Research Article

Worldwide Increasing Incidences of Cutaneous Malignant Melanoma

Dianne E. Godar

Center for Devices and Radiological Health, Food and Drug Administration, 10903 New Hampshire Avenue, WO64-4024, Silver Spring, MD 20993, USA

Correspondence should be addressed to Dianne E. Godar, dianne.godar@fda.hhs.gov

Received 20 November 2010; Revised 6 April 2011; Accepted 11 August 2011

Academic Editor: Arash Kimyai-Asadi

Copyright © 2011 Dianne E. Godar. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

The incidence of cutaneous malignant melanoma (CMM) has been increasing at a steady rate in fair-skinned populations around the world for decades [1–12]. Scientists are not certain why CMM has been steadily increasing over the decades, but strong intermittent UVB (290–320 nm) exposures, especially sunburn episodes, evidently initiate CMM [13]. The UV A (321–400 nm) passing through glass windows in offices and cars has been proposed to promote CMM [14]. In support of those possibilities exists the paradox between indoor and outdoor worker’s UV exposures and their incidences of CMM. Although outdoor workers get three to ten times the annual UV dose that indoor workers get [15, 16], they have similar or lower incidences of CMM [17]. Scientists think the increasing incidence of CMM is linear based on surveillance epidemiology and end results (SEER) data in the USA that only dates back to 1973 [12], but it may actually be exponential in the USA and in some other regions of the world.

To understand what factor(s) may be responsible for the increasing incidence of CMM, one must know the temporal incidence for as many decades as possible. Because whatever the causative agent is, or agents are, it must have entered or left our environment some time (∼10–30 yrs) before the increasing trend was first documented back in 1935 [1]. Thus, one can analyze the CMM incidence data for fair-skinned people around the world and plot it temporally by each country in the northern and southern hemispheres and by latitude. Further analysis of $R^2$ values can determine if the curves are linear or exponential. This paper will analyze the CMM incidences of fair-skinned populations all over the world, test if the increases are exponential or linear, and show that the increasing incidence decreases with increasing latitude until ∼50°N, where it reverses and begins to increase with increasing latitude in Northern Europe.

1. Introduction

The incidence of cutaneous malignant melanoma (CMM) has been increasing at a steady rate in fair-skinned populations around the world for decades [1–12]. Scientists are not certain why CMM has been steadily increasing over the decades, but strong intermittent UVB (290–320 nm) exposures, especially sunburn episodes, evidently initiate CMM [13]. The UVA (321–400 nm) passing through glass windows in offices and cars has been proposed to promote CMM [14]. In support of those possibilities exists the paradox between indoor and outdoor worker’s UV exposures and their incidences of CMM. Although outdoor workers get three to ten times the annual UV dose that indoor workers get [15, 16], they have similar or lower incidences of CMM [17]. Scientists think the increasing incidence of CMM is linear based on surveillance epidemiology and end results (SEER) data in the USA that only dates back to 1973 [12], but it may actually be exponential in the USA and in some other regions of the world.

To understand what factor(s) may be responsible for the increasing incidence of CMM, one must know the temporal incidence for as many decades as possible. Because whatever the causative agent is, or agents are, it must have entered or left our environment some time (∼10–30 yrs) before the increasing trend was first documented back in 1935 [1]. Thus, one can analyze the CMM incidence data for fair-skinned people around the world and plot it temporally by each country in the northern and southern hemispheres and by latitude. Further analysis of $R^2$ values can determine if the curves are linear or exponential. This paper will analyze the CMM incidences of fair-skinned populations all over the world, test if the increases are exponential or linear, and show that the increasing incidence decreases with increasing latitude until ∼50°N, where it reverses and begins to increase with increasing latitude in Northern Europe.

2. Materials and Methods

One can obtain CMM incidence’s throughout the decades from 1935 to 2007 for Australia, New Zealand, USA (except Connecticut 1935–1940 [1] and 1945 [2] and New York State 1955 [2]), middle Europe, Canada, and Northern Europe from the International Agency for Research on Cancer (IARC) [3–11]. The CMM incidence data from IARC was
Table 1: Averaged CMM age-adjusted incidences among whites per 100,000 people (world population) in each country of the northern and southern hemispheres (plotted in Figure 1); data is from [3–11] except 1940 [1], 1945, and 1955 [2] for the USA.

| Latitude/place          | 1940  | 1945  | 1950  | 1955  | 1960  | 1965  | 1970  | 1975  | 1980  | 1985  | 1990  | 1995  | 2000  |
|-------------------------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
| 30°S Australia          |       |       |       |       |       |       | 16.33 | 17.89 | 21.43 | 25.90 | 32.20 | 32.13 |       |
| 40°N USA                | 1.00  | 1.48  |       |       |       |       | 6.10  | 7.65  | 9.55  | 15.55 | 18.50 | 20.80 | 27.40 |
| 49°N Middle Europe      |       |       | 2.24  | 2.67  | 3.73  | 4.56  | 7.23  | 9.95  | 10.85 | 12.75 | 14.67 | 17.74 |       |
| 52°N Canada             |       |       |       | 2.00  | 2.12  | 2.64  | 3.18  | 4.73  | 5.97  | 7.08  | 7.95  | 9.74  |       |
| 60°N Northern Europe    | 1.80  | 2.24  | 3.28  | 4.18  | 5.09  | 5.99  | 7.68  | 9.20  | 10.29 | 11.33 |       |       |       |

Figure 1: Temporal exponential increase in the incidence of CMM by latitude worldwide. Note that only New Zealand (40°S) and Northern Europe (60°N) have linear rates of increase.

averaged from the following states/provinces/territories to get a mean value for each country at average latitude (Table 1 and Figure 1):

(1) Australia (~30°; range 19.5°–42.5°S)—Queensland (19.5°S), Western (24°S), South (32°S), New South Wales (33°S), Capital Territory (35.5°S), Victoria (36.5°S), and Tasmania (42.5°S).

(2) New Zealand (~40°S).

(3) USA (~40°N; range 20–47°N)—Hawaii (20°N), Los Angeles, California (34°N), Atlanta, Georgia (34°N), New Mexico (34°N), San Francisco, California (38°N), Utah (39°N), Connecticut (41.5°N), Iowa (42°N), Michigan (43.5°N), New York State (43°N; excludes New York City), and Washington State (47°N).

(4) Middle Europe (~49°N; range 46–52°N)—Switzerland (46°N), Slovenia (46°N), Romania (46°N), Hungary (47.5°N), Slovakia (48.5°N), France (48.5°N; Bas-Rhin), Germany (49.5°N; Saarland), all of Poland (51°N), The Netherlands (52°N), and all of England (52°N).

(5) Canada (~52°N; range 45–65°N)—Nova Scotia (45°N), New Brunswick (46.5°N), Prince Edward’s Island (46.5°N), Ontario (51°N), Newfoundland (53°N), Quebec (53°N), Albert (54°N), British Columbia (54°N), Manitoba (54°N), Saskatchewan (54°N), and Northwest Territories (65°N).

(6) Northern Europe (~60°N; range 53–65°N)—Ireland (53°N), Denmark (56°N), Scotland (57°N), Sweden (62°N), Iceland (63°N), Norway (64°N), and Finland (65°N).

Figure 2 has the same CMM incidence data for the year 2000 as shown in Figure 1, only plotted by latitude [1–11].

Figures 3(a) and 3(b) show CMM incidence data from [1–11], which includes 10 registries, the same 9 registries as SEER 9 along with Los Angeles. Figure 4(a) CMM incidence data for the USA (1973–2007) is from the SEER website at (http://seer.cancer.gov/faststats/selections.php) [12]. The data type is “SEER incidence,” the statistic type is “age-adjusted,” the year range is 1975–2007 (SEER 9), and the race/ethnicity is “white (includes Hispanics),” for both sexes of all ages. SEER 9 (white includes Hispanic) compared to SEER 17 (non-Hispanic white) has ~12-100,000 people lower incidence of CMM. The SEER 9 registries are Atlanta, Georgia; Connecticut; Detroit, Michigan; Hawaii; Iowa;
New Mexico; San Francisco-Oakland, California; Seattle-Puget Sound, Washington; Utah. Data are available for cases diagnosed from 1973 and later for these registries with the exception of Seattle-Puget Sound and Atlanta. The Seattle-Puget Sound and Atlanta registries joined the SEER program in 1974 and 1975, respectively. Seer 11 includes Los Angeles and San Jose-Monterey, California starting in 1992. Figure 4(b) shows the same data [12] as in Figure 4(a) only extended back to 1940 using data from [1–11].

3. Results

Table 1 contains the averaged CMM incidences of fair-skinned populations around the world from 1940 to 2000 for each country or region of the world [1–11]: Australia (∼30°S), New Zealand (∼40°S), USA (∼40°N), Middle Europe (∼49°N), Canada (∼52°N), and Northern Europe (∼60°N). New Zealand has the highest incidence of CMM closely followed by Australia while Middle Europe has the lowest incidence.

Figure 1 shows a temporal plot of the CMM incidence from 1940 to 2000 [1–11]. Note that Australia, USA, Middle Europe, and Canada all have exponential increases except New Zealand and Northern Europe, which have linear increases. In fact, all the countries of Northern Europe have linear increases in CMM except Iceland, which is exponential.

Figure 2 shows the incidence of CMM in 2000 decreases with increasing latitude up to ∼50°N where it changes and begins to increase with increasing latitude in Northern Europe [1–11]. Notice that the CMM incidence increases with decreasing latitude; however, near 50°N in Northern Europe the incidence begins to increase with increasing latitude. Note that the CMM incidence data from 1960 to 2000 all show the same change near 50°N (see Table 1).

Figure 3 shows the CMM incidence data in the USA analyzed in different ways to know whether or not the increase is exponential or linear [1–11]. Figure 3(a) shows how the CMM incidence data at 39°N can be linear if we only average San Francisco and Utah together. The data at 43°N is exponential because Connecticut is included in that data set. Figure 3(b) shows that when the Connecticut CMM incidence data is included with the 39°N data, it becomes exponential, while the data at 43°N becomes linear. Connecticut data extends a couple of decades further back in time (to 1935), which may be necessary to know if the trend is truly linear or exponential but might represent underestimates in CMM during that earlier time frame that would make the data appear to be exponential when it is really linear.

Figure 4(a) shows the SEER 9 data from 1975 to 2005 [12], which appears to be almost perfectly linear ($R^2$ is 0.9954), possibly because it does not extend far enough into the past as the data presented in Figures 1, 3(a), and 3(b). Figure 4(b) shows the same SEER 9 data [12] in Figure 4(a) only extended back in time to 1940 using references [1] through [11]. This changes the USA data from a linear ($R^2$ is now 0.9189) to an exponential ($R^2$ is 0.9755) increase in the incidence of CMM.

4. Discussion

The incidences of CMM in fair-skinned, indoor-working people have been increasing worldwide for decades (Figure 1). The countries with the highest incidences per annual erythemally weighted UV dose [15] are closest to
the equator: Australia (∼30°S), New Zealand (∼40°S), and the USA (∼40°N). The regions of the world with lowest fair-skinned incidences of CMM are in Middle Europe, around 49°N; however, the decreasing trend with increasing latitude changes around ∼50°N so that Canada (∼52°N) and Northern Europe (∼60°N) have higher incidences than Middle Europe (Figure 2). In fact, Northern Europe at ∼60°N has a higher incidence than Canada at 52°N and almost equates the incidence in the USA at ∼40°N. The increasing incidence of CMM appears to be exponential in most regions except Europe; however, as analysis in Figures 3(a) and 3(b) show, data in the USA extending to 1940 can make the CMM incidence appear to be exponential. The SEER data from 1973 to 2007 suggests that the CMM incidence is linearly increasing in the USA (R2 is 0.9954, Figure 4(a)). However, when the Connecticut [1, 2], New York State [2], and IARC data [3–11] extend the SEER data back to 1940, the incidence of CMM in the USA increases in an exponential manner (Figure 4(b)), possibly indicating that longer periods are needed to know if the increase is truly linear or if it is really exponential. Thus, we cannot be completely certain if the increase is linear or exponential in other countries because it may be that the data has to be collected for five decades or more to be conclusive.

Whether or not the incidence of CMM is increasing linearly or exponentially does not change the fact that it is increasing at the alarming rate of about 4-5% per year. In order to slow or stop this increasing trend, one must know what is causing it and change it. Based on the temporal plot shown in Figure 1, we know whatever started the increasing incidence of CMM either entered or left our environment before 1935, because that is when we have documented data for the first increases in CMM in the USA. Fluorescent lights (mid-1940's; [18]), sunscreens (late 1950's for UVB absorbing and 1988 for UVA and UVB absorbing; [19]), and tanning devices (∼1978; [20]) all entered our environment after the increasing incidence of CMM was first documented in the USA back in 1935 [1]. Thus, one should analyze what happened before 1935—during the early 20th century—to discover what may have really affected the incidence of CMM.

In the early 20th century, people went against evolution by going indoors during the day to work, which drastically decreased their daily amount of cutaneous vitamin D3 and exposed them to only UVA radiation passing through glass windows [14]. The artificial UV barrier created by windows divided UVB from UVA, so that the vitamin-D-making UVB wavelengths [21] were excluded and only the vitamin-D-breaking [22] and DNA-mutating UVA wavelengths [23–25] were included in our indoor-working environment. Possibly because this unnatural UV environment existed for decades in buildings and later in cars [26], CMM was promoted by UVA, after being initiated by UVB sunburns and began to steadily increase in the mid-1930’s.

Along these lines of reasoning, we now also have the increasing incidence of CMM with increasing latitude above ∼50°N (Figure 2). People living above 50°N go to the beach during the summer and get sunburned at lower latitudes to initiate CMM and then return home to northern latitudes that have primarily UVA for most of the year to promote CMM. The higher latitudes also allow the sun to aim more often at a perpendicular angle to the window glass allowing more UVA to pass through and directly expose people’s skin during their workday. In addition, above ∼50°N there is little UVB to make cutaneous vitamin D3 most of the year [27]. Further, in the northern regions of the world (above 37°N), a vitamin D3 “winter” occurs from at least November to February, which extends from October to March at higher latitudes, when the dose rate of UVB is too low to make any previtamin D3 even if an office worker goes outside during peak hours [27]. On the other hand, UVB exposure during peak hours occurs to outdoor workers to some extent during their workweek, so that they can maintain adequate levels of vitamin D3 in their skin and blood (as 25-hydroxyvitamin D) for most of the year. Note that the blood levels of vitamin

Figure 4: (a) The SEER 9 data from 1975 to 2007 indicates a linear increasing incidence in CMM (R2 is 0.9954). Five years of data was averaged to get each data point, for example, the data point for 1975 includes averaged data from 1973 to 1977. If the data is plotted out year by year the trend line and R2 values indicate it is a linear increase since 1973. Even if we add the IARC data for 1960, 1965, and 1970 to the SEER data, the incidence is still increasing in a linear manner from 1960 to 2005. (b) However, if one adds in the data for Connecticut and New York State extending it back 6 decades to 1940, then the increase appears to be exponential (R2 is 0.9755) rather than linear (R2 is now 0.9189).
D (measured as 25-hydroxyvitamin D in serum) in outdoor workers (gardeners), who get about five times the solar dose that indoor workers get, are about twice as high as indoor workers [28]. The reason vitamin D is important for controlling melanoma is because it can be converted to the hormone calcitriol inside melanoma cells [29]. Calcitriol can control the growth [30–32] and apoptotic cell death [33] of responsive melanoma cells, while it also affects the immune system [34, 35] and inhibits tumor promotion [36], which may all be responsible for increasing the survival of melanoma patients who get regular, moderate sun exposures [37]. Thus, intermittent, strong UVB-induced sunburns may initiate CMM, while low concentrations of vitamin D3 [38] in the skin and UVA-induced DNA damage may promote CMM [14].

5. Conclusions

The incidence of CMM is increasing at an alarming rate around the world in fair-skinned, indoor-working populations, and may be increasing at an exponential rate. The CMM incidence decreases with increasing latitude up to ~50°N where it changes and increases with increasing latitude. This inverse may occur because there is more UVA relative to UVB for most of the year at higher latitudes compared to lower latitudes. If windows, allowing UVA to enter our indoor-working environment and cars, are at least partly responsible for the increasing incidence of CMM, then UV filters can be applied to office and car windows to help reduce the rate of increase in the incidence of CMM worldwide.

Abbreviations

CMM: Cutaneous malignant melanoma
IARC: International Agency for Research on Cancer
SEER: Surveillance epidemiology and end results
USA: United States of America
UVA: 321–400 nm
UVB: 290–320 nm.

References

[1] Environmental protection agency, Ultraviolet radiation and Melanoma Vol. IV: Appendix A—Assessing the risks of stratospheric ozone depletion. US EPA 400/1-87/001D. Table 4-1, p. 4-2, December 1987.
[2] G. C. Roush, M. J. Schymura, and T. R. Holford, “Patterns of invasive melanoma in the Connecticut tumor registry. Is the long-term increase real?” Cancer, vol. 61, no. 12, pp. 2586–2595, 1988.
[3] R. Doll, P. Payne, and J. A. H. Waterhouse, Eds., Cancer Incidence in Five Continents, Vol. I, Union Internationale Contre le Cancer, Geneva, Switzerland, 1966.
[4] R. Doll, C. S. Muir, and J. A. H. Waterhouse, Eds., Cancer Incidence in Five Continents, Vol. II, Union Internationale Contre le Cancer, Geneva, Switzerland, 1970.
[5] J. Waterhouse, C. Muir, P. Correa, and J. Powell, Eds., Cancer Incidence in Five Continents, Volume III, IARC Scientific Publications No. 15, IARC, Lyon, France, 1976.
[6] J. Waterhouse, C. S. Muir, K. Shanmugaratnam, and J. Powell, Eds., Cancer Incidence in Five Continents, Vol. IV, IARC Scientific Publications No. 42, IARC, Lyon, France, 1982.
[7] C. S. Muir, J. Waterhouse, T. Mack, J. Powell, and S. L. Whelan, Eds., Cancer Incidence in Five Continents, Vol. V, IARC Scientific Publications No. 88, IARC, Lyon, France, 1987.
[8] D. M. Parkin, C. S. Muir, S. L. Whelan, Y. -T. Gao, J. Ferlay, and J. Powell, Eds., Cancer Incidence in Five Continents, Vol. VI, IARC Scientific Publications No. 120, IARC, Lyon, France, 1992.
[9] D. M. Parkin, S. L. Whelan, J. Ferlay, L. Raymond, and J. Young, Eds., Cancer Incidence in Five Continents, Vol. VII, IARC Scientific Publications No. 143, IARC, Lyon, France, 1997.
[10] D. M. Parkin, S. L. Whelan, J. Ferlay, L. Teppo, and D. B. Thomas, Cancer Incidence in Five Continents, Vol. VIII, IARC Scientific Publications No. 155, IARC, Lyon, France, 2002.
[11] M. P. Curado, B. Edwards, H. R. Shin et al., Eds., Cancer Incidence in Five Continents, Vol. IX, IARC Scientific Publications No. 160, IARC, Lyon, France, 2007.
[12] SEER Database, http://seer.cancer.gov/.
[13] J. M. Elwood, R. P. Gallagher, G. B. Hill, and J. C. G. Pearson, “Cutaneous melanoma in relation to intermittent and constant sun exposure—the Western Canada melanoma study,” British Journal of Cancer, vol. 35, no. 4, pp. 427–433, 1985.
[14] D. E. Godar, R. J. Landry, and A. D. Lucas, “Increased UVA exposures and decreased cutaneous Vitamin D3 levels may be responsible for the increasing incidence of melanoma,” Medical Hypotheses, vol. 72, no. 4, pp. 434–443, 2009.
[15] D. E. Godar, S. P. Wengratis, J. Shreffler, and D. H. Sliney, “UV doses of Americans,” Photochemistry and Photobiology, vol. 73, no. 6, pp. 621–629, 2001.
[16] D. E. Godar, “UV doses worldwide,” Photochemistry and Photobiology, vol. 81, no. 4, pp. 736–749, 2005.
[17] S. Gandini, F. Sera, M. S. Cattaruzza et al., “Meta-analysis of risk factors for cutaneous melanoma: II. Sun exposure,” European Journal of Cancer, vol. 41, no. 1, pp. 45–60, 2005.
[18] V. Beral, S. Evans, and G. Milton, “Malignant melanoma and exposure to fluorescent lighting at work,” The Lancet, vol. 2, no. 8293, pp. 290–293, 1982.
[19] http://en.wikipedia.org/wiki/Sunscreen.
[20] http://en.wikipedia.org/wiki/Tanning_bed#History.
[21] J. A. MacLaughlin, R. R. Anderson, and M. F. Holick, “Spectral character of sunlight modulates photosynthesis of previtamin D3 and its photoisomers in human skin,” Science, vol. 216, no. 4549, pp. 1001–1003, 1982.
[22] A. R. Webb, B. R. deCosta, and M. F. Holick, “Sunlight regulates the cutaneous production of vitamin D3 by causing its photodegradation,” Journal of Clinical Endocrinology and Metabolism, vol. 68, no. 5, pp. 882–887, 1989.
[23] J. G. Peak and M. J. Peak, “Comparison of initial yields of DNA-to-protein crosslinks and single-strand breaks induced in cultured human cells by far- and near-ultraviolet light, blue light and X-rays,” Mutation Research, vol. 246, no. 1, pp. 187–191, 1991.
[24] C. A. Jones, E. Huberman, M. L. Cunningham, and M. J. Peak, “Mutagenesis and cytotoxicity in human epithelial cells by far- and near-ultraviolet radiations: action spectra,” Radiation Research, vol. 110, no. 2, pp. 244–254, 1987.
[25] G. M. Halliday, N. S. Agar, R. S. Barnetson, H. N. Ananthaswamy, and A. M. Jones, “UV-A fingerprint mutations in
human skin cancer,” Photochemistry and Photobiology, vol. 81, no. 1, pp. 3–8, 2005.

[26] M. Moehrle, M. Soballa, and M. Korn, “UV exposure in cars,” Photodermatology Photoimmunology and Photomedicine, vol. 19, no. 4, pp. 175–181, 2003.

[27] A. R. Webb, L. Kline, and M. F. Holick, “Influence of season and latitude on the cutaneous synthesis of vitamin D3: exposure to winter sunlight in Boston and Edmonton will not promote vitamin D3 synthesis in human skin,” Journal of Clinical Endocrinology and Metabolism, vol. 67, no. 2, pp. 373–378, 1988.

[28] M. S. Devgun, C. R. Paterson, B. E. Johnson, and C. Cohen, “Vitamin D nutrition in relation to season and occupation,” American Journal of Clinical Nutrition, vol. 34, no. 8, pp. 1501–1504, 1981.

[29] J. Reichrath, M. Rech, M. Moeini, E. Meese, W. Tilgen, and M. Seifert, “In vitro comparison of the vitamin D endocrine system in 1,25(OH)2D3-responsive and -resistant melanoma cells,” Cancer Biology and Therapy, vol. 6, no. 1, pp. 48–55, 2007.

[30] J. A. Eisman, D. H. Barkla, and P. J. M. Tutton, “Suppression of in vivo growth of human cancer solid tumor xenografts by 1,25-dihydroxyvitamin D3,” Cancer Research, vol. 47, no. 1, pp. 21–25, 1987.

[31] K. Colston, M. J. Colston, and D. Feldman, “1,25-Dihydroxyvitamin D3 and malignant melanoma: the presence of receptors and inhibition of cell growth in culture,” Endocrinology, vol. 108, no. 3, pp. 1083–1086, 1981.

[32] R. J. Frampton, S. A. Omond, and J. A. Eisman, “Inhibition of human cancer cell growth by 1,25-dihydroxyvitamin D3 metabolites,” Cancer Research, vol. 43, no. 9, pp. 4443–4447, 1983.

[33] C. Danielsson, K. Fehsel, P. Polly, and C. Carlberg, “Differential apoptotic response of human melanoma cells to 1α,25-dihydroxyvitamin D3 and its analogues,” Cell Death and Differentiation, vol. 5, no. 11, pp. 946–952, 1998.

[34] S. Yang, C. Smith, and H. F. DeLuca, “1α,25-dihydroxyvitamin D3 and 19-nor-1α,25-dihydroxyvitamin D3 suppress immunoglobulin production and thymic lymphocyte proliferation in vivo,” Biochimica et Biophysica Acta, vol. 1158, no. 3, pp. 279–286, 1993.

[35] S. Yang, C. Smith, J. M. Prahl, X. Luo, and H. F. DeLuca, “Vitamin D deficiency suppresses cell-mediated immunity in vivo,” Archives of Biochemistry and Biophysics, vol. 303, no. 1, pp. 98–106, 1993.

[36] K. Chida, H. Hashiba, M. Fukushima, T. Suda, and T. Kuroki, “Inhibition of tumor promotion in mouse skin by 1α,25-dihydroxyvitamin D3,” Journal of Cancer Research, vol. 45, no. 11, pp. 5426–5430, 1985.

[37] M. Berwick, B. K. Armstrong, L. Ben-Porat et al., “Sun exposure and mortality from melanoma,” Journal of the National Cancer Institute, vol. 97, no. 3, pp. 195–199, 2005.

[38] A. E. Millen, M. A. Tucker, P. Hartge et al., “Diet and melanoma in a case-control study,” Cancer Epidemiology Biomarkers and Prevention, vol. 13, no. 6, pp. 1042–1051, 2004.