Antineutrophil Cytoplasmic Antibody–Associated Vasculitides: Could Geographic Patterns Be Explained by Ambient Ultraviolet Radiation?

PAUL A. GATENBY,1 ROBYN M. LUCAS,1 OLA ENGelsen,2 ANNE-LOUISE PONSONBY,3 AND MARK CLEMENTS1

Objective. This ecological study describes and quantifies the association between ambient ultraviolet (UV) radiation levels, including daily winter vitamin D effective UV radiation levels and the incidence of the 3 antineutrophil cytoplasmic antibody–associated vasculitides (AAVs): Wegener’s granulomatosis (WG), microscopic polyangiitis (MPA), and Churg-Strauss syndrome (CSS). Latitudinal variation in occurrence of the AAVs, especially WG, has been previously reported. For other autoimmune diseases such as multiple sclerosis and type 1 diabetes mellitus, inverse associations with latitude are hypothesized to indicate a causative role for low UV radiation exposure, possibly acting via vitamin D status.

Methods. Published epidemiologic studies provided data on incident cases, total population of study regions, age-specific incidence rates, and study location. From these data and online age-specific population data, we calculated crude incidence rates, the expected number of cases (to control for possible age confounding), and measures of ambient UV radiation. Negative binomial regression models were used to calculate the incidence rate ratio (IRR) for a 1,000 joules/m² increase in ambient UV radiation.

Results. The incidence of WG and CSS increased with increasing latitude and decreasing ambient UV radiation, with a stronger and more consistent effect across different UV radiation measures for WG, e.g., for average daily ambient clear sky erythemal UV radiation (WG: IRR 0.64 [95% confidence interval (95% CI) 0.44–0.94], P = 0.02; CSS: IRR 0.67 [95% CI 0.43–1.05], P = 0.08; MPA: IRR 1.16 [95% CI 0.92–1.47], P = 0.22). There was no apparent latitudinal variation in MPA incidence.

Conclusion. Our findings are consistent with a protective immunomodulatory effect of ambient UV radiation on the onset of WG and CSS. We discuss possible mechanisms, including the effect of vitamin D on the immune system.

INTRODUCTION

The antineutrophil cytoplasmic antibody (ANCA)–associated vasculitides (AAVs) are an important group of inflammatory vascular diseases of small blood vessels, with both overlapping and distinct features united by the association with ANCs (1). Included are Wegener’s granulomatosis (WG), microscopic polyangiitis (MPA), and Churg-Strauss syndrome (CSS) (2). The etiology of the AAVs is largely unknown, although they are widely believed to be autoimmune in origin, triggered by environmental events on an as-yet poorly characterized background of genetic susceptibility (3,4). No single dominant environmental factor has emerged with the exception of particular medications in a minority of cases (5,6). Epidemiologic studies have indicated clues, including silica exposure, infection, seasonal variation in occurrence, and the subject of this study, a latitudinal gradient. The latter is best documented for WG, which in both the Northern and Southern hemispheres appears to be more common at a higher latitude (7,8), although the magnitude of the latitudinal gradient has not been formally quantified. Some studies suggest an inverse relationship between latitude and MPA (9), although there are exceptions to this (7). Latitudinal gradients are seen in the autoimmune dis-
cases multiple sclerosis and type 1 diabetes mellitus, and the chronic inflammatory Crohn’s disease (10–12). All 3 are diseases in which Th1 cells play a predominant role (13,14). This type of T cell also appears very important in the pathogenesis of WG (15–18). More recently, it has become clear that another subset of T cells, Th17 cells, may contribute to the pathogenesis of diseases that were until now considered to be predominantly mediated by Th1 cells (19). This includes multiple sclerosis and granulomatous diseases such as Crohn’s disease and WG (20).

The most plausible hypothesis that has been raised to explain the latitude effect in immune diseases such as multiple sclerosis has been related to the interaction between vitamin D and the immune system. The active form of vitamin D, 1α,25-dihydroxyvitamin D₃, has particular immunomodulatory effects involving inhibition of certain T lymphocytes (21), with the reduction in a number of important cytokines derived from both mononuclear phagocytes and T cells; of particular relevance to this study, one of its effects is to suppress Th1 cells, and more recently, Th17 cell responses (20,22,23).

Sun exposure of the skin is the principal source of vitamin D for humans (24), and unless it is supplemented by diet, humans tend to be vitamin D deficient in higher latitude locations (25). It is important to recognize, however, that ultraviolet (UV) radiation exposure may have effects on the immune system independent of vitamin D, and any mechanisms remain speculative at this stage (26).

The objective of this ecological study was to describe and quantify the association between regional ambient UV radiation levels, including vitamin D effective UV radiation levels, and the incidence of the 3 AAVs: WG, MPA, and CSS. Contrary to previous studies that searched only for latitude dependencies (9,27), we compared worldwide incidence data with estimates of ambient UV radiation levels in addition to latitude, providing more pertinent descriptors of UV radiation exposure.

MATERIALS AND METHODS

Published epidemiologic studies of AAVs and each of the 3 diseases separately were sourced from PubMed using the following search strategy. We assessed studies that included the key words of the specific disease, e.g., “Wegener’s granulomatosis” and “incidence,” that were published before December 2008. Initial articles were used to check the completeness of the search by comparing the references with the articles obtained. Further unpublished work was obtained from the published abstracts of the 13th Vascularitis and Antineutrophil Cytoplasmic Antibody conference held in Cancun, Mexico in 2007.

We included studies that 1) used the diagnostic criteria developed by the American College of Rheumatology (ACR) (28) or the Chapel Hill Consensus Conference (CHCC) (29), 2) indicated a well-defined source population from which the incident cases arose and underlying population data were provided (at least total or adult-only population), 3) used complementary methods to ensure thorough detection of cases, and 4) did not restrict cases to only those with a particular phenotype subgroup of disease (e.g., only cases with renal disease). We restricted our analysis to published series where the population denominator was clear. Therefore, we did not use the study from Kuwait (30), a hospital-based study from Sweden (31), or the report from The Glomerular Disease Collaborative Network (32). We also excluded an important study from New Zealand that used discharge codes in a national data set that was not strictly comparable with the other studies (8). Three other studies did give rise to some concerns: the study from Lithuania (33) was hospital based and may have missed community cases, the diagnostic criteria used by Zeft et al (34) were not stated in their abstract, and the study from Japan (35) was based on referral to a renal unit and may have missed those with nonrenal disease. However, the denominator population in this case was clearly defined (35), and other sources (36) attest to the rarity of WG in this region. We extracted the number of cases of each disease entity, noting the location, year of publication, and the source population, e.g., general, adult-only.

We used online census data (see Supplemental Appendix A, available in the online version of this article at http://www3.interscience.wiley.com/journal/77005015/ for each study region (using the population estimate closest to the time period of the study, and if regional data were not available, applying the age distribution of the smallest available unit to the population figures in the published article) to derive the total population of the study region (by 5-year age groups), ensuring that this was consistent with the population data cited in the published study.

Where studies were reported in separate articles over several time periods, we used the total time period in the person-time denominator to provide the most stable incidence estimate. We then recalculated the crude incidence and 95% confidence intervals (95% CIs) at each location using a consistent method (Poisson distribution). For one study (34) that did not provide a total source population, we back calculated from the total number of cases (n = 36) with AAV and the reported annual incidence rate of 11.4 per million for 12 years to derive a population estimate. We extracted the available age-specific incidence rates for each disease (37–39). We then calculated the mean incidence rate for each age group and multiplied this by the population of the relevant age group for each study region. This provided the number of cases of each disease that would be expected in each age group and in each region if all of the regions had the same age-specific incidence rates. By summing the number of cases across the age groups, we calculated the expected total number of cases in the region, removing any confounding effect of any differences in the age structures of the populations (indirect age standardization). Because only a few studies noted the observed sex distribution of cases with each form of vasculitis, it was not possible to standardize for both age and sex.

Using the latitude and longitude of the largest city in each study and the time period during which the study was performed, we used satellite data to derive the mean daily ambient UV radiation level and the mean daily winter UV radiation level for each study (see Supplemental Appendix A, available in the online version of this article at http://www3.interscience.wiley.com/journal/77005015/
home). We estimated both the ambient UV radiation level weighted to erythemally effective wavelengths (UV<sub>ery</sub>) (40) and to vitamin D effective wavelengths (UV<sub>vitD</sub>) (41) under both clear sky and cloud-inclusive conditions. We used winter UV radiation levels in addition to mean daily ambient UV radiation levels (across the time period of the study) because there is a stronger latitudinal gradient in winter UV radiation compared with summer UV radiation (42). Additionally, it is the lower vitamin D effective UV radiation during the winter that limits vitamin D production from skin irradiation at high latitudes (25).

**Statistical analysis.** We first examined the correlation between the crude incidence rates and each measure of ambient UV radiation level using Pearson’s correlation coefficients (P value test if the correlation was different from zero or if there was a trend). We then used a negative binomial regression model (to account for the differences in precision between studies and the different age distributions of the source populations) (43) with the observed number of cases as the dependent variable, a measure of ambient UV radiation level as the independent variable, and the log of the expected number of cases as the independent variable. This resulted in only very small adjustments to the total number of cases and therefore to the estimated parameters. Sensitivity analyses. For all 3 diseases, the Lima, Peru data were influential (CSS: Cook’s distance 0.60; WG: Cook’s distance 0.22 and P = 0.02, respectively). In the negative binomial regression model, there was a modest increase in the incidence of CSS and WG per higher degree of latitude of 3.4% (IRR 1.04, 95% CI 0.99–1.08; P = 0.11) and 3.5% (IRR 1.04, 95% CI 1.00–1.07; P = 0.03), respectively. MPA incidence again did not appear to be associated with latitude (IRR 0.99, 95% CI 0.97–1.01; P = 0.23).

**Ambient UV radiation.** There was a strong inverse correlation between ambient UV radiation level and WG incidence (UV<sub>ery</sub>[clear]: r = −0.62, P = 0.02; winter UV<sub>vitD</sub>[cloud]: r = −0.60, P = 0.03) and a more modest inverse correlation with CSS incidence (UV<sub>ery</sub>[clear]: r = −0.43, P = 0.21; UV<sub>vitD</sub>[cloud]: r = −0.48, P = 0.16). In contrast, MPA incidence was not correlated with any measure of ambient UV radiation level (UV<sub>ery</sub>[clear]: r = −0.02, P = 0.95; UV<sub>vitD</sub>[cloud]: r = −0.13, P = 0.73). In the negative binomial regression model, the incidence of both WG and CSS was inversely associated with all of the measures of ambient UV radiation level (Figure 1). For an increase in UV<sub>ery</sub>[clear] of 1,000 joules/m<sup>2</sup> (e.g., the difference between Vilnius, Lithuania [1,733.11 joules/m<sup>2</sup>] and Lugo, Spain [2,665.61 joules/m<sup>2</sup>]), the CSS incidence decreased by 33% (IRR 0.67, 95% CI 0.43–1.05; P = 0.08), and decreased by 23% (IRR 0.77, 95% CI 0.62–0.95; P = 0.02) for a 1,000 joules/m<sup>2</sup> increase in winter UV<sub>vitD</sub>[cloud]. WG incidence decreased by 36% (IRR 0.64, 95% CI 0.44–0.94; P = 0.02) for a 1,000 joules/m<sup>2</sup> increase in UV<sub>ery</sub>[clear], and decreased by 25% (IRR 0.75, 95% CI 0.62–0.89; P = 0.001) for a 1,000 joules/m<sup>2</sup> increase in winter UV<sub>vitD</sub>[cloud]. For both CSS and WG, the strongest inverse associations were with measures of winter ambient UV radiation levels. There was no statistically significant variation in MPA incidence (Figure 1) for any measure of ambient UV radiation level when all of the data points were included (UV<sub>ery</sub>[clear]: IRR 1.16, 95% CI 0.92–1.47; P = 0.22 and winter UV<sub>vitD</sub>[cloud]: IRR 1.05, 95% CI 0.94–1.18; P = 0.37).

**Sensitivity analyses.** For all 3 diseases, the Lima, Peru data were influential (CSS: Cook’s distance = 0.87, MPA: Cook’s distance = 0.79, WG: Cook’s distance = 0.60). In addition, there was some evidence that the Miyazaki, Japan and Canberra, Australia data were influential for WG, using measures for change in model fit and change in estimated parameters.

When excluding the Lima, Peru data, although point estimates continued to show decreasing CSS and WG in-
cidence with increasing ambient UV radiation levels, these associations were no longer significant (CSS: IRR 0.87 [95% CI 0.44 – 1.74, \( P = 0.69 \)) and winter UVvitD [cloud] IRR 0.67 [95% CI 0.31 – 1.43, \( P = 0.30 \]). There was little change to the estimates of IRR for MPA, but notably, the point estimates were consistently 1, i.e., increasing incidence with increasing ambient UV radiation levels (UVery [clear]: IRR 1.28, 95% CI 0.78 – 2.10; \( P = 0.33 \) and winter UVvitD [cloud]: IRR 1.29, 95% CI 0.62 – 2.69; \( P = 0.49 \)).

In excluding the Miyazaki, Japan data (WG only), there was no appreciable change in the IRR estimates of decreasing WG incidence with measures of increasing ambient UV radiation levels per 1,000 joules/m² increase (UVery [clear]: IRR 0.71, 95% CI 0.53 – 0.95; \( P = 0.02 \) and winter UVvitD [cloud]: IRR 0.77, 95% CI 0.68 – 0.87; \( P < 0.001 \)).

When excluding the Canberra, Australia data (WG only), there was an increase in the estimates of IRR (i.e., a stronger association between WG incidence and ambient UV radiation), and these estimates were no longer influenced by exclusion of the Lima, Peru data (WG incidence: UVery [clear]).

**Table 1.** Studies reporting the population-based incidence of antineutrophil cytoplasmic antibody–associated vasculitides used in this analysis*

<table>
<thead>
<tr>
<th>Author, year (ref.)</th>
<th>Time period of study</th>
<th>Location (latitude, longitude)</th>
<th>Crude incidence per million population (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sanchez-Torres et al, 2006 (61)</td>
<td>1990–2004 Lima, Peru (12.00°S, 76.58°W)</td>
<td>CSS 0.11 (0.01–0.40) MPA 3.13 (2.36–4.06) WG 0.39 (0.16–0.81)</td>
<td></td>
</tr>
<tr>
<td>Fujimoto et al, 2006 (35)</td>
<td>2000–2004 Miyazaki, Japan (31.90°N, 131.43°E)</td>
<td>CSS 0.00 MPA 0.00</td>
<td></td>
</tr>
<tr>
<td>Ormerod and Cook, 2008 (64)</td>
<td>1995–2004 Canberra, Australia (35.30°S, 149.15°E)</td>
<td>CSS 1.81 (0.78 – 3.57) MPA 3.63 (2.07–5.89) WG 8.61 (6.09–11.80)</td>
<td></td>
</tr>
<tr>
<td>Gonzalez-Gay et al, 2003 (39)</td>
<td>1988–2001 Lugo, Spain (43.00°N, 7.50°W)</td>
<td>CSS 1.20 (0.33 – 3.07) MPA 10.18 (7.05 – 14.2) WG 3.59 (1.86–6.28)</td>
<td></td>
</tr>
<tr>
<td>Reinhold-Keller et al, 2002 (27)</td>
<td>1998–1999 South Germany (48.00°N, 7.85°E)</td>
<td>CSS 0.95 (0.26 – 2.42) MPA 1.66 (0.67–3.41)</td>
<td></td>
</tr>
<tr>
<td>Reinhold-Keller et al, 2005 (65)</td>
<td>1998–2002 North Germany (53.87°N, 10.70°E)</td>
<td>CSS 1.01 (0.55 – 1.69) MPA 2.66 (1.88–3.67)</td>
<td></td>
</tr>
<tr>
<td>Darodiniene et al, 2005 (33)</td>
<td>1990–1999 Vilnius, Lithuania (54.67°N, 25.27°E)</td>
<td>CSS 1.11 (0.41 – 2.41) MPA 2.58 (1.41–4.33)</td>
<td></td>
</tr>
<tr>
<td>Knight et al, 2006 (59)</td>
<td>1975–2001 Sweden (60°N, 15°E)</td>
<td>CSS 2.66 (0.32 – 9.61) MPA 6.65 (2.16–15.5)</td>
<td></td>
</tr>
<tr>
<td>Koldingsnes and Nossent, 2000 (38)</td>
<td>1984–1998 Northern Norway (69.67°N, 19.00°E)</td>
<td>CSS 0.39 (0.05–1.41) MPA 2.15 (1.07–3.85)</td>
<td></td>
</tr>
</tbody>
</table>

* 95% CI = 95% confidence interval; CSS = Churg-Strauss syndrome; MPA = microscopic polyangiitis; WG = Wegener’s granulomatosis.

**Figure 1.** Variation in the crude incidence rate of the antineutrophil cytoplasmic antibody–associated vasculitides with change in ambient erythemal ultraviolet radiation (UVR). The shaded area shows the 95% confidence limits. J = joules.
In our second sensitivity analysis, we recalculated case numbers where, for example, only adult cases were included, using the proportion of childhood cases compared with adult cases where the entire population was the source. Because the AAVs are rare in childhood, this had little effect on the total case numbers. For example, in the study from Lithuania (33), the number of cases with WG increased from 10 to 10.18, increasing the incidence from 1.84 per million to 1.88 per million. Repeating the analyses with the adjusted values did not change the results above.

DISCUSSION

These results confirm a preliminary observation that WG is latitude related (7,9), and quantify the magnitude of the gradient. We found a strong inverse association between measures of erythemal and vitamin D effective ambient UV radiation level and WG incidence, with a decrease in incidence of 38% (95% CI 6–56%) for every 1,000 joules/m² increase of UV_{ora}(clear). There was a similar, but with a lower magnitude, association for CSS, with a decrease in incidence of 23% (95% CI 4–38%) per 1,000 joules/m² increase in winter UV_{ora(cloud)}. In contrast, MPA incidence did not appear to be associated with latitude or ambient UV radiation when all data points were considered, although point estimates were consistently greater than 1, with some strengthening of effect with exclusion of the Lima, Peru data. Absence of a latitudinal gradient in MPA is at odds with previously published work (9). This may reflect our exclusion of the Kuwait data (30) because the precise population from which the subjects were derived is not clear, and the incidence estimates are potentially overestimated. As previously noted, 1,000 joules/m² of UV_{ora}(clear) is approximately the difference in this measure between Vilnius, Lithuania (1,733.11 joules/m²) and Lugo, Spain (2,665.61 joules/m²) during their respective study periods.

There is some uncertainty in the UV radiation estimates related to the evaluation of cloud effects. However, our results were consistent whether clear sky or cloud inclusive, erythemal, or vitamin D effective UV radiation estimates were used. The effects appeared to be stronger only for winter measures of ambient UV radiation. To put our findings into some perspective, our results indicate that the incidence of WG increases by 3.4% per higher degree of latitude and CSS by 3.5% per higher degree of latitude. The single comparable meta-analytic study of multiple sclerosis (46) found a nonsignificant latitudinal increase in age-adjusted incidence. From the data provided in their Figure 2, we estimate that this is of the order of a −4% increase in multiple sclerosis incidence per higher degree of latitude.

Ambient UV radiation was used as a proxy for population-level exposure to UV radiation and vitamin D status. However, at the population level, UV radiation exposure is affected by cultural and religious beliefs, e.g., clothing requirements or preference of pale or tanned skin, whereas within this context, individual-level sun exposure is highly behavior dependent (47). Vitamin D status depends not only on UV radiation exposure of the skin (48,49), but on skin pigmentation (50), diet (51), and genetic factors (52). Furthermore, at higher latitudes, increased dietary vitamin D intake (53) may uncouple a direct association between UV radiation exposure and vitamin D status. We recognize the limitations of the ecological approach, but are not aware of any studies for which individual UV dosimetry data or blood vitamin D levels are yet available for the AAVs. Therefore, we could not unravel these components in this ecological analysis. The findings should be interpreted to reflect not only latitude-related ambient UV radiation levels, but also variation in these other factors, i.e., diet, skin pigmentation, and sun exposure behavior.

We chose to study the association between 2 latitude-related factors, ambient erythemal and vitamin D effective UV radiation levels. Other environmental factors such as infection and nutrition may also vary by latitude, as may genetic predisposition to these diseases. For example, HLA status, a strong influence on antigen-induced immune responses, varies by race (54). However, our analyses consistently show a much stronger effect size for measures of ambient UV radiation level compared with latitude, although we cannot rule out a role for regional factors that are strongly linked to ambient UV radiation level.

UV radiation exposure of the skin leads to both direct and indirect (via vitamin D) immunosuppression (reviewed by Ponsonby et al in 2005 [22]). Imputed vitamin D levels would give results parallel to our conclusions here in regard to UV radiation, but there are insufficient data about population- or subject-specific vitamin D levels to allow anything other than speculation. Nevertheless, there are plausible immune mechanisms involving vitamin D₃ inhibition of Th1 and Th17 cell proliferation and cytokine production, tilting the immune system toward Th2 cells (20) and enhancing the suppressive activity of the CD24+CD25+ so-called Treg cells (55,56). WG is considered a Th1 and/or a Th17 cell–mediated autoimmune disease (57). A role for vitamin D (largely induced by UV radiation) in specifically suppressing Th1/Th17 cells and up-regulating Treg cells provides a biologically plausible explanation for the findings reported here.

The pathogenesis of CSS has not been dissected as effectively as WG. The occurrence of eosinophilia and elevated IgE levels is very suggestive of a Th2 cell response, which is not consistent with the above hypothesis with regard to WG. However, like WG, CSS is characterized by granuloma formation and we postulate that this aspect of the condition is likely to be Th1 cell driven (16).

Most evidence suggests that although Th1/Th17 cells may play a role in MPA, this condition is far more dependent on Th2 cells or antibody-related mechanisms than Th1 cells (1). In MPA, there is substantially stronger evidence for a pathogenetic role of the autoantibody antitymeloxygenase than there is for anti–proteinase 3 in WG and no occurrence of granulomas (16,58). Consequently, the lack of a latitude or UV radiation effect mediated through vitamin D is not unexpected in this particular AAV.
Two major classification systems are used for vasculitis: the ACR and CHCC criteria. Either both of the systems or only the CHCC were used by most of the studies included in these analyses; 4 studies used only the ACR criteria (33,38,59,60). Although MPA is only clearly defined in the CHCC criteria, the incidence of WG in those studies that used only the ACR criteria are consistent with that at other similar (59,60) or lower (33) latitude locations. Indeed, for the latter, the incidence of both WG and MPA is considerably lower than other similar latitude sites. The most likely misclassification when not using the CHCC criteria would be to include cases in the WG group that would now appropriately be designated MPA. This in effect would dilute the WG group at some of the sites and would weaken the association of WG and latitude and ambient UV radiation that we have demonstrated. Indeed, for a combined WG and MPA group, the associations are weaker than for WG alone and no longer statistically significant, although in the same direction. Therefore, for the combined data, the negative binomial regression with latitude demonstrates an increased incidence by 0.78% per degree of latitude (IRR 1.01, 95% CI 0.99–1.11) and the correlation with UV$_{\text{ory}}$(clear) and winter UV$_{\text{vid}}$(cloud) (r = −0.43, P = 0.22 and r = −0.56, P = 0.10, respectively).

To our knowledge, this study is the first to quantify the magnitude of the latitudinal variation in the incidence of these AAVs and to examine the association with levels of both average and winter regional ambient UV radiation. Although there is modest latitudinal variation in average daily ambient UV radiation levels (across the entire year), this variation is increased for winter ambient UV radiation and may explain the stronger associations we observed between WG incidence and winter ambient UV radiation. In addition, we have standardized the data for age to ensure that the observed variation is not confounded by variations in the age structure of populations.

However, in this work, we have not controlled for possible ethnic and genetic differences in the underlying populations. Conceivably, the very low reported incidence of WG in Kuwait (30), Japan (35), and Lima, Peru (61) could be caused by differences in genetic susceptibility of the underlying populations. Nevertheless, a prevalence survey within a relatively genetically homogeneous population, the New Zealand European population, provides supportive evidence for a latitudinal gradient in WG that is not confounded by variation in ethnicity (8). Clearly, this current study provides no clues to explain the dearth of AAVs in individuals of African American descent (32,62); however, a similar pattern is seen in multiple sclerosis, another autoimmune disease with a striking latitudinal pattern of occurrence (63). One focus of future work could be to examine the role of ethnicity in the development of the AAVs.

Here we have used the available literature reporting the population-based incidence of AAVs to explore possible etiologic associations with measures of ambient UV radiation. We observed a strong inverse association between the incidence of WG and measures of ambient UV radiation, with a lesser association for CSS and no association for MPA. The association with UV radiation (by either direct or indirect pathways mediated by enhancement of vitamin D synthesis) is biologically plausible through effects on Th1 and Th17 cell immune function and Treg cells. We regard these findings as important and, as has occurred in multiple sclerosis and type 1 diabetes mellitus, they should lead to the generation of etiologic hypotheses that can be more directly explored in individual-level epidemiologic studies.

ACKNOWLEDGMENTS

We gratefully acknowledge access to Total Ozone Mapping Spectrometer and Ozone Mapping Instrument ozone and UV radiation data from the NASA Goddard Space Flight Center and to Dobson and Brewer ozone data from the World Ozone and Ultraviolet Radiation Data Centre.

AUTHOR CONTRIBUTIONS

All authors were involved in drafting the article or revising it critically for important intellectual content, and all authors approved the final version to be published. Dr. Gatenby had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

Study conception and design. Gatenby, Lucas.

Acquisition of data. Gatenby, Lucas, Engelsen.

Analysis and interpretation of data. Gatenby, Lucas, Engelsen, Ponsonby, Clements.

REFERENCES

11. Staples JA, Ponsonby AL, Lim LL, McMichael AJ. Ecologic analysis of some immune-related disorders, including type 1 diabetes, in Australia: latitude, regional ultraviolet radiation,