A Rebuttal to Chiropractic Radiologists’ View of the 50-year-old, Linear-No-Threshold Radiation Risk Model

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This discussion is in response to a letter-to-the editor in the form of a ‘Commentary’ by Bussieres, Ammendolia, Peterson, and Taylor1 concerning our original commentary: ‘On “phantom risks” associated with diagnostic ionizing radiation: evidence in support of revising radiography standards and regulations in chiropractic,2 published in the December 2005 issue of this journal.

Bussieres et al.1 have expressed that our original commentary lacked credibility, while they claimed that: 1) we have a vested financial interest in promoting routine and follow-up x-rays; 2) we provided a biased and unscientific evaluation of the evidence; 3) there is “no convincing evidence that the use of radiography for spinal biomechanical assessment (other than for scoliosis) is of any therapeutic value”; and 4) ‘unnecessary’ x-rays are associated with high health care costs.

Ad Hominem Attacks
First, we will address their1 “Ad Hominem” attacks on us. They referred to our paper as “self serving”, “professionally irresponsible”, and having a “vested financial interest”. An Ad Hominem attack has no place in a scientific debate. In fact, the Ad Hominem attack is one of the fallacies in scientific debates; instead of critiquing the science, attack the character of the individual.3 According to Stein,3 when an individual resorts to an Ad Hominem attack, they have lost credibility. Normally, we would ignore such insults, however, we note only two of the four authors (Harrisons) have any financial gain from CBP technique (by seminar attendance) – but how do doctors, in different States/Provinces/Countries, x-raying their own patients, transcribe the knowledge gained about spinal health into a financial benefit for any of our authors?

Biased and Unscientific Evaluation of the Evidence
Second, Bussieres et al.1 claimed that we presented a biased and unscientific evaluation of the evidence on cancer risks from exposure to ionizing radiation. We believe, while being brief, we presented the facts. We presented a review of 7 studies of human exposures to low-level radiation that resulted in health benefits for these populations.2 So, in support of our previous article, we will expand on our evaluation of the evidence. For our discussion, we direct the reader to Figure 1 as a visual representation of the Linear-No-Threshold Model (LNT) and the Radiation Hormesis Model of cancer risks from exposure to ionizing radiation.

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Controversy in the BEIR VII Conclusions

Bussieres et al.\textsuperscript{1} fall prey to a typical argument on cancer risks by citing references that neglect hormesis evidence. They\textsuperscript{1} cited the 2005 BEIR report\textsuperscript{6} and a few other references used in the BEIR report.\textsuperscript{7–10} We had read parts of the long BEIR report and noted that it often made its claims while assuming the LNT model. In fact, what Bussieres et al.\textsuperscript{1} failed to mention is that, in 2005, Aurengo et al.\textsuperscript{11} compared the Reports by the French Academy of Sciences and the French Academy of Medicine, both of which reached the same opposite conclusion to BEIR VII.

Aurengo et al.\textsuperscript{11} found that the BEIR report neglected hormesis evidence and neglected negative analyses of the studies that were cited. Also we note that the BEIR Report has not been officially issued yet, (only a preliminary draft on their web site) and is still subject to change. In contrast, the unanimous report by the French Academy of Sciences and National Academy of Medicine (2005) stated:

"In conclusion, this report doubts the validity of using the LNT in the evaluation of the carcinogenic risk of low doses (<100mSv) and even more for very low doses (<10mSv). ... the use of LNT in the low dose or dose rate range is not consistent with the current radiobiological knowledge; LNT cannot be used without challenge ... for very low doses (<10mSv). ... The eventual risks in the dose range of radiological examinations (0.1 to 5 mSv, up to 20mSv for some examinations) must be estimated taking into account radiobiological and experimental data. An empirical relationship which is valid for doses higher than 200 mSv may lead to an overestimation of risk associated with doses one hundredfold lower and this overestimation could discourage patients from undergoing useful examinations and introduce a bias in radioprotection measures against very low doses (<10 mSv)."\textsuperscript{11}

Cohen's Outline

In preparation for this rebuttal, we contacted one of the leading authorities on radiation exposure risks, Dr. B. L. Cohen,\textsuperscript{12–22} University of Pittsburgh, Pennsylvania, USA. What follows is the outline that Cohen provided to us:\textsuperscript{23}

1. Problems with the Basis for the Linear-No-Threshold Theory

2. Direct Experimental Challenges to the Basis for LNT

3. Effects of Low-Level Radiation on Biological Defense Mechanisms

4. Stimulation of the Immune System

5. Cancer Risks vs Dose in Animal Experiments

6. Cancer Risks vs Dose in Human Experiments
   a. Critique of Data Frequently Cited Supportive of LNT
   b. Data Contradictory to LNT

1. Problems with the Basis for LNT

The LNT model is theoretical and simple: A single particle of radiation hitting a single DNA molecule in a single cell nucleus of the human body can initiate cancer. Therefore cancer initiation probability is proportional to the number of events, which is proportional to the number of particles of radiation, which is proportional to the dose. Thus the LNT theory is "the risk is proportional to the dose".\textsuperscript{23} The problem with this simple theory is that other factors affect cancer risk, i.e., human bodies have biological defense mechanisms that prevent the vast majority of radiation events from becoming a cancer.\textsuperscript{24}

There are several defense mechanisms: (1) The most important cause of DNA injury is corrosive chemicals termed reactive oxygen species (ROS) and low-level radiation has been shown to stimulate the scavenging processes to eliminate these from cells;\textsuperscript{25} (2) There is abundant evidence that low-level radiation stimulates the immune system, while high levels/doses depress the immune response;\textsuperscript{26} (3) Radiation can alter cell timing, i.e., the time before the next cell division/mitosis and low-levels of radiation increase this time and allow for more possible DNA repair; (4) Low dose hypersensitivity and increased radiation radioresistance are affected by low-level radiation;\textsuperscript{27} and (5) It is now recognized that tissue response, whole organ response, and organism response, rather than just single cellular response, must be considered.\textsuperscript{11}

There is another obvious failure of the original LNT model. The theory predicts that the number of initiating events is roughly proportional to the mass of the animal being irradiated. However, research has shown that the cancer risk for a given radiation field is similar for a 30 gram mouse and a 70,000 gram human.\textsuperscript{26}

Interestingly, validity of the LNT model is based on double strand breaks (DSB) in DNA molecules. Howev-
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er, Feinendegen estimated that ROS causes about 0.1 DSB per cell per day, whereas 100 mSv (10 rem) of radiation causes about 4 DSB per cell. Using this information, a 100 mSv dose of radiation would increase the lifetime risk of cancer (28,000 days x 0.1 DSB/day) by only about 0.14% (4/28,000), but the LNT model predicts 7 times that much at 1%.

2. Direct Experimental Challenges to the Basis for LNT
A direct failure of the basis for the LNT model is derived from microarray studies, which determine what genes are up-regulated and what genes are down-regulated by radiation. It was discovered that generally different sets of genes are affected by low-level radiation as compared to high-level doses. In 2003, Yin et al. used doses of 0.1 Sv and 2.0 Sv applied to mouse brain. The 0.1 Sv dose induced expression of protective and repair genes, while the 2.0 Sv dose did not.

A similar study on human fibroblast cells was conducted in 2002 by Golder-Novoselsky et al. Using doses of 0.02 Sv and 0.5 Sv, they discovered that the 0.02 Sv dose induced stress response genes, while the 0.5 Sv dose did not. Several other microarray studies have demonstrated that high radiation doses, which serve as the “calibration” for LNT, are not equivalent to adding an accumulation of low radiation doses.30

In fact, in 2001, Tanooka studied tumor induction by irradiating the skin of mice throughout their lifetimes. For irradiation rates of 1.5 Gy/week, 2.2 Gy/week, and 3 Gy/week, the percentage of mice that developed tumors was 0%, 35%, and 100%, respectively. This data demonstrated a clear threshold response directly in conflict with predictions of the LNT model.

3. Effects of Low-Level Radiation on Biological Defense Mechanisms
In 1994, the United Nations Scientific Committee on Effects of Atomic Radiation (UNSCEAR) report defined “adaptive response” as a type of biological defense mechanism that is characterized by sequent protection to stresses after an initial exposure of a stress (like radiation) to a cell. For radiation experiments, this is studied by exposing cells to low-doses to prime the adaptive response and then later exposing it to a high radiation “challenge dose” to see what happens. There have been several experiments in this topic, and we report on just a few of these.

In 1990, Cai and Liu exposed mouse cells in 2 different ways: (1) a high dose of 65 cGy (65 rad), and (2) a low-dose of 0.2 cGy before the high-dose of 65 cGy. The number of chromosome aberrations reduced in the second group compared to the first group was 38% bone marrow cell aberrations reduced to 19.5% and 12.6% spermatocyte aberrations reduced to 8.4%.

In 1992, Shadley and Dai irradiated human lymphocyte cells, some with high doses and some with a low-dose a few hours before a high-dose. The number of chromosome aberrations caused by a high-dose was substantially reduced when a preliminary low-dose was given first.

In 2001, Ghiassi-nejad et al. studied this effect in humans. In Iran, residents of a high background radiation area (1 cGy/year) were compared to residents in a normal background radiation area (0.1 cGy/yea). When lymphocytes, taken from these groups, were exposed to 1.5 Gy (150 rad), the percentages of aberrations were 0.098 for the high background area versus 0.176 (about double) for the low background area. The radiation in the high background area protected its residents from the 1.5 Gy dose.

4. Stimulation of the Immune System
The effects of low-level radiation on the immune system are important since the immune system is responsible for destroying cells with DNA damage. Low doses of radiation exposure cause stimulation of the immune system while high doses reduce immune activity.

Contrary to expectations from the basic assumption of the LNT model (cancer risks depends only on total dose), effects on the immune system are quite different for the same total dose given at a low dose rate (summation of several small doses) versus one high dose rate, i.e., at low dose rates the immune system is stimulated, while at high doses, cancers are caused.

5. Cancer Risks vs Dose (Animal Experiments)
To test the validity of the LNT model, there have been numerous direct experiments of cancer risk versus dose, with animals exposed to various radiation doses. In 1979, Ullrich and Storer reported that exposed animals lived up to 40% longer than controls. In a series of animal studies in the 1950s and 1960s, review articles by Finkel
and Biskis\textsuperscript{52–54} reported, with high statistical significance, that the LNT model over-estimated the cancer risks from low-level radiation exposures and they reported a threshold not a linear response.

In a 2001 review of over 100 animal radiation experiments, Duport\textsuperscript{55} reported on studies involving over 85,000 exposed animals and 45,000 controls, with a total of 60,000 cancers in exposed animals and 12,000 cancers in control animals. In cases where cancers were observed in controls receiving low doses, either no effect or an apparent reduction in cancer risk was observed in 40\% of the data sets for neutron exposure, 50\% of the data sets for x-ray exposure, 53\% of the data sets for gamma rays exposure, and 61\% of the data sets for alpha particles exposure.

6. Cancer Risks vs Dose (Human Experiments):
A. Critique of Data Frequently Cited in Support of LNT
The principle data cited and used to support the LNT model are those for solid tumors (all cancers except leukemia) in the survivors of the Japanese atomic bomb explosions. Pierce’s 1996 paper\textsuperscript{56} reported data from 1945–1990. By ignoring the error bars, supporters of the LNT model claim that the data suggests an approximate linear relationship with intercept near zero. But there is no data that gives statistical significant indication of excess cancers for radiation doses below 25 cSv.\textsuperscript{57} Leukemia data from Japanese A-bomb survivors strongly suggest a threshold above 20 cSv and the contradiction to the LNT model is recognized by the author.\textsuperscript{57} In 1998, Cohen\textsuperscript{58} used the three lowest dose points in the Japanese data (0-20 cSv) to show that the slope of the dose-response curve has a 20\% probability of being negative (i.e., Hormesis = risk decreasing with increasing dose).

The next often cited evidence, by supporters of the LNT model, is the International Association for Research on Cancer (IARC) studies on monitored radiation workers. In 1995, Cardis et al.\textsuperscript{59} reported on 95,673 monitored radiation workers in 3 countries and in a follow-up study by the same authors in 2005,\textsuperscript{60} they reported on 407,000 monitored workers in 154 facilities in 15 countries. In the first study, for all cancers except leukemia (there were 3,830 deaths, but no excess over the number expected from the general population), the risks were reported as –0.07/Sv with 90\% confidence limits of (−0.4,+0.3), i.e., there is NO support for LNT from this data! However, for leukemia (146 deaths), they reported a positive correlation, but their data had no indication of any excess cancers (risks) below 40 cSv. Most importantly, these authors discarded 3/7 of their data points when observed/expected was less than unity. In fact, Cohen\textsuperscript{23} noted that (1) no information on such confounding factors as smoking was given, (2) if data from just one of the 15 countries was eliminated (Canada), the apparent excess is no longer statistically different from zero, (3) the authors did not consider non-occupational exposure (natural background radiation) and if they had, they would have noticed that their excess “signal” was much smaller than the “noise” from background radiation.

Often critics of Radiation Hormesis use the “Healthy Worker effect” to discredit what is found. When studying mortality rates for employed workers compared to the general population, it is found that workers have lower mortality rates. In Sweden in 1999, Gridley\textsuperscript{61} compared 545,000 employed women to 1,600,000 unemployed women. He reported that the cancer incidence rate was slightly higher for employed women (1.05 ± 0.01). This eliminated the claims of the “Healthy Worker effect”. For an example of improper use of this effect, in 2005 Rogel\textsuperscript{62} studied 22,000 monitored workers in the French nuclear power industry. The cancer mortality rate was only 58\% of the general French population. Instead of concluding a Hormesis effect, Rogel claimed that this large difference was due to the “healthy worker effect”.\textsuperscript{62}

B. Data Contradictory to LNT
There is much data contradictory to the LNT model. There are multiple human studies which show a radiation Hormesis effect.\textsuperscript{16,22,64–72}

For breast cancer in Canadian women, Miller\textsuperscript{63} reported a decrease risk with increasing dose up to 25 cSv. Howe\textsuperscript{64} (for lung cancer in Canadian women) and Davis\textsuperscript{65} (for 10,000 people in Massachusetts) separately reported a decrease in cancers in the low-dose region up to 100 cSv. There is a difference between lung cancer rates in Japanese A-bomb survivors and the data from Howe and Davis: the Japanese survivors show a much higher risk at all doses. This indicates that one must not accept A-bomb survivor data (one large dose) to predict risks from low-dose rates where low-level doses are summed. It is known that risks from summing low doses (such as spinal radiography use in chiropractic) does not
equal the risks from one large dose (Tubiana).30
Kostyuchenko66 reported on a follow-up of 7,852 vil-
lagers exposed in the 1957 radioactive storage facility ex-
plosion in Russia. The cancer mortality rate was much
lower in these villagers than in unexposed villagers in the
same area supporting a hormetic effect. However, the ex-
posure of the workers directly at the facility was quite
high in one dose and these workers were found to have an
increase in cancers indicating a dose threshold for in-
crease cancers (Koshurnikova 2002).67
In 1997, Sakamoto68 reported on radiation treatments
in non-Hodgkin’s lymphoma. Patient groups were ran-
domly separated into radiation treatment and non-radia-
tion treatment. After 9 years, 50% of the control group
died but only 16% of the irradiated group died.
The conclusion from Cohen’s outline23 is that the LNT
theory fails badly in the low dose region. It grossly over-
estimates the cancer risks from low-level radiation. The
cancer risk from the vast majority of normally encoun-
tered radiation exposures (background radiation, medical
x-rays, etc.) is much lower than estimates given by sup-
porters of the LNT model, and it may well be zero or
even negative.

Critique of Bussieres et al.1 Cancer Risk References
As previously discussed, Aurengo11 reported on two
groups who came to the opposite conclusions compared to
the 2005 BEIR report.6
In their risks argument, Bussieres et al.1 passionately
present (their table 1) the number of estimated cancer
deaths per year as calculated from known x-ray usage in
the Berrington de Gonzalez study.8 This study8 relied
solely on the LNT model and has been criticized by many
for several reasons. First, as several critics noted73–76 and
for which the authors8 admitted in their reply,77 they
failed to weigh the benefits of diagnostic x-rays in their
study, which only guarantees an overestimate of death
calculations. Another criticism was their assumption of the
LNT to make their cancer death estimations. Tubiana
et al.74 pointed out the “speculative nature” of the LNT
hypothesis, and along with Simmons,78 noted that the
LNT is only compatible with exposures greater than
200mSv (significantly more radiation than any medical
x-rays).
Another criticism is that the Japanese survival data has
significant limitations to extrapolate its use for x-ray risk
estimates from γ rays. The Japanese exposure was a one
time high dose, which is entirely different from accumu-
lated small dose rates. Herzog and Rieger73 note that this
data will overestimate cancer risk because the Japanese
were exposed to γ rays from bombs, a different energy
spectrum than x-rays, but also the additional exposures of
β radiation, radionuclides emitting β, and high-energy α
radiation from contaminated water, food, and dust.
Yet another criticism of the study was that there was no
mention of the complexity and effectiveness of the hu-
man cell’s defenses against ionizing radiation. Tubiana et
al.74 noted there are hundreds of enzymes devoted to pro-
tect a cell from these effects and that “there is no single
defense mechanism but a variety…. an adaptive effect ex-
ists and a hormetic effect has even been seen in more than
half of experimental studies after low or moderate dos-
es.”79–80 “Extrapolation from high doses to low doses
with LNT is unlikely to be able to assess the risks accu-
rately.”74
For Bussieres et al.1 to use this study as ‘evidence’ for
rationale for x-ray guidelines is dubious. In fact, without
mentioning the scientific peer concerns surrounding the
study,73–76,78 (it is readily observed on pub-med that a
number of letters to the editor were published) we can
only conclude that they1 are the ones presenting a ‘bi-
ased’ evaluation of the evidence.

No Convincing Evidence for the Use of
Radiography for Spinal Assessment
Next, we arrive at Bussieres et al.1 third complaint about
our original article. Using 3 references, (Ammendolia et
al.,81 Mootz et al.,82 Haas et al.83) Bussieres et al.1 state x-
rays are not useful due to lack of clinical relevance. The
Ammendolia article81 is a questionnaire study with a fo-
cused group interview about clinicians’ opinions and
practices regarding acute low back pain only. This paper
disregarded an entire body of evidence in opposition to
the papers’ conclusions that was not addressed.84
Further, as in their commentary at hand,1 medical
‘diagnostic red flag only’ references are used for a
chiropractic argument against x-ray usage. That is, chi-
ropractic doctors differ fundamentally in assessment,
diagnosis, and treatment than their medical counterparts.
Therefore, Medical references apply to the profession of
drugs and surgery, NOT chiropractic, which uses physical
forces applied to spines. Not only is the spinal structure
intricately examined by chiropractors but as Mootz et al. states: “identifying contraindications to ... manipulation, however, is a purpose that belongs only to practitioners of manual therapies, especially DCs.”

Bussieres et al. commit a common error by citing Haas et al. without acknowledging that this is the middle paper of a 3-part debate. The Haas reference has been clearly rebutted.

The Mootz et al. review is a decade old, and even they stated “using plain film imaging, single studies result in relatively insignificant radiation dosages and expense.” At that time, (1997) Mootz et al. suggested that changes in patient misalignment had yet to be determined to impact clinical progress. Despite even being arguable at that time, in the following decade, there has been a variety of good quality, biomechanical and outcome studies defining this relationship published by the CBP group. In fact, in the recent ICA X-ray Guidelines (PCCRP), there are hundreds of Chiropractic outcome studies cited.

Furthermore, Bussieres et al. failed to acknowledge the recent randomized trial by Khorshid et al. where a clinically and statistically significant improvement in autistic children treated with upper cervical technique (using x-rays) was found compared to those treated with standard full spine technique (no x-rays).

Previous authors have stated that guidelines for chiropractic clinicians’ and manual therapists’ utilization of x-ray should be different from those of a medical practitioner who does not use spinal adjustments/forces and rehabilitation procedures as treatment for spinal subluxations. In studies specifically considering the role of chiropractic treatment interventions, spinal radiographic views indicate that between 66%–91% of patients can have significant abnormalities affecting treatment; 33% can have relative contraindications and 14% can have absolute contraindications to certain types of chiropractic adjustment procedures.

**Radiography Is Not Cost Effective**

Bussieres et al. mention that patients receiving radiography are more satisfied with care, but then discard this finding in favor of a ‘cost reducing model of health care’. We feel it is important to present this information in proper context.

For example, in a randomized trial comparing the intervention of lumbar radiography to no radiography in patients with at least 6 weeks duration of low back pain, Kendrick et al. found no differences in outcomes between the groups. Problematically, the intervention used for treatment did not specifically address any structural spinal displacements as chiropractic clinicians would readily apply. Importantly, patients receiving radiography were more satisfied with the care they received. Furthermore, patients allocated to a preference group, where the decision to receive lumbar radiography was made by them, achieved clinically significant improved outcomes compared to those randomized to a non-radiography or a radiography group.

Thus, undercutting patient choice by ‘red flag’ only guidelines, as Bussieres et al. would have us do for chiropractic practice, limits a patient’s right to choose and may impair or slow recovery. However, just as importantly, we caution Bussieres et al. for applying results from clinical studies with pharmacology and PT treatments to Chiropractic situations where the treatments are vastly different (physical forces applied to spinal structures).

While cost-effectiveness analysis may favor limited x-ray utilization in a volume 3rd party payer scenario where maximization of profits is the goal, in the individual patient, case specific circumstances can lead to a different conclusion. It is our perspective that, in chiropractic clinical practice, the needs of the one outweigh the needs of the many or the managed care organization; our duty is to identify the spinal problem of the individual and develop solution strategies where possible.

Lastly, there is an expectation by the consumer to have a thorough spinal evaluation when seeing a DC for a health problem and this includes an x-ray evaluation for alignment of the spine and the state of health of the spine.

**Conclusion**

In summary, Bussieres et al.’s Ad Hominem attacks on us and their main arguments against routine use of radiography in common practice, radiation risks, and lack of clinical usefulness are without scientific support. In fact, Luckey stated, “for every thousand cancer mortalities predicted by the linear models (LNT), there will be a thousand decreased cancer mortalities and ten thousand persons with improved quality of life.”
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