A New Fighter in the World

Pradip Mahato
Vill-Khudiatar, P.O.-Kaluhar, P.S.-Para, Dist-Purulia, Pin-723126, West Bengal, India

I. INTRODUCTION

Recently maximum cancer patients have died for wrong treatment. Otherwise antibiotics are losing its power of activity. So anti inflammatory anti receptors drugs will be an important part to treat of living mammals. Mahajyotisalaramine is an anti inflammatory anti receptors agent which can cure from cancer (pre-cancerous and cancerous both stages). The drug prevents and destroy DNA, RNA viruses and blocks the path way of metastasis formation. It can be chief fighter against beta mRNA COVID 19 virus. Moreover it has no side effects like other treatment chemotherapy and immunotherapy etc.

A normal cell =100%. 99.5%=abnormal cell. Decrease from normal cell 100%-0.5%=99.5% abnormal cell, 99.5%+0.5%=100% abnormal cell. So factor is 0.5% beta and +0.5% alpha. These two chains are responsible for cancer. Beta proteins decrease some essential proteins from the cell and alpha proteins increase concentration of proteins in cell. Beta proteins are belong to denatured family. Otherwise alpha proteins join to complex family. The denatured proteins create scarcity of demand in cell. Alpha proteins increase of demand in cell to produce complexity.

Why is a normal cell and a cancer cell have difference 0.5%? By searching for long term found two types of cell in a straw fungi and 0.5% difference between the two cells. I observed that the specific denatured proteins are responsible for it. These proteins inhibited the receptors proteins to produce normal cell. The denatured proteins are bind with alpha /beta receptors to produce first Gene mutation. Thereafter kinases/enzyme have been involved with the proteins produce complex proteins and it begins first single cell carcinoma. Beta regions receptors carried viruses. The viruses enter the host cell with the help of beta receptors. Here V or C zone were identified. mRNA COVID 19 have best choice in this areas for it’s first shelter and mutated.

Mahajyotisalaramine is derived from three different cells.

Why are these proteins denatured? The proteins are denatured by (1) heat (2) chemicals (3) metal (4) pollution etc. It behave different sites different nature as lungs, bone marrow, thyroid glands, liver, uterus etc. Studies were observed Leukorrhea, Trichomonas, Monilial infection can increased white discharge. Significantly increased excretion of some proteins. Those events are repeating many times, thus it create heat. Contact of sex can more increase to heat in uterus. So normal proteins altered to denature by heat. When denatured then its conjugated some proteins to direct wrongly. Those conjugated proteins are belong to some different proteins. Its make inflammation in uterus. Inflammation could begin Gene mutation, change nuclear, cytoplasmic ratio, abundant mitosis and sometimes create tumour giant cells. The pre-cancer period found pathological changes:(1) increase specific enzyme,(2) defective differential count or blood count,(3) defective antibody formation .(4) loss of cell pigmentation,(5) unrestricted cell formation.

Why did I think Mahajyotisalaramine could be useful in treating cancer? It was subjected for evaluation of anti cancer activity in experimental 15 rats. They were divided into four groups. Group—(1) Number of rats—4(a) Vagina of the female rats were massage with asbestos once every two days. (b) Starting Mahajyotisalaramine injection on same day every 72 hours (c) After three months, no ulcer, no tumour, no discharge observed. (d) After six months, stopped massage. (e) Result—Normal

Group—(2) Number of rats—4. (a) Vagina of the female rats were massage with asbestos once every two days. (b) After three months, vagina of the female rats have ulcerated and started discharge.(c) After four months, ulcer have extended to cervix.(d) Stopped massage.(e) After six months, attack with cancer: a cervix sections showing feature of chronic cervicitis with focal moderate dysplasia CIN-II Basement membrane is intact.(f) Starting Mahajyotisalaramine injection every 72 hours. (g) Result—Normal/cure.
Group—(3) Number of rats-3, (a) Vagina of the female rats were massaging with asbestos once every two days, (b) After two months, started orally three antibiotics cifroflaxacin, erythromycin, metronidazole respectively for the rats, (c) After three months, vagina of the female rats have ulcerated and started discharge, (d) After four months, ulcer have extended to cervix, (e) After six months, attack with cancer, Result-dead, (f) Antibiotics have no role in the disease. Its have lost of certain bacteria and peptide bond produced complex protein.

Group—(4) Number of rats-4, Following the serial numbers a, b, c, d, e respectively of group-2, (f) When tumour produces I used of surgical procedure to detach the tumour from the body. Thereafter treated the rats by Mahajyotisaralamine injection. Finally the rats recovered from the disease. Somehow no spread again cancer cells.

Studied in humans: (female) History/disease--- White discharge, --chronic cervicitis/ cystic mass /uterine polyp/ bulky uterus-----bloody discharge-----moderately differentiated squamous cell carcinoma, moreover another syndrome observed in squamous cell carcinoma as (1) Pain or inflammation in lower abdomen specially uterus (2) Anemia (3) constipation (4) Weakness (5) Keratinization of squamous epithelium (6) Atrophy of cells (7) Spot/ulceration (8) loss of appetite and acidity.

Anti cancer activity were observed with total dose in rats and above three hundred people. Mahajyotisaralamine prevents keratinization of epithelium cells, thus it keeps the mucous membranes healthy and moist. The drug has important role in the body to control of nuclear, cytoplasmic ratio and restricted infections. DNA, RNA RAS, SAC etc. viruses present by the drug. It binds glycoproteins with the help of cell surface receptors and fought with viral infections. So viruses cannot build their empower without receptors, they become powerless. Mahajyotisaralamine can accept other proteins and carbohydrate. It can accept other proteins or even decompose some proteins. At the end of the separator, the recipient complete these two tasks in a reverse process. That is, the cancer cells, which occur at the end of the last stage, turn the cells into normal cells, thus reaching the first differentiated cells of the periodically transformed cells. Finally Mahajyotisaralamine develops cancer prevention by taking the first inflammatory agent that helps abnormal cell division and completely preventing cancer. This process takes three to six months to complete. But in the case of the pre-cancer patient, the time is less. In the case of process is completed within two to three months.

Many cancer patients have been treated by the drug for 17 years. Such as (1) squamous cell carcinoma, (2) adeno carcinoma, (3) breast cancer, (4) thyroid cancer, (5) lungs cancer, (6) liver cancer, (7) rectum cancer, (8) prostate cancer and leukemia. It has been shown that during the treatment of a cancerous patient, the first stop of the attack disease and the mother tumour does not give birth to a daughter tumour. Even histopathology don’t show cancer cells. So abnormal growth stop and cell death occurs in the normal course. Inflammatory agents that convert proto-oncogene to oncogene become powerless. Significantly, Mahajyotisaralamineworks properly by targeting cancer cells. In leukemia, it can first detached nucleus from immature RBC (normoblast). So WBC don’t kill RBC and platelets. I observed that the WBC count increased more than normal count of leukemic patients. But treated them by this drug they come back to normal and other count turn to normal. So blood ratio remain stable.

Cancer Genetic disease, oncogene is duplicate copy of proto-oncogene, both of which are part of the body. Mahajyotisaralamine blocks the pathway for Gene mutation and also blocks the pathway of metastasis formation. In addition, since cell death occurs in the normal way, theonco-viruses can’t be produced in the body.

Finally, Dr. M.N. Chatterjea, Dr. Rana Shinde spoke "Hence, in a real sence, if metastasis could be controlled, cancer could be controlled and for the most part cured”. Azra Raza said that the two choices were (1) block the path of metastasis formation, (2) celebrate anti inflammatory agent throughout cancer. Therefore, it is proven in the treatment of cancer by Mahajyotisaralamine.

References:
[1]. Test book of Medical Biochemistry Dr. (Brig) M.N. Chatterjea, Dr. Rana Shinde, 3rd Edition-1998
[2]. Why curing cancer is so hard Azra Raza, TEDx-1915
[3]. The complex Biology Glenn Begley.