NEW DISCOVERY ON VIRAL DISEASE AND CANCER IN MEDICAL

Dr.U.P.CHAURASIYA

HIV-Virus is a product of yeast candida albicans. All viruses are product of systemic fungus. In influenza virus is product of yeast aspergillus fumigats hiv virus is ova/spore of yeast candida albicans. HIV-virus reproduction has becomes in yeast cell not be host cell. All viral diseases can be treated by systemic fungal drugs – Itraconazole,Ketaconazole, Fluconazole, Amphotericin-B, nystatin etc. successfully most important medicine of AIDS is amphotericin-B in the world. Systemic fungal drugs are flu worse, which is in treatment of all types flu like-swine flu, human flu, bird flu, horse flu, Ebola, AIDS etc. viral diseases. But it is important that Hormonal drugs, Chelating agent be must together fungal drugs for treatment of viral diseases. Cancer is a systemic fungus. It can be treated by systemic fungal drugs- Itraconazole, Ketaconazole, Fluconazole, Amphotericin-B, nystatin etc.

In blood cancer the value of WBC has becomes above 50000 to 100000, they are not WBC but they are yeast cell. The structure of HIV virus is same structure of yeast cell of candida albicans. It is impossible that no any HIV patient can be possible without infection of candida albicans in their body. It is proof of HIV virus being candida albicans. Virus is cause of cancer and viruses are produces by systemic fungus. Tobacco MoJack disease is a fungal disease. Human Papiloma viras (HPV) is cause of mouth cancer, uterus cancer and wart diseases. Wart is a cyst and cyst is a tumour. Wart/cyst/tumour can be treated by systemic fungal drugs successfully. All proof of above statement have been save in my book – VIROCHEMISTRY Written by Dr. U.P. Chaurasiya and I have done test on human body practically which file I have safe. So sir I want to meet of scientist team in research centre. Kindly I pray to you sir please give me a chance for meeting scientist team.

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Research book cystology+Mycology +Oncology +Bio chemistry =VIRO CHEMISTRY –Dr.Umapati chaurasiya
Virus is ova of yeast.
Cyst=Tumour=Fungus
HIV-Virus is ova of yeast candida albicans.
HIV-virus is ova of yeast candida albicans which is systemic infection in blood when he spreads in vagina then produces luecorrhea in women.after some time its infection spreads in uterus then produses uterus tumour ans its infection in ovary then decay of ovum is called uterus cancer (HPV)and when he spreads in blood then he defects all organs of body and entered in all tissues of body.When the drugs uses for this treatment the immune system more defected and change in blood culture.The immune system (WBC,RBC)has lossed before the virus killing.so patient surrounds by a lot of disease and became very weak.This state is called Aids.So the main reason of Aids is yeast candida albicans.

Organs Involvement in Aids by cysts(Fungus)-
Lungs-pneumo cystis carinii pneumonia,Fungi-Invesis candidosis,Asthma-aspergillus fumugates,Pulmonary kaposis sar coma,Cryptosporidium
Gastro intestinal tract- oropharynx-candidosis,kaposis sarcoma
Stomach-Crypto sporidium,kaposis sarcoma
Small instestine-Crytosporidium,Aspergillus
Colon-Kaposis sarcoma,aspergillus,candidosis
Liver/Billary tract-Histoplasmosis,toxoplasmosis,Cryposporidium,
Nervous system-Toxoplas mosis,Cryptococcus,Aspergillus,candida
Skin-Candida albicans,Herpes Zoster,savoricdermatitis
We found final result to above statement that cysts infection are all organs in Aids patient candida albicans,crypto sporidium,histoplasma,toxoplasma,aspergillus etc.cysts are spreads in all organs in Aids patient.When the antibiotic drugs uses for killing cysts then finished immune system (WBC-RBC)hormones,enzymes,vitamins,before dies of thease cysts and patient surrounds a lot of diseases.
Proof of being yeast candida albicans in HIV-virus:-
1-The structure of HIV-virus is same structure as yeast candida albicans.
2-The skin cancer (kaposi’s sarcoma) has found in whose patient those are confermly infected yeast candida albicans.
3-The patient feels difficulti in talking, eating, drinking whose throat and mouth had infected by yeast candida albicans.
4-The infection of candida albicans has confermly founds in fluids of vaginal and Anal HIV-patient.
5-The spore of candida albicans have been seen in stool, urine, pus, saliva, cough, blood, sweets of HIV patient. 
Above facts proves that the HIV-virus is yeast candida albicans. Its nucleus is made by RNA in ova or spore form but when it becomes develop, then it becomes yeast cell as both nuclieus DNA, RNA form. Nucleic acid is liquid, which is not centre of any cell. Nitrogen and phosphorus is centre of all cell whose valency is five.

\[ -N=N- \quad \text{binding energy} = 225 \text{kcal} \quad N(7 = 2,5) \]

In, 1958, M.S. Messelson and F.W. Stahl allowed the bacteria E.colii, to growin a medium containing haevy nitrogen (N15). After reproduction, the DNA of bacteria became labelled with (N15). This haevy DNA could be seperated from light DNA (containing N14).

In cancer or dead body carbon (C) decay in nitrogen (N) as follows:-

\[ 6C14 \quad \text{------------------------} \quad 7N14 \quad +1e0 \]

The bad smell of dead and decaying bodies is due to putrefaction of protien change nitrogen and sulphur.
Animals is a converter of carbon in nitrogen.
\[
\text{C} \rightarrow \text{N}
\]
Plant is a converter of nitrogen change in carbon.
\[
\text{N} \rightarrow \text{C}
\]
After the animal or plant dies the $^{6}\text{C}^{14}$ undergoes as decay as follows:-
\[
^{6}\text{C}^{14} \rightarrow ^{7}\text{N}^{14} + -_{1}\text{e}^{0}
\]
In Animals-- carbon------------change---------------Nitrogen

In Plants---Nitrogen--------------change-------------Carbon

Plants is a maschine (converter) who change Inorganic compound in Organic compound. Hence plants is Nitrogen converter in Carbon.

And Animals is a maschine(converter) who change Organiccompound in Inorganic compound. Hence animals is Carbon converter in Nitrogen.

Hence definition of Cancer :- by chemistry-
Carbon change in Nitrogen.

The body decay(melting) in cancer not growing(increasing ) .

HPV—Human papiloma virus---Uterus cancer, Mouth cancer, Wart

HBV-Hepetitis-B virus—Liver cancer
HIV, Ebola virus, Rabies virus, Influenza virus, smallpox virus, etc. cancer virus are produces by systemic fungus.

Virus is the cause of Cancer.
Cancer is a systemic fungus—Dr. U.P.Chaurasiya
All viruses are product of systimic fungus.
HIV-virus is product of yeast candida albicans.
Nickel di terthalimiate dioxgin

The metal ions chelates four nitrogen and stay in centre of compound is called nucleus of compound. Cancer and viral diseases have been treated by chelating of metal in body with increasing immune

Nucleus of complex compound is Metal ions
co++, Fe++, Mg++, Zn++, Cu++, Ni++
Biliverdin (Cu++) and Zinc Insulin (Zn++) are metal complexes that chelate four nitrogen and other elements such as hydrogen, oxygen, carbon, etc. Metal is called the ligand and it is stayed in the center of the compound called nucleus of compound. The main role of metals is to build complex compounds as human fluids, such as Hemoglobin, Zinc insulin, Biliverdin, vitamin-B12, and plants, such as Chlorophyll and dioxin. Its depend on the immune system. Metals chelating is the base of treatment of cancer and viral diseases by increasing the immune system. Zinc (Zn2+) is an essential element crucial for growth and development and also plays a role in cell signaling for cellular processes like cell division and apoptosis. In the mammalian pancreas, Zn2+ is essential for the correct processing, storage, secretion, and action of insulin in beta (β)-cells. Insulin is stored inside secretory vesicles or granules, where two Zn2+ ions coordinate six insulin monomers to form the hexameric-structure on which maturated insulin crystals are based. The total Zn2+ content of the mammalian pancreas is among the highest in the body, and Zn2+ concentration reach millimolar levels in the interior of the dense-core granule. Changes in Zn2+ levels in the pancreas have been found to be associated with diabetes. Hence, the relationship between co-stored Zn2+ and insulin undoubtedly is critical to normal β-cell function. The advances in the field of
c Zn2+ biology over the last decade have facilitated our understanding of Zn2+ trafficking, its intracellular distribution and its storage. When exocytosis of insulin occurs, insulin granules fuse with the β-cell plasma membrane and release their contents, i.e., insulin as well as substantial amount of free Zn2+, into the extracellular space and the local circulation. Studies increasingly indicate that secreted Zn2+ has autocrine or paracrine signaling in β-cells or the neighboring cells. This review discusses the Zn2+ homeostasis in β-cells with emphasis on the potential signaling role of Zn2+ to islet biology.

Copper in bile has been shown by electrophoresis to occur neither as free ions nor complexed to protein but to be associated with a component of the micellar complexes of bile. Solvent fractionation studies suggest that the bile salt components of the lecithin-bile salt complexes are the active binding agents. The effects of specific bile salts on the behaviour of copper during electrophoresis supports this possibility.
The relationship of certain bile salts to the excretion of copper in man during the time that an external biliary fistula was functioning and to the intestinal absorption of copper in the rat was found to confirm this concept.

The results show that copper in bile is associated with taurochenodeoxycholate and suggest an explanation for the elevated tissue copper levels found in Wilson's disease.

Thus we get final result that all immune system depend upon metal chelation and their ligands. All types of Cancer and Viral diseases can be treated successfully by chelating metals in body.---Dr. U.P. Chaurasiya

Oropharyngeal candidiasis (OPC), caused primarily by *Candida albicans*, is the most common oral infection in HIV+ persons. Although Th1-type CD4+ T cells are the predominant host defense mechanism against OPC, CD8+ T cells and epithelial cells become important when blood CD4+ T cells are reduced below a protective threshold during progression to AIDS. In an early cross-sectional study, OPC+ tissue biopsied from HIV+ persons had an accumulation of activated memory CD8+ T cells at the oral epithelial–lamina propria interface, with reduced expression of the adhesion molecule E-cadherin, suggesting a protective role for CD8+ T cells but a dysfunction in the mucosal migration of the cells. In a subsequent 1-year longitudinal study, OPC− patients with high oral *Candida* colonization (indicative of a preclinical OPC condition), had higher numbers of CD8+ T cells distributed throughout the tissue, with normal E-cadherin expression. In OPC+ patients, where lack of CD8+ T cell migration was associated with reduced E-cadherin, subsequent evaluations following successful treatment of infection revealed normal E-cadherin expression and cellular distribution. Regarding epithelial cell responses, intact oral epithelial cells exhibit fungistatic activity via an acid-labile protein moiety. A proteomic analysis revealed that annexin A1 is a strong candidate for the
effect moity). The current hypothesis is that under reduced CD4$^+$ T cells, HIV$^+$ persons protected from OPC have CD8$^+$ T cells that migrate to the site of a preclinical infection under normal expression of E-cadherin, whereas those with OPC have a transient reduction in E-cadherin that prohibits CD8$^+$ T cells from migrating for effector function. Oral epithelial cells concomitantly function through annexin A1 to keep *Candida* in a commensal state but can easily be overwhelmed, thereby contributing to susceptibility to OPC.

**Keywords:** AIDS, *Candida albicans*, epithelial cells, T cells, mucosal immunity, cytokines, these are related to each others.

"Cancer is a fungus, called *candida albicans*, and it can be treated using sodium bicarbonate". So says Tulio Simoncini

Simoncini is a former Italian oncologist in Rome who developed a theory that all cancer is caused exclusively by a fungus called *candida albicans*. His alternative cancer treatment is simple: To alkalise the body and tumour by the use of a cheap, common compound, sodium bicarbonate, tackling the candida, shrinking the tumour and stopping metastases. Thus it is also a natural cancer cure.

At this point we would like to remind readers of our often repeated stance that we do not believe that any single thing is the cause of all cancer. In 2012 MD Anderson analysed the histology and biochemistry of all their recent DCIS breast cancer patients (over 1400 women) and concluded that every person was different.
We cover Simoncini purely because people write in and ask about him. Also, much of what is on the internet is inaccurate. We are not fans of Simoncini or his theory and approach.

Dr. Simoncini was a respected Italian oncologist until he developed his theory—his theory is all cancer is caused by the fungus, candida albicans. His cure is to alkalse the body to produce a condition in which the candida could not thrive. His main weapon is the inexpensive, sodium bicarbonate.

Cancer is a not fungus, but cancer is a systemic fungus. Many types of cancer become by many type yeast or fungus germs. Dr. Umapati Chaurasiya.

Cancer has been not treated by sodium bicarbonate but it can be treated alkali drugs and systemic anti-fungal drugs with chelating agent. Dr. U.P. Chaurasiya.

Fungus and candida infections may cause of cancer. Some doctors implicate fungi as a cause of cancer (leukemia).

Systemic anti-fungal drugs which is success in cancer as follows:

Ketaconazole, Itraconazole, Fluconazole, Amphotericin – B, Nystatin.

Amphotericin-B is most powerful drugs which react in Aids, while it is deeply fungal drugs.
Ketaconazole can be act successfully in prostate cancer.

Fluconazole and Itraconazole have been use in Neutropenic cancer and Aids successfully.

**Antibiotic activity in increasing power-**

\[ o > I > N > F > Br > P > S > Cl \]

1. \( N_2 \)  \( N=N \)  225Kcals(For structure of DNA RNA of germs).

2. \( O_2 \)  \( O=O \)  118 kcals........

3. \( Cl_2 \)  \( cl-cl \)  57.8 kcals(antibiotic activity).

4. \( S_8 \)  \( S-S \)  54 kcals (antibiotic activity).

5. \( P_4 \)  \( P-P \)  50.0 kcals (antibiotic activity).

6. \( Br_2 \)  \( Br-Br \)  46.1 kcals (antibiotic activity).

7. \( F_2 \)  \( F-F \)  38.0 Kcals (antibiotic activity).

8. \( N_2 \)  \( N-N \)  37.0 Kcals (antibiotic activity).

9. \( I_2 \)  \( I-I \)  35.5 Kcals (antibiotic activity).

10. \( O_2 \)  \( O-O \)  33.0 Kcals (for breathing).

It is clear that lo power binding energy has lead high power antibiotic vartue.All compounds has lead antibiotic activity which structure made of above any Eliments.Very low power binding energy eliments has very high power antibiotic activity.

Ergosterol binding antifungal-Amphotericin-B
Ergosterol decreases antifungal drugs-Itraconazole, Ketaconazole, Fluconazole.
Type of viruses which is involve in fungi-
Actinomyces phase Av-1, Alcelaphine herpes virus-1,2, Asparagus virus-1,2,3, Avian carcinoma mill hill virus-2, Avian encephalomyelitis virus, Avian hepatitis E virus, Candida albicans Tca2,5 virus, Citrus psorosis virus, Common cold virus, Contagious pustular dermatitis virus, Crypto virus, Cryptosporidium parvum virus, Cytomegalo virus, Dengu virus, Encephlitis virus Hcmv (human cytomegalo virus), Hepatitis A virus, Hepatitis B virus, Hepatitis C virus, Hepatitis D virus, Hepatitis E virus, Hepatitis F virus, Hepatitis G virus, Herpes B virus, Herpes simplex virus (HSV), Herpes zoster virus, Human herpes virus 1,2,3,4,5,6,6A,6B,7,8, Human immunodeficiency virus (HIV) 1,2,3,4,5,6,7,8,9, Human papiloma virus (HPV), Human para infleunza virus 1,2,3,4, Human polyoma virus, Cypo virus 1,2,3,4,5,6,7,8,9,10,11,12,13,14,15,16, Human rhino virus A, B, C, Human T cellleukemia virus 1,2,3, Human T cell lymphoma virus (HTcLV-1/2/3), Human T cell lymphotropic virus 1,2 (HTLV), Influenza virus A/B/C/D, Japanese incephlitis virus, Kaposis sarcoma associated herpes virus, Measels virus, Mammery tumor virus, Poliomyelitis virus, Rabies virus, RNA tumor virus, Simplex virus, Small pox virus, Trichomonas vaginalis virus 1,2,3, Tumor virus, Reo virus 1,2,3, Retro virus, Rhabdo virus, Rhino virus,
All viruses are produces by systemic fungi. Virus replication in fungicide, not host cell. In Blood cancer have increasing WBC above 50000 to 100000, they are not WBC but they are yeast cell-----
Dr. U.P. Chaurasiya

In Asthma patient has infected by Aspergillus Fumigates yeast. Influenza virus is a product of yeast Aspergillus fumigates. HIV-virus is a product of yeast candida albicans.

All viruses are produces by replication of fungi cell.
Cyst=fungi=tumor=Wart
Origin of Cancer and Virus:-

In 1897 Luwanskko has been discovered Tobacco mojac disease on Tobacco leaf, which surroundings made very bad area due to stool. Stool which leads ova, spore, larva of fungus, bacteria, worm, etc. The main pathogens that are commonly looked for in feces include:

- **Bacteroides** species
- **Salmonella** and **Shigella**
- **Yersinia** tends to be incubated at 30 °C (86 °F), which is cooler than usual
- **Campylobacter** incubated at 42 °C (108 °F), in a special environment
- **Aeromonas**
- **Candida** if the person is immunosuppressed (e.g., undergoing cancer treatment)
- **E. coli O157** if blood is visible in the stool sample
- **Cryptosporidium**
- **Entamoeba histolytica**

Intestinal parasites and their ova (eggs) can sometimes be visible to the naked eye.

When Knoll and Ruska has been discovered electron microscope in 1934, then Stenely in 1935 has been crystallization of germ of tobacco mojac disease. This crystallization was particle form, therefore it is called vireon. At past time it name forward virus. Then developed virus concept. But it was main guilty that was germ of micro-fungus. Thus Scientist had been left main pole of theory.

- Actually It is micro fungal cell. If we will study in deeply then we found virus cell are produces fungal cell. **Candida albicans**
- **Candida tropicalis**
- **Candida glabrata**
- **Candida parapsilosis**

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<th>PREDISPOSING FACTOR</th>
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<td>Hematological</td>
<td>Neutropenia, cellular immunodeficiency (leukemia, lymphoma, AIDS, aplastic anemia)</td>
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<tr>
<td>Other</td>
<td>Intravenous drug addiction, malnutrition, malabsorption, thymoma</td>
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- **Candida krusei**
  Histoplasma, Toxoplasmata, Neoplasmata, Mycoplasma
- **Candida lusitaniae** Blastomysis, actinomycosis, aspergillus fumigates, Cryptococcus,

- **Candida kefyr**
- **Candida guilliermondii**
- **Candida dubliniensis**
- **Other species of Candida**
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