and Banjerdpongchai, 2018). On the other hand, caspase-8 and 9 play key roles in initiating the cascade of extrinsic apoptotic (death receptor) and intrinsic apoptotic (mitochondrial) pathway, respectively (Fulda and Debatin, 2006). Our study showed the PJM-treated HCCLM3 and MDA-MB 231 cells to demonstrate markedly enhanced levels of all three afore-mentioned caspases in addition to suppressing tumor promoting and survival proteins (Bcl-xl, survivin, and XIAP), and activating pro-apoptotic proteins (BAX, and BAK), hence strengthening the hypothesis that PJM induces HCCLM3 and MDA-MB 231 cells to regulate cell death by undergoing apoptosis.

Furthermore, the anti-metastatic effects of PJM were evaluated via wound-scratch and matrigel invasion chamber assays. Metastasis process involves many signaling pathways and is considered a multi-step process promoting the migration of cancer cells to distant organ sites. Matrix metalloproteinases (MMPs) are a family of structurally and functionally related Zinc-dependent enzymes, which play vital roles in promotion of metastasis and tumor growth. Amongst the MMPs, MMP2 and MMP9 are extensively involved in the facilitation of cancer metastasis (Luo et al., 2019). The inhibitory effect of PJM on MMP2 and MMP9 was validated via western blotting, which revealed the expression of both MMPs to decrease considerably after PJM treatment at 24 hrs dose-
dependently. Moreover, the protein expression level of pSrc was down-regulated in PJM-treated HCCLM3 and MDA-MB 231 cells, as reported elsewhere that the signal transducers (i.e. FAK and Src) contribute in the secretion of above MMPs and their expression levels are relatively found higher in cancer cells. This specifies the anti-metastatic effect of PJM to be mediated through inhibition of Src (Sigstedt et al., 2008).

Several hallmarks of cancer including uncontrollable cellular proliferation, metastasis/invasion, and apoptotic resistance are linked with dysregulated STAT3 signaling. STAT3 has been reconciled and validated as a drug target and is considered an important cell-signaling pathway in cancer therapeutics. Constitutively expressed STAT3 has been documented to trigger deregulated transcription of genes leading to gene products that consequently promote tumor progressions. STAT3 along with its downstream targets not only facilitate proliferation, survival, angiogenesis, and metastasis, but also suppress antitumor immune responses (Arora et al., 2018). Our study showed the suppression of JAK/STAT3 signaling pathway in HCCLM3 and MDA-MB 231 cells after treatment with PJM. Activation of STAT3 may be deteriorated in these cancer cells via interrupting with upstream kinases i.e. c-Src, JAK1, and JAK2. Their inhibition prompts apoptosis, increases cell-aggregation in subG1 phase of cell-cycle, and activates caspase 3 resulting in cell death (Kannaiyan et al., 2011). Our results also present the downregulation of p-Src, p-JAK1, and p-JAK2 following PJM treatment, which may further support Src and JAK-mediated inhibition of STAT3. The inhibitory effects displayed by PJM on STAT3 expression may be ascribed to its antioxidant nature (Ali et al., 2017) keeping in view the induction of STAT3 activation by ROS (reactive oxygen species) through Src, JAK, and NF-kB signaling and its reversal by antioxidant reagents (Song et al., 2017). Previous studies have also associated the abrogation of JAK/STAT3

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**Fig. 9.** Western blot analysis showing the effect of PJM on (A) Metastatic and cell-cycle proteins (B) pro-apoptotic and anti-apoptotic proteins against MDA-MB 231 cell line. MDA-MB 231 cells were treated with 100, 150, and 200 μg/ml for 24 hrs. Proteins were equally loaded onto the wells, also confirmed by β-actin loading control. Fold change was calculated using Fiji ImageJ software.
signaling pathway to induce cell-cycle arrest and apoptosis in vitro (Mackenzie et al., 2008), inhibit growth of tumors, and attain tumor elimination accompanied with improved survival rate in human HL murine models (Ju et al., 2016).

It is well-documented previously that inhibition of STAT3 activation requires the activity of PTP, SOCS, SHP-1/2, and PIAS, all of which negatively regulate STAT3 cell-signaling cascade and can effectively prevent the progression of cancer (Lee et al., 2017; Wu et al., 2019). Our study shows the up-regulation of PIAS1/3, SHP-1/2, and PTP-1β proteins in HCCLM3 and MDA-MB 231 cells after PJM treatment. PIAS protein(s) may regulate transcription via several mechanisms i.e. blocking DNA-binding activity of transcriptional factors, recruitment of transcription repressors, and promotion of protein SUMOylation. PIAS-1/3 binds to STAT-1/3 and can therefore inhibit their transcriptional activity. Several studies have shown the upregulation of PIAS-3 expression to inhibit cell-proliferation and enhance drug chemo-sensitivity in various tumors (Wu et al., 2019). For example, the inhibition of constitutively expressed STAT3 via curcumin attenuated tumor growth by over-expressing PIAS-3 in endometrial and ovarian cancer (Saydmohammed et al., 2010), the over-expression of PIAS-3 in prostate cancer induced cellular apoptosis in in vitro and in vivo assays (Wible et al., 2002), and the over-expression of PIAS-3 contributed to suppression of lung cancer and restored drug chemo-sensitivity. The above findings indicate that PIAS-3 may serve as an attractive candidate for JAK/STAT3 pathway targeting and restore the sensitivity to chemo-therapeutic drugs in cancer therapy (Wu et al., 2019). Apart from this, the tumor-suppressor activity of SHP-1 is mediated via negative regulation of STAT and JAK. Tai et al., (2012) reported dovitinib; a receptor-kinase inhibitor in sorafenib-resistant HCC to overcome sorafenib-resistance and induce apoptosis via SHP-1 mediated STAT3 signaling inhibition. Another study included plumbagin (Vitamin K3 analogue) to induce SHP-1 over-expression in human myeloma cells leading to prevention of STAT3 phosphorylation through the inactivation of c-Src, JAK-1, and JAK-2 (Sandur et al., 2010). In light of the above findings which suggest STAT3 activity to be vital to tumorigenesis and drug-resistance, the activating of SHP-1 by PJM may be a valuable applicant in cancer therapy.

Finally, we hereby report the STAT3 inhibition following PJM treatment as well as the suppression of STAT3 inhibition by using a specific inhibitor of STAT3 (Stattic; SC-203282), which blocked PJM-induced apoptosis. The inhibitory effect of PJM and SC-203282 was visualized through MTT, cell-cycle arrest analysis, and western blot analysis which suggested the sole treatment of PJM and SC-203282 to inhibit STAT3 activation whereas their combined treatment cancelled the inhibitory action of each other leading to the same phenotype as that of untreated. These observations clearly suggest that PJM most likely regulates cell death through the suppression of JAK/STAT3 pathway.

5. Conclusion

Overall our results direct PJM to inhibit the growth of wide-variety of cancer cells and induce apoptosis by down-regulating anti-apoptotic gene products, modulating cell-cycle proteins, activating caspases, and suppressing JAK/STAT3 cell-signaling pathway (Fig. 10). Owing to the significant medicinal value shown by PJM, the plant extract should further be explored for its anti-cancer aptitude at advanced level regarding its role in metastatic inhibition, immune response modulation for reducing tumor, and inducing apoptosis in suitable animal models.

Ethics approval
Not applicable.

Consent for publication
Not applicable.

Availability of data and materials
The data presented in this manuscript belongs to the research work done by Ms. Saima Ali under the supervision of Dr. Muhammad Rashid Khan and has not been submitted/published elsewhere.
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