**VERY LARGE AMOUNTS OF RADIATION ARE REQUIRED TO PRODUCE CANCER**

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1. Introduction

There is a public misconception regarding the relationship between radiation and cancer. This is the belief that any amount of radiation can cause cancer. This radio-phobia is unjustified since the real risk for radiation-induced cancer is very low. It takes a very large amount of radiation to cause cancer.

The biological damage that can be caused by high doses of ionizing radiation, including increased cancer, is well documented and justifiably feared [1]. However, the public perception of the ability of radiation to induce cancer is much greater than is supported by scientific data [2].

2. Where is the radiation-induced cancer?

If cancer is readily caused by every ionization of radiation, it should be possible to detect. However, radiation-induced cancer is difficult to detect. There are three major reasons for this. Variable background radiation dose makes increased cancer risk from small exposures impossible to detect. Cancer rate and cancer mortality are highly variable in different human populations making it very difficult to pinpoint excess radiation induced cancer. At the present time there are no specific biological markers for radiation-induced cancer so they cannot be identified or assigned a cause.

3. Why is cancer attributed to radiation exposure?

The public accepts the perception promulgated and enforced by the linear-no-threshold hypothesis which states that there is an increase in cancer risk for every unit of radiation exposure, or that any amount of radiation may cause cancer [3, 4]. This suggests that there is no “safe level” of radiation and all radiation exposure must be avoided.

The basis for this hypothesis is the extrapolation of risk from cancers produced by high doses into low dose regions where no significant increase in radiation-induced cancer can actually be detected. By suggesting that there is a “firm” link between dose and cancer, regardless of the total dose, it becomes possible to multiply small radiation doses delivered at low or high dose rates to very large populations and calculate an “excess” in cancers for any radiation exposure. However, this link is not “firm” because of the multiple extrapolations that are needed to go from measured changes in cancer frequency to predicted cancers which are extrapolated to exist following very low doses of radiation.

4.0 Multiple extrapolations are needed to move data from one scenario to another. This makes the extrapolated numbers of excess cancers uncertain, since they are not based on measured cancer frequency. The fact that the risk estimates from the LNTH are based on multiple extrapolations compounds the problems associated with uncertainty. The increased cancers are calculated numbers rather than real disease, and must be regarded as such.

4.1 Extrapolation from high to low dose

The extrapolation of biological effects over large dose ranges are not applicable. At low doses of radiation, many cell and molecular switches are activated which involve gene expression, DNA repair, cell death and cell transformation. Each of these biological processes modifies the shape of the
dose-response curve [5]. These biological processes make a linear extrapolation across even one order of magnitude of dose (10-100 mSv) not biologically acceptable.

### 4.2 Extrapolation from high to low dose-rate

The linear extrapolation of dose-rate from the A bomb survivors to environmental exposures is over more than 7 orders of magnitude! Assuming linearity over such a wide range of dose-rates is not reasonable. To accommodate this dramatic dose-rate extrapolation, the LNTH model uses a dose-dose-rate effectiveness factor (DDREF) of 1.5-2.0.

### 4.3 Extrapolation from whole body to partial body radiation exposures

There is extensive literature on the influence of internally deposited radioactive materials in experimental animals (6). The effectiveness of these protracted exposures to limited numbers of organs for the production of cancer is reduced relative to that following whole body acute exposure. Careful experimental animal studies have demonstrated that the risk to an organ per unit of dose from single acute whole body exposure is higher than the same dose delivered only to that organ [7].

### 4.4 Extrapolation across levels of biological organization

At every level of biological organization there are many non-linear dose relationships that have been shown to exist. To ignore all the non-linear data that has been derived at every level of organization and assume that the extrapolation between dose and biological response is linear is not acceptable.

#### 4.4.1 Molecular effects (8, 9, 10, 11)

#### 4.4.2 Cell and tissue effects (12, 13, 14, 15, 16)

#### 4.4.3 Matrix effects (17, 18)

#### 4.4.4 Whole animal effects (19, 20)

#### 4.4.5 Human population effects (21, 22)

### 4.5 Extrapolation between different human populations

The final extrapolation that is used in the a-bomb data is to extrapolate between different populations and ethnic groups, for example from Japanese to the U.S. population. Studies must attempt to control all other exposures and confounding factors and limit the differences between the selected populations to the difference in radiation exposure. In reality, this is almost impossible and therefore, divergent conclusions may often be drawn from the same data, depending on how the populations are defined.

All these extrapolations suggest that it is not possible to establish a linear link between dose and cancer. Using the LNTH, which suggests a “firm” link between dose and cancer risk makes it possible to multiply small radiation doses delivered at low or high dose rates to any exposed population and calculate “excess” cancers for any radiation exposure. These calculations may or may not reflect real risk.

### 5.0 How much radiation is required to increase cancer incidence?

Using the LNTH, it is possible to extrapolate or calculate the number of “excess” cancers in any exposed population without any data on cancer frequency. We have discussed several of the
difficulties associated with extrapolation of cancer frequency to determine the amount of radiation required to produce a cancer.

In many studies the “exposed” population has less cancer than the controls. To quote from BEIR VII (3), “In most of the nuclear industry workers studies, death rates in the worker populations were compared with national or regional rates. In most cases, rates for all causes and all cancer mortality in the workers were substantially lower than in the reference populations. Possible explanations include the healthy worker effect and unknown differences between nuclear industry workers and the general population”. However, when the cancer frequency is higher in the “exposed” population, risk estimates for radiation exposure are calculated even though an “unhealthy worker effect” or other unknown differences between the exposed and control population may be responsible for the increased cancer frequency seen in the exposed population. Thus, a causative link between cancer and radiation dose has not and can not be established following low doses.

Many people argue that just because a cancer increase cannot be detected in a population doesn’t mean that an excess in radiation-induced cancer doesn’t exist. This is theoretically true, but if you cannot detect an increase in cancer, the risk has to be relegated to a lower level of concern relative to other environmental insults such as life style, diet, smoking or asbestos where clear cut cancer increases can be demonstrated. BEIR VII (3) acknowledges that cancer risk following low doses is small. However, it is important to expand this observation to determine just how much radiation is required to produce an excess in total cancer in a population.

To illustrate the amount of radiation that is required to produce a cancer, two tables have been prepared. In these tables, the exposure has been assumed to be an acute whole-body exposure to low-LET radiation. The data for the tables comes from the BEIR VII (3) committee and relates the background cancer frequency, not mortality, to radiation doses. These tables use the data for “solid cancers” and the “conservative” Linear-No-Threshold model. However, as illustrated in this paper, there is a large body of scientific data that suggest that at low doses of low-LET radiation the response to radiation is less than predicted by the LNTH model [5, 15, 20, 23]. The LNT hypothesis is used in these tables to illustrate the point that, even with the conservative LNTH, very large amounts of radiation are required to produce a calculated excess in the cancer incidence.

5.1. Amount of radiation to increase cancer frequency: Radiation in the population held constant amount in the individual varied

In Table 1, the total amount of low LET-radiation delivered acutely to a population is held constant at 700 joules. The sum of the amount of radiation (energy in joules) delivered to each individual provides the best dose metric to estimate the cancer risk to a population [24]. This amount of radiation was selected as a starting point to illustrate that when 700 joules of radiation are delivered to a single 70 Kg man it results in a lethal dose of 10 Gy. Without medical intervention 100% of the people exposed to this large amount of radiation die of acute radiation sickness. Therefore, because this is a high “dangerous” individual dose, it is often assumed by the public that if this same amount of radiation, 700 joules, were delivered to a population it must cause a great deal of death and cancer. If this same large amount of radiation (700 J) were distributed to 10 people, it would result in a high dose of 1.0 Gy per person, and no acute radiation deaths would be expected at this high dose [1]. If 700 J was distributed to increasing numbers of people, as seen in Table 1, increased cancer risk, rather than death, becomes the major concern. Using the LNT hypothesis and the data from the BEIR VII [3] report, if 100 people were exposed to 700 J, the dose to each individual would be 0.1 Gy or 100 mGy. Using these most conservative estimates, one extra cancer is predicted in this exposed population in addition to the 42 cancers normally observed without radiation. Because of linearity, one extra cancer would be predicted in any population size given this amount of radiation energy. If this amount of radiation was distributed to 10,000 people the level of dose to each person would be similar to the amount received from low-LET background radiation in one year or if the dose and risk from background is added over a life time the risk would be 1/70th of that from background radiation.
It can be seen from this table, that this large, lethal amount of radiation can not produce a statistically significant or detectable increase in cancer frequency, regardless of the population size exposed.

<table>
<thead>
<tr>
<th>Number of People</th>
<th>Dose/Person (Gy)</th>
<th>Amount/Person (J)</th>
<th>Amount (J)</th>
<th>Background Cancer</th>
<th>Excess Cancer</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>10</td>
<td>*700</td>
<td>700</td>
<td>.42</td>
<td>0.0</td>
</tr>
<tr>
<td>10</td>
<td>1</td>
<td>70</td>
<td>700</td>
<td>4.2</td>
<td>1.0</td>
</tr>
<tr>
<td>100</td>
<td>0.1</td>
<td>7</td>
<td>700</td>
<td>42</td>
<td>1.0</td>
</tr>
<tr>
<td>1,000</td>
<td>0.01</td>
<td>0.7</td>
<td>700</td>
<td>420</td>
<td>1.0</td>
</tr>
<tr>
<td>10,000</td>
<td><strong>0.001</strong></td>
<td>0.07</td>
<td>700</td>
<td>4,200</td>
<td>1.0</td>
</tr>
<tr>
<td>100,000</td>
<td>0.0001</td>
<td>0.007</td>
<td>700</td>
<td>42,000</td>
<td>1.0</td>
</tr>
</tbody>
</table>

*This is a large lethal amount of radiation given to once person. Cancer can never be detected with this amount of radiation regardless of population size!**

**Background low LET dose/person

Table 1: The total amount of radiation held constant at a level that results in 100% lethality when given to one person (700 J or about 10 Gy) and the population size increased.

<table>
<thead>
<tr>
<th>Number of People</th>
<th>Dose/Person (Gy)</th>
<th>Amount/Person (J)</th>
<th>Amount (J)</th>
<th>Background Cancer</th>
<th>Excess Cancer</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.1</td>
<td>7</td>
<td>7</td>
<td>.42</td>
<td>0.01</td>
</tr>
<tr>
<td>10</td>
<td>0.1</td>
<td>7</td>
<td>70</td>
<td>4.2</td>
<td>0.1</td>
</tr>
<tr>
<td>100</td>
<td>0.1</td>
<td>7</td>
<td>700</td>
<td>42</td>
<td>1</td>
</tr>
<tr>
<td>1,000</td>
<td>0.1</td>
<td>7</td>
<td>7,000</td>
<td>420</td>
<td>10</td>
</tr>
<tr>
<td>10,000</td>
<td><strong>0.001</strong></td>
<td>0.07</td>
<td>7,000</td>
<td><strong>4,200</strong></td>
<td><strong>100</strong></td>
</tr>
<tr>
<td><strong>22,506</strong></td>
<td>0.1</td>
<td>7</td>
<td>158,000</td>
<td>9,450</td>
<td>increased</td>
</tr>
<tr>
<td><strong>86,611</strong></td>
<td>0.14</td>
<td>10</td>
<td>894,557</td>
<td>10,127</td>
<td>572</td>
</tr>
<tr>
<td>100,000</td>
<td>0.1</td>
<td>7</td>
<td>700,000</td>
<td>42,000</td>
<td>1000</td>
</tr>
</tbody>
</table>

*Amount of energy per person and the population size are below the level to detect cancer

Cancer is detectable in this range of population, dose, exposure

**Total amount of radiation, A-bomb and observed response

Table 2. The amount of radiation/person held constant at a level that results in a calculated 1% increase in cancer frequency (7 J/person or about 0.1 Gy).

*The population size and the total amount of radiation (J) required to detect a change in cancer frequency.

5.2. Amount of radiation to increase cancer frequency: Amount of radiation in each individual held constant amount in the population varied.

In the second table, each and every person is assumed to be exposed to a constant, acute exposure of radiation, 7.0 J or 0.1 Gy. This amount of radiation per person is twice the exposure allowed per year for radiation workers. As the number of people exposed to this amount of radiation increases, the amount of energy in the whole population becomes very large. Since the population is the unit measured to detect cancer, the amount of radiation energy in the whole population is assumed to be the important variable [24]. The table shows the background cancer rate and the predicted [3] excess cancer from the radiation exposure. The table illustrates, as has been published that it requires a very large population [25], each with this 7.0 J of radiation, to be able to detect an increase in cancer. If
10,000 people each get 7.0 J of radiation energy the total amount in the population would be 70,000 joules. However, the number of spontaneous cancers also increases as the population size increases. Since the frequency of radiation-induced cancers is small at this population size and level of radiation, it would still not be possible to detect an increase in the cancer frequency above the background level of cancer.

Using statistical methods, it can be calculated that it would require 22,606 people each exposed to 7.0 J to detect an increase in cancer frequency (with a confidence level of 5% false negatives and 5% false positives). This would result in deposition of more than 158,000 joules into the population. A quantity of 700 J delivered as an acute exposure to single person results in 100% lethality. This illustrates how much total radiation energy is required (158,000 J) to produce a significant increase in cancer frequency. From this discussion, it is obvious that although large amounts of radiation energy delivered to a single person are lethal, distribution of that amount over a large population is not lethal and doesn’t produce a detectable increase in cancer frequency. Only after very large amounts of radiation is it possible to detect excess cancer. This highlights the fact that radiation is a very good cell killer, (this is why we use it in radiation therapy), but that it is a poor carcinogen.

5.3 Real life examples

Although it is true that the extrapolations at the heart of public radio-phobia are based on real life examples, the perception of the radiation risk is much greater than the real risk [2]. The risk estimates used in the previous tables comes from extrapolation of the data from the A-bomb survivors. In contrast to calculations the next section reports the raw numbers of cancers. These may or may not be radiation-induced, but they are the excess cancers in the exposed populations compared to carefully selected controls. The use of these numbers help to put the perception of risk and the real risk into prospective so that the public can make their own decisions on the acceptability of radiation risk.

5.3.1 A-Bomb data

The A-bombs were the most terrible radiation events in the history of the world each of these bombs killed about 100,000 people from blast, burn and radiation-induced sickness. The impact of these nuclear weapons must never be trivialized. Of those that survived the bomb in 1945 (60 years ago), there has been a large follow-up study to determine the cause of death in the 86,611 people that were exposed to graded doses of radiation from the bomb. As of 2004, 47,685 of these people have died, leaving about 45% of the exposed population alive for future studies [26,27]. This is a very important population that must continue to be evaluated until their death. However, it can be seen from the survival in this group (45%) and the number of cancers induced (572 in a background cancer frequency of 10,127) that if an individual did not die from the blast, burns and acute radiation exposures, the risk for cancer induction is small and for most of the population the individuals will live out normal life spans. Of course, if a serious radiation catastrophe occurs (atomic bomb) where there will be very high levels of radiation exposure delivered to large populations. There will be large numbers of people killed by the bomb and there will be excess cancer produced. However, cancer should not be the primary concern for this catastrophe. Instead of thinking that each and every ionization results in cancer the reality is that large amounts of radiation can be delivered to human populations and result in very little, or no detectable increase in cancer.

5.3.2. Chernobyl

The exposure to the radiation from the A-bomb was delivered over a very short time. Most environmental radiation exposures are delivered at a low dose-rate protracted over long periods of time. These low dose-rate exposures come from natural background, fallout from atomic bomb tests, nuclear accidents, nuclear waste clean-up or other types of accidents involving radioactive material. These exposures involved both external radiation and radiation from inhaled or ingested radioactive materials which result in very low radiation dose-rates.
With this in mind, it is useful to review the worst nuclear accident in history. In 1986 there was an explosion in Chernobyl’s number four reactor that resulted in wide distribution of radioactive materials which contaminated the entire northern hemisphere. More than 600,000 people involved in trying to control the event and in the clean-up receiving varied and high levels of exposure [28]. Most of these that received high doses were reactor staff, emergency and recovery personnel. The very high doses resulted in 50 deaths after the accident from acute radiation syndromes. There were very high doses to the thyroid glands of children who drank milk from cows that had eaten grass contaminated with radioactive iodine. This resulted in high doses to the thyroid glands which have produced an excess of about 4,000 thyroid cancers “To date there have been 9 deaths from these cancers”. Therefore, in a population of several million exposed people there are to date a total of 59 deaths from this accident. Using the linear-no-threshold extrapolation and the calculated doses to this large population, it can be postulated that a 3,490 additional people will die from cancer as the result of this exposure. This can be related to the 252,000 cancers that will occur spontaneously in a population of this size or to the 5000 known deaths from coal mining accidents that occur each year or 100,000 since Chernobyl.

Chernobyl is a worst case example of a “dirty bomb”. The Chernobyl data are useful in estimating the impact of a terrorist radiation dispersal device where the amount of radiation involved, the population exposed, the radiation doses and thus, the cancer outcome would be much, much less than observed at Chernobyl.

### 5.3.3 Fallout

A prime example of extrapolation from high doses to low doses is seen in estimating “excess” cancers from radioactive fallout from Nuclear Weapons [29]. In this manuscript the authors multiply small doses times the linear risk per unit dose times a huge population and generate a predicted number of “excess cancers”. Using fallout doses and the LNTH, the frequency of “excess” leukemia in the United States was calculated to be 1,800. Even using the LNTH, which may not be true for leukemia, calculated leukemia frequency is a very, very small fraction of the spontaneous frequency (1,500,000) in this population.

This helps us to understand that these are indeed calculations and very large extrapolations. Such small changes in cancer frequency relative to the very high cancer background can never be detected. The use of the words, “might eventually occur” [29], could be replaced with “will never be detected” and demonstrate that this calculated excess risk is a very small public health concern.

### 6. Summary

This paper does not focus on calculated risk associated with radiation [29,30], but on detectable statistically significant increases in cancer frequency and the amount of radiation (energy in Joules) required to produce these changes regardless of the population size exposed. The risks for radiation-induced cancer are low relative to other potential causes of cancer. This paper has demonstrated that it takes a very large amount of radiation to produce an increase in cancer incidence, in contrast to the LNTH, which promotes the public misconception that any amount of radiation causes cancer.

Data from A-bomb, Chernobyl and fallout can and should be used by the scientific community to construct models, make risk estimates and predict cancer frequency. However, these calculations need to be complemented with the available clearly understandable raw data. When this is done it is obvious that it takes a lot of radiation to make cancer and that excess cancers are not the prime concern from a nuclear event.
7. References


[3] BIER VII-Phase 2, Health risks from exposure to low levels of ionizing radiation, Committee to assess health risks from exposure to low levels of ionizing radiation, National Research Council (National Academy of Sciences, Washington, D.C.), 2005


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