INTRODUCTION

Colon cancer is one of the most primitive living seed plants, cycas revoluta, which is used by the hilly people of Northeast India [10]. It is reported that seeds contain cycasin which is a neurotoxin when consumed orally due to the production of methyl adducts with DNA bases and causes point mutations [5]. This adduct interferes with normal cell growth by altered normal gene transcription. Over the past few years, the first-line clinical treatments for colon cancer patients are cytoreductive surgery and combined chemotherapy [3]. Drug resistance results in poor overall survival rate [4]. Therefore, the development of effective and less toxic drugs is urgent for colon cancer patients.

The colon-specific carcinogen 1,2-dimethylhydrazine (DMH) induces the formation of methyl adducts with DNA bases and causes point mutations [5]. This adduct interferes with normal cell growth by altered normal gene transcription. After the DNA damage, ROS production crossing the level of endogenous antioxidant increases, leading to colon cancer and neoplastic transformation. By Phase II biotransformation, these electrophilic intermediates are detoxified by enzymes such as glutathione (GSH) S-transferase [6]. The pro-carcinogen DMH undergoes hydroxylation in Phase I reactions catalyzed by cellular microsomal monooxygenases to produce strong electrophiles. The produced strong electrophiles are capable of interacting with cellular nucleophiles such as DNA to form adducts causing mutagenesis and neoplastic transformation. Phase II detoxification of carcinogenic metabolites is done by Phase I enzymes. The pro-carcinogen DMH undergoes hydroxylation in Phase I reactions catalyzed by cellular microsomal monooxygenases to produce strong electrophiles. The produced strong electrophiles are capable of interacting with cellular nucleophiles such as DNA to form adducts causing mutagenesis and neoplastic transformation. As a result, DNA damage occurs leading to colon cancer and less production of endogenous antioxidant. After the DNA damage, ROS is produced in fewer amounts for the survival of colon cancer cells.

Cycas revoluta, one of the most primitive living seed plants, contains edible starch in pith, and is used for making sago. Seeds of this plant contain cycasin which is a neurotoxin when consumed orally due to the production of methyl adducts with DNA bases and causes point mutations [5]. This adduct interferes with normal cell growth by altered normal gene transcription. After the DNA damage, ROS production crossing the level of endogenous antioxidant increases, leading to colon cancer and neoplastic transformation. By Phase II biotransformation, these electrophilic intermediates are detoxified by enzymes such as glutathione (GSH) S-transferase [6]. Due to binding of DMH metabolite to DNA, gene transcription is modified to create oxidative stress through the production of reactive oxygen species (ROS) inside the cells. The ROS has pathological status by inducing oxidative stress-mediated inflammatory response in several tissues [7]. Increase in ROS level is an important factor to develop colitis-associated colon cancer [8]. There are several endogenous antioxidant such as superoxide dismutase (SOD), CAT, and GSH increase in normal colon cell in response to small increase of ROS, but in DMH-induced rats due to modification gene transcription, excessive ROS is produced crossing the level of endogenous antioxidant production. As a result, DNA damage occurs leading to colon cancer and less production of endogenous antioxidant. After the DNA damage, ROS is produced in fewer amounts for the survival of colon cancer cells.

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MATERIALS AND METHODS

Materials

DMH, methanol, 5-fluorouracil (5-FU), phenazine methosulphate, NBT, reduced nicotinamide adenine dinucleotide, glacial acetic acid,
n-butanol, pure SOD, H$_2$O$_2$, pure catalase (CAT), phosphate buffer, ethylendiaminetetraacetic acid, trichloroacetic acid, Tris-HCl, Dithio- bis (2-nitrobenzoic acid), pure reduced GSH, and dd sodium dodecyl sulfate were purchased from Sigma Laboratories, Germany.

Animals
The experimental protocols involving animals were approved by the Institutional Animal Ethics Committee (NCPT/IAEC-15/2015), and the experiments on animals were performed in accordance with the Committee for the Purpose of Control and Supervision of Experiments on Animals guidelines. Experiments were performed on male Wistar rats (5 weeks old) procured from Indian Institute of Chemical Biology, Kolkata, India. The animals were fed standard balanced diet and drinking water and were maintained with 24 h day and night cycle. The animals were acclimatized for 2 weeks before starting of the experimentation.

Plant material
Fresh male cone was collected from the village of Srirampur under East Midnapur district and authenticated (Voucher Ref. No.: BSI/Pharma/SD/Tech./2016) by botanist, Dr. A B D Selvam of Botanical Survey of India (BSI), Shibpur, Howrah (West Bengal). The cones were cut into small pieces and dried in the shade below 50°C. Then, dried cone pieces were powdered in mixture grinder and stored in airtight container.

Extraction of plant extract
The cones were cut into small pieces and were shade dried and then milled into a coarse powder. Then, the air dried and powdered cones (175 g) were first defatted with petroleum ether (60–80°C) and then extracted with 2.5 L of methanol (90%) using Soxhlet apparatus. The solvent was then removed with repeated lyophilization. After drying, 11.7 g (6.6% yield) extract was obtained.

Determination of total polyphenolic compounds
UV spectrophotometric method was used to determine the concentration of phenolics in the plant extracts [17]. The content of total phenolics in extracts was expressed as mg gallic acid equivalent (GAE) per g of dry weight of extract (mg GAE/g DW).

Determination of total flavonoid content
The content of flavonoids in plant extracts was determined using spectrophotometric method [17]. The content of flavonoids in extracts was expressed as mg of rutin equivalent (RUE) per g of dry weight of extract (mg RUE/g DW).

Acute toxicity study
MECR in olive oil was administered orally to the animals in increasing doses up to 2000 mg/kg b.w. These animals were observed for 2 h for behavioral, neurological, and autonomic profiles and mortality and toxicity for 72 h.

In vivo experimental design for induction of colon cancer
Animals were divided into five groups (n=6). Group I serves as a vehicle control which received 0.25% carboxymethyl cellulose solution. Groups II-V were treated with DMH at the dose of 20 mg/kg b.w., s.c once a week for 4 weeks. Aqueous suspension of MECR at a dose of 200 mg/kg/day and 400 mg/kg was administered orally to the animals in Groups III to IV every day for 16 weeks. Group V received 5-FU as a standard drug at a dose of 10 mg/kg b.w., per day s.c. for 16 weeks [18].

Macrosopic evaluation of the incidence of polyps
At the end of the experiment, rats were sacrificed and colons were taken out and flushed with phosphate-buffered saline. The colons were cut to open longitudinally without disturbing the polyps and carefully counted through visual macroscopic examination. Then, the colons were verified histopathologically.

In vivo antioxidant status determination
After 16 weeks of treatment, rats were kept one night fasting and then sacrificed to collect colons. After washing in ice-cold saline, colons were kept in deep freeze (~20°C) to conduct the different types of biochemical tests. Then, the colons were taken out separately for each time just before the specific biochemical test to be conducted. Colon tissues were homogenated using homogenizer. For homogenization, 0.5 g of colon was taken in 5 ml (w/v) ice-cold saline followed by centrifugation at 2000 g for 10 min. The supernatant was used for evaluating the level of enzymatic antioxidant such as SOD and catalase (CAT) activity and non-enzymatic antioxidant such as reduced GSH [19].

Histological observation
Histological evaluation was performed in colon tissues. A portion of the specimen was fixed in 10% formalin and embedded in paraffin wax, sectioned at 4 µm thickness, and was stained with hematoxylin and eosin. Morphological changes of colon were evaluated by light microscopy method in control and experimental groups of animals [20].

Statistical analysis
All the data were evaluated with GraphPad Prism version 5 software (GraphPad Software Inc, La Jolla, CA). Hypothesis testing methods included one-way analysis of variance (ANOVA) followed by Dunnett's t-test to correct for multiple comparisons with acceptable statistical level significance (p<0.05). Each experiment was presented as the mean±SEM.

RESULTS
Total polyphenolic and total flavonoid content
MECR contains total polyphenols (6.3±0.09 mg of GAE/g) and total flavonoids (4.6±0.06 mg of RUE/g) (Table 1).

Acute toxicity study
In the acute toxicity assay, it was found that no mortality was observed up to doses 2000 mg/kg b.w. orally and was considered as safe, and no lethality or any toxic reaction was found up to the end of the study period. By keeping 1/5th (400 mg/kg) dose as highest, 200 mg/kg was selected as working doses for the present study.

Effect of DMH and MECR on polyps incidence
In the DMH-alone-treated rats (Group II), 100% incidence of polyps was found (Table 2). On supplementation with different doses of MECR (200 and 400 mg/kg b.w.) to DMH-induced rats (Groups III-IV), the incidence of polyps was significantly reduced as compared to DMH-alone induced rats (Group II). No specific changes were noticed control rats (Groups I), 400 mg/kg b.w, MECR was found to be effective, the incidence of polyps being 33.33%, but it was slightly less effective than the synthetic standard drug 5-FU (10 mg/kg b.w.), the incidence of polyph being 13.33%.

Effect of MECR on different antioxidant enzyme activity
SOD, catalase, and reduced GSH activity were decreased in DMH Group. All these parameters were restored significantly (p<0.05) toward the near normal value on supplementation with MECR (200 and 400 mg/kg b.w.) to DMH-treated rats (Groups III and IV). In Group V, the synthetic drug 5-FU (10 mg/kg b.w.) also increases the activities of SOD, CAT, and GSH significantly (p<0.05) more in DMH-treated rats (Fig. 1).

Table 1: Phytochemical content of MECR

<table>
<thead>
<tr>
<th>Phytochemical constituents</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total phenolic content (mg of GAE/g of DW extract)</td>
<td>6.3±0.09</td>
</tr>
<tr>
<td>Total flavonoid content (mg of rutin/g of DW extract)</td>
<td>4.6±0.06</td>
</tr>
</tbody>
</table>

Values were expressed as mean±SEM (n=3). MECR: Methanolic extract of Cycas revoluta, C. revoluta: Cycas revoluta.
Histopathological evaluation
The colon of control group showed normal mucosa and submucosal layers and normal colonic architecture without apparent abnormality in (Fig 2a). DMH-induced group showed clear degeneration as the size of the cells smaller than that of normal (Fig 2b). In DMH+MECR200 group, mucosal and submucosal layers were less ruptured compared to DMH-induced group (Fig 2c). In DMH+MECR400 group, the mucosal epithelial architecture along with sub-mucosal layer's integrity was better than that of DMH+MECR200 group (Fig 2d). Tubular glands gradually appeared clearly as the doses of MECR increased. However, DMH+5-FU-treated group showed better results as compared to DMH+MECR400 group as tubular gland appeared clearly with arranged manner (Fig 2e).

DMH group shows that tubular glands in mucosal and submucosal layer are not observed prominently. Tubular glands size is smaller in DMH Group than that of control group. Degenerative changes was observed at mucosal and submucosal layer in DMH Group. Tubular glands gradually appeared clearly as the doses of MECR increased. In DMH+5-FU group, degenerative changes are less than those of DMH+MECR 200 group and DMH+MECR 400 group (magnification, ×40).

DISCUSSION
Colon cancer was induced with DMH in Wistar rats through its metabolite azoxymethane which is potent genotoxic agents to trigger oxidative stress through DNA methylation of colonic epithelial cells [21]. This adduct interferes with normal cell growth by altered normal gene transcription. Due to binding of DMH metabolite to DNA, gene transcription is modified to create oxidative stress through the production of ROS inside the cells. Increase in ROS level is an important factor to develop colitis-associated colon cancer [8]. There are several endogenous antioxidants such as SOD, CAT, and GSH increase in normal colon cell in response to small increase of ROS, but in DMH-induced rats due to modification gene transcription, excessive ROS is produced crossing the level of endogenous antioxidant production. As a result, DNA damage occurs leading to colon cancer and less production of endogenous antioxidant. After the DNA damage, ROS is produced in fewer amounts for the survival of colon cancer cells. In the in vivo model, the levels of endogenous antioxidants SOD, CAT, and GSH were restored after administration of MECR in DMH-induced rats but not restored in only DMH-consumed rats. Endogenous antioxidant restoration by MECR occurs due to replacement of cancer cell in colon causing death in consequence of ROS increase by polyphenolic and flavonoid in MECR [22]. On the other hand, endogenous antioxidant increases in normal cell in response to ROS to reduce its level [23]. Polyphenolics and flavonoids of MECR are antioxidant [24] which perform anticancer [25] activity ultimately in rat colon. Hence, colon cancer polyps are less observed in MECR-treated group and polyp number decreases dose-dependent manner.

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### Table 2: Effect of MECR on the incidence of colonic polyps in the different groups

<table>
<thead>
<tr>
<th>Groups</th>
<th>Number of rats</th>
<th>Number of polyps bearing rats</th>
<th>Total number of polyps</th>
<th>Average number of polyps bearing rats*</th>
<th>Percentage incidence of polyps*</th>
<th>Percentage of polyps inhibition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control (I)</td>
<td>6</td>
<td>0</td>
<td>Nil</td>
<td>Nil</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>DMH (II)</td>
<td>6</td>
<td>6</td>
<td>15</td>
<td>2.5</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>DMH+MECR200 (III)</td>
<td>6</td>
<td>5</td>
<td>9</td>
<td>1.8</td>
<td>60</td>
<td>40</td>
</tr>
<tr>
<td>DMH+MECR400 (IV)</td>
<td>6</td>
<td>3</td>
<td>5</td>
<td>1.67</td>
<td>33.33</td>
<td>66.67</td>
</tr>
<tr>
<td>DMH+5-FU (V)</td>
<td>6</td>
<td>2</td>
<td>2</td>
<td>1</td>
<td>13.33</td>
<td>86.67</td>
</tr>
</tbody>
</table>

DMH: 1,2-dimethylhydrazine (20 mg/kg b.w.), MECR200: Methanolic extract of Cycas revoluta (200 mg/kg b.w.), MECR400: Methanolic extract of Cycas revoluta (400 mg/kg b.w.), 5-FU: 5-Fluoro-uracil (10 mg/kg b.w.).

*Total number of polyps/number of polyps-bearing rats in each group. *[(Total number polyps/total number of polyps in DMH group) ×100]
Furthermore, our histological observation revealed that clusters of abnormal degenerated glands and cells in the mucosal lining of the colon and rectum were observed in animals of DMH group along with severe mucosal and submucosal damage. However, animals treated with orally administered MECR show replacement of abnormal cell in the DMH-treated animals, justifying the anticancer potential of MECR.

CONCLUSION

From this study, it can be concluded that MECR might be a colon cancer protective agent. The present study opens many new areas of research work. This work can be continued in the future to study and to make it clinically applicable for colon cancer protective activity in different experimental models and also to isolate, identify, characterize, and standardize the active principle(s) that are responsible for this activity.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

REFERENCES