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Non-melanoma skin cancers and glucocorticoid therapy

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Summary Non-melanoma skin cancer (NMSC) is an important cause of morbidity and long-term mortality in organ transplant recipients receiving immunosuppressive drugs such as azathioprine and cyclosporin, often combined with adrenocortical steroids (glucocorticoids). At lower doses, glucocorticoids alone are prescribed for other conditions including musculoskeletal, connective tissue and respiratory disorders. Presently, it is unknown whether patients taking glucocorticoids are at an increased risk of skin malignances. In a population-based case-control study in New Hampshire, USA, we compared use of glucocorticoids in 592 basal cell carcinoma (BCC) and 281 squamous cell carcinoma (SCC) cases and in 532 age and gender matched controls; neither cases nor controls had a history of organ transplantation. Participants underwent a structured personal interview regarding history of medication use and skin cancer risk factors. We used unconditional logistic regression analysis to compute odds ratios associated with glucocorticoids (adjusted odds ratio = 2.31; 95% Cl = 1.27, 4.18), and risk of BCC was elevated modestly (adjusted odds ratio = 1.49; 95% Cl = 0.90, 2.47). In contrast, risk of both SCC and BCC were unrelated to use of inhaled steroids. Our data suggest that use of oral glucocorticoids may increase risk of NMSC, and SCC in particular, among patients other than organ transplant recipients. We hypothesize that immunosuppression induced by oral glucocorticoids may allow these cancers to emerge from immunosurveillance. © 2001 Cancer Research Campaign http://www.bjcancer.com

Keywords: non-melanoma skin cancer; squamous cell carcinoma; basal cell carcinoma; glucocorticoids; immunosuppressive therapy; case-control study

Organ transplant recipients taking immunosuppressive drugs are at an estimated 65-fold greater risk of squamous cell carcinoma (SCC) and a 10-fold higher risk of basal cell carcinoma (BCC) (McCann, 1999). Agents used to prevent allograft rejection include cyclosporine or azathioprine, often combined with glucocorticoids. Systemic glucocorticoids alone (e.g., prednisone, prednisolone, dexamethasone and cortisone) are widely used for a variety of medical conditions including connective tissue diseases, musculoskeletal disorders and allergies. However, it is presently unknown whether these patients are at an increased risk of NMSC. Therefore, we examined the potential risks of SCC and BCC associated with glucocorticoid use in non-transplant recipients, as part of a population-based case-control study of NMSC conducted in New Hampshire, USA.

MATERIALS AND METHODS

Study group

To identify cases for our study, we enlisted the collaboration of dermatologists and pathology laboratories throughout New Hampshire and bordering regions (Karagas et al, 1999). We selected a random sample of BCC cases (for efficiency) and all cases of invasive SCC diagnosed from July 1, 1993 through June 30, 1995 among New Hampshire residents, aged 25–74 years. The sample of BCC cases was drawn concomitantly with the SCC

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cases and controls, and stratified on anatomic site, age and gender to ensure representation of the entire group of BCC diagnoses. As of March 1996, we identified 1143 potential participants. One patient was not contacted at the physician's request, 31 (3%) were reported as deceased by a household member or physician, 10 (1%) lived in households in which no one answered after 40 attempts to telephone distributed over days, evenings and weekends, 178 (16%) declined participation and 27 (2%) were deemed mentally incompetent or too ill to take part. We interviewed a total of 603 BCC and 293 SCC cases.

We chose controls from New Hampshire residents aged 25-74 years who were frequency-matched on age (25-34, 35-44, 45-54, 55-64, 65-69, 70-74 years) and gender to represent the combined distribution of the SCC and BCC cases. We selected controls (roughly equal in number to the number of BCC cases) from lists of New Hampshire residents provided by the New Hampshire Department of Transportation (for those less than 65 years old) and Health Care Financing Administration's Medicare Program (for those 65 years and older). For interviewing purposes, controls were randomly assigned reference dates comparable to the cases' diagnosis dates. Of the 820 potential controls, 12 (2%) were reported as deceased by a member of the household, for 12 (2%), no one answered in the household after 40 attempts to telephone distributed over days, evenings and weekends, 228 (28%) declined participation and 28 (3%) were deemed mentally incompetent or too ill to take part. We thus interviewed 540 controls for the study.

Personal interview

All participants provided informed consent in accordance with the Committee for the Protection of Human Subjects at Dartmouth

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College. Study participants completed a structured personal interview, usually at their homes. Questions included sociodemographic information (level of education), skin sensitivity to the sun after first exposure in the summer (i.e., tendency to sunburn), previous radiation treatment, use of tobacco, and time spent outdoors during working and non-working hours in the summer and the rest of the year. We asked participants if their doctor had ever prescribed corticosteroids or steroids as pills, injections or inhalers for 1 month or longer. We then asked the age they were first treated, condition for which the corticosteroids were prescribed, the name of the drug, dose, and the duration of treatment. To aid recall, we developed a list of glucocorticoid drugs (trade name, generic name and description) and a pictorial guide showing the most commonly used drugs grouped by pill colour, size and shape. To minimize potential reporting bias, we did not reveal the specific hypotheses of interest to either the interviewer or participant, and we did not inform the interviewers of the casecontrol status of participants.

Statistical analysis

We computed the odds ratio (OR) and 95% confidence interval (CI) of BCC and SCC associated with use of glucocorticoids prior to the reference date using unconditional logistic regression, taking into account multiple confounding factors (Breslow and Day, 1980). In addition to age and gender, we assessed the potentially confounding effects of skin reaction to the sun (severe sunburn with blistering, painful sunburn, mild sunburn with some tanning and tanning with no sunburn), radiation treatment (no, yes), cigarette smoking history (never, former, current), and level of education (less than college, college, graduate/professional school). All relative risk estimates of SCC risk were adjusted for age, gender and sun sensitivity, and estimates of BCC for age and gender. No other factors appreciably influenced the results.

RESULTS

A total of 592 BCC, 281 SCC and 532 controls had available information on previous glucocorticoid use and no known history of an organ transplantation (4 BCC, 8 SCC and no controls had an organ transplant). Of the remaining subjects, SCC cases were older on average than BCC cases or controls (Table 1). A higher proportion of BCC and SCC cases than controls reported a tendency to sunburn, a history of radiation treatment, and graduate or professional school education (Table 1). The overall average number of hours spent outdoors during the summer months either recreationally or occupationally was roughly similar for SCC and BCC cases and controls (Table 1).

Overall, 13% of SCC cases, 10% of BCC cases and 8% of controls reported using inhaled or oral glucocorticoids for one month or longer (Table 1). The distribution of medical indications for glucocorticoid therapy did not differ notably between cases and controls (Table 1). Women reported using glucocorticoids more often than men (12.7% of women versus 8.9% of men); however, age, sun sensitivity, time spent outdoors, and level of educational attainment varied little according to glucocorticoid use (data not shown). Prednisone accounted for more than 80% of reported use of oral glucocorticoids.

Use of oral, but not inhaled glucocorticoids, was associated with an increased risk of both SCC and BCC. For SCC, risk was elevated more than two-fold (adjusted odds ratio = 2.31; 95% CI = 1.27, 4.18; P = 0.01) and was stronger for current than prior use (Table 2). We did not detect a trend in SCC risk with duration of use, but relatively few subjects indicated using glucocorticoids for more than 6 months (Table 2). Risk of BCC risk was modestly, and not clearly, elevated (adjusted odds ratio = 1.49; 95% CI = 0.90, 2.47; P = 0.12). Cases and controls reported a similar prevalence of inhaled glucocorticoid use (adjusted odds ratio for BCC = 0.76, 95% CI = 0.39, 1.47; P = 0.49; adjusted odds ratio for SCC = 1.44, 95% CI = 0.68, 3.05; P = 0.36).

DISCUSSION

Long-term immunosuppressive therapy to prevent allograft rejection markedly increases the occurrence of NMSCs. In organ transplant recipients, reported relative risks of NMSC range from 7 to as high as 250 (Blohme and Larko, 1984; Dyall-Smith and Ross, 1995: Gupta et al. 1986: Hartevelt et al. 1990: Hoxtell et al. 1977: Jensen et al, 1999; Kinlen, 1996; O'Connell et al, 1986; Sheiner et al, 2000). Treatment to prevent allograft rejection usually includes a multi-agent regimen of a cytotoxic drug (e.g., azathioprine), an agent affecting signal transduction in T-lymphocytes (e.g., cyclosporin), often combined with a glucocorticoid (e.g., prednisone), which suppresses T-cell proliferation and immune response. Other groups of patients use glucocorticoids alone, but at lower doses and generally for shorter durations. In our study, these individuals had an increased risk of SCC, and possibly of BCC, albeit to a lesser extent than that experienced by organ transplant recipients.

There are a few reports of NMSC among patients using immunosuppressive drugs for conditions other than organ transplantation. Kinlen and colleagues observed three cases of SCC versus 0.60 expected in a series of 1634 patients without transplants who were treated with azathioprine, cyclophosphamide, or chlorambucil (Kinlen, 1996). In a cohort of rheumatoid arthritis patients, skin cancer occurred in 19 out of 119 of the cyclophosphamide-treated patients (3 SCC cases) compared to 6 out of 119 patients not receiving cyclophosphamide (one SCC case) (Radis et al, 1995). In our study, we did not specifically ask subjects whether they took cyclophosphamide or other immunosuppressive agents. Nonetheless, our results for patients who reported taking glucocorticoids (and potentially combined with other agents), are consistent with prior studies on cyclophosphamide and other agents used in non-transplant recipients.

In contrast to the enhanced NMSC risk associated with oral glucocorticoids, we found no association between inhaled glucocorticoids and risk of either SCC or BCC. This result is not surprising, since inhaled glucocorticoids have limited systemic effects (Hardman et al, 1996). The fact that cases and controls were about equally likely to report using inhaled glucocorticoids suggests that NMSC cases do not report more use of drugs in general, or that increased surveillance for skin cancer among patients with chronic diseases can explain the observed association between oral glucocorticoids and NMSC. We did not note any particular condition for which NMSC patients reported taking glucocorticoids more than controls; however, the number of cases in each category was small.

The excess incidence of NMSCs seen in organ transplant recipients is likely due to the immunodeficient state itself. In a cohort study of 1098 renal transplant recipients, the excess NMSC risk occurred irrespective of the type of immunosuppressive therapy used (Bouwes Bavinck et al, 1996). Further, in a report of 2562 Table 1 Selected characteristics of basal cell carcinoma and squamous cell carcinoma cases and controls

Variable*	Controls <i>n</i> = 532 (%)	BCC cases <i>n</i> = 592 (%)	SCC cases <i>n</i> = 281 (%)
Gender			
Male	319 (60.0)	336 (56.8)	177 (63.0)
Female	213 (40.0)	256 (43.2)	104 (37.0)
Age (years)			
<60	202 (38.0)	268 (45.3)	64 (22.8)
60–64	70 (13.2)	86 (14.5)	42 (15.0)
65–69	137 (25.8)	118 (19.9)	71 (25.3)
70–74	123 (23.1)	120 (20.3)	104 (37.0)
Skin reaction to sun exposure for 1 hour the first time in summer			
Severe sunburn with blistering	35 (6.6)	26 (4.4)	30 (10.7)
Painful sunburn followed by peeling	135 (25.5)	212 (35.9)	99 (35.4)
Mildly burnt with some tanning	264 (49.9)	307 (52.0)	126 (45.0)
Tan without sunburn	95 (18.0)	46 (7.8)	25 (8.9)
Mean (SD) lifetime hours/week outdoors in summer			
Recreational	14.9 (6.8)	14.8 (6.6)	14.4 (6.5)
Occupational	10.2 (8.6)	9.3 (8.9)	10.5 (9.8)
Radiation treatment			
No	492 (92 5)	514 (86 8)	245 (87 5)
Yes	40 (7.5)	78 (13.2)	35 (12.5)
Smoking status			
Never	177 (33 3)	254 (42.9)	80 (31 7)
Former	251 (47 3)	254 (42.3)	142 (50 5)
Current	103 (19 4)	87 (14 7)	50 (17.8)
		0. ()	00 (110)
High school or technical school	250 (40 4)	220 (28 7)	100 (40 4)
	200 (40.1)	229 (30.7)	80 (21 7)
Graduate or professional school	103 (31.0)	213 (30.3)	70 (24 9)
	107 (20.1)	140 (20.0)	10 (24.3)
Any steroid use (innaled or oral)	404 (02.2)	524 (00.2)	245 (87.2)
None	491 (92.3)	534 (90.2)	245 (87.2)
Inhaled only	21 (4.0)	40 (0.8)	23 (0.2)
Oral and Inhaled	5 (0.9)	3 (0.5)	3 (1 1)
	5 (0.5)	3 (0.0)	3(11)
Reason for oral steroid use (users only) [†]	0 (00 0)	0 (40 0)	0 (22.2)
Respiratory conditions and astrima	9 (29.0)	8 (18.2)	9 (33.3)
Musculoskeletal and connective tissue disease	6 (19.4)	12 (27.3)	8 (29.6)
Alloray	I (J.Z) 7 (22.6)	4 (9.1) 6 (12.6)	2 (1.4) 1 (2.7)
Alleryy Gastrointestinal disease	1 (22.0) 1 (3.2)	5 (11 4)	(3.7)
Other	7 (22 6)	9 (20 5)	2 (1.4) 5 (18 5)
Villo	1 (22.0)	3 (20.3)	5 (10.5)

*Five individuals had missing data on skin reaction to sun exposure (1 BCC, 1 SCC, and 3 controls); one SCC case had missing data on radiation treatment; one control had missing data on smoking status; two individuals (1 BCC and 1 SCC) had missing data on indication for steroid use. [†]Four cases (2 BCC and 2 SCC) were treated for conditions in more than one category.

heart and kidney recipients, the magnitude of the NMSC risk related to the degree of drug-induced immunosuppression (Jensen et al, 1999). In animal experiments, immunosuppression produced by exposure to ultraviolet light plays a mechanistic role in skin carcinogenesis (Kripke, 1994).

Physicians prescribe glucocorticoids for patients with a variety of medical conditions because of their immunosuppressive and anti-inflammatory effects (Hardman et al, 1996), and in our study, about 8% of population controls reported using glucocorticoids for one month or longer. NMSC is a frequent and growing problem in Caucasian populations (Coebergh et al, 1991; Gallagher et al, 1990; Kaldor et al, 1993; Karagas et al, 1999; Levi et al, 1995; Staples et al, 1998). Our data along with others require further confirmation and indicate that glucocorticoids may contribute to NMSC occurrence.

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Table 2 Odds ratios (95% confidence intervals) for oral glucocorticoid use and basal cell carcinoma and squamous cell carcinoma*

Variable	Controls Number (%)	Basal cell carcinoma		Squamous cell carcinoma	
		Number (%)	Adjusted odds ratio (95% CI) [†]	Number (%)	Adjusted odds ratio (95% CI) [‡]
Any oral use					
No steroid use	491 (95.0)	534 (92.6)	1.00 -	245 (90.4)	1.00 -
Yes	26 (5.0)	43 (7.5)	1.49 (0.90, 2.47)	26 (9.6)	2.31 (1.27, 4.18)
Current or past use of oral glucocorticoid use§					
No steroid use	491 (95.2)	534 (92.7)	1.00 -	245 (90.4)	1.00 -
Current use	10 (1.9)	18 (3.1)	1.66 (0.76, 3.64)	15 (5.5)	3.67 (1.57, 8.63)
Past use	15 (2.9)	24 (4.2)	1.42 (0.73, 2.74)	11 (4.1)	1.59 (0.69, 3.68)
Total duration of oral glucocorticoid use§					
No steroid use	491 (95.2)	534 (92.7)	1.00 -	245 (90.4)	1.00 -
≤ 6 months	17 (3.3)	24 (4.2)	1.13 (0.53, 2.43)	17 (6.3)	2.45 (1.05, 5.72)
7 months–3 years	5 (1.0)	11 (1.9)	1.70 (0.71, 4.06)	4 (1.5)	2.26 (0.78, 6.56)
> 3 years	3 (0.6)	7 (1.2)	2.15 (0.74, 6.25)	5 (1.9)	2.52 (0.73, 8.69)

*Excludes individuals who used only inhaled steroids. †Adjusted for age and sex. ‡Adjusted for age and sex and skin reaction to sun exposure. SOne control and one BCC have missing data.

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