

Risk of Cancer in 767 Patients with Giant Cell Arteritis in Western Norway: A Retrospective Cohort with Matched Controls

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ABSTRACT. Objective. To determine the risk of cancer in a large Norwegian cohort of patients with giant cell arteritis (GCA).

Methods. This is a hospital-based, retrospective, observational cohort study including patients diagnosed with GCA in the Bergen Health Area during 1972–2012. Patients were identified through computerized hospital records using the International Classification of Diseases coding system. Medical records were reviewed. Each patient was randomly assigned population controls matched on age, sex, and geography from the Central Population Registry of Norway. Data on the occurrence of cancer were obtained from the Cancer Registry of Norway. The cumulative risk of malignancy was estimated using Kaplan-Meier methods and potential differences were analyzed using the Gehan-Breslow and log-rank tests.

Results. We identified 881 cases with a clinical diagnosis of GCA, of which 792 fulfilled the American College of Rheumatology (ACR) 1990 classification criteria and 528 were biopsy-verified. Cases with no registered cancer prior to GCA diagnosis were included in a time-to-event analysis, with first cancer as the event ($n = 767$ with clinical GCA diagnosis, 686 fulfilling ACR criteria for GCA, 463 biopsy-verified). These cases were matched with previously cancer-free population controls ($n = 1437$, 1284, 895, respectively). We found no significant difference in the risk of malignancy after time of diagnosis/matching for GCA patients compared to population controls ($p > 0.05$).

Conclusion. In this study of a large and well-characterized cohort of patients with GCA, there was no difference in the risk of malignancy in patients with GCA compared to matched population controls. (First Release January 15 2020; J Rheumatol 2020;47:722–9; doi:10.3899/jrheum.190147)

Key Indexing Terms:

VASCULITIS

GIANT CELL ARTERITIS

TEMPORAL ARTERITIS

MALIGNANCY

CANCER

EPIDEMIOLOGY

Giant cell arteritis (GCA) is the most common systemic vasculitis in adults. The immunopathogenesis of GCA, with

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This work was supported by unrestricted research grants from the Norwegian Association of Heart and Lung Patients; The Norwegian Rheumatism Association; Marit Hansens Memorial Fund; Merck Sharp & Dohme; Odd Fellow Medical Research Fund; and The Raagholt Foundation.

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Accepted for publication June 25, 2019.

its complexity and still many unknown factors, has been outlined in a review by Weyand, *et al* and more recently by Hid Cadena, *et al*^{1,2}. These reviews describe several lineages of dysfunctional immune cells as well as age-related remodeling of the immune system in GCA. The balance between the expression of coinhibitory and costimulatory immune checkpoint molecules appears to be crucial in fine-tuning the immune response and preventing autoimmunity. However, activation of specific inhibitory signals also allows cancer cells to avoid recognition and destruction^{3,4}. Immune checkpoint inhibitors, used in cancer treatment, have been followed by a range of immune-related adverse events, including polymyalgia rheumatica (PMR) and large-vessel vasculitis (LVV)^{5,6,7,8,9}. The increasing use of these drugs, which generate antitumor activity but also enhance autoimmunity, draws new attention to the “old” question of whether there is an association between GCA and cancer. Several investigators have addressed this question, but published reports have shown conflicting results^{10–18}. A metaanalysis from 2014 reported a low but statistically significant increased malignancy risk for patients with GCA¹⁹. However, most

studies on the risk of cancer in GCA have been limited by small sample sizes, possible misclassification bias, and/or short periods of followup. We report a large cohort study in which GCA diagnoses have been thoroughly verified and cancer diagnoses were obtained from a national registry with mandatory reporting, providing virtually no loss to followup. This study may contribute to clarifying the association between GCA and the risk of malignancy.

MATERIALS AND METHODS

This is a retrospective, observational cohort study including patients diagnosed with GCA in the Bergen Health Area during 1972–2012. Our material represents a predominantly white referral cohort from mixed rural and urban areas. The study setting was the Bergen Health Area, consisting of 3 somatic hospitals: Haukeland University Hospital, Haralds plass Deaconess Hospital, and Voss Hospital. Together these hospitals provide specialist healthcare services to the inhabitants of 22 municipalities in Hordaland County in western Norway. By population, Hordaland is the third largest county in Norway, and home to 10% of Norway's total population (Statistics Norway 2016). In this region there is only 1 laboratory for pathology, and no private hospitals that care for rheumatology patients. The majority (>90%) of rheumatologists and internal medicine specialists in the area are hospital-based, with only 3 private rheumatologists in the region, all publicly funded and collaborating closely with the hospital departments. Patients were identified through hospital records using the International Classification of Diseases (ICD) coding system: ICD-8 (446.4) for 1972–1987, ICD-9 (446.5) for 1987–1998, and ICD-10 (M31.5-6) for 1999–2012. Electronic medical records of hospital diagnoses were available from 1972. Medical records that were completely electronic were available from 2001. For patients diagnosed in 1972 through 2001, we also obtained access to archived paper-based medical records to complete the extraction of clinical data. We note that cancer data were obtained separately, because this information was provided by the Cancer Registry of Norway (CRN), in which all new cases of cancer in Norway have been registered since 1952. We collected other data by reviewing medical records of every patient registered with the diagnosis of GCA following an outpatient visit or admission to any ward in one of the 3 study hospitals between January 1, 1972, and December 31, 2012 (41-yr period). We excluded patients if their GCA diagnosis originated prior to the beginning of our study, if data were unavailable, or if the review of records from the time of diagnosis concluded that GCA was an implausible diagnosis. We subclassified patients according to the American College of Rheumatology (ACR) 1990 criteria for GCA and according to temporal artery biopsy (TAB) result. Further details about the selection process including characteristics of the excluded cases have been published previously²⁰. Every Norwegian is given a unique 11-digit identification number at birth or time of immigration. Each patient was initially matched for age (date of birth \pm 1 month), sex, and county of residence to 3 control subjects randomly selected from the Central Population Registry of Norway. The controls were required to be alive at the time of GCA diagnosis for their matched case and this date was defined as the start of the observation period for the control. The observation period ended with the diagnosis of cancer, death, or end of study (December 31, 2012), whichever came first. We excluded duplicate control subjects, control subjects who were also among the cases, and all subjects with precancerous lesions or unverified malignancy (e.g., tumor with uncertain topography and/or unverified histopathology). For persons with \geq 2 registered cancers, we included only the first cancer. Only individuals with no registered cancer before the time of GCA diagnosis or corresponding date among matched controls (i.e., outcome-/event-free on entry to the cohort) were included in the time-to-event analysis. The original target of 3:1 matching was designed with the intent to analyze also cause-specific mortality, including rare causes of death. However, the high prevalence of cancer in the community created an imbalance in this matching, and to avoid unnecessary exclusion of previ-

ously cancer-free cases, we dropped from 3:1 to 2:1 matching prior to the cancer-specific analyses. This was performed by random selection/inclusion of 2 of the original 3 controls when none of the controls had previous cancer. When only 1 of the 3 original controls had previous cancer, this control was excluded but the 2 cancer-free controls were kept in the analysis along with the cancer-free matched case. If 2 or all 3 original controls had previous cancer, we excluded all controls and the corresponding case from the time-to-event analysis.

Extensive demographic and clinical data were collected for the cases but for the population controls, we had no available information on potential risk factors, comorbid conditions, or other clinical data. Information on the occurrence of cancer was obtained from the CRN. The CRN registration is based on mandatory reporting by physicians involved in the investigation, treatment, or followup of cancer patients in Norway. CRN also receives information about all cancer deaths registered by the Norwegian Cause of Death Registry. This ensures near-complete ascertainment of incident cancers in the Norwegian population and the CRN is among the oldest and most complete cancer registries in the world²¹. Every cancer diagnosis is required to be reported, except basal cell carcinoma (BCC) in adults. Precancerous lesions, benign tumors, and tumors with uncertain/unknown potential for malignancy are also reported. When the study was performed, the registration of cancers in CRN was complete for the entire study period. Variables received from the CRN include date and ICD code of the cancer diagnosis, and the diagnostic basis/accuracy of the malignancy. Cancer diagnoses were grouped according to the European Shortlist for Causes of Death, 2012 version (COD-SL-2012). The interpretation and reporting of these data are the sole responsibility of the authors, and no endorsement by the CRN is intended nor should be inferred. The study was approved by the REK sør-øst B regional ethics committee (REC), which approved the study for all the hospitals involved (study reference number 2012/643/REK sør-øst B). REC granted permission to access records without obtaining consent from patients or their relatives owing to the long duration of the study and late onset of the disease. REC also granted permission to obtain a control cohort (3:1) matched on age, sex, and geography from the Central Population Registry of Norway.

Statistical analysis. Descriptive statistics were used to characterize the sample. The cumulative risk of cancer in cases and controls was estimated using Kaplan-Meier plots with the diagnosis of cancer as the event. P values comparing the Kaplan-Meier curves were calculated using the Gehan-Breslow and log-rank tests. Followup time was estimated using the reverse Kaplan-Meier method. HR were estimated using Cox regression with adjustment for time of diagnosis/matching (both linearly and categorized by decades). The significance level was set to 0.05. The computing was done using the Statistical Package for the Social Sciences (SPSS) software version 24 (IBM Corp.) and R software version 3.4 (R Foundation for Statistical Computing).

RESULTS

We identified 881 patients with a clinical diagnosis of GCA, of whom 792 fulfilled the ACR 1990 classification criteria for GCA and 528 had biopsy-verified GCA based on TAB. Incidence numbers for GCA in the cohort were comparable to previous reports, with a mean annual cumulative incidence of 18.4 per 100,000 persons aged 50 years or more for clinically diagnosed GCA, 16.7 for cases fulfilling the ACR 1990 criteria, and 11.2 for biopsy-verified cases alone²⁰. After excluding persons with registered cancer prior to GCA diagnosis, persons with registered cancer with uncertain diagnostic accuracy, controls that were also among the cases, and duplicate control subjects, we included the following numbers of cases and controls: 767 cases with a clinical diagnosis of GCA, of which 686 cases fulfilled the ACR 1990

criteria for GCA and 463 were biopsy-verified. These were matched (2:1) with 1437/1284/895 previously cancer-free population controls, respectively. Core characteristics of the included cases and controls are presented in Table 1.

At the end of study (December 31, 2012), a total of 120/107/69 patients with GCA and 227/204/141 population controls had been registered with a verified first malignancy in CRN after the time of GCA diagnosis/matching (Table 1). We found no significant difference in the risk of malignancy after time of diagnosis/matching for any subgroup of patients with GCA compared to population controls (Figure 1, Table 2). Followup times ranged from 0 to 35 years, with a median followup time of 3 years for cases and 5 years for controls. We did not compare the risk of cancer prior to GCA diagnosis/matching because of missing data on cancers diagnosed before 1972. Key features of our study compared to previous reports evaluating the risk of cancer in GCA cohorts from various populations are presented in Table 3.

The overall distribution of various cancer types, grouped by codes in COD-SL-2012, showed no difference between cases and controls ($p = 0.768$; Figure 2). The numbers for each cancer diagnosis are small and our study lacks sufficient power to analyze the risk of different cancer diagnoses separately. However, the combined lymphoid and hematopoietic malignancies (COD-SL-2012 codes 2.1.19–2.1.21) accounted for almost one-quarter of all registered cancers in both cases and controls in our study, afflicting 3.8% of all GCA cases and 3.5% of their corresponding controls. Malignant neoplasms of the colon, rectum, and anus (COD-SL-2012 code 2.1.4) accounted for another quarter of the registered cancers in cases and about 20% of the registered cancers in controls. Cancers of the breast (COD-SL-2012 code 2.1.10) followed, accounting for 10.8% of the registered malignancies in the patients with GCA and 12.8% in the controls.

DISCUSSION

This 41-year study of 881 Norwegian patients with GCA showed no difference in the risk of cancer compared to age-, sex-, and geographically matched population controls. This fits

well with epidemiological data showing little or no difference in overall mortality rates of patients with GCA compared to the general population^{22,23,24}. However, our result differs from that of some previous reports, in which an increased risk of cancer was seen in patients with GCA (Table 3).

Ungprasert, *et al*, in a metaanalysis from 2014, concluded with a low but statistically significant increased cancer risk, reporting a pooled risk ratio (RR) of 1.14 overall, 2.16 in the first 6–12 months after GCA diagnosis, and 1.35 for biopsy-proven GCA¹⁹. However, excluding a study with potential selection bias reduced the pooled RR to a nonsignificant level of 1.08. The excluded study was a large retrospective cohort study from Sweden reporting on 35,918 patients registered with a diagnostic code of GCA or PMR in a nationwide hospital discharge database¹⁶. The inclusion of only hospitalized patients may have led to a selection bias, favoring patients with severe disease who seem to have a higher cancer risk. A study by Michet, *et al* from the Mayo Clinic examined hospitalizations in the years 1996–2012 for patients with known GCA to evaluate whether hospitalization-related diagnoses accurately identified patients with this disease²⁵. They found that the GCA diagnosis was mentioned in only 31% of 502 hospitalizations among patients with GCA. This illustrates the potential failure of identification that may bias any study based on hospital-administrative diagnostic coding, ours included.

The highest risk of cancer in the Swedish study was noted during the first year after hospitalization for GCA or PMR¹⁶. This study was based solely on ICD coding without verification of the diagnosis through journal review. Consequently, incident cancers may have been misdiagnosed as GCA or PMR initially and thus contributed to the excess rate of cancer reported in the study. Symptoms similar to PMR and the sometimes-nonspecific presentation of GCA may mimic the presenting features of a variety of cancer diagnoses^{26–32}. In the selection process for our study, 35% of the initially selected patients coded as GCA were excluded to ensure a cohort of correctly diagnosed GCA²⁰. We verified GCA diagnoses by reviewing patient records from the time of diagnosis and we excluded patients if the reviewing rheuma-

Table 1. Core characteristics of cases and controls.

Characteristics	Clinical Diagnosis		ACR 1990 Criteria		Biopsy-verified	
	Cases, n = 767	Controls, n = 1437	Cases, n = 686	Controls, n = 1284	Cases, n = 463	Controls, n = 895
Mean age at time of GCA diagnosis/ matching, yrs (SD)	72.5 (8.5)	72.4 (8.5)	72.5 (8.4)	72.4 (8.4)	73.1 (7.8)	73.1 (7.8)
Female sex, n (%)	551 (71.8)	1030 (71.7)	497 (72.4)	928 (72.3)	339 (73.2)	656 (73.3)
Median observation time*, yrs (95% CI)	3 (2–3)	5 (4–5)	3 (2–3)	4 (3–4)	3 (1–3)	5 (4–5)
Events (first cancer), n (%)	120 (15.6)	227 (15.8)	107 (15.6)	204 (15.9)	69 (14.9)	141 (15.8)
Deaths during observation, n (%)	352 (45.9)	680 (47.3)	309 (45.0)	589 (45.9)	216 (46.7)	432 (48.3)

* Observation time according to reverse Kaplan-Meier. ACR: American College of Rheumatology; GCA: giant cell arteritis.

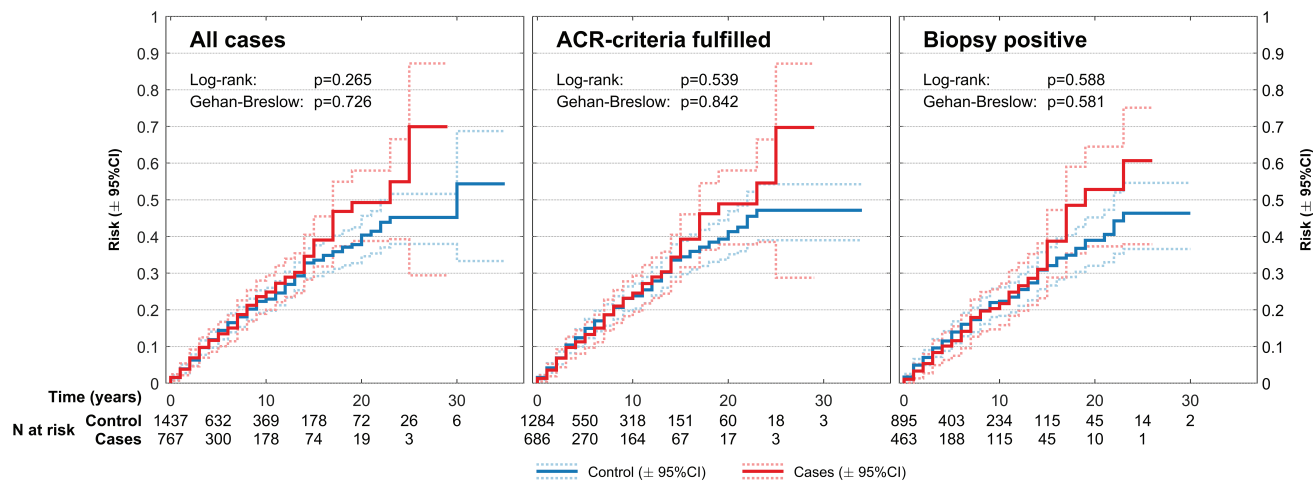


Figure 1. Cumulative risk of first malignancy after time of GCA diagnosis or matching. GCA: giant cell arteritis; ACR: American College of Rheumatology.

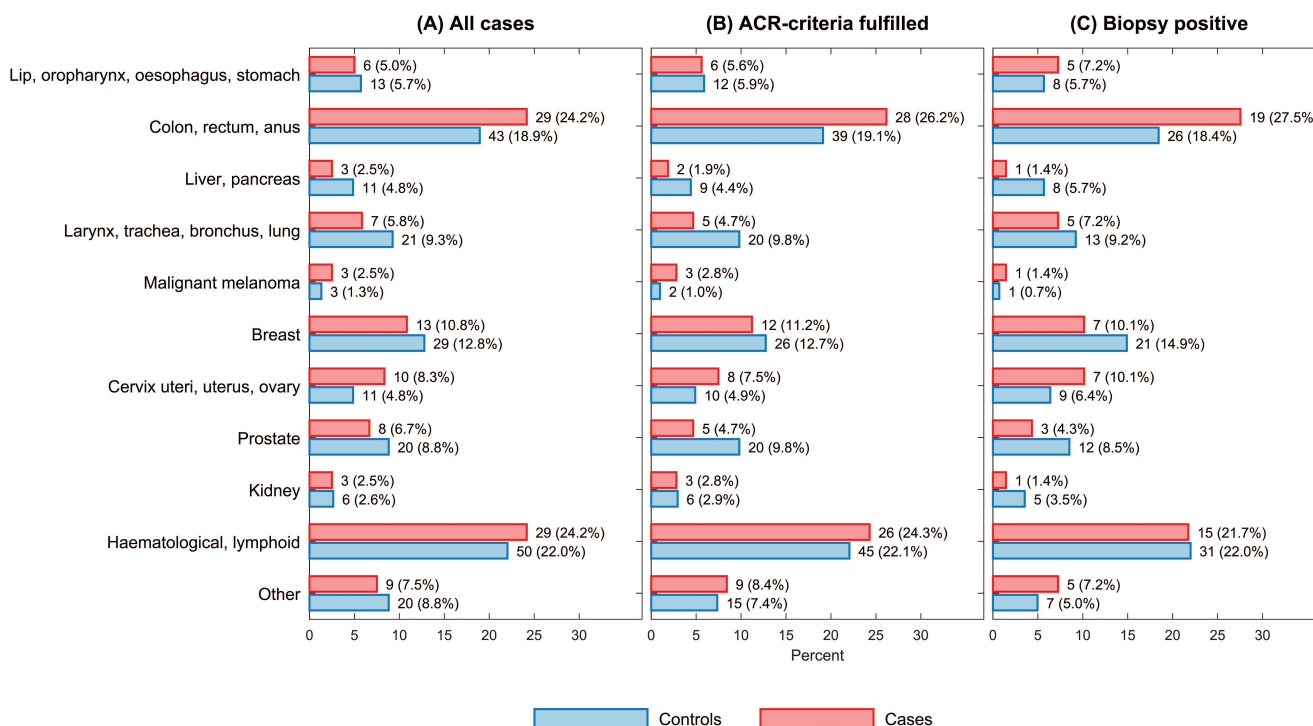


Figure 2. The distribution of different cancers in patients with GCA and matched controls in the Bergen Health Area (1972–2012). All values represent the number (%) of persons with registered cancer. A. Incident cancers for 767 patients with clinically diagnosed GCA and 1437 matched controls. B. Incident cancers for 686 patients with GCA diagnosis fulfilling the ACR 1990 classification criteria for GCA and their 1284 matched controls. C. Incident cancers for 463 biopsy-proven patients with GCA and 895 matched controls. Cancers are grouped according to COD-SL-2012: lip, oropharynx, esophagus, stomach (codes 2.1.1–3); colon, rectum, anus (code 2.1.4); liver, pancreas (codes 2.1.5–6); larynx, trachea, bronchus, lung (codes 2.1.7–8); malignant melanoma of skin (code 2.1.9); breast (code 2.1.10); cervix uteri, uterus, ovary (codes 2.1.11–13); prostate (code 2.1.14); kidney (code 2.1.15); hematological, lymphoid (codes 2.1.19–21); and other (codes 2.1.16–18, 2.1.22). ACR: American College of Rheumatology; GCA: giant cell arteritis; COD-SL-2012: European Shortlist for Causes of Death (2012 version).

tologist concluded that GCA was an implausible diagnosis based on clinical information available at that time. Characteristics of the excluded cases have been published previously²⁰. A similar proportion of erroneous diagnostic

coding in the Swedish hospital discharge study would represent a significant limitation of their results.

The excellent completeness of the CRN ensures that nearly all incident cancers are registered²¹. A study conducted

Table 2. HR for the risk of cancer in patients with GCA versus population controls.

	HR	Lower CI	Upper CI
All Cases, n = 2204			
Unadjusted	1.13	0.91	1.42
Adjusted for time of diagnosis/matching	1.16	0.93	1.45
Adjusted for time period (decades*)	1.16	0.93	1.45
ACR 1990 criteria fulfilled, n = 1970			
Unadjusted	1.12	0.85	1.36
Adjusted for time of diagnosis/matching	1.12	0.87	1.39
Adjusted for time period (decades*)	1.12	0.87	1.39
Biopsy-positive, n = 1358			
Unadjusted	1.09	0.81	1.45
Adjusted for time of diagnosis/matching	1.12	0.84	1.49
Adjusted for time period (decades*)	1.11	0.83	1.49

* Decades defined as the time periods 1972–1982, 1983–1992, 1993–2002, and 2003–2012. ACR: American College of Rheumatology; GCA: giant cell arteritis.

in a population where registration of cancers is not mandatory might be biased by an increased registration of cancers in patients followed within specialist care, compared to previously healthy individuals. Being monitored for a chronic disease might thus increase the probability of a subsequent cancer being registered. If so, this could partly explain the increased risk of cancer noted in patients with GCA in some studies.

To date no definite association between GCA/PMR and cancer has been agreed on because of conflicting epidemiological evidence, as illustrated in Table 3. The majority of previous studies have been retrospective cohort studies. However, 2 of the studies had a prospective design although differently executed^{11,18}. Prospective design is often considered superior as it allows an evaluation of the temporal sequence between exposure and outcome, and possibly reduces the risk of loss to followup. However, prospective studies are cost- and time-consuming, and often difficult to maintain for a long observation period, making them less suitable for the study of seldom and potentially late outcomes such as malignant diseases. Also, patients included in a prospective study are regularly evaluated and more likely to undergo extensive medical investigations, possibly leading to earlier detection and even an increased risk of cancer. As such, prospective studies run the risk of overestimating the incidence of cancer in the patient population compared to the controls. In the smaller prospective study, they found a standardized incidence ratio of malignancy of 4.61¹⁸. However, the number of patients was rather low (n = 107) and the number of recent and successive cancers was low (n = 7), rendering these numbers uncertain. Also, there were no age- and sex-matched controls, only the use of crude incidence rates.

It may be argued that a surveillance bias applies to any patient with GCA, often presenting with elevated inflammatory markers and nonspecific symptoms and signs. This clinical presentation may start an extensive diagnostic inves-

tigation, in which concomitant cancers are detected earlier than they otherwise would have been, thus shifting the timing of cancer diagnosis from post-GCA to pre-GCA. However, a large population-based study from the UK found no difference in the frequency of prior cancer between patients with and without GCA¹⁷, and Kermani, *et al* found that patients with GCA in a cohort from Olmsted County (USA) had significantly fewer malignancies prior to diagnosis compared to controls³³.

The 2 previously published reports on cancer risk in Norwegian patients with GCA concluded differently^{10,11}. Myklebust, *et al* concluded that there was no difference in risk in their cohort of 398 patients with GCA and 1592 controls¹¹. Haga, *et al* published a case-control study based on prospectively included cases of GCA or PMR during 1978–1983 with subsequent retrospective analysis of cancer risk based on cancer (registered in the CRN) from the inception of the registry through 1988¹⁰. The inclusion criteria were biopsy-proven GCA or fulfillment of study-specific clinical criteria for either GCA or PMR, and 185 cases and 925 controls were included (Table 3). While Haga, *et al* found no difference in overall cancer risk, they found an increased risk in biopsy-proven GCA with HR of 2.35 for cases versus controls. They also reported an increased HR of 7.25 for biopsy-positive compared to biopsy-negative cases. The Haga study included patients from hospitals within our catchment area and within our study period. However, they also included isolated PMR (91 of the 185 cases) and the study-specific inclusion criteria were somewhat different from ours. Further, the number of biopsy-proven GCA was low (n = 65), and the finding of 16 cancers among these might have been coincidental because we found no such difference in our 528 biopsy-proven cases. Two other previously published reports limited to biopsy-proven GCA cases have not confirmed the finding of increased risk associated with positive biopsy. In disagreement with Haga, *et al*¹⁰, they found no overall difference in

Table 3. Key features of previous studies on cancer risk in patients with giant cell arteritis (GCA).

Country and time period	Ungrasert, 2014 ¹⁹	Brekke, 2018 (PS)	Jese, 2018 ¹⁸	Li, 2017 ¹⁷	Ji, 2010 ¹⁶	Kermani, 2010 ¹⁵	Hill, 2010 ¹⁴	Gonzalez-Gay, 2007 ¹³	Liozon, 2006 ¹²	Myklebust, 2002 ¹¹	Haga, 1993 ¹⁰
Study design	NA	Norway, 1972–2012	Slovenia, 2011–2015	UK, 1990–2014	Sweden, 1965–2006	USA, 1950–2006	Australia, 1991–2006	Spain, 1981–2005	France, 1976–2005	Norway, 1987–1998	Norway, 1978–1988
Inclusion criteria	Metaanal.	Retro GCA defined by clinical diagnosis, ACR 1990 criteria, or positive biopsy	Prosp ACR 1990 criteria for cranial GCA, imaging criteria for purely extracranial GCA	Retro Diagnostic code GCA in nationwide general practice database	Retro Diagnostic code GCA or PMR in nationwide hospital discharge database	Retro ACR 1990 criteria for GCA	Retro Biopsy-proven GCA	Retro Biopsy-proven GCA	Retro ACR 1990 criteria for GCA	Prosp Biopsy-proven GCA, Bird or Hamrin criteria for PMR	Retro Biopsy-proven GCA or study-specific clinical criteria for PMR
Controls	NA	Age-, sex-, and geographically matched controls	General population > 50 yrs	Age-, sex-, and geographically matched controls without GCA	General population (database subjects without the disease)	Age-, sex-, and geographically matched controls without GCA	General population (age- and sex-specific)	NR	General population (same age group)	Age-, sex-, and geographically matched controls	Age-, sex-, and geographically matched controls
No. cases / controls	Pooled cases: 39,808	767 / 1437	107 / NR	9778 / 92,268	35,918 / NR	204 / 407	226 / NR	255 / NR	271 / NR	398 / 1592	185 / 925
Method of cancer identification	NA	Norwegian Cancer Registry	Prosp data + Slovenian cancer registry	Diagnostic code in general practice database	Swedish cancer registry	Review of complete medical records	South Australia cancer registry	Review of medical records	Review of medical records + regional cancer registry	Norwegian Cancer Registry	Norwegian Cancer Registry
Main conclusion	Increased risk	No difference	Increased risk	No difference	Increased risk	No difference	No difference	No difference	Increased risk	No difference	Increased risk (if TAB-positive)

ACR: American College of Rheumatology; GCA: giant cell arteritis; NA: not applicable (metaanalysis reporting pooled data from several cohort studies^{10,11,14,15,16}); Metaanal.: metaanalysis; Retro: retrospective; Prosp: prospective; NR: not reported; PMR: polymyalgia rheumatica; PS: present study; TAB: temporal artery biopsy.

risk of malignancy in biopsy-proven GCA compared to the general population^{13,14}.

Published reports have varied regarding the distribution of different types of cancers that patients with GCA experience. Kermani, *et al* found that nonmelanoma skin cancers were the most common cancers in both cases and controls¹⁵. This finding has not been confirmed by other studies, although Ji, *et al* also found an excess of skin cancers¹⁶. However, several studies have not included BCC of the skin as a malignancy^{10,11,12,14}. BCC in adults are not reported to the CRN and were thus not included among cancers observed in our study or in the previous Norwegian studies.

Liozon, *et al* found a strong temporal association between GCA and myelodysplastic syndromes and reported a concurrent (occurring 1 yr before or after GCA diagnosis) hematological malignancy in 3.6% of their cases¹². The combined malignancies of lymphoid and hematopoietic tissues accounted for almost one-quarter of the registered cancers in our study, afflicting 3.8% of all GCA cases and 3.5% of the matched controls. Overall, both frequency and types of cancers found in patients with GCA and population controls in our study were similar. This is in agreement with the findings reported by Myklebust, *et al* and largely with the sizeable Swedish study, except for their finding of more skin cancers and leukemia^{11,16}. The numbers for each cancer diagnosis are small and most studies lack sufficient power to analyze the risk of different cancer diagnoses separately.

We have discussed factors that may have contributed to our findings being different from those of some other studies. Our data are limited by the retrospective design and have incomplete (for cases) and missing (for controls) data on important cancer risk factors such as smoking, use of cytotoxic medications or hormones, and other potential confounders such as body mass index and family history of malignancy. For example, if the prevalence of smokers is reduced in patients with GCA compared to the controls, our study could underestimate the true cancer risk associated with GCA. We also note that our cohort consists of cases with predominantly cranial GCA (65% with positive TAB). Thus, our results may not be representative for cases with purely extracranial LVV. One strength of our study is the completeness and high quality of the CRN. It has been mandatory to register every new case of cancer in Norway in this registry since 1952. Another strength is the large and well-defined cohort of GCA cases resulting from a thorough review of clinical data, excluding misclassified cases, and including hospitalized patients, as well as those only treated in outpatient clinics. The study also included a large cohort of population controls that were tightly matched regarding the most significant of all cancer risk factors — age. The large sample size of both cases and controls rendered a well-powered analysis, allowing us to detect relevant differences between the groups. The long duration of our study (41

years) is a strength that reduces the risk of missing late-occurring cancers as well as diminishes the risk of erroneous conclusion based on variations through time. However, the long duration also poses a challenge because the diagnostic criteria for both cancer and GCA might have changed substantially during such a long period. Even so, time-dependent changes are controlled for by including the control population on which such factors presumably would have a similar effect.

Based on our findings, patients with GCA are not at increased risk of cancer following the diagnosis of their vasculitis. From a clinical point of view, our results indicate that patients with an unequivocal diagnosis of GCA do not need specific screening for malignancy.

ACKNOWLEDGMENT

The authors thank Dr. Rabab Adam, MD, for contributions in the early phase of the study. The authors also thank Alf Aksland (IT consultant, Haukeland University Hospital), Anita Mellingen (Department of Rheumatology, Haukeland University Hospital), Bjørg Sandvik (Department of Pathology, Haukeland University Hospital), Robinson Lazcano (Central archive, Haukeland University Hospital), Lisbeth Thorsen (Haralds plass Deaconess Hospital), Laila Kvåle (Voss Hospital), and their coworkers, who aided with the identification of patients or the extensive retrieval of patient records.

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