Incidence rates of systemic lupus erythematosus in the USA: estimates from a meta-analysis of the Centers for Disease Control and Prevention national lupus registries

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ABSTRACT

Objective To estimate the annual incidence rate of SLE in the USA.

Methods A meta-analysis used sex/race/ethnicity-specific data spanning 2002–2009 from the Centers for Disease Control and Prevention network of four population-based state registries to estimate the incidence rates. SLE was defined as fulfilling the 1997 revised American College of Rheumatology classification criteria. Given heterogeneity across sites, a random effects model was employed. Applying sex/race/ethnicity-stratified rates, including data from the Indian Health Service registry, to the 2018 US Census population generated estimates of newly diagnosed SLE cases.

Results The pooled incidence rate per 100 000 person-years was 5.1 (95% CI 4.6 to 5.6), higher in females than in males (8.7 vs 1.2), and highest among black females (15.9), followed by Asian/Pacific Islander (7.6), Hispanic (6.8) and white (5.7) females. Male incidence was highest in black males (2.4), followed by Hispanic (0.9), white (0.8) and Asian/Pacific Islander (0.4) males. The American Indian/Alaska Native population had the second highest race-specific SLE estimates for females (10.4 per 100 000) and highest for males (3.8 per 100 000). In 2018, an estimated 14 263 persons (95% CI 11 563 to 17 735) were newly diagnosed with SLE in the USA.

Conclusions A network of population-based SLE registries provided estimates of SLE incidence rates and numbers diagnosed in the USA.

INTRODUCTION

The heterogeneity of clinical manifestations of SLE, lack of a singular diagnostic test, racial/ethnic disparities in SLE susceptibility and mortality, differing case definitions, heterogeneous sources for case ascertainment, possible inaccuracy of self-report, unreliability in coding in health system databases, and variable access to healthcare for high-risk populations have made SLE difficult for epidemiologists to study in the USA and resulted in widely varying epidemiological estimates.1–3
The Centers for Disease Control and Prevention (CDC) supported a network of five population-based SLE registries using standardised active surveillance methods to determine SLE prevalence and incidence in populations reflecting the broad distribution of racial/ethnic demographics in the USA. These regional registries provided overall prevalence and incidence of SLE, as well as estimates that focused on the major US demographic groups. A meta-analysis using these registries provided overall estimates of SLE prevalence and the number of individuals diagnosed with SLE in the USA in 2018. Further leveraging these registries, a meta-analysis to estimate the general and sex-specific and race/ethnicity-specific incidence rates of SLE was performed and used to provide an estimate of the number of newly diagnosed SLE cases in the USA in 2018.

METHODS

Data sources and study selection

The methodology for this paper mirrored that used in our meta-analysis of the prevalence of SLE. The CDC-supported lupus registries included two sites with large black and white populations (Georgia Lupus Registry (GLR), Fulton and DeKalb counties); Michigan Lupus Epidemiology and Surveillance Program (MILES), Washtenaw and Wayne counties); two sites with large Hispanic and Asian/Pacific Islander populations (California Lupus Surveillance Program (CLSP), San Francisco county; Manhattan Lupus Surveillance Program (MLSP), New York County); and one system to capture the American Indian/Alaska Native (AI/AN) population (Indian Health Service (IHS), including facilities in Alaska, Phoenix and Oklahoma City areas). These registries evaluated incident and prevalent cases occurring in 2002–2009 by performing active surveillance at various times between 2003 and 2015 as previously described. The case definitions for SLE incidence varied slightly by the time period evaluated in each registry, with all taking place between 2002 and 2009. The primary case definition for SLE was the 1997 revised American College of Rheumatology (ACR) classification criteria for SLE. The registries used harmonised methods including screening for potential SLE cases using the same core set of International Classification of Diseases, Ninth Revision (ICD-9) codes and a variety of case-finding sources and a consistent approach to capture the relevant clinical and demographic information and core definitions from a standardised data dictionary. Medical abstractors, who underwent routine training and quality assurance monitoring, collected the data. Denominators were based on intercensal population estimates for the respective source populations. Sex-specific and race/ethnicity-specific incidence estimates were calculated per 100,000 person-years and age-adjusted to the 2000 US standard population. Data were extracted by two authors independently (PI and HP) from published manuscripts, who agreed on all data used.

Data synthesis and analysis

A meta-analysis derived pooled incidence estimates using data from four state-based, CDC-funded registries (GLR, MILES, CLSP and MLSP) for age-standardised incidence rates and for rates stratified by sex and race/ethnicity categories, with the exception of the AI/AN population. Given that the IHS-based registry was different, focusing on one demographic population (AI/AN), it was handled separately. For the meta-analysis, heterogeneity across the four state-based sites was tested by Cochran’s Q and I² statistics. Due to significant heterogeneity, a random effects model, weighted by the population denominator for each site, was used to calculate the pooled incidence, which allowed for an underlying distribution of the effect sizes across different studies. Pooled sex-specific and race/ethnicity-specific estimates were calculated, except for the AI/AN estimate, which used the previously published female age-adjusted incidence estimates and newly calculated male age-adjusted estimates to the 2000 US standard population that were not previously age-adjusted due to the small number of cases. The MLSP rates were initially presented as combined race and ethnicity categories (eg, non-Hispanic white), but for the meta-analysis race and ethnicity rates were recalculated separately for consistency. The number of new SLE cases in the USA was estimated using the pooled age-adjusted sex-specific and race-specific incidence rates from the four states and the age-adjusted AI/AN population incidence rates from the IHS, which were extrapolated to 2018 US Census population data.

Figure 1 Meta-analysis results of the SLE incidence based on four CDC population-based registries, overall and by registry site. Overall incidence estimates for the SLE registry sites in Michigan (MI), Georgia (GA), New York (NY) and California (CA) are represented by circles, with the size of the circle corresponding to the weight of the contribution to the meta-analysis, and the diamond representing the results from the meta-analysis and the lines corresponding to 95% CI. SLE cases were defined according to the 1997 revised American College of Rheumatology criteria. CDC, Centers for Disease Control and Prevention.
The sex-specific and race-specific estimates were then summed for the total population count.

**Patient and public involvement**

Neither patients nor the public were involved in the design, or conduct, or reporting or dissemination plans of our research.

**RESULTS**

**Incidence**

The five registries contributed 1057 SLE cases fulfilling the ACR classification. The random effects model for the meta-analysis of estimates from the four state-based registries yielded an overall SLE incidence rate of 5.1 per 100,000 person-years (95% CI 4.6 to 5.6) (figure 1). The incidence among females was about seven times higher than among males (8.7 vs 1.2) (table 1). From the race-specific and ethnicity-specific pooled estimates from the four state-specific registries, the incidence rate was highest among black females (15.9, 95% CI 12.5 to 20.3), followed by Asian/Pacific Islander (7.6, 95% CI 5.5 to 10.4), Hispanic (6.8, 95% CI 6.2 to 7.6) and white females (5.7, 95% CI 4.9 to 6.7) (figure 2A and table 1). Among males, the incidence rate was highest among black males (2.4, 95% CI 1.8 to 3.0), followed by Hispanic (0.9, 95% CI 0.4 to 1.9), white (0.8, 95% CI 0.6 to 1.1) and Asian/Pacific Islander males (0.4, 95% CI 0.2 to 0.6) (figure 2B and table 1). The age-adjusted incidence rate estimates for the AI/AN population from the IHS registry (which were not included in the pooled meta-analysis estimates from the four state-based registries) were the second highest of all the race/ethnicity categories for females (10.4, 95% CI 6.6 to 14.6) and highest for males (3.8, 95% CI 1.6 to 7.8) (figure 2A,B and table 1).

**Number of newly diagnosed SLE cases in the USA**

Applying our sex-specific and race-specific incidence estimates to the corresponding population denominators from 2018 Census data, we estimated that 14263 (95% CI 11563 to 17735; 12560 females and 1703 males) persons in the USA were newly diagnosed with SLE in 2018 (table 1).

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**Table 1** Estimated number of new persons diagnosed with SLE in the USA in 2018

<table>
<thead>
<tr>
<th>Race/ethnicity (number of sites in the analysis)</th>
<th>Female</th>
<th></th>
<th></th>
<th></th>
<th>Male</th>
</tr>
</thead>
<tbody>
<tr>
<td>Race</td>
<td>SLE incidence* per 100 000 (95% CI)</td>
<td>Population denominator</td>
<td>Estimated SLE cases in the USA, n (95% CI)</td>
<td>SLE incidence* per 100 000 (95% CI)</td>
<td>Population denominator</td>
</tr>
<tr>
<td>Black (4)</td>
<td>15.9 (12.5 to 20.3)</td>
<td>24 880 722</td>
<td>3956 (3110 to 5051)</td>
<td>2.4 (1.8 to 3.0)</td>
<td>22 961 129</td>
</tr>
<tr>
<td>White (4)</td>
<td>5.7 (4.9 to 6.7)</td>
<td>130 137 989</td>
<td>7418 (6377 to 8719)</td>
<td>0.8 (0.6 to 1.1)</td>
<td>127 942 583</td>
</tr>
<tr>
<td>Asian/PI (2)</td>
<td>7.6 (5.5 to 10.4)</td>
<td>12 544 896</td>
<td>953 (690 to 1305)</td>
<td>0.4 (0.2 to 0.6)</td>
<td>11 660 533</td>
</tr>
<tr>
<td>AI/AN (1)</td>
<td>10.4 (6.6 to 14.6)</td>
<td>2 238 966</td>
<td>233 (148 to 327)</td>
<td>3.8 (1.6 to 7.8)</td>
<td>2 134 870</td>
</tr>
<tr>
<td>Total†</td>
<td>8.7 (8.1 to 9.4)</td>
<td>169 802 573</td>
<td>12 560 (10 325 to 15 402)</td>
<td>1.2 (0.9 to 1.6)</td>
<td>164 699 115</td>
</tr>
<tr>
<td>Ethnicity</td>
<td>Hispanic‡ (2)</td>
<td>6.8 (6.2 to 7.6)</td>
<td>30 689 083</td>
<td>2087 (1903 to 2332)</td>
<td>0.9 (0.4 to 1.9)</td>
</tr>
</tbody>
</table>

SLE cases were defined according to the 1997 revised American College of Rheumatology criteria.

*Estimates for black and white persons are based on pooled estimates from the four state-based registries; Asian/PI and Hispanics are based on pooled estimates from California and New York; AI/AN estimates are based on the Indian Health Service registry.

†The pooled "total" incidence estimate includes black, white and Asian/PI. Since the AI/AN incidence was based on one registry and was significantly higher, it was not included in the pooled incidence per 100,000.

‡Hispanic ethnicity is not mutually exclusive from the race categories, that is, all Hispanic persons are included in one of the race categories. Thus, the pooled estimates do not incorporate the Hispanic rates since that would lead to duplicate counting. Estimates for Hispanics are based on pooled estimates from California and New York.

AI/AN, American Indian/Alaska Native; Asian/PI, Asian/Pacific Islander.
DISCUSSION

The overall SLE incidence in the USA was estimated to be 5.1 per 100,000 person-years (95% CI 4.6 to 5.6) during calendar years 2002–2009. Incidence rates were approximately seven times higher in females compared with males and highest among black and AI/AN females. Extrapolating sex-specific and race-specific estimates to 2018 US Census data, we estimated that 14,263 (95% CI 11,563 to 17,735) persons (12,560 and 1,703 females and males, respectively) in the USA were newly diagnosed with SLE fulfilling ACR classification criteria.

Our analyses have several limitations as previously described.3–8 There were minor differences in the ICD-9 criteria and case-finding sources used to identify possible cases by the different registries. Capture-recapture analyses conducted by the state-based registries demonstrated small numbers of missing cases meeting the ACR criteria.3–6 Undiagnosed cases were not counted, nor were ‘incomplete’ lupus, drug-induced lupus or primary cutaneous lupus.15–16 Race and ethnicity were abstracted from medical records, which may not accurately represent the patient’s own racial or ethnic identification. The different races and Hispanic ethnicity encompass heterogeneous subgroups, and SLE rates among these subgroups may differ. The AI/AN incidence rates were based only on the IHS registry7 and were not used for the meta-analysis pooled incidence calculations given it focused on one demographic population. However, this registry included three geographical regions with different population characteristics to improve the generalisability of results, and these results were used for the national estimate calculation of new SLE cases. Other SLE case definitions used by the five registries3–7 resulted in slightly higher estimates in most instances, although greater sensitivity may have occurred with lower specificity. Finally, our incidence estimates from 2002 to 2004 and from 2007 to 2009 were applied to the 2018 Census population, which
provides a companion to our 2018 prevalence estimate; both might be slightly affected if lupus incidence or prevalence changed significantly during that period.

Despite these limitations, our analysis has several strengths. The case-finding was population-based and thus likely captured a wider spectrum of SLE than previous studies. The public health surveillance exemption additionally facilitated broad case-finding and data collection. Cases were also validated through standardised and quality-controlled abstracting, with rigid reviews of all available medical records by trained abstractors. Additionally, we used harmonised methods and data dictionaries and defined cases using the 1997 revised ACR SLE classification criteria. The CDC registries included SLE cases from diverse populations across the country with substantial representation of the major racial and ethnic groups found in the USA, allowing for a better understanding of SLE by race/ethnicity as well as among understudied males.

Most importantly, this is the first study to use representative registry data to estimate the incidence of SLE in the USA. We were able to use information from a large, coordinated network of population-based active surveillance registries to estimate an SLE incidence of 5.1 per 100,000 person-years in the USA, corresponding to 14,263 newly diagnosed persons in 2018. Both likely represent a lower bound for SLE incidence but provide new information to inform future research and policy efforts addressing the burden of this rare disease.

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Competing interests None declared.

Patient and public involvement Patients and/or the public were not involved in the design, or conduct, or reporting, or dissemination plans of this research.

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