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Clinical associations of the metabolic syndrome in systemic lupus erythematosus: data from an international inception cohort

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ABSTRACT

Background The metabolic syndrome (MetS) may contribute to increased cardiovascular risk in systemic lupus erythematosus (SLE). We aimed to examine the association of demographic factors, lupus phenotype and therapy exposure with the presence of MetS.

Methods The Systemic Lupus International Collaborating Clinics Registry for Atherosclerosis Inception cohort enrolled recently diagnosed (<15 months) SLE patients from 30 centres across 11 countries from 2000. Clinical, laboratory and therapeutic data were collected according to a standardised protocol. MetS was defined according to the 2009 consensus statement from the International Diabetes Federation. Univariate and backward stepwise multivariate logistic regression were used to assess the relationship of individual variables with MetS.

Results We studied 1686 patients, of whom 1494 (86.6%) had sufficient data to determine their MetS status. The mean (SD) age at enrolment and disease duration was 35.2 years (13.4) and 24.1 weeks (18.0), respectively. MetS was present at the enrolment visit in 239 (16%). In backward stepwise multivariable regression analysis, higher daily average prednisolone dose (mg) (OR 1.02, 95% CI 1.00 to 1.03), older age (years) (OR 1.04, 95% CI 1.03 to 1.06), Korean (OR 6.33, 95% CI 3.68 to 10.86) and Hispanic (OR 6.2, 95% CI 3.78 to 10.12) ethnicity, current renal disease (OR 1.79, 95% CI 1.14 to 2.80) and immunosuppressant use (OR 1.81, 95% CI 1.18 to 2.78) were associated with MetS.

Conclusions Renal lupus, higher corticosteroid doses, Korean and Hispanic ethnicity are associated with MetS in SLE patients. Balancing disease control and minimising corticosteroid exposure should therefore be at the forefront of personalised treatment decisions in SLE patients.
The Systemic Lupus International Collaborating Clinics (SLICC) group has developed an international registry of patients with newly diagnosed SLE to facilitate prospective, longitudinal studies of risk factors for the development of atherosclerosis in SLE. The aim of the present cross-sectional study was to investigate the prevalence of, and factors associated with, MetS in patients with SLE at enrolment into the SLICC inception cohort.

METHODS
SLICC registry for atherosclerosis
SLICC comprises 30 centres from 11 countries in North America, Europe and Asia. An inception cohort was recruited between 2000 and 2009 to investigate risk factors for atherosclerosis in SLE. Data were submitted to the coordinating centre at the University of Toronto at enrolment, and patients were reviewed annually in their local centre. Laboratory tests (fasting or non-fasting) necessary to evaluate disease activity, CHD risk factors, and to define MetS were performed locally. The study was approved by the University Health Network Research Institute, Research Ethics Committee, Toronto, Canada and by the institutional research ethics boards of participating centres in accordance with the Declaration of Helsinki’s guidelines for research in humans.

Patients
Patients were enrolled when four or more American College of Rheumatology (ACR) classification criteria for SLE were recognised. All patients were enrolled within 15 months of the date of their diagnosis. There were no specific exclusion criteria other than failing to meet four or more ACR criteria and being more than 15 months since diagnosis. Clinical and laboratory features, including CHD risk factors and therapeutic exposures, were recorded according to a standard protocol. SLE disease activity and damage were assessed using the SLE disease activity index (SLEDAI-2K) and the SLICC/ACR damage index (SLICC-DI), respectively. Active nephritis was defined as: haematuria greater than five red blood cells/high power field, excluding other causes; pyuria greater than five white blood cells/high power field, excluding infection; new/recent increase of more than 500 mg/24 h protein; casts including granular or red blood cells; or consistent renal biopsy. Nephrotic syndrome was defined as proteinuria greater than 3 g/24 h, oedema and increased blood pressure (BP). All patients provided written informed consent.

Definition of MetS
MetS was defined according to the 2009 definition in the joint interim statement from the International Diabetes Federation Task Force on Epidemiology and Prevention; National Heart, Lung, and Blood Institute; American Heart Association; World Heart Federation; International Atherosclerosis Society and International Association for the Study of Obesity. This harmonising statement requires three or more of the following five criteria to be present: (1) elevated waist circumference (MetS WC) using population/country-specific thresholds; (2) elevated triglycerides (MetS TG) of 1.7 mmol/l or greater (≥150 mg/dl) or drug therapy for hypertriglyceridaemia; (3) reduced high-density lipoprotein (HDL) cholesterol (MetS HDL) less than 1.3 mmol/l (<50 mg/dl) in women and less than 1.0 in men or drug therapy for reduced HDL-cholesterol; (4) elevated BP (MetS BP) of 150/85 mm Hg or greater or drug therapy for hypertension; and (5) elevated fasting glucose (MetS Glu) of 5.6 mmol/l or greater (≥100 mg/dl) or drug therapy for hyperglycaemia.

Statistical analysis
SLE factors implicated in MetS development in SLE were defined a priori. These represented inflammatory disease activity (SLEDAI-2K), disease phenotype including active renal disease, low complement or high anti-double-stranded DNA antibodies and therapeutic exposures including several measures of corticosteroid exposure. All corticosteroid doses were converted to milligrams (mg) of prednisolone equivalent. The cumulative oral prednisolone dose received before enrolment (g) was calculated for each individual. A comparison of continuous data was performed using Wilcoxon’s rank sum test, and of categorical data using the χ² test. Univariate logistic regression was used to assess the relationship between the presence of MetS at enrolment into the SLICC registry for atherosclerosis (RAS) and individual variables. Results presented are adjusted OR and 95% CI. Analysis was adjusted for age, ethnicity and gender. Those factors associated with MetS on univariate analysis (p<0.2) were entered into a multivariable model. Backward stepwise multivariate logistic regression was performed with significance set at 5%, resulting in a final model. All statistical analyses were performed using STATA 10.0.

RESULTS
Patients
By 2009, 1686 patients were enrolled into SLICC–RAS. Sufficient data were available to define the presence or absence of MetS in 1494 (88.6%) patients, of whom 1356 (59.4%) were women. The mean (SD) age at enrolment into the study was 35.2 years (13.4) and the mean (SD) disease duration was 24.1 weeks (18.0). There was a wide ethnic variation, reflecting the participating centres; 660 (44.2%) were Caucasian, 240 (16.1%) were Hispanic, 228 (15.3%) were black African, African-American or Afro-Caribbean and 303 (20.3%) were south-east Asian. At enrolment the mean (SD) SLEDAI was 5.5 (5.4) and 287 (19.5%) had very active disease with a SLEDAI of 10 or greater (table 1). There were 192 (11.4%) patients with insufficient data to determine the MetS status. This group had similar demographics and SLICC-DI to the rest of the cohort. However, they had a lower mean SLEDAI, a lower prevalence of renal disease, less immunosuppressant use and were more likely to be receiving antimarial therapies. This subset also had a lower frequency of corticosteroid exposure (see supplementary table S1, available online only).

Prevalence of MetS
MetS was present in 239/1494 (16%) patients. The individual MetS criteria met were: MetS WC (686/1354, 48.4%), MetS BP (686/1491, 46%), MetS TG (619/1342, 46.1%), MetS HDL (486/822, 59.1%) and MetS Glu (271/1344, 20.2%). MetS was more common in men than women (22.2% vs 15.2%; p=0.03) and those with MetS were older than those without (mean (SD) age 36.9 years (13.3) vs 34.9 years (14.7); p<0.04). Patients of Hispanic and Korean ethnicity had the highest prevalence of MetS, compared to the rest of the cohort (31.3% and 50.1% vs 10.5%; p<0.0001).

Factors associated with MetS at enrolment
In an age, ethnicity and gender-adjusted analysis we assessed the strength of the relationship between the presence of MetS and our predefined variables related to inflammation, disease

Clinical and epidemiological research

To test whether the effect of corticosteroids was dose depend-ent, the multivariable model was run using only current oral cor-

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Characteristics of patients at enrolment into SLICC–RAS</th>
</tr>
</thead>
<tbody>
<tr>
<td>No of patients</td>
<td>1494</td>
</tr>
<tr>
<td>Age, years (mean (SD))</td>
<td>35.2 (13.4)</td>
</tr>
<tr>
<td>Gender (%)</td>
<td>1336 (89.4) Female 158 (10.6) Male</td>
</tr>
<tr>
<td>Ethnicity (%)</td>
<td>660/1492 (44.2) Caucasian 228/1492 (15.3) Black African/Afro-Caribbean 303/1492 (20.3) SE Asian 240/1492 (16.1) Hispanic 61/1492 (4.1) Other</td>
</tr>
<tr>
<td>Region (%)</td>
<td>358/1477 (24.2) Canada 194/1477 (13.1) Mexico 374/1477 (25.3) USA 168/1477 (11.4) Asia 383/1477 (25.3) Europe</td>
</tr>
<tr>
<td>CHD risk factors (mean (SD))</td>
<td>BP systolic, mm Hg 119.5 (16.8) BP diastolic, mm Hg 75.3 (11.0) On AHT medication, % 435 (29.1) Total cholesterol, mmol/l 4.93 (1.49) Triglyceride, mmol/l 1.79 (1.19) HDL-cholesterol, mmol/l 1.39 (0.60) On lipid-lowering medication, % 171 (11.5) Glucose, mmol/l 5.03 (1.63) Diabetes, % 50 (3.4) Smoker current, % 225 (15.1) Premenopausal, % 1244 (83.3) BMI 25.1 (5.9) WC, cm 82.9 (14.0) 5-Year % Framingham risk Women 0.57 Men 5.03</td>
</tr>
<tr>
<td>Disease duration, weeks (mean SD)</td>
<td>24.1 (18.0)</td>
</tr>
<tr>
<td>SLEDAI (mean SD)</td>
<td>5.5 (5.4)</td>
</tr>
<tr>
<td>SLICC/ACR-DI=0</td>
<td>528 (81.9%)</td>
</tr>
<tr>
<td>Disease phenotype (%)</td>
<td>Active renal disease 314 (22.9) Anti-dsDNA positive 541/1347 (40.2) Low complement 519/1349 (38.5) Thrombocytopenia 44/1313 (3.4)</td>
</tr>
<tr>
<td>Oral CS use (median (IQR))</td>
<td>1043 (69.8) Average CS dose, mg 20 (10, 30) Highest CS dose, mg 40 (20, 60) Cumulative CS dose, g 2.6 (1.1, 5.0)</td>
</tr>
<tr>
<td>Pulse IV CS (%)</td>
<td>70/1423 (4.9)</td>
</tr>
<tr>
<td>Immunosuppressant use (%)</td>
<td>Azathioprine 599/1491 (31.0) Methotrexate 262 (43.7) Mycophenolate mofetil 104 (17.4) IV cyclophosphamide 98 (16.4) 95 (15.9) Ciclosporin 21 (3.5) Other 19 (3.2) Antimalarial use (%) 971 (65.0)</td>
</tr>
</tbody>
</table>

ACR, American College of Rheumatology; AHT, antihypertensive; BMI, body mass index; BP, blood pressure; CHD, coronary heart disease; CS, corticosteroid; DI, damage index; HDL, high-density lipoprotein; IV, intravenous; RAS, Registry for Atherosclerosis; SLEDAI, systemic lupus erythematosus disease activity index; SLICC, Systemic Lupus International Collaborating Clinics; WC, waist circumference.

phenotype and therapeutic exposure (table 2). SLE features associated with MetS included active renal lupus (OR 2.87, 95% CI 2.05 to 4.02), SLEDAI-2K greater than 10 (OR 1.73, 95% CI 1.22 to 2.44) and thrombocytopenia (OR 2.10, 95% CI 1.05 to 4.29). Several corticosteroid variables showed an association, including current oral (OR 1.53, 95% CI 1.05 to 2.25) and past intravenous (OR 3.22, 95% CI 1.55 to 7.68) corticosteroid use. SLE patients with MetS received oral corticosteroids at higher average daily doses and with higher cumulative and peak doses compared to those without MetS. The use of immunosuppressive agents was also associated with MetS (OR 2.21, 95% CI 1.65 to 3.00), and a negative association with antimalarial use was observed (OR 0.51, 95% CI 0.38 to 0.67).

In a backward stepwise multivariable regression analysis, factors independently associated with MetS were higher average daily oral prednisolone dose (mg) (OR 1.02, 95% CI 1.00 to 1.03), older age at study entry (years) (OR 1.04, 95% CI 1.03, 1.06), Korean (OR 6.35, 95% CI 3.68 to 10.86) and Hispanic (OR 6.2, 95% CI 3.78 to 10.12) ethnicity, active renal lupus (OR 1.79, 95% CI 1.14 to 2.80) and immunosuppressant use (OR 1.51, 95% CI 1.18 to 2.78). These remained unchanged when overlapping variables (such as proteinuria and active renal disease) were excluded. When active renal involvement was removed from the multivariable model in favour of the renal components scored on SLEDAI, the presence of haematuria became significant (OR 1.70, 95% CI 1.03 to 2.82) (with other variables in the final model unchanged).

Exploratory analyses

We further examined the two highest risk ethnicities to explore potential differences in SLE features or therapeutic exposures that may influence the presence of MetS. One hundred and sixty-five of 169 (97.6%) patients of Korean ethnicity resided in South Korea; 192 of 240 (80%) patients of Hispanic ethnicity resided in Mexico and 16.3% in the USA. Korean and Hispanic patients demonstrated distinct and contrasting MetS phenotypes compared to each other and the rest of the cohort (table 3). Patients of Korean ethnicity had a lower prevalence of central obesity (MetS WC 20.1% vs 51.3%; p<0.0001) and lower mean (SD) body mass index (BMI) (21.6 (4.3) vs 25.8 (6.1) kg/m2; p=<0.0001), but a significantly increased prevalence of hyperglycaemia and dyslipidaemia. Korean patients also had more active laboratory features in the SLEDAI, such as positive anti-dsDNA antibodies (66% vs 36.0%; p<0.0001), hypercomplementaemia (75.2% vs 31.1%; p<0.0001) and thromboctopenia (11.2% vs 2.7%; p<0.0001). Oral corticosteroid use in the Korean cohort was almost universal (95.3%), although the average daily, peak and cumulative doses were similar or lower than the rest of the cohort.

In the Hispanic cohort, MetS was contributed to by significantly more dyslipidaemia (MetS TG 64.3% vs 40.3%; p<0.0001), but a similar prevalence of central obesity. Hispanic patients also had more active renal disease at enrolment (40.4% vs 15.5%; p<0.0001) and similar prevalence of hypertension and active serological indicators (ie, elevated anti-dsDNA antibodies and low complement) to other ethnicities. Hispanic patients were also exposed to higher average, peak and cumulative doses of corticosteroids but less antimalarial agents than the rest of the SLICC–RAS cohort (table 4).
Table 2  Significant factors associated with MetS at enrolment into SLICC–RAS in age, ethnicity and gender-adjusted analyses

<table>
<thead>
<tr>
<th>MetS</th>
<th>Yes</th>
<th>No</th>
<th>p Value</th>
<th>OR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Current CS (%)</td>
<td>193/239 (80.8)</td>
<td>8250/1255 (67.7)</td>
<td>&lt;0.001</td>
<td>1.53 (1.05 to 2.25)</td>
</tr>
<tr>
<td>Average CS dose, mg (median (IQR))</td>
<td>30 (15, 45)</td>
<td>20 (10, 30)</td>
<td>&lt;0.001</td>
<td>1.02 (1.01 to 1.04)</td>
</tr>
<tr>
<td>Highest CS dose, mg (median (IQR))</td>
<td>50 (30, 60)</td>
<td>30 (20, 50)</td>
<td>&lt;0.001</td>
<td>1.00 (1.00 to 1.01)</td>
</tr>
<tr>
<td>Cumulative CS dose, g (median (IQR))</td>
<td>3.1 (1.5, 5.4)</td>
<td>2.3 (1.0, 4.0)</td>
<td>0.006</td>
<td>1.05 (1.00 to 1.09)</td>
</tr>
<tr>
<td>Past IV CS (%)</td>
<td>18/230 (7.8)</td>
<td>52/1193 (4.4)</td>
<td>0.03</td>
<td>1.88 (1.35 to 2.62)</td>
</tr>
<tr>
<td>Antimalarial (%)</td>
<td>123/239 (51.5)</td>
<td>648/1255 (67.7)</td>
<td>&lt;0.001</td>
<td>0.51 (0.38 to 0.67)</td>
</tr>
<tr>
<td>Immunosuppressant (%)</td>
<td>140/238 (58.8)</td>
<td>459/1253 (36.6)</td>
<td>&lt;0.001</td>
<td>2.21 (1.63 to 3.00)</td>
</tr>
<tr>
<td>SLICC-DI ≥1 (%)</td>
<td>26/95 (27.4)</td>
<td>91/550 (16.6)</td>
<td>0.01</td>
<td>1.99 (1.16 to 3.40)</td>
</tr>
<tr>
<td>SLEDAI-2K (mean (SD))</td>
<td>7.79 (14.0)</td>
<td>5.24 (5.25)</td>
<td>&lt;0.001</td>
<td>1.05 (1.02 to 1.07)</td>
</tr>
<tr>
<td>SLEDAI ≥10 (%)</td>
<td>66/239 (27.6)</td>
<td>221/1255 (17.7)</td>
<td>&lt;0.001</td>
<td>1.73 (1.22 to 2.44)</td>
</tr>
<tr>
<td>High anti-dsDNA (%)</td>
<td>14/206 (6.8)</td>
<td>30/1031 (2.9)</td>
<td>0.02</td>
<td>2.10 (1.03 to 4.29)</td>
</tr>
<tr>
<td>Leucopenia (%)</td>
<td>7/203 (3.5)</td>
<td>92/1108 (8.3)</td>
<td>0.02</td>
<td>0.33 (0.14 to 0.75)</td>
</tr>
<tr>
<td>Active renal disease (%)</td>
<td>94/239 (39.3)</td>
<td>220/1255 (17.5)</td>
<td>&lt;0.001</td>
<td>2.87 (2.05 to 4.02)</td>
</tr>
<tr>
<td>Past renal disease (%)</td>
<td>22/239 (9.2)</td>
<td>69/1255 (5.5)</td>
<td>0.03</td>
<td>1.67 (1.00 to 2.88)</td>
</tr>
</tbody>
</table>

Variables reflect current exposures, recorded at enrolment.
CS, corticosteroid; DI, damage index; IV, intravenous; MetS, metabolic syndrome; RAS, Registry for Atherosclerosis; SLEDAI, systemic lupus erythematosus disease activity index; SLICC, Systemic Lupus International Collaborating Clinics.

Table 3  Characteristics of patients of Korean and Hispanic ethnicity compared to all other ethnicities

<table>
<thead>
<tr>
<th></th>
<th>Korean</th>
<th>p Value*</th>
<th>Hispanic</th>
<th>p Value†</th>
<th>Other ethnicities</th>
</tr>
</thead>
<tbody>
<tr>
<td>No of patients</td>
<td>169</td>
<td>240</td>
<td>1085</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age, years (mean (SD))</td>
<td>28.8 (9.7)</td>
<td>&lt;0.0001</td>
<td>29.2 (10.3)</td>
<td>&lt;0.0001</td>
<td>37.5 (14.0)</td>
</tr>
<tr>
<td>Gender (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>150 (88.8)</td>
<td>0.90</td>
<td>218 (90.8)</td>
<td>0.45</td>
<td>968 (89.2)</td>
</tr>
<tr>
<td>Male</td>
<td>19 (11.2)</td>
<td>0.90</td>
<td>22 (9.2)</td>
<td>0.45</td>
<td>117 (10.8)</td>
</tr>
<tr>
<td>MetS (%)</td>
<td>52 (30.8)</td>
<td>&lt;0.0001</td>
<td>72 (31.3)</td>
<td>&lt;0.0001</td>
<td>112 (10.3)</td>
</tr>
<tr>
<td>MetS phenotype (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MetS WC</td>
<td>33/164 (20.1)</td>
<td>&lt;0.0001</td>
<td>129/228 (56.6)</td>
<td>0.15</td>
<td>483/942 (51.3)</td>
</tr>
<tr>
<td>MetS BP</td>
<td>74 (43.8)</td>
<td>0.63</td>
<td>117 (48.8)</td>
<td>0.40</td>
<td>495/1082 (45.8)</td>
</tr>
<tr>
<td>MetS TG</td>
<td>100/153 (65.4)</td>
<td>&lt;0.0001</td>
<td>108/168 (64.3)</td>
<td>&lt;0.0001</td>
<td>411/1021 (40.3)</td>
</tr>
<tr>
<td>MetS HDL</td>
<td>110/144 (76.4)</td>
<td>&lt;0.0001</td>
<td>97/149 (65.1)</td>
<td>0.007</td>
<td>279/529 (52.7)</td>
</tr>
<tr>
<td>MetS Glu</td>
<td>45/168 (26.8)</td>
<td>0.04</td>
<td>41/236 (17.4)</td>
<td>0.42</td>
<td>185/940 (19.7)</td>
</tr>
<tr>
<td>BMI (mean(SD))</td>
<td>21.6 (4.3)</td>
<td>&lt;0.0001</td>
<td>24.5 (5.0)</td>
<td>0.002</td>
<td>25.8 (6.1)</td>
</tr>
<tr>
<td>WC (cm (mean (SD))</td>
<td>74.7 (8.1)</td>
<td>&lt;0.0001</td>
<td>82.9 (10.9)</td>
<td>0.17</td>
<td>84.3 (14.9)</td>
</tr>
<tr>
<td>Disease duration, weeks (mean (SD))</td>
<td>18.5 (15.9)</td>
<td>&lt;0.0001</td>
<td>23.6 (16.9)</td>
<td>0.15</td>
<td>25.1 (18.4)</td>
</tr>
<tr>
<td>SLEDAI (mean (SD))</td>
<td>7.45 (6.09)</td>
<td>&lt;0.0001</td>
<td>6.46 (5.75)</td>
<td>&lt;0.0001</td>
<td>5.0 (5.2)</td>
</tr>
<tr>
<td>SLICC/ACR-DI (mean (SD))</td>
<td>0.24 (0.69)</td>
<td>0.50</td>
<td>0.28 (0.69)</td>
<td>0.87</td>
<td>0.30 (0.74)</td>
</tr>
<tr>
<td>Disease phenotype (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Active renal disease (%)</td>
<td>49 (29.0)</td>
<td>&lt;0.0001</td>
<td>97 (40.4)</td>
<td>&lt;0.0001</td>
<td>168 (15.5)</td>
</tr>
<tr>
<td>Anti-dsDNA positive (%)</td>
<td>105/159 (66.0)</td>
<td>&lt;0.0001</td>
<td>84/211 (39.8)</td>
<td>0.30</td>
<td>352/977 (36.0)</td>
</tr>
<tr>
<td>Low complement (%)</td>
<td>121/161 (75.2)</td>
<td>&lt;0.0001</td>
<td>74/208 (35.6)</td>
<td>0.46</td>
<td>324/980 (33.1)</td>
</tr>
<tr>
<td>Thrombocytopenia (%)</td>
<td>16/143 (11.2)</td>
<td>&lt;0.0001</td>
<td>2/210 (1.0)</td>
<td>0.13</td>
<td>26/960 (2.7)</td>
</tr>
<tr>
<td>Medication (median (IQR))</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oral Cs, %*</td>
<td>161 (95.3)</td>
<td>&lt;0.0001</td>
<td>211 (87.9)</td>
<td>&lt;0.0001</td>
<td>671 (61.8)</td>
</tr>
<tr>
<td>Average Cs dose, mg</td>
<td>20 (10, 35)</td>
<td>0.26</td>
<td>30 (15, 42.5)</td>
<td>&lt;0.0001</td>
<td>20 (10, 30)</td>
</tr>
<tr>
<td>Highest Cs dose, mg</td>
<td>30 (15, 55)</td>
<td>0.07</td>
<td>50 (30, 60)</td>
<td>&lt;0.0001</td>
<td>40 (20, 60)</td>
</tr>
<tr>
<td>Cumulative Cs dose, g</td>
<td>1.4 (0.4, 3.1)</td>
<td>&lt;0.0001</td>
<td>3.9 (1.8, 6.2)</td>
<td>&lt;0.0001</td>
<td>2.5 (1.2, 4.8)</td>
</tr>
<tr>
<td>Pulse intravenous Cs, %</td>
<td>26 (15.4)</td>
<td>0.001</td>
<td>5/232 (2.2)</td>
<td>0.26</td>
<td>39/1031 (3.8)</td>
</tr>
<tr>
<td>Immunosuppressant (%)</td>
<td>86 (50.9)</td>
<td>&lt;0.0001</td>
<td>146 (60.8)</td>
<td>&lt;0.0001</td>
<td>367/1082 (33.9)</td>
</tr>
<tr>
<td>Antimalarial (%)</td>
<td>120 (71.0)</td>
<td>0.29</td>
<td>125 (52.1)</td>
<td>&lt;0.0001</td>
<td>705 (65.0)</td>
</tr>
</tbody>
</table>

*Korean versus all other (non-Hispanic) ethnicities.
†Hispanic versus all other (non-Korean) ethnicities.
ACR, American College of Rheumatology; BMI, body mass index; BP, blood pressure; CS, corticosteroid; DI, damage index; Glu, glucose; HDL, high-density lipoprotein; MetS, metabolic syndrome; SLEDAI, systemic lupus erythematosus disease activity index; SLICC, Systemic Lupus International Collaborating Clinics; TG, triglycerides; WC, waist circumference.
ticosteroid use (yes/no), with dose-related variables excluded. In this model, current corticosteroid use was not significantly associated with MetS although past intravenous corticosteroid use was (OR 2.45, 95%CI 1.01 to 5.97), suggesting that increasing corticosteroid doses have a greater impact on MetS susceptibility than simple exposure status. Immunosuppressive therapies were not predicted a priori to be mechanistically involved in MetS development but rather act as a marker of disease severity, higher disease activity and/or corticosteroid use. The majority of patients receiving immunosuppressive therapies were also receiving oral corticosteroids (91.5%), and at higher doses than those not on immunosuppressive therapies. However, use of immunosuppressive therapies remained a significant predictor of MetS even after adjusting for all clinically correlating factors, such as SLEDAI, renal disease and corticosteroid use/dose (fully adjusted OR 2.15, 95%CI 1.15 to 4.00).

Finally, those with renal involvement had more hypertension (MetS BP 72.9% vs 38.8%; p<0.0001) and hypertriglyceridaemia (MetS TG 74.1% vs 38.8%; p<0.0001) than those without renal lupus. Central obesity parameters (BMI and MetS WC) were, however, lower in those with renal disease, despite significantly higher corticosteroid exposures (eg, median (IQR) average oral prednisolone dose 30 mg (20, 50) vs 16 mg (10, 30)) (see supplementary table S2, available online only). However, use of immunosuppressive therapies remained a significant predictor of MetS even after adjusting for all clinically correlating factors, such as SLEDAI, renal disease and corticosteroid use/dose (fully adjusted OR 2.15, 95%CI 1.15 to 4.00).

DISCUSSION

We report that MetS was common (16%) in SLE despite our cohort being young, predominantly female and early in the course of their disease. The relatively high prevalence of MetS at enrolment suggests that the metabolic derangements that contribute to long-term CHD risk characterised by MetS appear early in the course of the disease. Smaller studies of established lupus cohorts have reported MetS rates of 18–32%, which were consistently higher than control populations. For example, Parker et al. found a prevalence of 50% in a UK SLE cohort, compared to 20% in controls, and Sabio et al. reported a prevalence of 20%, compared to 15% in controls. These results suggest the hypothesis that rapid control of inflammatory disease activity with the lowest doses of corticosteroid possible is likely to be beneficial to long-term cardiovascular risk. The role of inflammation in the development of atherosclerosis has been increasingly recognised, and SLE is associated with higher circulating levels of high sensitivity C-reactive protein, interleukin 18 and tumour necrosis factor α, which are associated with insulin resistance and endothelial dysfunction and have been implicated in the development of CHD in the general population. Long-term and high-dose corticosteroid use is associated with the pro-atherogenic metabolic disturbances characterised by MetS. and the weight of evidence in SLE suggests they are pro-atherogenic. Others have, however, suggested that their anti-inflammatory activity in lupus may exert an atheroprotective role. The present study suggests that in early disease, higher doses of corticosteroids play a pivotal role in the development of MetS.

The strong association of MetS with Korean and Hispanic ethnicities partly reflects a higher background population prevalence of MetS. In 2006 a Mexican study found the prevalence of MetS in adults was 37–50%, depending on the definition used, and 24–36% in adults aged 20–59 years. Similarly, the San Antonio Heart Study demonstrated that people of Mexican Hispanic descent had a higher prevalence of MetS than Caucasian individuals, an observation especially pronounced in women (30.9% vs 16.8%). A recent study of South Korean adults described a MetS prevalence in women of 16–31% depending on the definition used, with low levels of central obesity, as we found in our study. While there is therefore variability in the background prevalence of MetS, MetS remains more prevalent in SLE than controls. The ethnic gradient may therefore reflect an increased susceptibility to MetS and enhanced sensitivity to the adverse effects of inflammation and corticosteroid exposure in Korean and Hispanic patients. There were, however, important differences in the SLE phenotype observed in these two subsets, as well as differences in the clinical and inflammatory pattern of disease observed to be associated with MetS. Whether these differences translate into a differential effect on future cardiovascular endpoints is the subject of prospective study.

Antimalarial agents showed a significant negative association with MetS on univariate analysis; however, this did not remain significant in our multivariable model. A protective effect of antimalarial use against MetS has been demonstrated by other groups with more stable, mild disease. Our study included recently diagnosed patients with a shorter exposure to the drug in the context of a more severe, active disease. It will be of interest to observe how longer exposure to antimalarial agents in the context of disease stabilisation will influence the MetS phenotype over time in this cohort. The significant association between immunosuppressant use and MetS may represent confounding, an indication of disease severity rather than mediating metabolic derangement. However, the relationship persists even after adjusting for all measured potential confounders (such as SLEDAI and corticosteroid use). This suggests that either immunosuppressive therapies have direct adverse metabolic effects, or there is residual confounding related to inadequately measured exposures, such as disease activity. Apart from cyclosporin A, immunosuppressive therapies commonly used in SLE are not associated with metabolic derangements, and may play a role in preventing atherosclerosis. Therefore, immunosuppressant use is likely to reflect residual confounding, and additional biomarkers and/or indices of disease activity may improve the estimation of exposure to systemic inflammation.

This is the largest study to date examining MetS in SLE and has many advantages over previous studies. The cohort is young, with a range of disease activity that allows us to explore more effectively the impact of inflammation on MetS development. We also studied an international cohort recruited from 11 countries with a range of ethnicities, and therefore the results can be generalised to a range of populations. Our data also included a breadth of corticosteroid data, allowing us to...
explore their effects in more detail. Finally, the prospective nature of the cohort limits many potential sources of bias associated with retrospective studies.

The analysis does, however, have several limitations. First, there are missing MetS data on 11.4% of the cohort. While the demographics of this cohort are broadly similar they appear to have less severe disease, which may bias the analysis towards disease severity markers. Our data may therefore overestimate the true prevalence of MetS in this population. The most common missing variable was HDL-cholesterol, which was not universally performed locally in all centres. Second, a cross-sectional study is not the ideal study design to dissect the interplay between inflammation, corticosteroid exposure and MetS, although it can provide important data with which to inform prospective studies. Future work will examine how these factors influence MetS over time and investigate the influence of genetic factors on MetS prevalence and susceptibility. Finally, the use of MetS as a CHD risk prediction tool for adverse cardiovascular events has yet to be validated in SLE and the SLICC–RAS cohort is an ideal setting in which to examine this further.

Our study confirms that MetS is common in young patients with recently diagnosed SLE. This clustering of CHD risk factors and the observed ethnic variation in MetS susceptibility should help inform risk stratification in the management of early disease. MetS is associated with a more severe disease phenotype and higher doses of corticosteroids, therefore balancing disease control while minimising corticosteroid exposure should be at the forefront of personalised treatment decisions in these patients.

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**Contributors**
The study was conceived by IB, MU, DG and ML, and all SLICC investigators contributed to its design. All authors contributed to data collection. Data analysis, interpretation and manuscript preparation was performed by BP, IB, ML, MU and DG. All authors have critically reviewed and edited the manuscript, and approved its publication.

**Competing interests**
None.

**Ethics approval**
The study was approved by the University Health Network Research Institute, Research Ethics Committee, Toronto, Canada and by the institutional research ethics boards of participating centres in accordance with the Declaration of Helsinki’s guidelines for research in humans.

**Patient consent** Obtained.

**Provenance and peer review**
Not commissioned; externally peer reviewed.

**Correction notice**
This article has been corrected since it was published Online First. The spelling of one of the authors’ surnames has been corrected.

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Clinical and epidemiological research


