

Table 1: Comparison of tail length of mice before and after treating with mineral oil.

Types of mice	Tail length in cm	
	Before treating with mineral oil	After treating with mineral oil
Normal mice	9.6cm	9.6cm
Mouse with plasmacytomas alone	9.7cm	5.4cm
Mouse with metastasize tumor in the head	9.6	5.8cm
Mouse with metastasize tumor in the throat	9.5cm	5.9cm

This de-fibrinated blood was taken and diluted with equal volume of physiological saline. 1ml of lymphoprep solution was taken in a vial to which added the diluted blood in slanting position using pipette. The content of the vial was centrifuged at 1400 rpm for 20 minutes. The interphase was removed using pipette. The cells were centrifuged at 1400 rpm for 20 minutes and the pellets were washed with 1ml saline. The pellet was resuspended in 100 µl of RPMI 1640 medium.

12-14 cm of drinking straw was cut. One end of the straw was slantly cut and sealed by slightly heating the tip in a flame. Nylon wool fibres were finely teased using a pair of forceps and the teased fibres were packed (loosely) into the straw. Washed the straw using 5ml of physiological saline and made a small opening at the sealed end of the straw to drain the saline. It was then filled with 3ml of RPMI1640 medium in a horizontal position. The column was now kept in incubator at 37°C for 30 minutes in horizontal position. Resuspended lymphocytes were loaded into the activated nylon wool column. Then the column was held vertically above an eppendorf tube, now hot saline (about 60°C) was added. The hot saline passing out of the column was collected. To this 0.2 ml of 1% SRBC (Sheep Red Blood Corpuscles) was added and the mixture was centrifuged for 12 minutes at 1600 rpm. After centrifugation the samples were incubated in the refrigerator at 4°C for 5 minutes. The pellet was resuspended in the 100 µl of RPMI medium. Then a drop of it was taken in a clean dry slide, observed and enumerated T cells under the microscope (20x/40x) for rosettes. Number

of T cell rosettes formed were observed among 100 lymphocytes and tabulated.

Results and discussion:

Anti cancer study

The anticancerous activity of *A. aspera* extracts were checked with the cancer mice. The present results show that ether extract of the plant leaves has efficient anticancerous activity compared to other extracts. The anticancerous activity, phytochemical screening of *A. aspera* and its impact on cancer cells was analyzed and recorded.

Reduction in the tail length of cancer mice

Reduction in the tail length of the mineral oil induced mice confirms the presence of cancer in those mice. The tail length of the normal mice was 9.6 cm whereas the mice with metastasize tumor in the head had tail length of 5.8 cm, the metastasize throat cancer mice had 5.9 cm of tail length and the mice with plasmacytoma alone had the tail length of 5.4 cm. The comparison of the tail length of mice before and after the mineral oil injections was shown in the Table 1.

T cell count

The increase in T cell count further confirms the presence of cancer in the mice. The comparison of T cell count for normal and cancer mice before and after giving extracts is given in the Table 2.

The anticancerous activity of *A. aspera* leaves was tested against mineral oil induced swiss albino mice (group V). It was found that none of the mice got the symptoms of cancer. This result is in agreement with the earlier reports of Careonal, (2000). Careonal, (2000) stated that the *A. aspera* leaves contain Vitamin

Table 2. Comparison of T cell count for normal mouse and cancer mouse before and after giving extracts.

Types of mouse	Tcell count before giving extracts	Tcell count after giving extracts
Normal mouse	62/100 lymphocytes	62/100 lymphocytes
Mouse with plasmacytoma and metastasize tumor in the head	96/100 lymphocytes	81/100 lymphocytes
Mouse plasmacytoma alone	84/100 lymphocytes	83/100 lymphocytes
Mouse with plasmacytoma and metastasize tumor in the throat	93/100 lymphocytes	94/100 lymphocytes

C and beta-carotene that has the property of anti-tumoral activity. Similar findings were obtained in *Mallotus peltatus* leaf extracts administered animals (Bhattacharya *et al.*, 2002), from this phytochemical obtained in plants were used as an anti cancerous agent.

The group II mice developed plasmacytoma, among the six, one of the mice was found to develop metastasize tumor in the head and one got metastasize tumor in the throat. The mouse with the metatasize tumor in the head, throat and plasmacytoma were treated with ether extract for 15 days. There was reduction in the cancer symptoms for the ether extract treated cancer mouse.

The present work clearly indicates that the ether extract at the concentration of 3 mg/ml is very effective in reducing the cancer symptoms. These results are in agreement with the earlier reports of Careonal, (2000) and Bhattacharya *et al.*; (2002). Because of the easily soluble nature of this plant in organic solvents and of its anticancerous activity, suggests that *A. aspera* leaves can be used in the anticancerous drug preparation in alternative medicine. This suggests that using the *A. aspera* leaves is cheap and the economic herbal drugs can be prepared for cancer treatment. The use of *A. aspera* leaves was highly promising for cancer treatment.

A. aspera leaves were traditionally used by the tribal people for the treatment of tumors (Joy, 1998). Hence in the present

study different extracts of *A. aspera* leaves were tested with the cancer mice. The treatment showed that the ether extracts of *A. aspera* leaves has the greater anti cancerous activity and it is highly promising for the further phytochemical evaluation and this will be continued in the further studies.

The free radicles present in the mineral oil have the capability to bind covalently with the DNA and interact with purine; pyrimidines and phosphodiester groups of DNA. Thus the normal cells turn to cancerous cells. If the cancer cell spread throughout the body along with tumor site, that type of cancer is malignant tumor. *A. aspera* leaves has been used traditionally by tribal people for the treatment of inflammation, swellings, wounds and tumors. But hitherto, no work has been carried out scientifically to prove the efficacy of this plant to inhibit the growth of cancer cells. Hence, in the present study mineral oil induced swiss albino mice with plasmacytoma and metastasized plasmacytoma with head and throat tumors were chosen to find out the anticancerous properties of *A. aspera* leaves. The antioxidant property of *Achyranthes aspera* Linn leaves prevents the damage caused by the carcinogens to the DNA that disturbs the cell function (Murthy and Aleyamma Mathew, 2004). Hence, the present study promotes the utilization of plant-derived substances to prevent the growth of cancer cells.

In spite of tremendous development in the field of synthetic drugs during recent era, they are found to have some or other side effects, whereas plants still hold their own unique place, by the way of having no side effects (Jayakar *et al.*, 2003). Therefore, a systematic approach should be made to find out the efficacy of plants against cancer. Hence, the present study strongly suggests that the utilization of plant decoction to prevent the growth of cancer cells and inflammatory disorders. Further studies involving the purification of the chemical constituents of the plant and the investigation in the bio chemical pathways may result in the development of a potent anti-cancerous agent with better therapeutical index.

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