Characterization and risk estimate of cancer in patients with primary Sjögren syndrome

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Abstract

Background: The purpose of this study is to characterize the risk of cancer in a large cohort of patients with primary Sjögren syndrome (SjS).

Methods: We had analyzed the development of cancer in 1300 consecutive patients fulfilling the 2002 SjS classification criteria. The baseline clinical and immunological characteristics and systemic activity (ESSDAI scores) were assessed at diagnosis as predictors of cancer using Cox proportional hazards regression analysis adjusted for age at diagnosis and gender. The sex-and age-specific standardized incidence ratios (SIR) of cancer were estimated from 2012 Spanish mortality data.

Results: After a mean follow-up of 91 months, 127 (9.8%) patients developed 133 cancers. The most frequent type of cancer was B-cell lymphoma (including 27 MALT and 19 non-MALT B-cell lymphomas). Systemic activity at diagnosis of primary SjS correlated with the risk of hematological neoplasia and cryoglobulins with a high risk of either B-cell or non-B-cell lymphoma subtypes. Patients with cytopenias had a high risk of non-MALT B-cell and non-B-cell cancer, while those with low C3 levels had a high risk of MALT lymphomas and those with monoclonal gammapathy and low C4 levels had a high risk of non-MALT lymphomas. The estimated SIR for solid cancer was 1.13 and 11.02 for hematological cancer. SIRs for specific cancers were 36.17 for multiple myeloma and immunoproliferative diseases, 19.41 for Hodgkin lymphoma, 6.04 for other non-Hodgkin lymphomas, 5.17 for thyroid cancer, 4.81 for cancers of the lip and oral cavity, and 2.53 for stomach cancer.

Conclusions: One third of cancers developed by patients with primary SjS are B-cell lymphomas. The prognostic factors identified at SjS diagnosis differed according to the subtype of B-cell lymphoma developed. Primary SjS is also associated with the development of some non-hematological cancers (thyroid, oral cavity, and stomach).

Keywords: Sjögren syndrome, Cancer, Lymphoma
Background

Sjögren syndrome (SjS) is a systemic autoimmune disease that principally affects women between the fourth and sixth decades of life who present with sicca symptomatology caused by dryness of the main mucosal surfaces [1]. The clinical spectrum of SjS extends from dryness to systemic involvement. Since 1978 [2], SjS has been closely associated with an enhanced risk of lymphoma, one of the most severe complications a patient may develop. Primary SjS patients have a 10–44-fold greater risk of lymphoma than healthy individuals, higher than that reported for systemic lupus erythematosus (sevenfold) and rheumatoid arthritis (fourfold) [3].

The close link between lymphoma and SjS is clearly exemplified by the very specific type of lymphoma arising in SjS patients, mainly low-grade B-cell lymphomas (predominantly marginal zone histological type) with primary extranodal involvement of the major salivary glands (overwhelmingly parotid) [4, 5]. The lymphoma-genesis hypothesis in primary SjS suggests a key role for the continued stimulation of B cells in the exocrine glands and organs containing mucosa-associated lymphoid tissue (MALT); in some patients, this autoimmune process is further altered by a combination of oncogenic factors that promote abnormal increases in B-cell survival, thus putting the patient at high risk of a malignant B-cell transformation [6].

Studies that have analyzed the association between primary SjS and cancer have overwhelmingly been centered on the characterization and risk estimation of lymphoma [7]. Little data is available on the association between primary SjS and non-B-cell lymphomas or solid cancer [11]. The aims of this study were to analyze the development of solid and hematological cancer, identify related baseline SjS-prognostic factors (including systemic activity), and estimate the corresponding standardized incidence ratios (SIR) per type of cancer in a large, well-characterized cohort of Spanish patients with primary SjS.

Methods

Patients

The GEAS-SS Study Group was formed in 2005 with the aim of collecting a large series of Spanish patients with primary SjS. Both incident and prevalent cases were included; for incident cases, the diagnosis of primary SjS was confirmed after January 2005. By January 2016, the database included 1300 consecutive patients (513 prevalent cases) who fulfilled the 2002 classification criteria for primary SjS [12]. Exclusion criteria were chronic HCV/HIV infection and associated systemic autoimmune diseases. Diagnostic tests for SjS (ocular tests, parotid scintigraphy, and salivary gland biopsy) were performed according to the European Community Study Group recommendations [12]. Clinical and laboratory data were collected and computerized according to a standard previously reported protocol [13].

Definition of variables

The date of disease diagnosis was defined as the date when the attending physician confirmed fulfillment of the 2002 criteria [12]. Systemic involvement was defined according to the ESSDAI [14]. Disease activity states (DAS) were categorized according to the global ESSDAI score as low activity (ESSDAI <5), moderate activity (5 ≤ ESSDAI ≤ 13), and high-activity (ESSDAI ≥ 14) [15].

Cancers were determined through medical chart review of the hospital discharge summaries by the corresponding oncology/hematology departments. We included only cancers occurring after the confirmed diagnosis of primary SjS by the attending physician. Solid cancers were classified using the nomenclature included in the GLOBOCAN Project (http://globocan.iarc.fr/) and hematological cancers using the 2016 WHO classification [16, 17].

Statistical analysis

Descriptive data are presented as means and standard deviation (SD) for continuous variables and numbers and percentages (%) for categorical variables. Time-to-event analyzes for cancer are presented as Kaplan-Meier curves. The baseline clinical and immunological characteristics and systemic activity (ESSDAI scores) were assessed at diagnosis as predictors of cancer using univariate Cox proportional-hazards regression analysis adjusted for age at diagnosis and gender. The level of systemic activity was recoded as no activity vs. any type of activity (low/moderate/high) in the analysis. Multivariate Cox proportional-hazards regression analysis allowed adjustment for age at diagnosis and gender and variables that were statistically significant (p < 0.05) in the univariate analysis, in order to establish independent variables associated with hematological and solid cancers. The hazard ratios (HRs) and their 95% confidence intervals (CIs) obtained in the adjusted regression analysis were calculated. The person-years of follow-up were calculated from the date of diagnosis to the last visit, cancer, or death (whichever occurred earliest). The standardized incidence ratios (SIR) were computed as the ratio of observed to expected cancers. Expected cancers were determined by multiplying person-years by the corresponding sex- and age-specific incidence rates of cancer in the general Spanish population in 2012 (the most recent data available from the International Agency for Research on Cancer) provided by the GLOBOCAN project (http://globocan.iarc.fr/old/method/method.asp?country=724) and summing overall person-years. The sex- and age-specific incidence rates of cancer for Spain were estimated from national mortality data for 2012 by
modeling, using a set of age-, sex-, and site-specific incidence:mortality ratios obtained by the aggregation of recorded data from the 12 Spanish cancer registries [18]. The incidence rates of several types of cancers could not be estimated for population belonging to the age category “0–14” by the fact that these types of cancers were not detected in this population. Thus, this category was excluded when SIR were computed. The 95% confidence intervals (CI) of the SIR were also calculated. All significance tests were two tailed, and values of \( p < 0.05 \) were considered significant. All analyses were conducted using the R version 3.2.3 for Windows statistical software package.

**Results**

**Baseline prognostic factors**

Baseline characteristics of primary SjS patients are summarized in Table 1. The cohort consisted of 1300 patients, including 1201 (92.4%) women and 99 (7.6%) men (female: male ratio, 12:1), with a mean age at diagnosis of 55.1 (SD 15.4) years (range, 15.3–92.9 years).

After a median follow-up of 66.1 months (range 1–560.3 months; 9922.3 person-years), 127 (9.8%) patients developed 133 cancers: 64 patients developed a solid cancer, 57 patients hematological cancer, 4 patients both solid and hematological cancers, and 2 patients two different types of solid neoplasia. The most frequent types of cancers included B-cell MALT lymphomas (\( n = 27, 20\% \)), other B-cell lymphomas (\( n = 19, 14\% \)), breast (\( n = 14, 11\% \)), colorectal (\( n = 9, 7\% \)), myeloid neoplasia/leukemia (\( n = 8, 6\% \)), lung (\( n = 6, 5\% \)), and stomach (\( n = 5, 4\% \))
cancers (Additional file 1: Table S1). In comparison with patients who developed solid cancer, those who developed hematological cancer had a higher frequency at diagnosis of cryoglobulins (\( p = 0.002 \)), low C3 levels (\( p = 0.018 \)), high ESSDAI score (\( p = 0.001 \)), and high DAS (\( p < 0.001 \)) (Additional file 1: Table S1). The median time to the development of cancer was 59.9 months. Figure 1 shows the Kaplan-Meier curves for the development of the first solid or hematological cancer. Survival (free of cancer) at 5, 10, 20, and 30 years was 93.9, 87.9, 76.9, and 70.3%, respectively. The following baseline

### Table 1 Baseline SjS-related features and risk of development of solid and hematological cancer

| Baseline features | Values at diagnosis (\( n = 1300 \)) | Solid cancer (\( n = 70 \)) | Hematological cancer (\( n = 61 \)) |
|-------------------|--------------------------------------|----------------------------|-------------------------------------|
| Age at diagnosis  | 54.8 ± 15.4                          | †                          | †                                   |
| Gender (male)     | 99 (7.6)                             | †                          | †                                   |
| Ethnicity (white) | 1246 (95.8)                          | 0.41 [0.10–1.71]           | 0.99 [0.14–7.29]                    |
| Dry mouth         | 1299 (99.9)                          | –                          | –                                   |
| Dry eye           | 1234 (94.9)                          | 0.48 [0.19–1.20]           | 1.37 [0.33–5.63]                    |
| Altered ocular tests | 1013/1156 (87.6)                     | 1.29 [0.47–3.58]           | 5.38 [0.74–39.03]                   |
| Altered parotid scintigraphy | 835/992 (84.2) | 0.87 [0.41–1.86] | 1.34 [0.53–3.41] |
| Positive salivary gland biopsy | 497/629 (79) | 0.92 [0.35–2.44] | 1.01 [0.42–2.45] |
| Anemia (Hb < 110 g/L) | 216/1297 (16.7) | 1.05 [0.59–1.86] | 2.07 [1.20–3.58]‡ |
| Leukopenia (<4000/mm\(^3\)) | 227/1297 (17.5) | 1.49 [0.87–2.55] | 0.96 [0.50–1.86] |
| Thrombocytopenia (<150,000/mm\(^3\)) | 87/1297 (6.7) | 1.18 [0.54–2.57] | 1.17 [0.50–2.72] |
| Neutropenia (<1500/mm\(^3\)) | 137/1296 (10.6) | 1.41 [0.74–2.71] | 1.66 [0.83–3.32] |
| Lymphopenia (<1000/mm\(^3\)) | 141/1295 (10.9) | 1.80 [0.96–3.35] | 1.43 [0.71–2.86] |
| Antinuclear antibodies+ | 1124/1295 (86.8) | 0.95 [0.49–1.85] | 1.05 [0.50–2.21] |
| Rheumatoid factor+ | 583/1245 (46.8) | 1.13 [0.71–1.82] | 1.08 [0.65–1.80] |
| Anti-Ro/SS-A+ | 981/1294 (75.8) | 1.43 [0.81–2.53] | 1.28 [0.70–2.36] |
| Anti-La/SS-B+ | 626/1289 (48.6) | 0.99 [0.62–1.60] | 1.17 [0.69–1.97] |
| Monoclonal gammopathy | 114/1018 (11.2) | 1.94 [0.99–3.80] | 2.13 [1.08–4.19] |
| Cryoglobulins+ | 70/929 (7.5) | 0.70 [0.25–1.94] | 4.37 [2.32–8.22]‡ |
| Low C3 levels (<0.82 g/L) | 144/1233 (11.7) | 0.59 [0.21–1.62] | 2.07 [1.06–4.03] |
| Low C4 levels (<0.11 g/L) | 163/1218 (13.4) | 1.07 [0.51–2.24] | 2.23 [1.17–4.26] |

Values are represented as HRs [95% CIs]
Set in italics, statistically significant (\( p < 0.05 \)) SjS-related features associated with cancer in the univariate Cox proportional hazards regression analysis adjusted for age at diagnosis and gender
SjS Sjögren syndrome
†Adjusting variable
‡Statistically significant (\( p < 0.05 \)) SjS-related features associated with cancer in the multivariate Cox proportional hazards regression analysis adjusted for age at diagnosis and gender
variables at diagnosis were associated with hematological cancer in the Cox regression analysis (Table 1): anemia (HR 2.07; \( p = 0.009 \)), C3 levels <0.82 g/L (HR 2.07; \( p = 0.033 \)), C4 levels <0.11 g/L (HR 2.23; \( p = 0.015 \)), monoclonal gammopathy (HR 2.13; \( p = 0.028 \)), and cryoglobulins (HR 4.37; \( p < 0.001 \)). Multivariate analysis identified anemia and cryoglobulins as variables independently associated with hematological cancer. No variables were significantly associated with the development of solid cancer.

**Systemic activity**

Table 2 summarizes the relationship between organ-by-organ baseline ESSDAI activity at diagnosis and the risk of hematological and solid cancer. The mean ESSDAI score at diagnosis of the entire cohort was 6.00 (SD 6.6) and correlated with the risk of hematological cancer (HR 1.06; \( p < 0.001 \)). The highest risk was found in patients with a high baseline DAS (HR 4.34; \( p < 0.001 \)). Baseline activity in the constitutional (HR 2.44; \( p = 0.011 \)), lymphadenopathy (HR 8.03; \( p < 0.001 \)), glandular (HR 3.04; \( p < 0.001 \)), and biological (HR 2.12; \( p = 0.011 \)) domains was associated with a higher risk of hematological cancer. Multivariate analysis showed that the lymphadenopathy and glandular domains were independently associated with the risk of hematological cancer. A sensitivity analysis comparing no/low activity vs. moderate/high activity showed similar results (although with higher HRs), with the addition of the hematological domain, which was significantly associated with the development of hematological cancer (HR 2.72; \( p = 0.001 \)) (Additional file 1: Table S3). Additional file 1: Table S4 summarizes the mean baseline ESSDAI scores of patients classified according to the type of cancer.

Additional file 1: Table S5 summarizes the potential influence of SJS-related therapies (pilocarpine, hydroxychloroquine, corticosteroids, immunosuppressive agents, intravenous immunoglobulin, and rituximab) in the development of cancer. In the univariate analysis, we found a higher risk of developing hematological cancer in patients receiving corticosteroids (HR 2.67; \( p < 0.001 \)) or an immunosuppressive agent (HR 2.83; \( p < 0.001 \)), although the multivariate analysis adjusted for the main prognostic factors identified (anemia, monoclonal gammopathy, cryoglobulins, low C3, low C4, and ESSDAI) showed only non-significant differences.

**Prognostic factors for hematological cancer subtypes**

According to the WHO classification, we classified hematological cancers into three main groups: B-cell MALT lymphoma \( (n = 27) \), B-cell non-MALT lymphoma \( (n = 19) \), and non-B-cell \( (n = 15) \) cancers (Table 3). Baseline prognostic factors associated with B-cell MALT lymphomas included cryoglobulins (HR 6.32; \( p < 0.001 \)) and low C3 levels (HR 3.25; \( p = 0.010 \)). For B-cell non-MALT lymphomas, the prognostic factors included anemia (HR 2.58; \( p = 0.047 \)), monoclonal gammopathy (HR 3.45; \( p = 0.024 \)), cryoglobulins (HR 3.34; \( p = 0.028 \)), and low C4 levels (HR 3.83; \( p = 0.014 \)). For non-B-cell hematological cancers, prognostic factors included anemia (HR 4.02; \( p = 0.009 \)), neutropenia (HR 5.86; \( p = 0.002 \)), thrombocytopenia (HR 4.85; \( p = 0.004 \)), and cryoglobulins (HR 4.36; \( p = 0.034 \)).

Baseline systemic activity (global ESSDAI score) and high DAS were associated with a higher risk of B-cell MALT and non-B-cell cancers (but not B-cell non-MALT lymphomas). Distribution per organ showed that baseline systemic activity in the lymphadenopathy domain was linked to a higher risk for the three subtypes of hematological cancer, activity in the glandular domain with a higher risk of MALT lymphomas, activity in the biological domain with a higher risk of non-MALT B-cell lymphomas, and activity in the constitutional, pulmonary, and hematological domains with a higher risk of non-B-cell hematological cancer (Table 3).

**Standardized incidence ratios (SIRs) for cancer**

For all cancers combined, the SIR estimate was 1.91 (95% CI 1.60 to 2.28) and was higher in men than in women (2.29 vs. 1.87) (Table 4). The SIR was 1.13 (95% CI 0.88 to 1.46) for solid cancer and 11.02 (95% CI 8.35 to 14.54) for hematological cancer. With respect to solid cancers, we found an increased risk for thyroid cancer (SIR 5.17; 95% CI 1.94 to 13.79), cancers of the lip and oral cavity (SIR 4.81; 95% CI 1.81 to 12.83), and stomach cancer (SIR 2.53; 95% CI 1.05 to 6.07) in women. We analyzed potential predictive factors for the development of thyroid, lip, and oral cavity, and stomach cancers.
(Additional file 1: Table S6) and found that only ethnicity was a predictive factor for the development of any of these cancers, since non-white patients had a HR of 10.44 \((p = 0.004)\).

With respect to hematological cancers, we had to adapt our classification of hematological cancer (based on the WHO 2016 nomenclature) to the four categories of hematological cancer used in the GLOBOCAN database (which are based on ICD codes); the correspondence between the two classifications is shown in Additional file 1: Table S7. We found an increased risk for three of the four GLOBOCAN categories corresponding to hematological cancer: multiple myeloma and immunoproliferative diseases \((SIR 36.17; 95\% CI 25.44 \text{ to } 51.43)\), Hodgkin lymphoma \((SIR 19.41; 95\% CI 7.29 \text{ to } 51.72)\), and non-Hodgkin lymphoma \((SIR 6.04; 95\% CI 3.43 \text{ to } 10.64)\).

**Discussion**

In non-specialized medical settings, primary SjS is often considered a chronic, non-life threatening disease that causes dryness, fatigue, and pain. However, systemic involvement has increasingly been recognized as a key part of the disease spectrum with a significant weight in dictating the prognosis and survival \([13]\). Among the systemic manifestations of SjS, lymphoma is one of the worst complications that physicians should expect. In 1978, Kassan et al. \([2]\) estimated a SIR for NHL of 44.4 in patients with primary SjS. The SIRs estimated by subsequent studies, even though the great majority have been lower, have confirmed that primary SjS patients are at higher risk of lymphoma, with a pooled 14-fold higher risk reported by Liang et al. \([19]\) in a recent meta-analysis. However, this meta-analysis included cohorts of patients in whom the diagnosis of primary SjS was based on the 1993 criteria, whose fulfillment allows the inclusion of patients with negative Ro/La antibodies and negative biopsy. This creates a significant bias, since several studies have reported a lower risk of lymphoproliferation in immunonegative patients \([8, 20]\). Table 5 summarizes the studies based on the fulfillment of the 2002 criteria \([8–11, 21–27]\). The SIRs for B-cell lymphoma range between 7 and 9 in population-based studies and between 16 and 48 in hospital-based studies. The majority of these studies were focused only on B-cell lymphoma, and most of them used an old terminology (NHL).
The present study is the first to analyze the risk of the different types of cancer (including solid cancer and hematological cancers other than lymphoma) and identify the corresponding baseline predictive factors in a hospital-based cohort of patients with primary SjS. Only three previous studies have estimated the SIR for all type of cancers in primary SjS. The risk was not significant in a population-based study [11] but was significant in the two hospital-based studies [8, 21] (although the significance was at the limit in the study by Theander et al [8]). We also found a higher significant risk with a SIR of nearly 20, although when we separated solid and hematological cancers, the SIR for solid cancers was not significant, while the SIR for hematological cancer was 11-fold higher (10-fold higher in women and 20-fold higher in men).
Although the vast majority of cells infiltrating the salivary glands of patients with primary SjS are T cells [6], the majority of lymphomas reported are of B-cell origin (in our study, the ratio between B- and T-cell lymphomas was 15:1). Among B-cell lymphomas, the different subtypes not only have differing clinical presentations but also have a different frequency and, logically, a different prognosis. Therefore, the identification of predictive factors for the development of the different subtypes of B-cell lymphoma seems rational. Three subtypes of lymphoma alone account for more than 90% of reported cases in primary SjS (Table 5): MALT lymphoma (58%), DLBC (20%), and MZ lymphoma (13%). Plasma-cell myeloma was reported in only four cases (2%), although a recent study [28] has described an increased risk in primary SjS patients presenting with monoclonal gammapathy. Among non-B-cell hematological cancers, we found 8 patients with myeloid neoplasia/leukemia (not reported in previous studies), four cases of Hodgkin disease, and three cases of T/NK-cell lymphoma (4 cases and 2 cases previously reported, respectively) [8–10, 21], representing 25% of the hematological cancers observed.

Recent studies have confirmed severe parotid involvement, purpura, leukopenia, anti-La antibodies, raised levels of BAFF and beta2-microglobulin, cryoglobulins, monoclonal band, and hypocomplementemia as risk factors for B-cell lymphoma [24, 25, 29, 30].

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Table 4  Standardized incidence ratios (SIRs) for cancer classified according to the GLOBOCAN categories

| Cancer categories | Total (n = 1239)* | Women (n = 1145) | Men (n = 94) |
|-------------------|------------------|-----------------|-------------|
|                   | Obs† | Exp | SIR | LCI | UCI | Obs | Exp | SIR | LCI | UCI | Obs | Exp | SIR | LCI | UCI |
| All cancers†      | 121  | 63.77 | 1.9 | 1.59 | 2.27 | 105 | 56.77 | 1.85 | 1.53 | 2.24 | 16  | 7  | 2.29 | 1.4 | 3.73 |
| Solid cancers§    | 60   | 53.1 | 1.13 | 0.88 | 1.46 | 55  | 47.42 | 1.16 | 0.89 | 1.51 | 5   | 5.68 | 0.88 | 0.37 | 2.12 |
| Thyroid           | 4    | 0.79 | 5.05 | 1.89 | 13.45 | 4   | 0.77 | 5.17 | 1.94 | 13.79 | 0   | 0.02 | 0    | –  | –   |
| Lip, oral cavity  | 4    | 0.99 | 4.05 | 1.52 | 10.8  | 4   | 0.83 | 4.81 | 1.81 | 12.83 | 0   | 0.16 | 0    | –  | –   |
| Stomach           | 5    | 2.24 | 2.23 | 0.93 | 5.36  | 5   | 1.98 | 2.53 | 1.05 | 6.07  | 0   | 0.26 | 0    | –  | –   |
| Kidney            | 3    | 1.67 | 1.8  | 0.58 | 5.57  | 2   | 1.44 | 1.39 | 0.35 | 5.55  | 1   | 0.23 | 4.39 | 0.62 | 31.14 |
| Brain, nervous system | 2  | 1.17 | 1.72 | 0.43 | 6.86  | 2   | 1.07 | 1.87 | 0.47 | 7.48  | 0   | 0.1  | 0    | –  | –   |
| Gallbladder       | 1    | 0.77 | 1.3  | 0.18 | 9.2   | 1   | 0.72 | 1.39 | 0.2  | 9.83  | 0   | 0.05 | 0    | –  | –   |
| Lung              | 6    | 4.66 | 1.29 | 0.58 | 2.87  | 4   | 3.43 | 1.17 | 0.44 | 3.11  | 2   | 1.23 | 1.63 | 0.41 | 6.5 |
| Prostate          | 2    | 1.66 | 1.2  | 0.3  | 4.81  | 2   | 1.66 | 1.2  | 0.3  | 4.81  | 2   | 1.66 | 1.2  | 0.3  | 4.81 |
| Melanoma of skin  | 2    | 1.73 | 1.16 | 0.29 | 4.62  | 2   | 1.63 | 1.23 | 0.31 | 4.92  | 0   | 0.1  | 0    | –  | –   |
| Bladder           | 2    | 2.1  | 0.95 | 0.24 | 3.8   | 2   | 1.48 | 1.35 | 0.34 | 5.42  | 0   | 0.63 | 0    | –  | –   |
| Colorectal        | 9    | 10.08| 0.89 | 0.46 | 1.72  | 9   | 9.02 | 1    | 0.52 | 1.92  | 0   | 1.06 | 0    | –  | –   |
| Breast            | 14   | 15.68| 0.89 | 0.53 | 1.51  | 14  | 15.68| 0.89 | 0.53 | 1.51  | 0   | 0    | 1    | –  | –   |
| Pancreas          | 2    | 2.29 | 0.87 | 0.22 | 3.5   | 2   | 2.1  | 0.95 | 0.24 | 3.8   | 0   | 0.18 | 0    | –  | –   |
| Cervix uteri      | 1    | 1.39 | 0.72 | 0.1  | 5.1   | 1   | 1.39 | 0.72 | 0.1  | 5.1   | 0   | 0.1  | 1    | –  | –   |
| Corpus uteri      | 2    | 3.72 | 0.54 | 0.13 | 2.15  | 2   | 3.72 | 0.54 | 0.13 | 2.15  | 0   | 0    | 1    | –  | –   |
| Ovary             | 1    | 2.16 | 0.46 | 0.07 | 3.28  | 1   | 2.16 | 0.46 | 0.07 | 3.28  | 0   | 0.18 | 0    | –  | –   |
| Hematologicalb    | 50   | 4.54 | 11.02| 3.85 | 14.54 | 42  | 4.14 | 10.14 | 7.49 | 13.72 | 8   | 0.39 | 20.34 | 10.17 | 40.67 |
| Leukemia          | 3    | 1.49 | 2.02 | 0.65 | 6.26  | 2   | 1.34 | 1.49 | 0.37 | 5.95  | 1   | 0.14 | 7.02  | 0.99 | 49.8 |
| MM/ID and lymphoma| 47   | 3.05 | 15.41| 11.58| 20.51 | 40  | 2.80 | 14.29 | 10.48| 19.48 | 7   | 0.25 | 27.91 | 13.31 | 58.54 |
| Hodgkin lymphoma  | 4    | 0.21 | 19.41| 7.29 | 51.72 | 3   | 0.19 | 15.87| 5.12 | 49.21 | 1   | 0.02 | 58.64 | 8.26 | 416.27 |
| MM/ID             | 31   | 0.86 | 36.17| 25.44| 51.43 | 28  | 0.79 | 35.59| 24.58| 51.55 | 3   | 0.07 | 42.58 | 13.73 | 132.03 |
| Non-Hodgkin lymphoma | 12 | 1.99 | 6.04 | 3.43 | 10.64 | 9   | 1.82 | 4.94 | 2.57 | 9.49  | 3   | 0.16 | 18.37 | 5.93 | 56.96 |

*Excluding patients diagnosed with cancer before fulfillment of primary SjS criteria (n = 55) and patients with an age at SjS diagnosis <15 years (n = 6) (the age category “0–14” was excluded when SIR were computed). From 6 patients with an age at SjS diagnosis <15 years, one was diagnosed with cancer bExcluding 7 patients with hematological neoplasia not included in the GLOBOCAN categories (1 refractory cytopenia with multilineage dysplasia, 1 refractory anemia with unilineage dysplasia, 1 refractory anemia with excess blasts, 1 myelodisplastic syndrome undiagnosable, 1 mastocytosis, 1 essential thrombocythemia, and 1 angioimmunoblastic T-cell lymphoma) (see Additional file 1: Table S5) and 4 patients who developed first a solid cancer and subsequently a hematological cancer

†Excluding patients with non-melanoma skin cancer (n = 4, not included in the GLOBOCAN classification) and a non-classifiable skin cancer (n = 1, uncertain whether was a melanoma or non-melanoma cancer)

§Excluding 4 patients with neoplasia not included in the GLOBOCAN categories (2 endocrine, 2 ocular)
| Author          | Year | Country | Setting | Primary SjS patients (n) | Mean follow-up (yrs/pat-year) | Population cancer registry/classif | Cancer (n) | All-cancer SIR (95% CI) | Specific solid cancer SIR | Hemat neoplasia classif | NHL (n) | MALT (n) | DLBC (n) | MZ (n) | Myel (n) | Other B-cell | Non-B-cell | NHL SIR (95% CI) | MM SIR (95% CI) |
|-----------------|------|---------|---------|--------------------------|-----------------------------|----------------------------------|------------|------------------------|--------------------------|--------------------------|---------|---------|---------|-------|---------|-------------|-----------|-----------------|----------------|
| Theander 2006   | Sweden | Hosp    | 286     |                          | jul-64                      | Swedish Cancer Reg/ICD7          | 33         | 1.42 (0.98–2.00)       | NA                       | WHO 2001                  | 12      | 11      | 7       | 0     | 2       | 1           | T-cell (1) | 15.6 (7.8–27.8) | 3.27 (0.1–18.2) |
| Baimpa 2009     | Greece | Hosp    | 536     |                          | 2.6/ND                      | NA                               | ND         | NA                     | NA                       | ND                       | 40      | 38      | 21      | 7     | 5       | 0           | 0         | 1.35 (0.7–3.9)  | ND             |
| Zhang 2010      | China  | Hosp    | 1320    |                          | 4.4/ND                      | Shanghai Reg/ICD10              | 29         | 3.25 (2.12–4.52)       | NA                       | WHO 2001                  | 10      | 8       | 2       | 2     | 0       | 0           | 3         | T-cell (1)     | 48.1 (20.7–94.8) | 37.9 (4.58–136.7) |
| Weng 2012       | Taiwan | Pop     | 7852    |                          | ND/27.24                     | NHIV/ICD9                         | 227        | 1.01 (0.74–1.35)       | Colon 0.22 (0.05–0.6); Thyroid 2.56 | ICD9                    | 31      | 23      | ND      | ND    | ND      | ND          | ND         | 7.1 (4.2–10.3)  | 6.1 (2.0–14.2)  |
| Baldini 2012    | Italy  | Hosp    | 563     |                          | 6/ND                         | NA                               | ND         | NA                     | NA                       | WHO 2001                  | 12      | 8       | 3       | 1     | 0       | 0           | ND         | NA             | ND             |
| Hemminki 2012   | Sweden | Pop     | 1516    |                          | ND/16.700                    | Swedish Cancer Reg/ICD            | NA         | NA                     | Breast 0.46 (0.26–0.75) | NA                      | NA      | ND      | ND      | ND    | ND      | ND          | ND         | ND             | ND             |
| Johnsen 2013    | Norway | Pop     | 443     |                          | ND/38.13                     | Norway Reg/ICD10                 | NA         | NA                     | NA                       | ICD10                   | 7       | 6       | 0       | 1     | 0       | 0           | ND         | 9.0 (7.1–26.3)  | ND             |
| Risselada 2013  | Netherl | Hosp   | 195     |                          | 7.7/ND                       | NA                               | NA         | NA                     | NA                       | WHO 2001                  | 21      | 10      | 8       | 3     | 0       | 0           | ND         | NA             | ND             |
| Quartuccio 2014 | Italy  | Hosp    | 661     |                          | ND                           | NA                               | NA         | NA                     | NA                       | ND                      | 40      | ND      | ND      | ND    | ND      | ND          | ND         | NA             | ND             |
| Papageorgiou 2015 | Greece | Hosp | ND     |                          | ND                           | NA                               | ND         | ND                     | ND                       | ND                      | 77      | 51      | 12      | 8     | 0       | 6           | ND         | ND             | ND             |
| Nocturne 2016 | France | Hosp    | ND     |                          | ND                           | WHO                               | ND         | ND                     | ND                       | ND                      | 99      | 58      | 17      | 18    | 0       | 6           | ND         | ND             | ND             |
| Present study 2016 | Spain | Hosp   | 1300    |                          | 7.6/ND                       | GLOBOCAN/ICD10                   | 122        | 1.91 (1.6–2.28)        | Thyroid 5.17 (1.94–13.79); Lip/oral 4.81 (1.81–12.83); Stomach 2.53 (1.05–6.07) | WHO 2016                     | 61      | 12      | 27      | 7     | 2       | 6           | HD (4); T-cell (3); myeloid/leuk (8) | 6.04 (3.43–10.64) | 36.17 (25.44–51.43) |

ND not detailed, NA non-applicable, MALT mucosa-associated lymphoid tissue, DLBC Diffuse large B-cell lymphoma, MZ marginal zone lymphoma, MM myeloma multiple, Netherl Netherlands, Hosp hospital, Pop population, Classif classification, Hemat hematological, Myel Myeloma
study is the first to report that the prognostic role of these risk factors may differ according to the subtype of hematological cancer. We found a higher risk of MALT lymphoma in patients presenting with systemic activity, positive cryoglobulins, and low C3 levels at SjS diagnosis, while the risk of non-MALT B-cell lymphomas was unrelated to systemic activity, with anemia, monoclonal gammapathy, cryoglobulins, and low C4 levels at SjS diagnosis being the main risk factors. For non-B-cell hematological cancers, risk factors were systemic activity, cytopenias (anemia, thrombocytopenia and leukopenia), and cryoglobulins at SjS diagnosis. With respect to the influence of immunosuppressive therapies in the development of cancer, we found a higher risk of developing hematological cancer in patients who ever received corticosteroids/immunosuppressants, although the differences were non-significant after adjustment for the main predictive factors in which systemic activity is included. The explanation for the disappearance of significant differences in the multivariate model is that the more active patients are those who are mainly treated with corticosteroids/immunosuppressants and, therefore, the adjustment of comparisons for systemic activity yielded non-significant differences.

Analysis of systemic organ-by-organ activity showed that activity in the lymphadenopathy domain at SjS diagnosis was associated with a high risk of all subtypes of hematological neoplasia, while activity in the glandular domain was associated with the development of MALT lymphoma, activity in the biological domain with non-MALT lymphoma, and activity in the constitutional, pulmonary, and hematological domains with non-B-cell cancers. These differentiated high-risk profiles (Additional file 2: Figure S1), which are present at the diagnosis of primary SjS, may help physicians identify which patients may be at high risk of developing a specific subtype of hematological cancer. Together with the differentiated clinical presentation and the predilection for specific lymphoma subtypes, these high-risk profiles may contribute to an as-early-as-possible diagnosis and, therefore, to earlier specific therapeutic management that could improve survival.

We also analyzed and characterized the development of solid neoplasia in our cohort of patients with primary SjS and found that the risk was not increased with respect to the control population, in contrast to the results reported in Chinese patients [21]. Only two previous studies analyzed the risk for specific types of solid cancer and reported a lower risk for the development of colon and breast cancers [11, 27] and a higher risk for thyroid cancer [11]. We found an enhanced risk for the development of thyroid, lip/oral cavity, and stomach cancers (SIRs of 5.17, 4.81, and 2.53, respectively), with an enhanced risk of developing any of these three cancers (HR 10.44) found in non-white patients. The highest risk was for thyroid cancer, in which several risk factors [31] are clearly shared with primary SjS, including the predominantly female involvement [32], the low frequency of smokers [33], and the high frequency of association with autoimmune thyroiditis [34, 35]. The finding of the higher risk for cancers of the oral cavity and stomach in women with primary SjS is interesting, since the oral cavity is overwhelmingly involved in primary SjS and the stomach is the most frequent extraglandular extranodal site of lymphoma involvement in primary SjS [36], although further specific studies of the potential influence of dietary and other lifestyle factors (not analyzed in our study) are necessary. We also explored a potential association between the degree of oral involvement and the risk of developing cancer of the lip/oral cavity, but found no significant association between dry mouth, the severity of parotid scintigraphy results, and the presence of lymphocytic sialoadenitis in the salivary biopsy (data not shown), although the number of patients was too small to make solid conclusions.

Several methodological considerations should be discussed with respect to our results. The first is the limitation of the database available to estimate the SIRs for hematological cancer in the Spanish population (GLOBOCAN), since the two main GLOBOCAN categories (NH and MM/ID) contain a mix of B-cell subtypes. In the case of primary SjS, the inclusion of the most frequent hematological cancer (MALT) in the MM/ID category and not in the NHL category may cause some misinterpretation of the corresponding SIRs (31 for MM/ID, 12 for NHL); in fact, the SIR for the combination of the three categories of lymphomas included in GLOBOCAN was 15.41 (95% CI 11.58–20.51). In addition, 11 of the 61 hematological cancers developed by our primary SjS patients were not included in the GLOBOCAN categories corresponding to the hematological cancers, leading to a probable underestimation of the global SIR for hematological neoplasia found in our study. This highlights the importance of detailed information in these types of studies on the correspondence between the classification of hematological neoplasia often used in clinical practice (WHO) and the classification used in the cancer database of the general population (overwhelmingly using ICD codes). In addition, a potential risk of selection bias between the different types of hematological cancers should be taken into account (tertiary care centers will probably report a wider spectrum of cancer subtypes than less specialized centers). Homogeneity in classifying SjS patients according to the most accepted criteria (2002 AECG) is important, a factor not applied in recent studies that used the older 1993 criteria [37, 38] or in which the diagnosis of SjS was self-reported by the patient [39–41].
Conclusions
Patients with primary SjS had an 11-fold higher risk of developing hematological cancers than the general Spanish population. One third of the cancers developed during the study follow-up period were B-cell lymphomas (of which MALT lymphomas accounted for 60% of cases); the main prognostic factors identified at SjS diagnosis included systemic activity, cytopenias, and cryoglobulin-related immunological markers, although the weight of these factors differed for each subtype of B-cell lymphoma. Primary SjS was also associated with a higher risk of some types of non-hematological cancers (thyroid, oral cavity, and stomach). Patients with primary SjS should be closely followed not only for the enhanced risk of hematological cancer (not only B-cell lymphomas), but also for some types of solid cancer.

Additional file

Additional file 1: Table S1. Frequency of solid and hematologic cancers. Table S2. Comparison between patients who developed solid cancer and those who developed hematological cancer. Table S3. Sensitivity analysis comparing no/low activity vs. moderate/high activity. Table S4. Mean baseline ESSDAI scores of patients classified according to the type of cancer. Table S5. Analysis of a potential influence of SJS-related therapies (pilocarpine, hydroxycarbamol, corticosteroids, immunosuppressive agents, intravenous immunoglobulin and rituximab) in the development of cancer. Table S6. Analysis of potential predictive factors for the development of these three non-hematological cancers (thyroid, lip and oral cavity, and stomach cancers). Table S7. Correlation between the WHO 2016 nomenclature and the four categories of hematological cancer used in the GLOBOCAN database (which are based on ICD codes). (DOCX 38 kb)

Additional file 2: Figure S1. Frequency of the main WHO subtypes of hematological cancer and the corresponding predictive factors identified at SjS diagnosis. (TIF 182 kb)

Abbreviations
DAS: Disease activity states; MALT: Mucosa-associated lymphoid tissue; SIR: Standardized incidence ratios; SjS: Sjögren syndrome

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Authors’ contributions
All authors listed have contributed sufficiently to the project to be included as authors, and all those who are qualified to be authors are listed in the author byline. Authors’ contributions were the next. PBZ and MRC contributed to the conception and design; all authors contributed to the acquisition of data; PBZ, BK, and MRC contributed to the analysis and interpretation of data; BK and MRC contributed to the statistical analysis; all authors contributed in drafting the article or revising it critically for important intellectual content; all authors gave the final approval of the version published.

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References

1. Ramos-Casals M, Brito-Zerón P, Sísó-Almirall A, Bosch X. Primary Sjögren syndrome. BMJ. 2012;344:e3821.
2. Kassan SS, Thomas TL, Moutsopoulos HM, Hoover R, Kimberley RP, Budman DR, et al. Increased risk of lymphoma in sicca syndrome. Ann Intern Med. 1978;89:888–92.
3. Zintzaras E, Voulgarelis M, Moutsopoulos HM. The risk of lymphoma development in autoimmune diseases: a meta-analysis. Arch Intern Med. 2005;165:2337–44.
4. Papageorgiou A, Voulgarelis M, Tzioufas AG. Clinical picture, outcome and predictive factors of lymphoma in Sjögren syndrome. Autoimmun Rev. 2015;14:641–9.
5. Ramos-Casals M, De Vita S, Tzioufas AG. Hepatitis C virus, Sjögren’s syndrome and B-cell lymphoma: linking infection, autoimmunity and cancer. Autoimmun Rev. 2005;4:8–15.
6. Nocturne G, Mariette X. Sjögren syndrome-associated lymphomas: an update on pathogenesis and management. Br J Haematol. 2015;168:317–27.
7. Brito-Zeron P, Baldini C, Bootsoa H, Bowman SJ, Jonsson R, Mariette X, et al. Sjögren syndrome. Nat Rev Dis Prim. 2016;2:16047.
8. Theander E, Henriksson G, LJungberg O, Mandt T, Manthorpe R, Jacobsson LT. Lymphoma and other malignancies in primary Sjögren’s syndrome: a cohort study on cancer incidence and lymphoma predictors. Ann Rheum Dis. 2006;65:796–803.
9. Baimpa E, Dahabreh I, Voulgarelis M, Moutsopoulos HM. Hematologic manifestations and predictors of lymphoma development in primary Sjögren syndrome: clinical and pathophysiologic aspects. Medicine (Baltimore). 2009;88:284–93.
10. Nocturne G, Vicone A, Ng WF, Le Guern V, Hachulla E, Cornez D, et al. Rheumatoid factor and disease activity are independent predictors of lymphoma in primary Sjögren’s syndrome. Arthritis Rheumatol (Hoboken, NJ). 2016;68:977–85.
11. Weng M-Y, HuAmy Y-T, Liu M-F, Lu TH. Incidence of cancer in a nationwide population cohort of 7852 patients with primary Sjogren’s syndrome in Taiwan. Ann Rheum Dis. 2012;71:524–7.
12. Vitali C, Bombardieri S, Jonsson R, Moutsopoulos HM, Alexander EL, Carsens SE, et al. Classification criteria for Sjögren’s syndrome: a revised version of the European criteria proposed by the American-European Consensus Group. Ann Rheum Dis. 2002;61:554–8.
13. Brito-Zeron P, Kostov B, Solans R, Fraile G, Suárez-Cuervo C, Casanova S, et al. Systemic activity and mortality in primary Sjögren syndrome: predicting survival using the EULAR-SS Disease Activity Index (ESSDAI) in 1045 patients. Ann Rheum Dis. 2016;75:548–55.
14. Seror R, Theander E, Brun JG, Ramos-Casals M, Valin V, Dörner T, et al. Validation of EULAR primary Sjögren’s syndrome disease activity (ESSDAI) and patient indexes (ESSPRI). Ann Rheum Dis. 2015;74:859–66.
15. Seror R, Bootsoa H, Saraux A, Bowman SJ, Theander E, Brun JG, et al. Defining disease activity states and clinically meaningful improvement in primary Sjögren’s syndrome with EULAR primary Sjögren’s syndrome disease activity (ESSDAI) and patient-reported indexes (ESSPRI). Ann Rheum Dis. 2016;75:1–8.
16. Swedlow SH, Campo E, Pilleri SA, Harris NL, Stein H, Siebert R, et al. The 2016 revision of the World Health Organization classification of lymphoid neoplasms. Blood. 2016;127:2375–90.
17. Arber DA, Odah A, Hasserjian R, Thiele J, Borowitz MJ, Le Beau MM, et al. The 2016 revision to the World Health Organization classification of myeloid neoplasms and acute leukemia. Blood. 2016;127:2391–405.
18. Ferlay J, Steillerova-Foucher E, Lorret-Tieulent J, Rosso S, Coebergh JW, Comber H, et al. Cancer incidence and mortality patterns in Europe: estimates for 40 countries in 2012. Eur J Cancer. 2013;49:1374–403.
19. Liang Y, Yang Z, Qin B, Zhong R. Primary Sjögren’s syndrome and malignancy risk: a systematic review and meta-analysis. Ann Rheum Dis. 2014;73:1151–6.
20. Quattrocchi L, Baldini C, Bartolini E, Priori R, Carubbi F, Corazza L, et al. Anti-SSA/SSB-negative Sjögren’s syndrome shows a lower prevalence of lymphoproliferative manifestations, and a lower risk of lymphoma evolution. Autoimmun Rev. 2015;14:1019–22.
21. Zhang W, Feng S, Yan S, Zhao Y, Li M, Sun J, et al. Incidence of malignancy in primary Sjögren’s syndrome in a Chinese cohort. Rheumatology (Oxford). 2010;49:571–7.
22. Balldini C, Pepe P, Luciano N, Ferro F, Talarico R, Grossi S, et al. A clinical prediction rule for lymphoma development in primary Sjögren’s syndrome. J Rheumatol. 2012;39:804–8.
23. Johnsen SJ, Brun JG, Goransson LG, Småstuen MC, Johannesen TB, Haldorsen K, et al. Risk of non-Hodgkin’s lymphoma in primary Sjögren’s syndrome: a population-based study. Arthritis Care Res (Hoboken). 2013;65:816–25.
24. Riselada AP, Krueze AA, Blijswa JW. Clinical features distinguishing lymphoma development in primary Sjögren’s syndrome—a retrospective cohort study. Semin Arthritis Rheum. 2013;43:171–7.
25. Quartuccio L, Isola M, Baldini C, Priori R, Bartolini Bocci E, Carubbi F, et al. Biomarkers of lymphoma in Sjögren syndrome and evaluation of the lymphoma risk in prelymphomatous conditions: results of a multicenter study. J Autoimmun. 2014;45:75–90.
26. Papageorgiou A, Zigos DC, Mavragani CP, Zintzaras E, Tsoulas AG, Moutsopoulos HM, et al. Predicting the outcome of Sjögren’s syndrome-associated non-hodgkin’s lymphoma patients. PLoS One. 2015;10:e0161869.
27. Hemminki K, Liu J, Ji J, Forsti A, Sundquist J, Sundquist K. Effect of autoimmune diseases on risk and survival in female cancers. Gynecol Oncol. 2012;127:180–5.
28. Tomi A-L, Belhki R, Nocturne G, Desmoïns F, Bërge E, Pavy S, et al. Brief report: monoclonal gammopathy and risk of lymphoma and multiple myeloma in patients with primary Sjögren’s syndrome. Arthritis Rheumatol (Hoboken, NJ). 2016;68:1245–50.
29. Retamozo S, Gheitasi H, Quartuccio L, Kostov B, Corazza L, Bövé A, et al. Cryoglobulinemic vasculitis at diagnosis predicts mortality in primary Sjögren syndrome: analysis of 515 patients. Rheumatology (Oxford). 2016;55:1443–51.
30. godtberg je, seror r, miceli-richard c, benessiano j, devauchelle-pensec v, dieude p, et al. Serum levels of beta-2microglobulin and free light chains of immunoglobulins are associated with systemic disease activity in primary Sjögren’s syndrome. Data at enrollment in the prospective ASSESS cohort. PLoS One. 2013;8:1–6.
31. Holmes D. Thyroid cancer: incidence trends in the USA. Nat Rev Endocrinol. 2012;16:12312.
32. Ramos-Casals M, Brito-Zerón P, Kostov B, Sísó-Almirall A, Bosch X, Buss D, et al. Google-driven search for big data in autoimmune geoepidemiology: analysis of 394,827 patients with systemic autoimmune diseases. Autoimmun Rev. 2015;14:670–9.
33. Bartolini E, Baldini C, Schillaci G, Quartuccio L, Priori R, Carubbi F, et al. Cardiovascular disease risk burden in primary Sjögren’s syndrome: results of a population-based multicentre cohort study. J Intern Med. 2015;278:185–92.
34. Ramos-Casals M, Garcia-Carrasco M, Cervera R, Gaya J, Halperin J, Buxo L, et al. Thyroid disease in primary Sjögren syndrome. Study in a series of 160 patients. Medicine (Baltimore). 2000;79:103–8.
35. Farrell E, Heffron C, Murphy M, O’Leary G, Sheahan P. Impact of lymphocytic thyroiditis on incidence of pathological incidental thyroid carcinoma. Head Neck. 2017;39:122–7.
36. Garcia-Carrasco M, Ramos-Casals M, Cervera R, Font J. Primary Sjögren’s syndrome and lymphatic proliferation. Med Clin (Barc). 2000;114:740–6.
37. Solans-Laque R, Lopez-Hernandez A, Bosch-Gil JA, Palacios A, Campillo M, Villard-Tares M. Risk, predictors, and clinical characteristics of lymphoma development in primary Sjögren’s syndrome. Semin Arthritis Rheum. 2011;41:415–23.
38. Lazarus MN, Robinson D, Mak V, Waller H, Isenberg DA. Incidence of cancer in a cohort of patients with primary Sjögren’s syndrome. Rheumatology (Oxford). 2006;45:1012–7.
39. Linet MS, Vajdic CM, Morton LM, de Roos AJ, Skibola CF, Boffetta P, et al. Medical history, lifestyle, family history, and occupational risk factors for follicular lymphoma: the InterLymph Non-Hodgkin Lymphoma Subtypes Project. J Natl Cancer Inst Monogr. 2014;2014:426–40.
40. Engels EA, Cerhan JR, Linet MS, Cozen W, Colt JS, Davis S, et al. Immune-related conditions and immune-modulating medications as risk factors for non-Hodgkin's lymphoma: a case-control study. Am J Epidemiol. 2005;162:1153–61.

41. Ekstrom Smedby K, Vajdic CM, Falster M, Engels EA, Martinez-Maza O, Turner J, et al. Autoimmune disorders and risk of non-Hodgkin lymphoma subtypes: a pooled analysis within the InterLymph Consortium. Blood. 2008;111:4029–38.