Introduction:

Cancer incidence and deaths continue to rise in Ohio (Figure 1) and around the world. In the last two decades there have been major changes in air pollution that could potentially affect many cancers. There has been a dramatic drop in lead (Pb) air pollution from the removal of Pb from gasoline combined with an increase in benzene, a carcinogen. Pesticide and fertilizer use have risen sharply. All of these factors have increased the toxic effects of cadmium (Cd) air pollution. Cd is a ubiquitous pollutant present in tobacco smoke, incineration fumes, phosphate fertilizers and sewage sludge. With the increase in global population there is also an increase in Cd air pollution.

Of all the metals Cd is the most easily absorbed and the most influenced by nutritional factors. It has a special affinity for multiple sulfhydryl groups, but it binds weakly to so many other compounds that it moves to all compartments of the cell. Very low Cd concentrations down to 100pM stimulate cell growth and DNA synthesis significantly (von Zglinicki 1992).

Cd's behavior in complex mixtures is difficult to predict. It plays a pivotal role in stress responses. The stress response provides a first line of defense against infections and cancer transformation (Young 1989). Cd exposure, however, is associated with tumor formation and coupling of leukocytes, effects not generally found with other stressors in a model system of the skin of the common carp (Iger 1994). High dose Cd alone produced tumors in planaria, while PCB's and Aroclor 1254, a
pesticide, did not initiate tumors but potentiated the tumorigenic effects of low dose Cd (Tehseen 1992, Schaeffer 1991).
**Interactions:**

Cd has complex interactions with metals, chemicals and hormones. Cd is antagonized by Zn and is synergistically toxic with copper (Cu), tin (Sn), and Ni (Geertz 1994). Antagonism and synergism were found with mercury (Hg) and Cd depending on concentration. Cd and Ni powders injected into different legs were far more potent in combination in inducing fibrosarcomas in female rats than males (Furst 1989). These 2 metals as sulfides stimulate $\text{H}_2\text{O}_2$ formation by human polymorphonuclear cells at 0.5 (Cd) and 1 nM (Ni) concentrations, comparable to effects of the tumor promoter 12-O-tetradecanoylphorbol-13-acetate. (Zhong 1990). In a model system using murine sarcoma virus, Ni produced a splicing defect mutation while Cd and chromium (Cr) produced frameshift mutations (Chiocca 1989). Deletions of up to 300 nucleotides were noted with Cd exposure in this system (Heckman 1992). Cd, Cr, and Ni all inhibit DNA repair in mammalian cells and are co-mutagens with ultraviolet rays (Hartwig 1989).

Substances like dioxins, pesticides, and herbicides bind to the estrogen receptor. Estrogen increases the uptake of Cd into the liver, kidneys, and mammary glands. Since Cd binds to the metal binding site of the steroid binding site of DNA (Freedman 1988), it can be expected to have a modulatory role on steroid mediated cellular control processes. This provides one molecular basis for the synergistic effects of these chemicals with Cd. In addition, Cd by increasing glutathione levels in the liver, increases blood lipid levels which increases the levels of
these fat soluble chemicals in the rest of the body (Leonzio 1992).
Gene Effects:
With exposure to Cd, organisms or cells increase their production of metallothionein (MT). Cd moves from the cell membrane to the nucleus where it binds to DNA, inducing gene transcription in as little as 4 hours (Fowler 1993). With chronic exposure gene duplications can occur (Maroni 1987). In Cd resistant cells the transcription of MT can increase 20-40 fold (Sequin 1987). In humans there are at least 14 genes that have been identified which control the production of MT (Gedamu 1987). A mutation that results in the loss of a charged amino acid at the hinge between the metal binding domains makes it ineffective (Cody 1993).

Starvation, infection, physical stress, inflammation, radiation, endotoxin, glucocorticoids, glucagon, catecholamines, calcium ionophores, estrogen, progesterone, interferons, interleukin-1, carbon tetrachloride, chloroform, ethanol, and alkylating agents are some of the stimuli that activate MT genes (Bremner 1987, Karin 1987). Interferon has been shown to protect cells from DNA damage caused by Cd (Vasil'eva 1989). Methylation of the MT gene affects gene expression (Bhave 1988). The promoters greatly modify the availability of free Cd.

Cd binds to a metal responsive element in the MT-1 promoter, activating it in Cd exposed and non-Cd exposed cells (Anderson 1990), suggesting that Cd plays a role in all stress responses, not just those caused by Cd exposure. Because it gets into cells readily, it can
modulate any stress response. If all goes well, Cd detoxifies itself and terminates the stress response to that the cell can return to normal housekeeping function.

**Carcinogenesis:**

Carcinogenic doses of Cd induce oxidative stress while impairing cellular defense mechanisms against such stress (Koizumi 1992). It is evident that co-exposure to a variety of chemicals result in this carcinogenic state (Yamada 1993).

By altering cell metabolism, Cd fosters cell proliferation. The cell surface receptor activated by Cd (Smith 1989), interacts with oncogenes. Cd is known to increase the expression of two oncogenes c-myc and c-jun (Jin 1990). C-myc is associated with aggressive tumors. Cd also alters the tumor suppressor protein p53, eliminating its suppressant effect on cancers in a variety of tumors (Hainaut 1993).

In a situation of Zn deficiency, which always accompanies stress, there is inhibition of DNA repair, which is also significantly associated with carcinogenesis (Nocentini 1987). In any stress situation which results in a loss of tight epithelial junctions, either in the GI tract or in the vasculature, Cd is taken up by these lining cells and the transfer of nutrients is inhibited, compromising the cellular defenses.

In cell culture systems Cd at certain doses increases cell proliferation, an effect blocked by Se (Webber 1985). By providing Se, some cancer blocking herbs block the
uncontrolled growth of cancer cells stimulated by Cd.

Moreover, Cd effects on the primary tumor may allow shedding of tumor cells into the blood and the breaks in the vascular lining allow metastatic cells access to tissue. Cd has effects on proteases which are necessary for basement-membrane degradation and invasion of tumor cells.
Cathepsin D:

There have been several studies on the effects of Cd on cathepsin D, an important acid protease in lysosomes. Cd increases cathepsin D mRNA in human breast cancer cells (Garcia-Morales 1993). Cathepsin B increases mature cathepsin D in hormone refractory prostate tumor cells, promoting tumor invasion (Weiss 1993), an effect inhibited by a protease inhibitor. Elevated levels of cathepsin D were found in breast secretions from women with breast cancer (Sanchez 1993).

In human ovarian carcinoma cell lines, 1 nM 17-beta-estradiol increases secretion of procathepsin D by 50% (Rowlands 1993), an effect blocked by 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD). An inhibitory effect of TCDD on estrogen stimulated formation of lactate from glucose in human breast cancer cells was reversed by a phorbol ester, an effect that could involve Cd but that was not addressed in the study (Moore 1993). The increased incidence of reproductive tumors in fish and humans around pollution sources of Cd and dioxins suggests a synergistic effect.
Resistant Tumors:

Air pollution with Cd can make cancers resistant to cancer treatments. Tumors resistant to radiation and chemotherapy are resistant to Cd and may contain an overexpression of the MT gene. Overexpression of MT in primary invasive ductal carcinoma of the breast is associated with metastases which are resistant to anti-cancer drugs and radiation and have a poor prognosis (Schmidt 1993). It is possible that the resistance blocks apoptosis.

Over-expression of heat shock protein 60 in ovarian cancer is associated with resistance to cis-platin therapy and a poor prognosis (Kimura 1993). In this study the researchers did not find an induction by Cd but they evaluated the cells at 4 hours. Hiramura et al (1993) found that induction by Cd of this protein was maximal at 18 hours after exposure.

Over-expression of heme-oxygenase in human lung adenocarcinoma cell line (CL3R) is associated with resistance to Cd, arsenite, and adriamycin (Lee 1993). Heme-oxygenase is induced by oxidant stress and Cd in conditions of low glutathione. Glutathione suppresses the induction of heme oxygenase (Sunderman 1987).
Anti-carcinogenesis:

Although low doses can be carcinogenic, especially in combination with other chemicals, high doses can induce apoptosis. Mammalian cells contain a Ca(2+)-dependent endonuclease which is a necessary step in apoptosis. Zinc inhibits this process, depending on the free Ca2+ concentration. It appears that a balance between Zn and Ca regulates this process (Lohmann 1993). Cd alone stimulates the endonuclease, replacing Ca2+, and is more inhibitory than Zn in blocking apoptosis. This ability to take the place of both Zn and Ca and to have higher potencies than either ion, helps explain the highly divergent effects that it can produce.

As a stress agent Cd can kill chemically or spontaneously initiated cancers of the lung, liver, and blood in rats (Waalkes 1991,1992). Lung tumors induced by chemical exposures in humans are inhibited in individuals who are heavy smokers. Unfortunately, they generally suffer from other toxic effects from Cd exposure.
Female Cancers:

With the reduction in atmospheric lead which antagonizes the effects of Cd and the increase in environmental chemicals that bind to the estrogen receptor combined with Cd pollution, one can predict a change in female morbidity and mortality. The increase in breast cancer could be in response to Cd. Infiltrating inflammatory cells in the stroma appear to be the source of cathespin D levels associated with enhanced metastatic potential of node negative cancers (Johnson 1993), an effect that Cd could promote. Cathespin D stimulates the release of transforming growth factor-alpha (TGF-alpha), which stimulates breast cell growth by binding to TGF-alpha/epidermal growth factor receptors on the cell surface (Henry 1992). Metastases from human breast tumor cells grown in mice without estrogen supplementation, are associated with elevated cell production of cathepsin D (Thompson 1993).

The behavior of dioxins are particularly problematic. 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) acts as liver tumor promoter in female rats but not male rats (Schrenk 1992). In the human breast cancer cell line MCF-7, TCDD acts as an anti-estrogen but when combined with a phorbol ester the anti-estrogen response is inhibited (Moore 1993).

Chronic cervicitis and cervical cancer are increased in smokers (Bartecchi 1994) and associated with human papilloma virus. It is likely that Cd facilitates the carcinogenic effects of the virus which it does with murine
retrovirus infections.
Male Tumors:

Prostate cancer has been associated with increased Cd levels in the soil in Spain (Garcia Sanchez 1992). It is one of the cancers that has been increasing in incidence and severity. Older patients in Utah with aggressive tumors had a higher intake of dietary fat (West 1991). Cd exposure from occupational exposure, high dietary intake, or smoking increased the risk for aggressive prostate tumors (Elghany 1990). Men employed as janitors had greatly increased risk for aggressive tumors in this study. The carcinogen 3,2'-dimethyl-4-aminobiphenyl when combined with Cd produced significantly more prostate carcinomas in rats than either chemical given alone, showing a synergistic effect (Shirai 1993). Janitors are exposed to Cd in dust and other toxic chemicals in cleaning supplies.

Experimentally Cd can produce testicular cancers in rodents.
Lung and Colon Cancer:

Synergy of Cd with other carcinogenic chemicals undoubtedly plays a role in the development of lung cancer in smokers and non-smokers exposed to passive smoke. The effects of such synergy have been studied in rats using Cd, n-nitrosoheptamethyleneimine, and asbestos fibers to produce lung cancer. Elevated levels of Cd, Ni and Cr are found in lung cancer and colorectal cancer and the metal levels are correlated with blood levels of tumor markers (Martin Mateo 1990).

Workers exposed to 4-aminobiphenyl, which is a potent bladder carcinogen, who developed chloracne from exposure to dioxin (TCDD) in 1949, had increased mortality from soft tissue sarcoma, bladder cancer and respiratory cancer (Collins 1993). Since 4-aminobiphenyl is a co-carcinogen with Cd it is likely that Cd could be a co-carcinogen with dioxin and 4-aminobiphenyl.
Clinical Evaluation of Cd Exposure and Toxicity:

1. Urinary-cotinine is correlated with blood Cd (Willers 1992). Co-exposure to physical, emotional, chemical, or biological stressors, nutritional deficiencies, exaggerated responses to stress are factors that suggest increased levels of free Cd. There is a wide variability in sensitivity to the toxicity of Cd ions (Sens 1994).

2. Free Cd is not measured by blood, urine and hair Cd levels. Mag-fura-2 can be used to measure free Cd levels in cells (Quamme 1992). The Cd concentration in blood collected at autopsy is several hundred times higher than values measured before death. Formalin fixative greatly reduces the level of Cd found in tissues (Koizumi 1994).

3. By exposing dispersed cancer cells to varying concentrations of Cd and evaluating their viability in 24 hours, one can identify Cd sensitivity before the administration of radiation or chemotherapy. Cd effects on over-expression of the MT gene, heat shock protein 60, or heme oxygenase can be used to identify Cd resistance in cancer cells using immunological stains.

4. Decreased anti-oxidants, lipid peroxidation, elevation of LDH, Alk Phosphatase, eosinophilia, lymphopenia, elevated monocytes, fever, cachexia, viral, bacterial and fungal infection, neurotoxicity, fatigue, and muscle
weakness are non-specific effects of increased free Cd.

**Therapeutics:**

1. Cd could be used therapeutically in cancer treatments to induce apoptosis in Cd sensitive tumor cells. Anti-sense oligonucleotides complimentary to the messenger RNA coding for MT-II increased the sensitivity of neuroblastoma cells to Cd (Iversen 1992).

2. Anti-fungal herbs have anti-carcinogenic effects and chelate and remove Cd, ie Garlic, Pau D'Arco, and licorice.

3. High doses of vitamin C eliminate plasmids in bacteria which confer cadmium resistance. It is conceivable that high doses of vitamin C could make cancer cells Cd sensitive as well.

4. Cd can be used therapeutically in gene therapy to incorporate plasmids and promote gene expression. These attributes could be used in new gene therapies to restore the very malfunctions it may have caused.

5. Mg and Zn protect normal cells from toxic effects of chemotherapeutic agents. Niacin as NAD, protects DNA from oxidative strand breaks (Zhang 1993). Zn deficiency prevents the mobilization of vitamin A, another cancer fighting vitamin from the liver.
6. N-acetyl-cysteine facilitates Cd excretion, increases intracellular glutathione, and prevents the production of interleukin-8 induced by Cd, as does EDTA, a metal chelator (Horiguchi 1993).
Summary:

As a stress agent, inducing apoptosis and blocking it, Cd can have both helpful and harmful effects. The atmosphere is a thin envelope which makes the world a global village. Cd is the most toxic metal in air. As both the first and second messenger of the stress response, it is synergistically toxic with all other stressors, including many other carcinogens. Elimination of Pb and its replacement with added benzene in gasoline appears to have increased the toxicity of atmospheric Cd. With scientific understanding of the molecular basis of Cd's role in carcinogenesis and anti-carcinogenesis, primary cancer prevention can be practiced by reducing Cd and chemical air pollution and educating the public on smoke cessation, healthy eating habits and stress reduction. Using the existing information on Cd and its effects, determinations could be made on established cancers so that individualized treatment protocols can be developed to improve patient care.