Systemic lupus erythematosus on the Caribbean island of Curacao: an epidemiological investigation

J C Nossent

Abstract
To determine the incidence, prevalence, and outcome of systemic lupus erythematosus (SLE) in a well delineated black population in the Caribbean basin data were collected on the disease course of all patients with definite SLE seen during a 10 year period (1980–9) using three different sources of information (hospital records, private practice records, and death certificates). Ninety four patients were identified giving an average annual incidence rate of 4.6/100 000 (95% confidence interval (CI) 1.6 to 8.8), which showed little variation during the study period. Twenty five patients (27%) died during the study period, giving a point prevalence of 47/100 000 (CI 34.1 to 51.1) in 1990. In women aged 15–44 years the annual incidence (12/100 000; CI 5.3 to 18.9) was highest, whereas in women aged 44–65 years the 1990 point prevalence rate (one in 526; CI 469 to 625) was highest. Annual mortality was 1.7/100 000 (CI 0.0 to 4.2) with a female to male ratio of 5.3. Renal disease was the most common complication, occurring in 73 (78%) patients. Thus the transatlantic movement from an area with a (presumably) low prevalence of SLE (Central Africa) has been accompanied by an increase in the prevalence of SLE in the black population of Curacao, indicating that environmental factors may prevail over genetic factors in the expression of this disease.


Reports on the incidence and prevalence of systemic lupus erythematosus (SLE) show a considerable variation between countries and sometimes within one country.1–9 Although a considerable part of these variations is probably due to differences in patient selection, environmental and genetic factors are also believed to influence disease expression.1 4 Ethnicity is considered to be one of the predominant determinants, as black subjects in the USA have a threefold increased incidence of SLE compared with white subjects,3 5 whereas in Chinese subjects the disease is even more common.6 7 On the other hand, SLE is still considered to be a rare disease (although sound epidemiological data from the area are not available) in Central Africa, which is the original homeland of many black Americans.8 8 At the same time as the black population was introduced into the USA, the island of Curacao in the Caribbean basin was also populated by black people from Central Africa. They now make up more than 95% of the population on the island. This study reports the epidemiology of SLE in this well delineated black population.

Patients and methods
PATIENTS
The island of Curacao is an autonomous part of the Dutch Kingdom, located 60 km off the coast of Venezuela with a (sub) tropical climate. Its population is estimated to be 146 500, of which less than 5% are white and which has a migratory rate of about 1%.10 There is one major 450 bed teaching hospital which serves the whole island. To ensure a maximum degree of completeness, information on all patients with SLE seen during the period 1980–9 was gathered from three different sources. Firstly, all medical records of patients discharged from the hospital during the study period with a diagnosis of SLE (ICD code 710.0) were reviewed; with this method 85 patients were identified. Secondly, all specialists in internal medicine (six) and dermatology (two) (there is no rheumatologist on the island) were asked in 1989 to provide information on all patients with lupus who had been in their care during that period; although most of the consultants did not keep an official diagnostic index, this provided another eight patients not identified by the hospital registry. Thirdly, all death certificates in the Public Health Department from that period containing a diagnosis of SLE were reviewed and medical records of the patients were retrieved; this provided one patient not identified by either of the other two sources.

METHODS
All patient records were reviewed by one investigator using a predefined data extraction form. Only patients fulfilling four or more American Rheumatism Association (ARA) criteria for the classification of SLE11 were entered into the study. Information was gathered on demographic features, (cumulative) ARA criteria, initial disease activity at the time of diagnosis using the validated SLEDAI score12 and subsequent disease course. Period prevalence rates were calculated using all patients with definite SLE seen during the study period, whereas incidence rates were calculated using only true incident cases—that is, those diagnosed during the 10 year study period. The time at which a patient fulfilled at least four ARA criteria was chosen as the time of diagnosis. For calculations of age specific rates we used the age distribution classes as given in the report by

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Siegel et al. Table 1 gives the breakdown figures for the general population of Curaçao, together with a correction factor that allows comparison of these rates with a European population which has a rather different age distribution.

Table 1  Epidemiological data for the population of Curaçao according to gender and age classes used in this study

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Men</th>
<th>Women</th>
<th>Total</th>
<th>Correction factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;14</td>
<td>22 000</td>
<td>21 500</td>
<td>43 500</td>
<td>0.7</td>
</tr>
<tr>
<td>15–44</td>
<td>34 200</td>
<td>37 900</td>
<td>72 100</td>
<td>0.91</td>
</tr>
<tr>
<td>45–64</td>
<td>10 050</td>
<td>11 250</td>
<td>21 300</td>
<td>1.45</td>
</tr>
<tr>
<td>&gt;65</td>
<td>4000</td>
<td>5600</td>
<td>9600</td>
<td>1.95</td>
</tr>
<tr>
<td>All</td>
<td>70 250</td>
<td>76 250</td>
<td>146 500</td>
<td></td>
</tr>
</tbody>
</table>

*This allows calculation of rates corrected for a European population, which has a different age group distribution with more elderly subjects.*

Results

INCIDENCE AND PREVALENCE

A total of 94 patients were identified during the 10 year period. All patients were of African descent; table 2 summarises their demographic features. There was no significant difference in these features between true incident cases and previously (before 1980) diagnosed cases, except for the follow up period (as expected). Twenty five patients died during this period, giving an overall prevalence rate for SLE on 1 January 1990 of 47/100 000 (CI 34.1 to 51.1) at risk. Figure 1 shows the course of prevalence rates for the study period. Analysis of age specific prevalence rates by gender at the end of the study period revealed a maximum prevalence rate of 186.6/100 000 (or one in 526) in women aged 45–64 (table 3). Thirty six eight true incident cases were identified during the 10 year study, giving a mean annual incidence rate of 4.6/100 000 at risk (CI 0.4 to 8.8). Annual incidence rates for the whole study period are also shown in figure 1 and show a relatively constant level of newly diagnosed cases over the 10 year period. Table 4 shows the age specific incidence rates by gender during that period. Considerable variations were found, with the lowest rate (0.57/100 000) for men aged 15–44 years and the highest rate (12.1/100 000) in women in the same age group.

MORTALITY

Twenty five patients (27%) died during the study period. There was no difference in the

Statistical analysis of data was performed with the Mann-Whitney U test for differences between continuous variables and with the χ² test for contingency tables. Ninety five per cent confidence intervals (CI) were calculated using the approximation formula from the standard error, whereas survival probabilities were estimated from Kaplan Meier curves. Figures represent mean values unless otherwise stated; p<0.05 was considered statistically significant.

Figure 1  Crude annual incidence and (period) prevalence rates for systemic lupus erythematosus in Curaçao between 1980 and 1989. Bars indicate numbers of patients.

Table 2  Demographic features of 94 black patients with systemic lupus erythematosus on Curaçao

<table>
<thead>
<tr>
<th>Demographic feature</th>
<th>No men/women (ratio)</th>
<th>83/11 (7.5)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Symptomatic period before diagnosis (years)</td>
<td>1-7</td>
<td>(0-11.8)</td>
</tr>
<tr>
<td>Mean (SD) age at diagnosis (years)</td>
<td>34.6 (14.8)</td>
<td></td>
</tr>
<tr>
<td>Mean (SD) SLEDAI score at diagnosis</td>
<td>13.3 (6.8)</td>
<td></td>
</tr>
<tr>
<td>Mean (SD) follow up period (months)</td>
<td>70.7 (69)</td>
<td></td>
</tr>
<tr>
<td>No (%) of deaths</td>
<td>25 (27)</td>
<td></td>
</tr>
</tbody>
</table>

Table 3  Age specific period prevalence rate of systemic lupus erythematosus in Curaçao by gender, 1980–90

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Men</th>
<th>Women</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>No of patients</td>
<td>Prevalence rate (CI)*</td>
<td>No of patients</td>
<td>Prevalence rate (CI)*</td>
</tr>
<tr>
<td>&lt;14</td>
<td>1 (4.5:0.3 to 8.7)</td>
<td>0</td>
<td>—</td>
</tr>
<tr>
<td>15–44</td>
<td>0 (—)</td>
<td>38 (0.5:0.3 to 0.7)</td>
<td>21</td>
</tr>
<tr>
<td>45–64</td>
<td>1 (10.0:5.6 to 16.2)</td>
<td>4</td>
<td>18.6 (15.9:21.3)</td>
</tr>
<tr>
<td>&gt;65</td>
<td>4 (8.5:2.8 to 16.2)</td>
<td>63</td>
<td>83.8 (85.8:101.8)</td>
</tr>
</tbody>
</table>

*Indicates rate per 100 000 subjects; CI denotes 95% confidence interval.

Table 4  Age specific incidence rates of systemic lupus erythematosus by gender in Curaçao, 1980–9

<table>
<thead>
<tr>
<th>Age group at diagnosis (years)</th>
<th>Men</th>
<th>Women</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>No of patients</td>
<td>Incidence rate (CI)*</td>
<td>No of patients</td>
<td>Incidence rate (CI)*</td>
</tr>
<tr>
<td>&lt;14</td>
<td>2 (0.91:0.1 to 2.8)</td>
<td>2</td>
<td>0.93 (0.1:0.4 to 2.8)</td>
</tr>
<tr>
<td>15–44</td>
<td>2 (0.57:0.9 to 2.1)</td>
<td>9</td>
<td>0.8 (0.2:1.7)</td>
</tr>
<tr>
<td>45–64</td>
<td>2 (0.06:0.6 to 9.4)</td>
<td>3</td>
<td>0.59 (0.08:9.8)</td>
</tr>
<tr>
<td>&gt;65</td>
<td>2 (1.13:0.9 to 3.1)</td>
<td>60</td>
<td>7.86 (0.3:13.2)</td>
</tr>
</tbody>
</table>

*Average annual rate per 100 000 subjects during study period; CI denotes 95% confidence interval.
proportion of deaths in true incident cases compared with patients diagnosed before 1980. The mean (SD) age at death was 42.0 (18.1) years, whereas the median (range) disease duration at death was 80.5 (9-214) months. Mean (SD) disease activity (by SLEDAI score) at the time of death was 14.4 (9.1; median 13.5). Table 5 gives the age specific annual death rates by gender. Mortality rates increased steadily with age in men, whereas in women this increase was also seen, but at a persistently higher rate (nearly six times higher in the age group 15–45 years). The mean annual death rate for all patients during the study period was 1.7/100 000.

### Survival

Life table analysis showed a survival probability for the 68 true incident cases in this study of 90.5%, 60.1, and 45.7% at one, five, and 10 years with a decreased survival probability for men (p<0.01 by log rank test), although age group was not a significant prognostic factor.

### Morbidity

Most patients had a prolonged symptomatic period before the diagnosis of SLE was made (table 1). Figure 2 gives the occurrence of initial and cumulative disease manifestations. The occurrence of overall mucocutaneous lesions was low at 20%, whereas clinical renal disease (defined as persistent proteinuria or cellular casts, or both, in the absence of other disease) was found in 73 (78%) patients (histological confirmation of lupus nephritis was obtained in only 11 patients). Patients with clinical lupus nephritis did not differ significantly from the other patients with respect to age, gender, symptomatic period before diagnosis or follow up, but only in patients with lupus nephritis did death occur (30% vs 0% in the patients without lupus nephritis). Renal failure requiring chronic renal replacement treatment developed in 10 (14%) of the patients with renal disease.

### Discussion

This is the first study to present epidemiologic data on SLE in a well delineated, black Caribbean population. Only the report by Wilson and Hughes in 1979 on rheumatic diseases in Jamaica has presented epidemiologic data on SLE from this area; they reported the number of patients with SLE seen in an outpatient clinic during a three year period. Rough estimates indicated that the prevalence of SLE in Jamaica at that time was about 5–17/100 000, depending on the estimated size of the population served by their clinic. Our data indicate a 1980 prevalence rate of 22/100 000 in Curacao. The exclusion of private patients from the study in Jamaica may account for the difference. In the period studied the highest prevalence of SLE in Curacao was found in women aged 15–64 years; for this whole age group the rate indicates that one in 689 black women have SLE (for the age group 15–44, one in 970; for the age group 45–64, one in 526) as opposed to men of the same age, in whom the disease affects one in 14 285. This prevalence rate is lower than that in black women, but similar to that in white women in the USA as reported by Fessel. This variance in prevalence and the fact that SLE seems to be such a rare disease in Central Africa (although this statement has been treated with caution because of the limited availability of diagnostic tools and the lack of good epidemiologic data from this area) indicates that ethnicity can be superseded by environmental factors as a major determinant for the expression of SLE. Also, the annual incidence rate in this black population (4.6/100 000) is almost identical to that found in a study in south Sweden within a study group with a entirely different genotypic make up, although it is lower than that in a group of black subjects in Baltimore. The environmental factors affecting these results are unknown; one theory relies on the immuno-
suppressive effect of ubiquitous parasitic infections to explain the low prevalence of SLE in Africa,6 but this finding has not been expanded. Others assume that various factors (toxic, psychological) in the Western world may predispose for the development of SLE,7 making SLE a disease of prosperity. Siegel, in his landmark study, found a lower prevalence of lupus in black and white subjects in rural areas compared with those in a large city, but definitive proof is lacking. The possibility of not determining all subjects affected was a major concern in this study, as the most critical part was the non-availability of diagnostic indices from the cooperating consultants, who thus had to rely on memory to identify patients. As SLE is a chronic disease in which exacerbations may occur at any time in the disease course, most patients with lupus remain under prolonged control (although the frequency of visits may vary) and, as the island population has a low migratory rate,10 it seems unlikely that the number of patients who may have been missed by the methods used (because they were discharged or had left the island) would have significantly altered the given rates. Also, Petri et al11 have found an average admission frequency of 4.4 over a five year period for patients with lupus in an inner city population, whereas Pistiner et al showed that more than half of their new (private) patients with lupus in the last decade needed at least one admission.12 Thus the admission rate for patients with SLE in the first years of their disease is high and socioeconomic factors seem to influence this rate. As the per capita annual income in Curacao is about US$5000,10 the high frequency of admitted patients in this study is not remarkable and lessens the probability of cases being under estimated. The good overlap of patient identification (only 10% of patients identified by one source only) also supports this view.

The occurrence of specific clinical manifestations (initially and overall) in SLE of our patients and the sex distribution was similar to that in reports from Jamaica,14 Sweden,15 the USA,16 17 and The Netherlands.20 As this study was based on patient record findings, however, it is conceivable that skin lesions (erythema may be hard to distinguish in black patients) and mouth ulcers may have been underestimated as these diagnoses are easily missed during routine investigations. Lupus nephritis was a common complication, present at diagnosis in 48% and finally in 78% of our patients. As histological confirmation was not always obtained these results may be an overestimation, but it has been reported that black patients with SLE tend to have more common and more serious renal disease.21 22 Furthermore, the clinical manifestations of lupus nephritis have been shown to be reliable indicators for the presence or absence of renal disease,23 though they do not accurately reflect its severity.24 As renal disease is still the principal prognostic factor in lupus, this high percentage of renal disease may explain the relatively poor survival in our true incident patients. Also, the median initial SLEDAI scores (table 1) indicated a high disease activity in our patients at diagnosis, which is another poor prognostic sign as shown by Ginzler et al25 and Chang et al.26 The mortality rate in male patients in this study was comparable with the rate (0.8) reported by Siegel et al2 and showed a steady increase with age, indicating that lupus did not alter the life expectancy of male patients. In female patients, however, the mortality rate was continuously higher than in male patients with a peak rate (5.1) in women aged 45–64 years and lower rates (3.16–3.5) in younger and older female patients. This indicates a significant negative influence of lupus on life expectancy overall, but foremost in the young female patients, as was also found in the study in Jefferson, USA.2 Some caution must be taken in accepting this interpretation as the number of (especially male) patients was small.

In summary, this study indicates that SLE is a prevalent disease in black Carribean women aged between 15 and 64 years, with a steady rate of incidence over the last decade. Renal disease is the most common complication of lupus in these patients, and is associated with a decreased life expectancy in young women.

10 Central Bureau Statistiek van Nederlandse Antillen, Yearbook, 1989.


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