INTRODUCTION

According to the Vedas, the cow was considered as the most valuable animal and called as the mother of all [1]. Cow’s urine was known as “GOMUTRA” has many advantages in curing several diseases [2]. It has a unique place in Ayurveda and was described as an effective medicinal substance or secretion of animal origin with innumerable therapeutic properties [3]. It is an important ingredient of panchagavya, a term used to describe five major substances such as urine, milk, ghee, curd, and dung obtained from cow [3,4]. All the five products possess medicinal properties and are used singly or in combination with some other herbs against many diseases [5,6]. This kind of alternative treatment, termed as “cowpat,” has been reported [7] to be beneficial even for threatening and clinical studies are required to confirm its therapeutic efficacy.

Various analyses done on cow urine, revealed the presence of substances such as nitrogen, sulfur, phosphate, sodium, manganese, iron, silicon, chlorine, magnesium, malic, citric, tartaric, succinic, carbolic acids, calcium salts, Vitamins A, B, C, D, E, lactose, enzymes, creatinine, hormones, and gold acids. Therefore, these are needed in smaller amounts by the human body to cure various diseases. The presence of certain volatile and nonvolatile components was also present in cow’s urine [12].

COW URINE DISTILLATE AS A BIOENHANCER FOR ANTIMICROBIAL & ANTIPROLIFERATIVE ACTIVITY AND REDISTILLED COW URINE DISTILLATE AS AN ANTICLASTGEN AGENT

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ABSTRACT

Objective: The objective of this study was to prove that cow urine distillate (CUD) is a bioenhancer for antimicrobial activity and antiproliferative activity, redistilled CUD (RCUD) as an anticlastogen agent.

Methods: The antimicrobial activity of rifampicin with CUD at different concentrations was determined against pathogenic Escherichia coli by well puncture method. The Penicillin and ciprofloxacin in combination with CUD at different increasing concentrations against pathogenic E. coli culture were also determined by disc diffusion method. Sulforaphane (ACA) as an anticancer agent was extracted from cruciferous vegetables and purified by high-performance liquid chromatography. The Breast cancer cell lines (MCF-7) were treated with anticancer agents along with CUD in increasing concentrations. The anticlastogenic activity of RCUD in human peripheral lymphocytes was tested with clastogens such as manganese dioxide and hexavalent chromium.

Results: CUD showed to enhance the antimicrobial activity of rifampicin with 20 µl concentration by well puncture method; penicillin with increasing concentration of up to 80 µl and ciprofloxacin up to 80 µl, respectively, by disc diffusion method. The rate of degeneration of breast cancer cell lines (MCF-7) was increased with increasing concentration of CUD. Clastogen (MnO₂) of 10 µl with 200 µl of RCUD showed effective anticlastogenic activity in agarose gel electrophoresis as the activity of clastogen decreased with increasing concentration of RCUD.

Conclusion: CUD acts as a bioenhancer to increase antimicrobial and antiproliferative activity. RCUD showed a high level of anticlastogenic activity toward clastogen. Thus, cow urine is found to have special properties that can be used in combination with different therapeutic agents to cure several diseases such as tuberculosis, leprosy, and cancer. Further in vivo and clinical studies are required to confirm its therapeutic efficacy.

Keywords: Cow urine distillate, Redistilled cow urine distillate, Bioenhancer, Antimicrobial activity, Antiproliferative activity, Anticlastogen activity.

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INTRODUCTION

The cow urine distillate (CUD) is used as a bioavailability facilitator for anticancer therapy directly or in combination with anticancer molecules [2]. The bioactive fraction enhances the activity of antibacterial agents, anticancer agents, and antituberculosis agents from 2 to 80 folds and helps the antibiotics and other molecules to act better on the target by transferring the compound across the membrane to the target site.

Sulforaphane and diindolylmethane (compounds from brassica vegetables) have been shown to synergize together in inhibition of cancer growth. The former compound has cancer chemopreventive activity and is also classified as an isothiocyanate [3]. Sulforaphane is a glycol breakdown product of the glucosinolate glucoraphanin (sulforaphane glucosinate). Young broccoli sprouts and cauliflower, sprouts are especially rich in glucoraphanin [6].

A clastogen is a mutagenic agent form mutagenesis, and it is human carcinogens as well [11], an anticlastogen is an agent that inhibits or suppresses the activity of clastogen [13]. Various research studies have already reported regarding the redistilled CUD (RCUD) as an anticlastogenic agent [14].

The present work was aimed at proving CUD as a bioenhancer for antimicrobial activity [15] and anticancer activity and RCUD as an anticlastogenic agent. To prove, the antimicrobial activity of CUD in various concentrations was tested against different antibiotics. Similarly, the anticancer activity of CUD with different anticancer agents.
against cancer cell lines was checked. The anticlastogenic activity of RCUD in human peripheral lymphocytes was tested with clastogens namely manganese dioxide and hexavalent chromium.

**METHODS**

**Collection of sample (CUD)**
CUD was collected from Asaram Gow-Shala, bypass tank road, Niwai, Rajasthan.

**CUD - bioenhancer for antimicrobial activity**

About 50 ml of Luria-Bertani broth was autoclaved at 121°C for 15 minutes and followed by inoculating with *Escherichia coli* (Diagnostic Centre at Mambalam, Chennai), it was incubated at 37°C in orbital shaker overnight.

**Well puncture method**
Mueller-Hinton agar (MHA) was autoclaved at 121°C for 15 minutes. Rifampicin was prepared using sterile distilled water at a concentration of 1 mg/ml wells were created in the medium using gel puncture. *E. coli* broth culture was uniformly spread over the medium using a sterile swab. Rifampicin added in the wells as standard concentration of 10 µl would act as a control. CUD was added in different concentrations of 10 µl, 20 µl. The plates were incubated at 37°C for 48 hrs [2].

**Disc diffusion method**
MHA was prepared and autoclaved at 121°C for 15 minutes. Using a sterile swab *E. coli* broth culture was uniformly spread over the media. Penicillin and ciprofloxacin discs were placed on the media. CUD was added in the concentration of 20 µl, 40 µl, 60 µl, and 80 µl. The plates were incubated at 37°C for 48 hrs [2].

**Extraction of anticancer agent from broccoli sprouts and cabbage seeds**
Broccoli sprouts and cabbage seeds were rinsed with distilled water. It was then ground using mortar and pestle. Ground sprouts and seeds were defeated thrice with hexane and allowed to dry overnight at room temperature. MIB-Q water was added in 3:1 ratio (w/w) to defatted meal, father was allowed to utilize for 8 hrs at room temperature. Sodium chloride:wet meal:sodium sulfate was added in the ratio 1:1:0.75 (w/w/w) and mixed thoroughly followed by extracting thrice with equal volumes of methylene chloride. It was centrifuged at 4,000 rpm for 10 minutes. Upper aqueous phase was collected in a separate centrifuge tube. The liquid was filtered using a Whatman filter paper. The filtrate was stored in the refrigerator [16-18].

**High-performance liquid chromatography (HPLC)**
Preparative HPLC separation was performed by injecting 0.5 ml of filtered aqueous extract onto a Waters Prep Nova-Pak (19 × 300 mm, 60 Å, 6 µm) HR C-18 reversed-phase HPLC column (Waters, Milford, MA). Elution was performed using Waters model 501 pumps to deliver a constant flow rate. The solvent system consisted of 50% acetonitrile in water. Sulforaphane was detected by absorbance at 254 NM using a ultraviolet (UV) detector (19).

**Maintenance of breast cancer cell lines**
Breast cancer cell lines (MCF-7) strains were ordered from National Centre for Cell Sciences (NCCS) from Pune. The confluent cells from NCCS were subcultured and maintained aseptically followed by examining under phase contrast microscope. They used up medium in a centrifuge tube followed by adding 2 ml of SE buffer and mixed by vortexing. 2 ml of 0.2% freshly prepared SDS was added and mixed thoroughly, the tubes were incubated at 65°C for 1 hr in dry bath. 400 µl of 2M NaCl was added to the tubes and mixed by inverting the tubes gently. Tubes were centrifuged at 10,000 RPM for 15 minutes at 4°C and the supernatant was carefully transferred to centrifuge tubes and the pellet was discarded. Double the volume of ice cold absolute ethanol was added to each tube, the tubes were gently inverted several times until the DNA precipitate was formed. Visible DNA strands were transferred to microfuge tube followed by adding 200 µl of 70% ethanol and tubes were centrifuged at 2,000 rpm for 5 minutes. The supernatant was discarded, the pellet was air dried, after drying it was re-suspended with 200 µl of TE buffer and stored at −20°C. Finally, the isolated DNA from clastogen treated lymphocyte was viewed by running gel electrophoresis [14].

**RESULTS**

**Antimicrobial activity**

The antimicrobial activity of rifampicin was increased when in combination with CUD in different concentrations 10 µl and 20 µl respectively. Rifampicin + CUD showed a clear zone of inhibition, whereas rifampicin alone shows a pale zone of inhibition. Therefore, the
microbial activity of rifampicin increased with increasing concentration along with CUD. 20 µL showed larger zone of inhibition compared with rifampicin + CUD (10 µL). Hence, it was proved that CUD is bioenhancer for the microbial activity of antibiotic rifampicin.

Disc diffusion method with penicillin + CUD showed the zone of inhibition is same as in control and even with 20 µl of CUD after 24 hrs of incubation. However, after 48 hrs of incubation the zone of inhibition with 40 µL, 60 µL, and 80 µL CUD concentrations were observed and were found to be effective (Fig. 1).

The zone of inhibition was found to increase with increasing concentration 20 µL, 40 µL, 60 µL, and 80 µL of the CUD + ciprofloxacin disc. Percentage of the zone of inhibition increases with CUD concentration, hence proving that CUD is an enhancer for antimicrobial activity (Fig. 2).

Two distinct layers as upper aqueous phase and lower organic phase, the former contains the anticancer agent. The HPLC peak (sharp peak) for the extracts from broccoli sprouts was seen in Fig. 3 and indicates that the compound has very less impurities, whereas in Fig. 4 shows that the cabbage seed extract has a high level of impurity when compared to the broccoli sprouts.

After 24 hrs of incubation, the degeneration of breast cancer cell lines was seen by a broccoli anticancer agent alone and therefore the increase in degeneration rate with the increase in the concentration of CUD with broccoli sprouts. Degeneration of cancer cells with cabbage seeds was less as the purity of sample was less found by HPLC results. Redistillation of CUD was done to check the anticlastogen activity Fig. 5 shows the normal lymphocyte culture and treated lymphocytes with clastogen and RCUD was seen in Fig. 6.

Isolation of DNA was done from lymphocyte and run at 1% and 0.8% agarose gel seen in Figs. 7 and 8. The lane 3, 4 with 10 µl MnO₂ + 200 µl of RCUD, showed the high dastrogenic activity as the concentration of RCUD was less in lane 5 and 6 with 40 µl MnO₂ + 400 µl of RCUD comparatively, this explains the anti-clastogenic activity of RCUD. Fig. 9 shows the activity of RCUD as an anticlastogenic agent.

**DISCUSSION**

Cow’s urine is a mobile medical dispensary, and it’s a panacea of all diseases stated by Pathak and Kumar [19]. Cow’s urine is believed to cure many diseases [20] as the CUD, bioenhancer for antimicrobial activity proved by disc diffusion method and well puncture method. In accordance to Wate et al., 2011 distillate cow’s urine, an activity enhancer and availability facilitator for bio active molecules such as antibiotic, antifungal, and anticancer drugs [21]. The CUD was added in increasing concentrations of 20 µL, 40 µL, 60 µL, and 80 µL and the results were found to be more effective in disc diffusion method, and therefore it shows an effective enhancing property to antibiotics rifampicin, penicillin, and ciprofloxacin within 48 hrs of incubation. Therefore, similar results were expressed by Khanuja et al., 2002 showed that the activity of antibiotic rifampicin and penicillin used alone and several folds activity increased when used in combination with CUD in addition to that Vijayalakshmi and Saranya stated that Gomutra has both antifungal [22,23] and antibacterial activity.

According to Vermeulan et al., 2006 extraction of anticancer agents from a plant source is the most economical method, the cruciferous vegetables such as broccoli, brussels sprouts, and cabbage, can lower the risk of developing pancreatic, lung, colorectal, and prostate cancers [18,24,25]. According to Kushad et al., 1999, the broccoli seeds were found to be rich in glucosinolates or isothiocyanate [26] compounds as HPLC was performed using HR C-18 reversed-phase column, the solvent system consisted of 10% acetonitrile in water and therefore his findings revealed the presence of compound sulforaphane at 254 nm absorbance using a waters 486 tunable absorbance detector (Waters, Milford, MA) [27]. In our present work, the anticancer agent
sulforaphane, a glucosinolate was extracted from broccoli sprouts and cabbage seeds, followed by preparative HPLC by injecting 0.5 ml of filtered aqueous extract onto a C-18 column with a solvent system consisting of 50% acetonitrile in water, their purity was detected by absorbance at 254 nm using a UV detector. The peaks of HPLC reveal the presence of impurities as the broccoli extract has less impurities when compared to the cabbage seeds.

Randhawa, 2010 stated that CUD as a bioenhancer for anticancer agents, in the present study anticancer agent was extracted from broccoli sprouts and cabbage seeds [26]. The MCF-7 was treated with sulforaphane as control (100 µl, 200 µl) and sulforaphane (ACA) along with CUD in increasing concentrations (100 µl ACA + 100 µl CUD, 100 µl ACA + 200 µl CUD, and 100 µl ACA + 300 µl CUD) followed by adding the ACA of about 100 µl. As the concentration of CUD increased with the rate of degeneration of breast cancer cells was also increased gradually, and degeneration was observed maximum in 100 µl ACA + 300 µl CUD. Hence, CUD acts as an enhancer of anticancer agent [26]. Dutta et al., 2005 reported that the anticlastogenic and antigenotoxic effect of RCUD in human peripheral lymphocytes, which has been challenged with manganese dioxide (MnO₂) and hexavalent chromium (Cr+6). Human peripheral lymphocytes in vitro were treated with manganese dioxide and hexavalent chromium as established genotoxicants and clastogens which could cause induction of DNA strand break, chromosomal aberration and micronucleus [29]. Our present work showed that three different concentrations of RCUD such as 1 µL/ml, 50 µL/ml, and 100 µL/ml were used and the manganese dioxide and hexavalent chromium caused statistically significant damages to DNA, which could be protected by RCUD which in turn confirmed the clastogenic effect by the clastogenic assay.

The anticlastogenic effect of RCUD revealed that the concentration of clastogen to that of RCUD, i.e., 10 µL MnO₂/Cr+6 + 200 µL of RCUD and 40 µL MnO₂/Cr+6 + 400 µL of RCUD, as latter causes DNA strand break and the difference in thickness of a DNA band showed the effect of RCUD as anticlastogenic agent and the control was treated with 30 µL of MnO₂/Cr+6 whereas the lymphocytes were pretreated (1 hr before the treatment of clastogens) with RCUD for better results.
CONCLUSION

The present work proved that the CUD serves as a bioenhancer for antimicrobial activity by testing against different antibiotics; anticancer activity was being tested in MCF-7 cell lines and RCU as an anticlastogenic agent toward clastogens. Therefore, cow’s urine has potential to cure several major adverse diseases, as this imparts the vital therapeutic effect of CUD. Further, in vivo study has to be conducted to prove the anticancer activity.

REFERENCES