Evaluation of Validated Herbal Medicine Formulation for Anti-Proliferative and Apoptotic Activities Used In the Tribal Medicine System

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Abstract: The current study was carried out to appraise the Anti-proliferative activity of Herbal Medicine Formulation (HMF) comprising different components of ethno-medicinal plant drugs which are practiced by the local healers in the Tribal Medicine System (TMS). The medicine formulation was subjected for validation with the authorized Ayurvedic Medical Practitioner in order to ascertain the active resources for novel lead constituents present in the drug. In addition, the validated HMF will also facilitate the movement of indigenous practices of traditional medicines to the public domain. The analysis was made on preliminary phyto-chemicals using fractions of HMF drug which reveals the active presence and were known to demonstrate both biomedical and physiological activities. Subsequently, the analysis of Thin Layer Chromatographic confirms the presence of diversified bioactive ingredients in the different components of HMF drug which indicates the efficacy of the HMF drug. The active fractions were prepared by serial extraction with ethanol, methanol, petroleum ether and ethyl acetate respectively based on the standard protocols and tested against MCF-7 (Mammarian Cancer cells) followed by HeLa cancer cells lines in vitro and their probable mechanism of action was critically analyzed. Further, the anti-proliferative activity was evaluated with these diversified HMF drug fractions against cancer cell lines by MTT assay, Trypan Blue assay and Hoechst’s staining methods respectively. The apoptotic effects was determined and correlated for its anti-cancer activity in the extracts of HMF drug. The TLC of extracts showed that, chloroform extract moved to the maximum distance of 15cm followed by ethyl acetate extract with a 14.7 cm as retention factor. Whereas, the ethanolic extract of HMF drug moved the least distance with 6.5cm retention factor and aqueous extract did not show any movement on the stationary phase. The analysis for MTT assay demonstrated that, the cell viability was diminished with the increased concentration of the HMF drug. The inhibition concentration value (IC₅₀) for the MTT assay at 24hrs was found to be 5.1µg/ml and for 48hrs at 5.4µg/ml. Later, the attentiveness of HMF drug against both MCF-7 and HeLa cancer cells was instituted to appraise the anti-proliferative activity. The arresting of cell proliferation and growth of these cancerous cells was observed even in very low concentration of the HMF drug, which may be due to the presence of bio-active constituents in the medicine formulation. Subsequently, the apoptotic assays showed that, the increased fractions of HMF drug (60, 75, 80 and 95%) induced mitochondrial depolarization in MCF cells correspondingly, at those fractions which could have triggered the apoptosis in mitochondrial pathway. Eventually, the ability of curbing and inhibition capability of the free radicals was assessed in the different fractions of HMF drug through biochemical method like, ABTS scavenging assays respectively. The result reveals; significant ABTS free radical scavenging activity of 74±0.74 was evident in the extract. Hence, the HMF drug was found to be most effective in deactivating these Cancer Cells and justifies the tested herbal formulation, HMF being practiced by the tribal healers in their traditional medicine system. Further, the formulation needs to be evaluated for its clinical trials through the absolute purification process in order to propose the medicine as most potential anticancer herbal drug.

Key Words: Anti-proliferative activity; Apoptosis; Antioxidant activity; Ethno-medicinal plant drugs, Herbal Medicine Formula; HeLa and MCF-7 Cell lines
I. Introduction

The natural products derived mainly from plants and propose a diverse sequence of active constituents which plays a key role by regulating the biological organization with these natural antioxidants. The remedy using chief constituents of herbal formulation has been significantly accomplished which can facilitate development of most desired drug by breakthrough technology for the benefit of mankind. The large interest is currently being paid by the researchers to explore most potent natural products for their interesting counterpart activities against different ailments apart from anticancer drug research (Van, 1993).

Furthermore, the implication of wide range of medicines towards management and prevention of cancer and associated ailments are explicitly derived either from indigenous or aboriginal plants contribute around 25% of its total effectiveness. It was estimated that, nearly 60-80% of the drugs approved for cancer related therapeutics are derived from plants only. This has protracted the pursuit of efficient antioxidant and anticancer agents from natural sources particularly medicinal plants which are practiced by some ethnic group (Mishra et al., 2008; Boopathy and Karthiresen, 2010). However, the investigations on lead constituents from pant drugs have been recuperated the status by means of getting superior perceptive of their biological importance such as antimicrobial, antioxidant, radical scavenging, anti-proliferative activities (Manian et al., 2008; Caamal, 2011; Gul et al., 2013).

The ethno-medicinal plants were appraise all the way through systematic screening protocols and possess an imperative position in the drug invention and many innovative drugs have been justified with respect to effectiveness of their formulations in different cultures of traditional medicine system. Even though, the beneficial aspects of both synthetic and chemo-typing profile along with molecular modeling are in the front line. Similarly, the medicinal plants remain an essential source of any new-fangled drugs, which further leads to new chemical lead constituents along with the vital role in the biological system. Therefore, bio-prospecting of antitumor drugs from natural products is escalated now and receiving outstanding interest globally due to its target specificity and have no side effects in the therapeutic strategies (Sharma et al., 2011; Sanaz et al., 2012).

Generally, cancer’ is a very serious health setback and patients are confronted with undesirable side effects resulting from conventional treatments. The complementary and alternative medicines are optional choices. Some herbal or traditional formulations from oriental medicines in some parts of India were found to be evidence for anti-cancer activities such as, anti-proliferation, anti-angiogenesis and apoptosis (Van, 1993; Manian et al., 2008; Atjansuppap et al., 2009; Al-Rashidi et al., 2011). In Karnataka, many herbal, folklore medicines, and traditional medicine formulas have been used extensively as complementary medicines without any proven evidence of their effectiveness in the biological system. Hence, after interviewing traditional or tribal medicine men in the province of Biligirirangana Hills (Chamarajanagara district of Karnataka, it was noticed practically that, an herbal formulation (pre-prepared) used to treat cancer related ailments such as, breast, lung and liver cancers was preferred in the study. This tribal herbal formulation merits investigation for the anti-proliferative activity of its crude-aqueous extracts and its each plant component of the Herbal Medicine formulation against human cancer cells (Ravishankar and Murthy, 2011).

Mammalian cancer or breast cancer is the most numerous malignancies among women are the leading cause of death due to cancer related ailments and the consequential of the metastatic development of primary stage of cell tumors are of great concern (Jemal et al., 2006). The possible exploration of plants based medicine formulations in the treatment of different diseases together with cancers is inevitable. This would be the very basis for innovations via modern medical science, as they are considered to be vast sources of new-fangled drugs (Jones et al., 2006; Kim, 2008; Aune et al., 2009; Hasan et al., 2011; Woo and Kim, 2011). The discovery and production of drugs has been dominated by the synthetic chemistry which further facilitate both target specific and non-target drugs. Therefore, the specific protocol was established to ascertain and production of the potential drugs from bioactive plants and their extracts of active fractions will create a platform to provide new and novel products for disease management via strategic treatments but, prevention is still sizeable. Besides, the antitumor area has the greatest impact by the active constituents derived from plant drugs, where drugs like vinblastine, vincristine, taxol, and camptothecin have improved the chemotherapy of some cancers in a remarkable way.

The plants based medicines are stable which have unlimited capacity to produce active constituents that in turn attract researchers in the quest for innovative, novel and active chemotherapeutics (Lampronti et al., 2003; Ravelo et al., 2004; Itharat, 2004; Jemal, 2006; Aune et al., 2009). The long-lasting search for new-fangled anticancer lead molecules in plant medicines and traditional foods is a realistic and promising strategy for its prevention (Anusha and Murthy, 2010) in the daily life. But, in accordance with this worldwide trend of cancer related ailments and increased mortality rate, the current study was undertaken to evaluate the extracts of validated Herbal Medicine Formulation (HMF) for anti-proliferative and apoptotic activities which are being practiced by Tribal healers in the Traditional Medicine System (TMS) for Cancer and related ailments.
Ethno-medicinal plants

The different parts of five ethno-medicinal plant drugs, *A. serphyllifolia* (leaves), *D. hispida* (tubers); *G. mauritiana* (leaves); *N. nimmoniana* (leaves) and *R. densiflora* (whole plant) were collected from different tracts/regions of B.R. Hills of Chamaraja Nagara districts of Karnataka (Fig. 1A-H).

**Materials and Methods**

**Ethno-medicinal plant drugs**

The different parts of five ethno-medicinal plant drugs, *A. serphyllifolia* (leaves), *D. hispida* (tubers); *G. mauritiana* (leaves); *N. nimmoniana* (leaves) and *R. densiflora* (whole plant) were collected from different tracts/regions of B.R. Hills of Chamaraja Nagara districts of Karnataka (Fig. 1A-H).

**Instruments**

The instruments such as, Electric blender, Microscope (Olympus), Muffle furnace (Meta lab, Scientific industries, Mumbai), Centrifuge (REMI R-4C & 8RC Centrifuge Machine Bengaluru), Soxhlets Apparatus (Multiple Units), Flash Evaporator, Shaker incubator, Laminar Air Flow (Meditech, Chennai), Spectrophotometer (UV/Visible, Elico Limited, Hyderabad) etc. were employed in the study.

**Chemicals**

In the study, the standard chemicals of analytical grade were used namely, Picric acid, α-napthol, Benedict’s reagent, 5% Ferric chloride, 1%Gelatin, 10% sodium hydroxide, Alcohol, Biuret’s reagent, Ninhydrine reagent, Lead acetate, NaOH, Conc.H2SO4, copper sulphate, dimethylsulfoxide (DMSO), di-sodium hydrogen orthophosphate (Na2HPO4), ethylenediamine tetracetic acid (EDTA), Folin Ciocalteu’s phenol (FC) reagent. Besides, the chemicals used in the study were of, 3(4,5-dimethylthiazol-2-yl)2,5-diphenyl-tetrazolium bromide (MTT), rutin, gallic acid followed by ferric chloride (FeCl3), hydrogen peroxide (H2O2), Hydroxylamine hydrochloride, sodium dodecyl sulphate (SDS), magnesium sulphate (MgSO4), potassium chloride (KCl), magnesium chloride (MgCl2), sodium hydroxide (NaOH), sodium chloride (NaCl), sodium dihydrogen-orthophosphate (NaH2PO4), trichloroacetic acid (TCA), and Tris-HCl were obtained from Sisco Research Lab., (Mumbai, India). Additionally, other specific solvents and reagents of analytical grade were used in the studies, which were procured from authorized S.D. Fine chemicals Pvt. Ltd., followed by Merck, India. The media like, agar, cholesterol, hypochlorite solution, proteose peptone and yeast extract were procured from Hi-Media, (Mumbai), India (Deng et al., 2006; Zakaria et al., 2011).

**Interaction with Tribal Medicine Men and collection of Ethno-medicinal plants**

The interactions were conducted intermittently with Tribal Medicine Men at the different locale of Biligirirangana Hills, Karnataka (during the period, 2013-2014) with a semi-structured questionnaire. The data on Medicine formulation and responsible plant components were documented and the Herbal Medicine Formulations (HMF) and individual ethno-medicinal plant materials were obtained from the Tribal Medicine Men (Fig. 3A-J). The particulars on atypical practiced medicine formulation comprising different parts of ethno-medicinal plants such as, *Andrographis serphyllifolia*, Vahl (leaves); *Discorea hispida*, Dennst, (tubers); *Glycosmis mauritiana* Tanaka, (leaves); *Notaphyodes nimmoniana* Blume (leaves) and *Rauwolfia densiflora* (Wall.) Benth & Hook (whole plant) respectively were explicitly collected during the interaction. The plants species were identified and authenticated by consulting a taxonomist followed by standard flora, correspondingly, the plant materials were deposited at Bhoomigeetha Institute of Research & Development, (Tumkur), Karnataka, India. The baseline informations of selected ethno-medicinal plants are represented in the Table 1.

**Validation of Tribal Medicine Formulation (HMF)**

The different plants samples of Tribal medicine formulation were scientifically validated based on their physical characteristics in association with an authorized Ayurvedic practitioner, Nisarga Ayurveda Research Foundation, Sakaleshpur, Hassan district, India (Table 1). The standard protocols were identified and the methodology was employed in the present study based on the descriptions of Chaithra (2013).

**Preparation and Processing of HMF**

The collected ethno-medicinal plant materials of HMF were subjected for unraveling different desirable parts like, leaves, stem, root/ tubers from the main plants or whole plant parts. The different parts of the selected ethno-medicinal plant drug materials were subjected for shade drying for 20 days to ensure that, the active constituents were free from decomposition and possibility of photo-chemical degradation was also monitored. The whole HMF formula and its components were extracted by following the traditional process i.e., boiling with water for 15 minutes followed by filtering the same using muslin cloth then, the filtrates were dried by means of lyophilization processes. Further, the dried extracts of HMF were examined to determine antioxidant, anti-proliferative and apoptotic activities as per the standard analytical procedures (Anonymous, 2002).

**Preparation of Solvent extracts**

The air-dried components of the HMF drug were powdered using a suitable mechanical grinder to obtain a coarse powder, which was then subjected to successive solvent extraction with ethanol, methanol and petroleum ether in a soxhlet apparatus. The material was dried in hot air oven at 40°C each time before extracting with the
next solvent. The extracts were then filtered through a Whatman No.1 filter paper and concentrated to the dry mass using rotary evaporator. The extraction process was repeated for three times at different time intervals. The yield of each extract was measured and residues were stored in dark glass tubes for further analysis.

**Phyto-chemical analysis of EMP and HMF**

The extracts from ethno-medicinal plant (EMP) drugs and Tribal Medicine Formulation (HMF) were used for the Phyto-chemical analysis qualitatively for the detection of carbohydrates, proteins followed by the secondary metabolites like alkaloids, flavonoids, terpenoids, steroids, tannins, saponins and total phenols etc. The aqueous extracts of the plant was subjected to qualitative chemical screening for the identification of the alkaloids, flavonoids and tannins using standard procedures (Trease, G. E.; and Evans –2004; Mondal et al., 2013).

**ABTS radical scavenging activity**

The ABTS assay (2, 2-azinobis 3-ethylbenzothiazoline-6-sulfonic acid) was performed by preparing a stock solution which is consisting of 7 mM ABTS solution and 2.45 mM potassium persulfate solution at equal proportion. This was subjected for incubation at room temperature for 12h during dark condition that was further yielded a dark colored solution which contains radicals of ABTS. Subsequently, to perform each assay, the fresh working solution was prepared by mixing stock solution with methanol (50%) and the initial absorbance was 0.700 (± 0.02) at 745 nm at 30°C temperature. Further, the extracts of ethno-medicinal plant drugs and tribal medicine formulation were used at variable concentrations (50-3000μg/ml). Then, these concentrations were subjected for reaction with known volume (3 ml) of ABTS solution and the absorbances were taken at 734 nm. Meanwhile, the ascorbic acid was taken as positive control. Finally, the radical scavenging activity was assessed based on the percent activity of ABTS and calculated the value as per the standard formula (Re et al., 1999).

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\text{ABTS radical scavenging activity (\%) = \frac{\text{Control OD} - \text{Sample OD} \times 100}{\text{Control OD}}}
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**Thin Layer chromatography (TLC)**

Thin layer chromatography (TLC) is a chromatography technique used to separate diverse concoction bioactive constituents. Thin layer chromatography is performed on a sheet of glass, plastic, or aluminum foil, which is coated with a thin layer of adsorbent material, usually silica gel, aluminium oxide, or cellulose. This layer of adsorbent is known as the stationary phase. After the sample has been applied on the plate, a solvent or solvent mixture (known as the mobile phase) is drawn up the plate via capillary action. Because different analytes ascend the TLC plate at different rates, separation is achieved and the compounds in the extracts gets separated based on their affinity. The fluorescence bands were observed at 254nm (short wavelength) in UV light followed by 366 nm (long wavelength) using UV light in the respective plates. After the run, plates were dried and sprayed with NP/PEG reagents were used to detect the bands on the TLC plates and the observation of chromatograms was done under long wavelength UV followed by visible light. The movement of the active compound was expressed by its retention factor (Rf), values were calculated for different samples (Wagner and Bladt, 2009).

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\text{Rf} = \frac{\text{Distance travelled by the solute}}{\text{Distance travelled by the solvent front TLC plates}}
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**Cell Lines and Cell Culture Preparation**

The Cell- lines used in this study were of estrogen receptor–positive ‘MCF-7’ breast cancer cells and cervical ‘HeLa’ cancer cells. The MCF-7 cancer cells were cultured in 89% DMEM and 10% FBS along with 1% penicillin/streptomycin. Similarly, the HeLa, cells were cultured in 89% RPMI 1640 with 10% FBS and 1% penicillin/streptomycin. All the cells were cultured at 37 °C at 95% humidity and 5% CO_2 for 3 days as they reached 80%-90% confluency. Subsequently, the spent medium was removed and replaced with fresh medium and incubated again for 24 h. The cell cultures were then washed with PBS 1–2 times and were suspended using trypsin-EDTA and finally, fresh medium was added to the cells (Evan and Vousden, 2001).

**Anti-proliferative activity**

Anti-proliferative activity was measured by MTT assay and that was performed using 96-well plate at a cell density of 3 x 103 cells/well. The cancer cells, MCF-7 and HeLa were subjected for analysis using pre-cultured by Minimum Essential Medium (MEM) containing 10% Fetal Bovine Serum (FBS) and as per their specificity described above. All the cell lines were pre-cultured for 24 h before confronting with variable concentration of the extracts i.e., 10, 30, 100, 300, and 1,000 μg/ml. Then, phosphate buffer saline (PBS) was used as negative control and adriamycin was used as positive control, finally subjected for incubation as per the
protocol explained above (Nesaretnam et al., 1998; Sun et al., 2002; Giri et al., 2006; Siripong et al., 2006; Hu et al., 2011). After the incubation period (72 h), the treated cells were added with MTT reagent then incubated again for 3h and the formozan salts were dissolved with DMSO. The absorption was measured at 550 nm. The concentration that inhibited 50% cell growth (IC₅₀) was calculated using curve fitting and the each experiment was done in 3 replicates and reported as IC₅₀ ± SD (Elumalai et al., 2012).

**MTT bioassay**

The cytotoxic effect in different active fractions of HMF drug was evaluated against MCF-7 using MTT bioassay (Sanaz et al., 2012). The human breast cancer MCF-7 Cell line was cultured explicitly as mentioned in the procedure. Accordingly, the cells were seeded in 96-well microtiter plate (200 μl/well) with concentration of 4x10⁴ cells/cm². The cultivated cells were exposed to various concentration of the methanolic extract (1, 0.75, 0.5, 0.25, 0.1, 0.075, 0.05, 0.025, 0.01 mg/mL) prepared in 1% dimethyl sulfoxide (DMSO) as they reached 40-50% confluency and were then incubated for different periods of time (24, 48 and 72 h). The control groups received the same amounts of DMSO with four wells remained untreated as control. After the treatment, normal culture medium was replaced with 200 μl fresh media and 50 μl MTT reagent (2 mg/mL in PBS), except the cell-free blank control wells. The cells were maintained vi incubation as per the procedure mentioned above subsequently, the MTT solution was substituted with 200 μl of DMSO and 25 μl sorenson buffer (0.1M NaCl, 0.1M glycine regulated to pH: 10.5 with 1M NaOH), incubated for 15 min at 37°C. Eventually, the optical density of the wells was measured at 570 nm by means of a spectrophotometric plate reader of standard firm. The growth of tumoral cells and viability of the cells was determined using the formula.

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\text{Viability} \% = \left(\frac{\text{optical density of sample}}{\text{optical density of control}}\right) \times 100
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Furthermore, the cyto-toxicity of the HMF extract was determined by plotting of the percent cytotoxicity index, CI % = [1-(optical density of sample/optical density of control)] × 100, versus concentrations of the fractions of Tribal medicine formulation.

**Evaluation of Anticancer activity from HMF drug formula**

The concept of making dilution which is a process of declining the concentration by adding of a solution such as water. The crude extracts of HMF drug was diluted according to the requirements and the extracts were syringe filtered to avoid contamination prior to use.

**Trypan Blue Assay**

Trypan blue is an imperative stain applied to examine the dead tissues or the cells which takes blue colour selectively. This assay is used to determine the dead cell count as well as the living cell count. The living cells will have an intact membrane which does not allow the dye to pass since the cells are very selective in compounds. The dead cell does not process an intact membrane and takes up the stain.

**Hoechst Stain Assay**

This assay was done to check that the cell death has occurred due to apoptosis (the cell death due to destruction in the actual functions of membrane followed by cells that in turn leads to inflammation) or not. Besides, Hoechst Stain assay was done to confirm that, the cell death has occurred explicitly due to apoptosis only.

**MTT Assay**

The MTT assay was performed to reduce the yellow composite called 3-(4,4-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide (MTT) under the influence of succinate dehydrogenase enzyme in the complex-II at the electron transport chain that occurs in Mitochondria. The turn down of MTT can occur only physiologically active cells so, the activity will be determined based on the viability status of the cells. The whole reaction comprises, the MTT which go into the cell subsequently to the mitochondria wherein, MTT was reduced to form an insoluble Formosan product which appears in dark purple color. Meanwhile, the cells are solubilized through DMSO solvent thus, the released Formosan product was assessed through spectrophotometric method (Mosmann, 1983).

**III. Results**

The present study was carried out at the Department of Engineering Chemistry, Maharaja Institute of Technology, (Visvesvaraya Technological University), Thandavapura, Nanjanagud taluk, Mysuru dist - 571302, (Karnataka), India and the analytical study was conducted at the department of Biotechnology and applied sciences, SIET, Tumkur. The herbal/tribal medicine formulation was validated in association with Bhoomigeetha Institute of Research and Development (BIRD), Tumkur. The anti-cancer study was executed in association with Sri Raghavendra Biotechnologies, Bengaluru. The HMF (Tribal Medicine Formula) or Herbal Medicine Formula
(HMF) was procured from traditional practitioner at Biligirirangana Hills, Chamarajanagara district, Karnataka (Table 1 and Fig 1).

Phyto-chemical analysis
The phyto-chemical screening of aqueous extract and solvent extracts of EMP and HMF demonstrated the presence of Carbohydrate, proteins, alkaloids, flavonoids, saponins, tannins, gums & mucilages, coumarins, terpenoids, tannins, steroids, glycosides, phyto sterols, fixed oils and fats, phenols, saponin etc in Cold water, hot water followed by solvent extracts. The presence of these phyto-chemicals suggested to taking part in synergistic role to exert the observed pharmacological activity. The fact that strong synergism of several constituents in the crude drug may prove more potent and effective than any single purified compound, is always overlooked. Moreover, this may help to nullify the toxic effects (if any) of individual constituents (Table 2).

Antioxidant activity
The method used, ABTS radical scavenging assay, gave the measure of antioxidant activity of the HMF drug extract determined by the decolorization of the ABTS*, through measuring the reduction of the radical cation as the percentage inhibition of absorbance at 734nm. The effects of HMF drug aqueous extract on ABTS free radical scavenging activities was assayed at various concentrations. The IC<sub>50</sub> value was found to be 07± 0.74µl/ml was evident in this extract (Table 3).

Chromatographic studies
This was done in order to purify the HMF drug solvent extracts and the TLC chromatograms were developed using the solvent system, toluene, chloroform and ethanol. The bioactive compound moved on the stationary phase and the retention factor was calculated. There were two to five bands of whole HMF drug extract at Rf values of 0.32 and 0.44 in A. serpyllifolia; 0.52 and 0.26 in D. hispida; 0.59 and 0.23 in G. mauritiana; 0.34, 0.14, 0.16, 0.62 and 0.95 in N.nimmoniana similarly, in case of R. densiflora 0.76 and 0.85 were present. The extracts showed variable Rf values as compared among these extracts (Table 4 and Fig.2A&B).

Anticancer activity
The partially purified extract of HMF drug was tested for anticancer property against both HeLa and MCF-7 cell lines respectively. The inhibition concentration (IC<sub>50</sub>) as observed from the graph lies between 3 and 4 in 24 hours of trypan blue assay (Table 5 and Fig. 3, 8, 10 & 11). The cells show positive Hoechst assay as the DNA has acquired the Hoechst stain and the same is seen under the fluorescent microscope as observed (Fig. 14 & 17).

The result analysis for MTT assay shows that, the cell viability decreases with the increase in concentration of the HMF drug. The inhibition concentration value (IC<sub>50</sub>) for MTT assay at 24 hours showed to be 3% and for 48 hours (Table 6 & 7) lies in between 3 and 4 (Fig. 4,5,6,7 and 9A &B). The positivity of the MTT Assay can be observed in the fig 5 and 7 by the formation of crystals.

Anticancer activity against MCF-7 Cell Lines
The cells were cultured on the suitable Animal cell media under controlled condition and the various concentrations of the formulation was taken and tested for its action on the growth of HeLa cells and Mammarian Cancer cells (MCF-7). The Anticancer activity was determined using the ‘Trypan Blue assay’ in Table 5. This result revealed that, profound activity was noticed by the end of 24 hours than 48 hours shown in Fig. 8, 10, 11,12 & 16. Then, the Cell viability count was determined through MTT assay along with a media control contained growing cells with no drug. Similarly, the vehicle control was set up in presence of the respective solvents. The ethanolic extract of HMF drug was tested against HeLa cells followed by MCF-7 cell lines (Fig. 4,5, 6,7 and 9A &B). The HMF drug extract was found to be superior as compared to other extracts. The result reveals the apoptosis of both cancer cells indicating the most positive action of the Herbal Medicine formulation (Fig.12,13 &14).

The ‘Hoechst staining’ technique was showed the apoptotic state of cells and DNA, through fluorescent microscopy. The remarkable activity was noticed in the HMF drug tested in its crude form compared to the purified form which confirms that, the HMF drug works best in its crude form which is being currently practiced by the Traditional healers (Fig. 15, 16 & 17).

IV. Discussion
The presence of active secondary metabolites in the extracts of ethno-medicinal plants may have profound activity and justifies the status for preparation of crude potential drug by the tribal people. The phytochemical screening of the ethno-medicinal plants showed the presence of alkaloids, flavonoids, terpenoids, saponins, tannins, phenolic compounds and reducing sugars. A. serpyllifolia and D. hispida did not contain
cardiac glycosides and coumarins while, *G. mauritiana*, *N. nimmoniana* and *R. densiflora* showed the presence of glycosides, tannins and alkaloids. The phyto-chemical analysis conducted on the plant extracts revealed the presence of constituents which are known to exhibit medicinal as well as physiological activities (Sofowora, 1993). The TLC result in the different components of HMF drug exhibited the presence of diversified bioactive constituents and confirms the overall potentiality of the HMF drug.

The MTT assay on the cytotoxic activity of the partially purified extract of HMF plant drug on MCF-7 cell line apart from experimentation with HeLa cells suggested that, the extract was fairly cytotoxic to MCF-7 cells in a dose and time dependent manner. The findings of the present study will provide an insight into a new implication of the traditional usage of HMF drug in the tribal medicine system found to be as a most potential, efficacious and novel cancer chemo-preventive agent, where integration of the different herbal compositions remedy may help in preventing or reducing the risk of breast cancer and associated oxidative stress diseases. The similar expressions were also made in traditional medicinal plants (Abdul et al., 2009; Althunibat et al., 2009; Rajesh et al., 2011; Cieckiewicz et al., 2012).

Several studies in this filed have shown that, the ethno-medicinal plants are of potential value for identifying anti-proliferative agents followed by determination of the anti-inflammatory and anti-proliferative activity of herbal medicine formulation where, the suppression of cell growth as well as induction of apoptosis in human breast cancer cells was observed (Conforti et al., 2008; Nurhanan et al., 2008; Hogan et al., 2010; Mbaveng et al., 2011; Lin et al., 2012). Elsewhere, metabolites like, flavonoids, di-terpenoids and poly phenols from Herbal Medicine formulation was appraised for their Cytotoxic effects against both MCF-7 and HeLa cancer cell lines respectively. This finding from the study revealing that, the HMF has strong tumor regression potentiality even for a broad range of tumor cells depicted in the ailment. This is in accordance with the reports of Sun et al., (2002); Matito et al., (2003); Giri et al., (2006); Hu et al., (2011); Elumalai et al., (2012).

In addition, this study has clearly shown that, HMF drug has relatively substantial antioxidant activity with regard to the potential constituents present in it that might be responsible for the chemo-preventive effects of the HMF drug extract. The protective role and consistency of these active compounds of the HMF drug might have connected to the ailment in controlling the various oxidative stress factors (Braca et al., 2002; Matito et al., 2003; Aiyegoro et al., 2009; Tyagi et al., 2010; Murthy et al., 2011; Hu et al., 2011; Hazirah et al., 2013). The different endogenous antioxidants produced by the human body have potential health benefits against oxidative stress and related ailments, but also naturally supplemented herbal antioxidant compounds such as, phenolic acids, polyphenols and flavonoids scavenge free radicals like peroxide, hydroperoxide or lipid peroxyl are most promising and inhibits the oxidative mechanisms to prevent the system from degenerative diseases (Bhandari and Kawabata, 2004; Barriera et al., 2008; Aiyegoro and Okoh, 2010; Verma et al., 2009; Hogan et al., 2010; Yang et al., 2010).

In addition, this study has clearly shown that, the active fractions of HMF drug has antioxidant activity and comparatively high with regard to the potential radical scavenging ability of its methanolic extract. Since, the HMF drug are normally comprising complex mixtures of different active metabolites i.e., flavonoids, alkaloids and other phenolic compounds present in the methanolic extract of HMF drug which further donates a hydrogen atom for scavenging the stable DPPH radical, would be beneficial in quantifying the presumed charge of them as antioxidant and anti-cancer agents (Oskoveian et al., 2011; Hazirah et al., 2013).

The exceptional fact that, anticancer activities of ethno-medicinal plants have been verified critically to be coupled with a diverse range of phyto-chemicals, such as polyphenols, flavonoids, terpenoids, steroids, catechins etc (Tung et al., 2009). Many researchers have also opined that, the more phenolic content of different plant components increases the antioxidant activity in the system since, there is a linear correlation between phenolic contents and antioxidant activity (Prasad et al., 2009; Hogan et al., 2010; Jain and Jain, 2011; Nagmoti et al., 2012). Although, the phenolics of plant sources are very basis for antioxidants, in which they have differential expressions along with a specific ability to quench various free radicals. As a result, verifying the role of these phyto-chemicals in HMF drug, certain kind of an oxidative stress related diseases is of substantial importance. In the study, there is a correlation between total active phyto-chemicals with special reference to phenolic and flavonoids contents as well as radical scavenging activity along with the anti-proliferative activity of breast cancer in the active fractions of HMF drug was explored.

By taking into account, the results on cyto-toxicity in the extracts of HMF drug at MCF-7 cells was found to be superior with increased contents of both phenolics and flavonoids and thereby, higher radical scavenging activity in higher concentrations was noticed. On the whole, this study suggests, the potentialities of both antioxidant and cytotoxic activities of HMF extract which could be helpful in preventing or curbing the progress of various oxidative stress-related diseases such as breast cancer. This was evident in the approaches on bioassay-guided fractionation in HMF drug, further it would be of value to purify and identify the foremost active constituents in the extract which is responsible for inhibiting the proliferation of MCF-7 cells in the system (Ksouri et al., 2009; Fu et al., 2010). Similarly, as some phenolic antioxidants have a specific role in suppressing the growth and proliferation of transformed or malignant cells through induction of programmed cell death or
apoptosis, it materializes to demand the need for additional investigation on cell cycle analysis and determination of the distinctive mechanism of action for providing anti-proliferative activity in the extract of Herbal Medicine formulation (Puneeth kumar et al., 2011; Oskoueiyan et al., 2011; Sanaz et al., 2012). The authors believe that, the current objectives of the study could furnish the appropriate background for detailed examination on anticancer properties of this HMF drug. Besides, it is the very first report on the analysis of the anti-proliferative (MCF-7) and antioxidant activities of Herbal Medicine formulation being practiced in the Tribal Medicine System (TMS).

V. Conclusion

Finally, it can be concluded that, the Herbal medicine formulation has potent antimicrobial and antioxidant activities apart from their antiproliferative activities. Hence, with this basis, the formulation showed the highest anticancer activity against MCF-7 Cell lines and justifies their practice of Herbal Formulation in their TMS. Further, the susceptibility of mammalian cancer cells has been influenced by HMF drug has been substantiated based on the outcome of the study. In addition, the significance of these observations and recommendations in the light of previous studies with asynchronous population of MCF-7 cells also has been discussed.

Acknowledgements

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References


Evaluation of validated herbal medicine formulation for Anti-proliferative and apoptotic activities...


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Evaluation of validated herbal medicine formulation for Anti-proliferative and apoptotic activities.


Fig. 1A-H: Ethno-Medicinal Plant Drugs (Emp) And Parts Of Tribal Medicinal Formulation (Hmf)  
A: Andrographis serphyllifolia, B: Dioscorea hispida, C & D: Tubers of Dioscorea hispida;  
And processing of Tubers, E: Glycosmis mauritiana, F: Nothopodytes nimontiana,  
G & H: Habit and Root samples of Rauwolfia densiflora,
Table 1: Validated Tribal Medicine formulation (HMF) and its components practiced for antiproliferative and anti-inflammatory related ailments at Biligirirangana Hill Tracts, Karnataka.

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Ethno-medicinal plants with Vernacular Name.</th>
<th>Family</th>
<th>Plant parts used</th>
<th>Quantity (powder) (g/kg)</th>
<th>Validated Quantity of HMF (g)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Andrographis serphyllifolia Vahl. (A) Vr. Name: Kasinasara</td>
<td>Acanthaceae</td>
<td>Whole plant</td>
<td>20</td>
<td>(A) 20+ (D) 15+ (G) 25+ (N) 25+ (R) 15+ (HMF) ADGNR = 100g</td>
</tr>
<tr>
<td>2</td>
<td>Dioscorea hispida Dennst. (D) Vr. Name: Noolana hambu</td>
<td>Dioscoreaceae</td>
<td>Tubers</td>
<td>15</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Glycosmis mauritiana (Lam) Tanaka. (G) Vr. Name: Orange berry</td>
<td>Rutaceae</td>
<td>leaves</td>
<td>25</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Nothapodytes nimoniana, Blume. (N) Vr. Name: Durvasane mara</td>
<td>Icacinaceae</td>
<td>Leaves</td>
<td>25</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Rauwolfia densiflora Benth &amp; Hook. R Vr. Name: Snake root</td>
<td>Apocynaceae</td>
<td>Leaves</td>
<td>15</td>
<td></td>
</tr>
</tbody>
</table>

**DOSAGE, DURATION AND MODE OF TREATMENTS OF TRIBAL MEDICINE FORMULATION**

**Paste of HMF**

It is applied on affected part of the wound, skin cut, infected region due to tumour formation with few drops of Lime juice as external application.

**Duration:**
Apply paste at wound area & cover with a thin cloth 4 times/week

**Decoction of HMF**

Ground & juice boiled with warm water & swallowed internally for tumour related problems. Decoction with warm water/ goat milk for inflammation, skin destructions and related ailments.

**Duration:**
One tsp two times a day for 7 days.

*HMF obtained from TMM was validated by Authorized Ayurvedic Practitioner*

Table 2: Phyto-chemical analysis in different extracts of HMF drug

<table>
<thead>
<tr>
<th>SL. No.</th>
<th>Phytochemicals</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
<th>F</th>
<th>G</th>
<th>H</th>
<th>I</th>
<th>J</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Carbohydrate</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>2</td>
<td>Proteins</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>3</td>
<td>Alkaloids</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>4</td>
<td>Flavonoids</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>5</td>
<td>Terpenoids</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>6</td>
<td>Tannins</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>7</td>
<td>Phenol</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>8</td>
<td>Saponin</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>9</td>
<td>Oils &amp; fats</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
</tbody>
</table>

A-Cold water, B-Hot water, C-Ethanol, D-Methanol, E-Acetone, F-Chloroform, G-petroleum ether, H-Ethyl acetate, I-Hexane, J-NaCl

DOI: 10.9790/5736-1001010524 www.iosrjournals.org 17 Page
Evaluation of validated herbal medicine formulation for Anti-proliferative and apoptotic activities...

Table 3. Antioxidant activity (ABTS) free radical scavenging activity in HMF drug

<table>
<thead>
<tr>
<th>Test Variables</th>
<th>Plant Extrait</th>
<th>IC50µg/ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample</td>
<td>HMF drug extract</td>
<td>7±0.74</td>
</tr>
<tr>
<td>Référence</td>
<td>Gallic acid</td>
<td>1.72±0.03</td>
</tr>
</tbody>
</table>

Table 4. Showing column fractions of Crude drug in Column Chromatography

<table>
<thead>
<tr>
<th>Column eluents of ethanol extract</th>
<th>Distance moved by the compound(cm)</th>
<th>Distance moved by the solvent(cm)</th>
<th>Retention factor(cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crude sample</td>
<td>14.7</td>
<td>19.5</td>
<td>0.754</td>
</tr>
<tr>
<td>Fraction 1</td>
<td>16.1</td>
<td>19.5</td>
<td>0.826</td>
</tr>
<tr>
<td>Fraction 2</td>
<td>16.1</td>
<td>19.5</td>
<td>0.826</td>
</tr>
<tr>
<td>Fraction 3</td>
<td>16.2</td>
<td>19.5</td>
<td>0.83</td>
</tr>
<tr>
<td>Fraction 4</td>
<td>15.8</td>
<td>19.5</td>
<td>0.81</td>
</tr>
</tbody>
</table>

Fig. 2A & B. The TLC chromatograms of the individual components of HMF drug extracts

A: A. serpyllifolia, B: D.hispida; C: G. mauritiana; D: N.nimmoniana; E: R. densiflora
Stationary phase: Silica gel GF254; Mobile phase: toluene: chloroform: ethanol = 5: 5:2
{(Observed under ultra violet light 254 nm (A) and 366nm (B))}

Evaluation of Anticancer activity

Table 5. Showing Trypan Blue assay values in different concentrations of HMF drug

<table>
<thead>
<tr>
<th>SL. No</th>
<th>Concentration</th>
<th>% Cell Death</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Media</td>
<td>8.39</td>
</tr>
<tr>
<td>2</td>
<td>Vehicle</td>
<td>17.46</td>
</tr>
<tr>
<td>3</td>
<td>10%</td>
<td>48.82</td>
</tr>
<tr>
<td>4</td>
<td>6%</td>
<td>41.72</td>
</tr>
<tr>
<td>5</td>
<td>5%</td>
<td>39.6</td>
</tr>
<tr>
<td>6</td>
<td>4%</td>
<td>30.26</td>
</tr>
<tr>
<td>7</td>
<td>3%</td>
<td>26.6</td>
</tr>
</tbody>
</table>
Evaluation of validated herbal medicine formulation for Anti-proliferative and apoptotic activities...

Fig. 3. Showing the % of cell death after Trypan blue assay in different concentrations of HMF drug.

Table 6. Showing IC$_{50}$ value of cells from MTT assay after 24hrs of incubation in different concentrations of HMF drug.

<table>
<thead>
<tr>
<th>Concentration of crude ethanol extract media</th>
<th>% viability of cells</th>
<th>IC$_{50}$µg/ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>vehicle</td>
<td>32.12</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>47.22</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>42.00</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>33.82</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>24.84</td>
<td>5.45</td>
</tr>
<tr>
<td>6</td>
<td>11.11</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>20.42</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>26.96</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>38.72</td>
<td></td>
</tr>
</tbody>
</table>
Table 7. Table showing IC50 value of cells from MTT assay after 48hrs of incubation in different concentrations of HMF drug

<table>
<thead>
<tr>
<th>Concentration of crude ethanol extract</th>
<th>% Viability of cells</th>
<th>IC50µg/ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>media</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vehicle</td>
<td>96.69</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>47.52</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>24.75</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>11.22</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>3.96</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>13.20</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>17.49</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>33.99</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>46.20</td>
<td></td>
</tr>
</tbody>
</table>

Fig. 5. Showing the IC50 value of MTT assay after 48 hrs of incubation in different concentrations of HMF drug

Fig. 6. MTT assay in different concentrations of aqueous extract of HMF drug
Fig. 7. MTT assay in different concentrations of HMF drug for 24hrs

Fig. 8. Trypan Blue assay in different concentrations of HMF drug for 48hrs Anticancer activity from different (Crude) concentrations of HMF drug

Fig. 9A & B. MTT Assay in different concentrations HMF drug (24 and 48 hrs)
Evaluation of validated herbal medicine formulation for Anti-proliferative and apoptotic activities..

**Fig. 10.** Trypan blue test in different concentrations HMF drug for 24 hours

![Trypan blue assay after 24 hrs](image)

**Fig. 11.** Trypan blue test in different concentrations HMF drug for 48 hours

![Trypan blue assay after 48 hrs](image)

Apoptosis of HeLa cells after the action of HMF drug extract

**Fig. 12.** Trypan Blue Assay -Lowest concentration: 20 to 30% Cell death

![Trypan blue assay result](image)
Evaluation of validated herbal medicine formulation for Anti-proliferative and apoptotic activities.

Fig. 13. MTT Assay showing crystallization of the cell matrix in 48 hours

Fig. 14. Hoechst's Assay – High Concentration: 100% Cell death Apoptosis of MCF-7 Cells, after the action of HMF drug extract

Fig. 15. MTT Assay-Crystal formation (24hrs) in Ethno-medicinal plants Formulation HMF drug
Evaluation of validated herbal medicine formulation for Anti-proliferative and apoptotic activities...

Fig. 16. Trypan Blue Assay-Lowest concentration: 40-50% Cell Death (24hrs) in Ethno-medicinal plants formulation HMF drug

Fig. 17. Hoechst staining: To visualize the chromosomes after the action of the test sample HMF drug