Association of neighbourhood socioeconomic status and diabetes burden using electronic health records in Madrid (Spain): the HeartHealthyHoods study

Usama Bilal,1,2,3 Felicia Hill-Briggs,1,4,5 Luis Sánchez-Perruca,6,7 Isabel Del Cura-González,7,8,9 Manuel Franco1,2

ABSTRACT

Objective To study the association between neighbourhood socioeconomic status and diabetes prevalence, incidence, and control in the entire population of northeastern Madrid, Spain.

Setting Electronic health records of the primary-care system in four districts of Madrid (Spain).

Participants 269,942 people aged 40 or older, followed from 2013 to 2014.

Exposure Neighbourhood socioeconomic status (NSES), measured using a composite index of seven indicators from four domains of education, wealth, occupation and living conditions.

Primary outcome measures Diagnosis of diabetes based on ICPC-2 codes and glycated haemoglobin (HbA1c %).

Results In regression analyses adjusted by age and sex and compared with individuals living in low NSES neighbourhoods, men living in medium and high NSES neighbourhoods had 27% (95% CI: 23% to 30%) and 50% (95% CI: 47% to 52%) lower prevalence of diabetes. Moreover, the hazard of diabetes in men living in medium and high NSES neighbourhoods was 13% (95% CI: 1% to 23%) and 20% (95% CI: 9% to 29%) lower, while the hazard of diabetes in women living in medium and high NSES neighbourhoods was 17% (95% CI: 3% to 29%) and 31% (95% CI: 20% to 41%) lower. Individuals living in medium and high SES neighbourhoods had 8% (95% CI: 2% to 15%) and 15% (95% CI: 9% to 21%) lower prevalence of lack of diabetes control, and a decrease in average HbA1c % of 0.05 (95% CI: 0.01 to 0.10) and 0.11 (95% CI: 0.06 to 0.15).

Conclusions Diabetes prevalence, incidence and lack of control increased with decreasing NSES in a southern European city. Future studies should provide mechanistic insights and targets for intervention to address this health inequity.

INTRODUCTION

The burden of diabetes has seen a large increase in Western countries in recent decades.1 Diabetes-attributable costs in the European Union have been estimated to be over $100 billion per year and are predicted to continue increasing in the following decades.2 Population preventive strategies are needed to decrease this burden,3 taking into consideration mass influences that differ across populations.3

Among these mass influences are neighbourhood characteristics. A large body of literature has explored contextual socioeconomic influences on health. In particular, the association between neighbourhood socioeconomic status (NSES) and several measures of diabetes (prevalence, incidence...
or control) is robust and has been replicated in the USA,4–10 other Anglo-Saxon countries11–19 and northern and central Europe20–26 including in experimental or quasi-experimental settings.21 27 Nonetheless, these influences have received scant attention in southern Europe.28 Moreover, previous studies have shown a strong social gradient in diabetes mortality in Spain, which warrants further mechanistic insights into its causes.29 Recent studies have shown that segregation patterns and neighbourhood selection phenomena is changing in southern Europe,30 necessitating a study of the health outcomes associated with these changes.

Finally, many of the studies outlined above use data from research-driven cohort studies. While these types of studies have the advantage of standardised and high-quality data collection, they may suffer from a number of biases derived from a non-random sampling of the study participants.31 In particular, the role that context plays in determining selection into a study may be particularly relevant in studies on the effect of context on health.32 With electronic health records (EHR) in a health system with universal health coverage, these drawbacks may be overcome by avoiding the necessity for sampling altogether.

Taking the above into consideration, we studied the association between NSES and diabetes prevalence, incidence and control in an electronic health record-based cohort of the entire population of northeastern Madrid that includes data on more than 640,000 people.

**METHODS**

**Study setting**

This study was conducted within the HeartHealthyHoods project (www.ihanna.eu) in the city of Madrid, Spain.32 We took data for 2013 and 2014 from all healthcare centres in four districts of the city of Madrid, all belonging to the same health district. These four districts contain around 20% of the total population of Madrid and are representative of the rest of the city of Madrid (online appendix figure 1). Our unit of analysis is the census section (n=427), which is the smallest area for which the census collects data and has around 1200 people (range: 583 to 3865). Individual-level data were obtained from EHR including 640,217 individuals registered in any health centre of the area. These EHR contain data on patient age, sex, residential location, clinical diagnoses and laboratory values (lipids and HbA1c).

Since this screening for cardiovascular risk factors is limited to people 40 years and older,32 we restricted our dataset to people born after January 1, 1973 (aged 40 or older by 2013). Our final study sample was composed of 270,660 individuals, of which 23,908 had a diagnosis of diabetes. Primary care EHR includes 99.5% of the individuals living in the area per the census.

**Neighbourhood socioeconomic status**

The main exposure of this study was NSES. To measure NSES, we considered the four domains of the Spanish Commission to Reduce Health Inequalities33: education, wealth, occupation and living conditions. To search for indicators to measure these four domains, we explored all available data sources, to our knowledge, on social, economic and contextual factors in Madrid, Spain. We looked for readily available indicators (to ease replicability) that were measured at the neighbourhood or census section level (to improve granularity) and that were available for several years (to allow for further studies looking at longitudinal changes). After this process we selected seven indicators that represent the four domains: education—(1) primary education (% people above 25 years of age with primary studies or below), (2) university education (% people above 25 years of age with university education or above); wealth—(3) average housing prices (per sq. m); occupation—(4) part-time employment (% workers in part-time jobs), (5) temporary employment (% workers in temporary jobs), (6) manual occupational class (% workers in manual or unqualified jobs); and living conditions—(7) unemployment rate (% registered unemployed individuals/people aged 16 to 64). Indicator data were obtained from the Padrón (a continuous and universal census collected for administrative purposes), the social security and employment services registries and the IDEALISTA report (a report from a large real estate corporation in Spain). All data were available by January 2013. The online resource contains a detailed description of the operationalisation of indicators.

We computed a weighted index of the seven indicators by: (1) making the directionality of the associations consistent, by reversing some of the indicators (primary education, part-time employment, temporary employment, manual occupational class and unemployment rate) so that all indicators had a consistent association with the final index; (2) for each indicator, we centred by the mean and divided it by the SD in order to obtain a Z-score of each indicator; (3) in each domain, we averaged the Z-score of each indicator, resulting in a Z-score for each domain (education, wealth, occupation and living conditions) and (4) finally, we calculated the composite index of NSES by averaging the Z-score of each of the four domains. This composite NSES index was then operationalised in separate analyses as a categorical variable (NSES in tertiles) or as a continuous variable.

**Diabetes prevalence, incidence and control**

Diabetes diagnoses were extracted from the EHR for all individuals, as recorded by primary care physicians during their usual clinical practice. A type-2 diabetes diagnosis was defined using the T90 diagnosis code of the ICPC-2 (‘diabetes non-insulin dependent’). A previous study has validated the diagnosis of diabetes in this dataset with a kappa of 0.99, with high sensitivity (99.5%) and specificity (99.5%).34 Prevalent cases were defined as diabetes diagnoses dated before 1 January 2013. Incident cases were
those occurring from 1 January 2013 to 31 December 2014 in people free of diabetes by baseline (1 January 2013). We operationalised lack of diabetes control as either a dichotomous variable (HbA1c>=7%) or a continuous variable (HbA1c %). If more than one value of HbA1c was available, we used the last available measurement of the year.

**Statistical methods**

The overall goal of this analysis is to study the association between NSES and diabetes prevalence, incidence and control. We computed descriptive statistics by tertile of NSES.

To study the association between NSES and diabetes prevalence or lack of control (binary indicator) we used a log-binomial regression model with robust standard errors clustered at the census section level using a sandwich Huber–White estimator. These models were adjusted for age (in five categories; 40 to 49, 50 to 59, 60 to 69, 70 to 79 and 80 and older) and sex. Continuous HbA1c (for diabetes control) was examined using a linear regression with robust standard errors clustered at the census section level using a sandwich Huber–White estimator. Around 21% of the sample that had prevalent diabetes had no HbA1c % measured in 2013 or 2014. To assess whether this missing data affected our inferences, we did a sensitivity analysis using a conditional mean imputation of HbA1c % in people with diabetes. In this model, we predicted the HbA1c % value using age, sex, healthcare centre, NSES index and diagnosis of other cardiovascular risk factors or conditions (hypertension, dyslipidaemia, prevalent cardiovascular disease, chronic kidney disease and retinopathy). We then compared the point estimates of the association between prevalent lack of control and average HbA1c % obtained with and without conditional mean imputation.

In the analysis of diabetes incidence, each individual entered the sample on 1 January 2013 and exited on the date of diabetes diagnosis (outcome), date of death (censored), date of moving out of a health centre in the area (censored) or study end by 31 December 2014 (administrative censoring). We used Kaplan–Meier survival estimates to explore the differences in the hazard of diabetes incidence by NSES tertile. Cox proportional hazards models were used to estimate the adjusted association, with clustered standard errors on the census section. Since we censored individuals at death, a potential competing risk, our estimates from the model are analogous to cause-specific hazard ratios, and can therefore be interpreted as the increase in the hazard of diabetes if people that do not die. We checked the proportionality of hazards assumption by plotting Schoenfeld residuals and by checking their trend over time.35

To graphically display the association between the exposure and the outcome variables, we also modelled the associations above using restricted cubic splines with four knots in the percentiles recommended by Harrell.36 A previous report in the Spanish setting highlighted a significant interaction by sex of contextual socioeconomic status and diabetes,28 so we explored whether this interaction existed in our analysis and displayed stratified results if this was the case. All analyses were conducted in R V.3.3.0 (R Software Foundation).

**RESULTS**

**Study population**

Table 1 shows a description of the study population by tertile of NSES and in the total population. The total sample size was 269,942 people, with around 25%, 30% and 45% of the population living in low, medium and high NSES areas. Overall, the median age was 56.5 (IQR=47.4 to 69.8) and 54.9% of the population were women. Of this, 8.8% of the population older than 40 years of age had diabetes, 1.0% developed diabetes during follow-up and the average HbA1c in diabetic people was 6.7 (IQR=6.2 to 7.5). Thirty-nine percent of all diabetic people had uncontrolled diabetes (HbA1c equal or above 7%). Stratifying the population by tertile of NSES revealed that younger people lived in neighbourhoods with higher SES. The prevalence of diabetes decreased sharply with NSES (11.9% in the lowest NSES, 9.6% in the medium NSES and 6.5% in the highest NSES), and the incidence of diabetes followed a similar gradient by NSES (1.3%, 1.1% and 0.9% in the lowest, medium and highest NSES areas).

**NSES and diabetes prevalence**

Table 2 shows the association between NSES and diabetes prevalence, control and incidence. Diabetes prevalence was associated in a dose–response manner to NSES. This association was significantly stronger in women as compared with men (P value for the interaction <0.001). In particular, compared with men living in low NSES neighbourhoods, those living in medium NSES neighbourhoods had 8% lower prevalence of having diabetes (PR=0.92, 95% CI 0.89 to 0.96), while those living in the highest NSES neighbourhoods had 24% lower prevalence of diabetes (PR=0.76, 95% CI 0.74 to 0.80). In the case of women, those living in medium and high NSES neighbourhods had 24% and 46% lower prevalence of diabetes, respectively, as compared with those living low NSES neighbourhoods (PR=0.76, 95% CI 0.73 to 0.79, and PR=0.54, 95% CI 0.52 to 0.57). These associations were consistent in models looking at continuous NSES: a one SD increase in NSES was associated with 14% and 26% lower prevalence of diabetes in men and women, respectively (PR=0.86, 95% CI 0.84 to 0.87, PR=0.74, 95% CI 0.72 to 0.75). Figure 1 shows the association using continuous NSES with restricted cubic splines, where the steeper pattern for women is evident.

**NSES and diabetes control**

Table 2 also shows the association between NSES and diabetes control, operationalised as a dichotomous variable (lack of diabetes control, or HbA1c>=7%) or a
continuous variable (HbA1c %). There was no significant interaction by sex in the NSES and diabetes control (P value for the interaction=0.219 and 0.358 in the dichotomous and continuous model). As compared with people with diabetes living in the lowest NSES neighbourhoods, those living in medium NSES areas had 5% lower prevalence of lack of diabetes control (PR=0.95, 95% CI 0.91 to 0.99), while those living in the highest NSES areas had 9% lower prevalence of lack of diabetes control (PR=0.91, 95% CI 0.87 to 0.95). Moreover, a one SD increase in NSES was associated with 4% lower prevalence of lack of diabetes control (PR=0.96, 95% CI 0.94 to 0.98). These associations were maintained when looking at continuous HbA1c: diabetic people living in medium and high NSES had a lower average HbA1c % (see table 2). Figure 2 shows the prevalence of lack of diabetes control and average HbA1c levels across levels of NSES using restricted cubic splines, showing a linear decrease both in lack of control and in average HbA1c % with increasing NSES. In the sensitivity analysis using conditional mean imputation of HbA1c %, we found no change in our inferences after accounting for missing HbA1c % (see online appendix figure 2).

### NSES and diabetes incidence

Overall, at 1 and 2 years of follow-up, the diabetes incidence was 5.7 per 1000 and 10.5 per 1000. Figure 3 shows the Kaplan–Meier estimate of diabetes incidence by tertile of NSES, showing a social gradient in diabetes incidence (lower NSES corresponding to higher diabetes incidence, P<0.001). Table 2 also shows the results of the adjusted Cox proportional hazards models. We found a significant interaction by sex (P value for interaction=0.004). The hazard of diabetes incidence in men living in medium and high NSES neighbourhoods was 13% and 20% lower compared with men living in low NSES neighbourhoods (HR=0.87, 95% CI 0.77 to 0.99, and HR=0.80, 95% CI 0.71 to 0.91). A stronger association was observed in women, as the hazard of diabetes incidence in women living in medium and high NSES neighbourhoods was 17% and

<table>
<thead>
<tr>
<th>Variable</th>
<th>Total</th>
<th>Tertile 1 (Lowest NSES)</th>
<th>Tertile 2 (Mid NSES)</th>
<th>Tertile 3 (High NSES)</th>
<th>P values*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample Size (N)</td>
<td>269 942</td>
<td>68 369</td>
<td>81 072</td>
<td>120 501</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Median Age (IQR)</td>
<td>56.5 (47.4;69.8)</td>
<td>58.6 (48.3;74.5)</td>
<td>58.1 (48.0;71.1)</td>
<td>54.7 (46.6;66.9)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>% Men</td>
<td>45.1%</td>
<td>44.6%</td>
<td>44.2%</td>
<td>45.9%</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>% Women</td>
<td>54.9%</td>
<td>55.4%</td>
<td>55.8%</td>
<td>54.1%</td>
<td></td>
</tr>
<tr>
<td>% Death during follow-up</td>
<td>1.2%</td>
<td>1.4%</td>
<td>1.3%</td>
<td>1.0%</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>% Moved during follow-up</td>
<td>0.8%</td>
<td>0.8%</td>
<td>0.8%</td>
<td>0.8%</td>
<td>0.673</td>
</tr>
<tr>
<td>% With prevalent diabetes</td>
<td>8.8%</td>
<td>11.9%</td>
<td>9.6%</td>
<td>6.5%</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>% With incident diabetes†</td>
<td>1.0%</td>
<td>1.3%</td>
<td>1.1%</td>
<td>0.9%</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Median HbA1c (IQR)</td>
<td>6.7 (6.2;7.5)</td>
<td>6.7 (6.2;7.5)</td>
<td>6.7 (6.2;7.5)</td>
<td>6.7 (6.2;7.4)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>HbA1c≥7%</td>
<td>38.8%</td>
<td>40.5%</td>
<td>38.7%</td>
<td>37.1%</td>
<td>0.237</td>
</tr>
<tr>
<td>HbA1c&lt;5%</td>
<td>0.3%</td>
<td>0.3%</td>
<td>0.4%</td>
<td>0.3%</td>
<td>0.285</td>
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<tr>
<td>HbA1c 5%–6.5%</td>
<td>41.1%</td>
<td>40.0%</td>
<td>40.5%</td>
<td>42.7%</td>
<td></td>
</tr>
<tr>
<td>HbA1c 6.5%–7%</td>
<td>20.1%</td>
<td>19.4%</td>
<td>20.6%</td>
<td>20.3%</td>
<td></td>
</tr>
<tr>
<td>HbA1c 7%–9%</td>
<td>32.4%</td>
<td>34.0%</td>
<td>32.2%</td>
<td>30.9%</td>
<td></td>
</tr>
<tr>
<td>HbA1c&gt;9%</td>
<td>6.1%</td>
<td>6.3%</td>
<td>6.3%</td>
<td>5.7%</td>
<td></td>
</tr>
<tr>
<td>Primary education, % (IQR)</td>
<td>24.6% (15.1;32.2)</td>
<td>36.3% (30.7;40.3)</td>
<td>24.7% (20.8;27.9)</td>
<td>24.7% (27.1;19.5)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>University education, % (IQR)</td>
<td>20.8% (13.0;33.7)</td>
<td>10.2% (7.4;13.0)</td>
<td>20.8% (16.8;24.7)</td>
<td>40.1% (29.9;52.5)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Unemployment rate, % (IQR)</td>
<td>12.6% (10.6;13.8)</td>
<td>13.8% (13.8;16.4)</td>
<td>12.6% (12.0;12.7)</td>
<td>8.9% (7.8;10.6)</td>
<td>&lt;0.001</td>
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<tr>
<td>Part-time workers, % (IQR)</td>
<td>23.4% (18.7;25.9)</td>
<td>26.7% (24.8;26.8)</td>
<td>23.4% (22.4;25.9)</td>
<td>16.5% (12.7;19.4)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Temporary workers, % (IQR)</td>
<td>19.0% (17.3;20.9)</td>
<td>20.5% (20.4;21.5)</td>
<td>20.4% (18.9;20.9)</td>
<td>16.7% (13.8;18.2)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Manual class, % (IQR)</td>
<td>37.1% (27.4;40.0)</td>
<td>40.3% (40.0;43.1)</td>
<td>37.1% (36.2;40.0)</td>
<td>22.4% (17.4;30.2)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Property value, EUR/m² (IQR)</td>
<td>2286.0 (1975.0;2659.0)</td>
<td>1776.0 (1561.0;1971.0)</td>
<td>2243.0 (2128.0;2398.0)</td>
<td>2832.0 (2608.0;3382.0)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>SES index (IQR)</td>
<td>0.0 (-0.6;0.6)</td>
<td>-0.8 (-1.2;-0.6)</td>
<td>-0.2 (-0.3;0.1)</td>
<td>1.0 (0.6;1.6)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

*P value - values for continuous individual-level characteristics were computed using a clustered Somers’ D comparison of medians; P-values for categorical individual-level characteristics were computed using Donner’s χ² adjusted for clustered data. P-values for contextual characteristics were conducted at the neighbourhood level using a Kruskal-Wallis test for the comparison of medians.
†Incident diabetes refers to new diagnoses of diabetes in 2013 or 2014 in people free of diabetes at baseline.
NSES, neighbourhood socioeconomic status index.
31% lower compared with women living in low NSES
neighbourhoods (HR=0.83, 95% CI 0.71 to 0.97, and
HR=0.69, 95% CI 0.59 to 0.80). These associations
were consistent in models looking at continuous NSES:
a one SD increase in NSES was associated with a 10% and 18%
decrease in the hazard of incident diabetes in men and
women, respectively (HR=0.90, 95% CI 0.85 to 0.94, and
HR=0.82, 95% CI 0.77 to 0.87). We tested the assumption
of proportionality of hazards and found no evidence to
reject the null hypothesis of proportionality (P value for
the global chi²-test=0.604 for the unadjusted model, and
0.365 for the fully adjusted model).

### DISCUSSION

This study has shown a strong association between NSES
and diabetes burden. In particular, there is a dose–
response association: as NSES increases, diabetes prevalence,
lack of control and incidence decrease in a linear
fashion. This association is seen for both a categorical

### Table 2  Association of neighbourhood socioeconomic status (NSES) and diabetes outcomes

<table>
<thead>
<tr>
<th>Variable</th>
<th>Total</th>
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<tbody>
<tr>
<td></td>
<td>Diabetes Prevalence</td>
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<td></td>
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<td></td>
</tr>
<tr>
<td></td>
<td>PR (95% CI)</td>
<td>P values</td>
<td>PR (95% CI)</td>
<td>P values</td>
<td>PR (95% CI)</td>
</tr>
<tr>
<td>Tertile 1 of NSES (Low)</td>
<td>1(Ref.)</td>
<td></td>
<td>1(Ref.)</td>
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<td>1(Ref.)</td>
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<tr>
<td>Tertile 2 of NSES (Middle)</td>
<td>0.84 (0.82 to 0.87)</td>
<td>&lt;0.001</td>
<td>0.92 (0.89 to 0.96)</td>
<td>&lt;0.001</td>
<td>0.76 (0.73 to 0.79)</td>
</tr>
<tr>
<td>Tertile 3 of NSES (High)</td>
<td>0.66 (0.64 to 0.68)</td>
<td>&lt;0.001</td>
<td>0.76 (0.74 to 0.80)</td>
<td>&lt;0.001</td>
<td>0.54 (0.52 to 0.57)</td>
</tr>
<tr>
<td>Continuous NSES</td>
<td>0.80 (0.79 to 0.81)</td>
<td>&lt;0.001</td>
<td>0.86 (0.84 to 0.87)</td>
<td>&lt;0.001</td>
<td>0.74 (0.72 to 0.75)</td>
</tr>
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| Variable                        | Lack of Diabetes Control (HbA1c ≥7%)
<table>
<thead>
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<tbody>
<tr>
<td></td>
<td>PR (95% CI)</td>
</tr>
<tr>
<td>Tertile 1 of NSES (Low)</td>
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<tr>
<td>Tertile 2 of NSES (Middle)</td>
<td>0.95 (0.91 to 0.99)</td>
</tr>
<tr>
<td>Tertile 3 of NSES (High)</td>
<td>0.91 (0.87 to 0.95)</td>
</tr>
<tr>
<td>Continuous NSES</td>
<td>0.96 (0.94 to 0.98)</td>
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| Variable                        | Lack of Diabetes Control (Continuous HbA1c %)
<table>
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<tbody>
<tr>
<td></td>
<td>Beta (95% CI)</td>
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<td>Tertile 2 of NSES (Middle)</td>
<td>−0.05 (-0.10 to −0.01)</td>
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<td>Tertile 3 of NSES (High)</td>
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<tr>
<td>Continuous NSES</td>
<td>−0.04 (-0.06 to −0.02)</td>
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<table>
<thead>
<tr>
<th>Variable</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>HR (95% CI)</td>
</tr>
<tr>
<td>Tertile 1 of NSES (Low)</td>
<td>1(Ref.)</td>
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<tr>
<td>Tertile 2 of NSES (Middle)</td>
<td>0.85 (0.77 to 0.95)</td>
</tr>
<tr>
<td>Tertile 3 of NSES (High)</td>
<td>0.75 (0.68 to 0.83)</td>
</tr>
<tr>
<td>Continuous NSES</td>
<td>0.86 (0.83 to 0.90)</td>
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</table>

*Models adjusted by age, sex and year and clustered on the census section. Results for diabetes prevalence and lack of diabetes control
(binary) are shown in prevalence ratios (95% CI); results for lack of diabetes control (continuous) are presented as changes in average HbA1c
% (95% CI); results for diabetes incidence are presented as hazard ratios (95% CI).
(...tertiles) and a continuous operationalisation of the exposure. There seems to be an interaction by sex in the association with diabetes prevalence and incidence, which is stronger in women as compared to men.

Previous studies have shown analogous results to ours. A report by Larrañaga found an increase in the prevalence of diabetes in more deprived neighbourhoods in the Basque Country (northern Spain), using a sample of primary care practices, displaying a similar interaction by sex as our study. Other studies using EHR in other countries have found significant associations between area-level poverty, deprivation or socioeconomic status and diabetes prevalence, incidence and control. A study by Cox using EHR from a Scottish region found increased diabetes prevalence in more deprived areas, as measured using the Carstairs index of deprivation. Studies by Mezuk and Sundquist showed a significant increase in diabetes incidence in the Swedish population living in medium and high deprivation neighbourhoods, measured using four indicators of NSES. Several studies in the UK, USA and Israel have studied the association of NSES with diabetes control as measured by HbA1c % in EHR, finding a consistent gradient similar to ours (lower NSES associated with lower likelihood of control or higher HbA1c %). Other studies using data from cross-sectional surveys or cohort studies, but with similar spatial units as ours have also found significant associations in the USA, France, and Sweden. 

![Figure 1](estimated-diabetes-prevalence-by-levels-of-neighbourhood-socioeconomic-status-index.png)

**Figure 1** Estimated diabetes prevalence by levels of neighbourhood socioeconomic status index.

![Figure 2](estimated-diabetes-control-by-levels-of-neighbourhood-socioeconomic-status.png)

**Figure 2** Estimated diabetes control by levels of neighbourhood socioeconomic status.
Third, HbA1c represents a robust measure of diabetes before and shown to have a very high validity with a kappa value. The diagnosis of diabetes in our EHR has been validated using models adjusted by age, sex and year and clustered on the census section. For prediction purposes age was set to the third category (60 to 70 years of age).

Our study is the first in Spain (and to our knowledge in southern Europe) to show an association between NSES and diabetes control.

**Strengths and limitations of this study**

Our study has several strengths. First, we study the entire population of an area of a very large city (Madrid) where almost 600,000 people live. This results in a very large sample size and decreased concerns for selection bias as compared to regular cohort studies or surveys. Second, the diagnosis of diabetes in our EHR has been validated before and shown to have a very high validity with a kappa value of 0.96. Third, HbA1c represents a robust measure of diabetes control and is the standard of care in clinical practice. Finally, we used an exposure constructed from publicly available indicators, increasing the replicability of our findings and the applicability to other health outcomes. Our study also has some limitations. First and foremost, while the validity of our measures of diabetes prevalence, incidence and control is high, we cannot achieve the standardisation of measurements that cohort studies do. While there exists the possibility of differential measurement error, we have no reason to suspect that the accuracy of the measure of diabetes prevalence varies by socioeconomic status, given that Spain has a universal healthcare system. Second, while our exposure is built from publicly available indicators, this also restricts our capacity to build a complex exposure that may capture socioeconomic status better. Third, the available data for individual level confounders were restricted to basic socio-demographic variables, age and sex, which opens the possibility for residual confounding in our inferences. In particular, we do not have data on individual-level socioeconomic status. Unmeasured confounding by neighbourhood selection may be an important source of bias in our study. However, whether adjusting for individual-level socioeconomic status brings estimates closer to the truth or induces overadjustment may depend on the level of social mobility of each country. Last, the generalisability of these results to other Spanish or European cities may be limited for cities that do not have similar segregation patterns. Recent research has shown increased segregation in Madrid, with levels similar to London.

The implications of our study are several. As this is the first study, to our knowledge, to show strong contextual gradients in diabetes burden in Spain, we believe these findings should be incorporated in the National Health Equity Strategy. Research wise, this study opens the possibility to study the connection between contextual factors (food, physical activity, tobacco and alcohol environment) and diabetes. Future studies may consider providing specific mechanistic insights into the contextual determinants of diabetes in southern Europe. For example, Auchincloss and Christine have reported over several studies increased prevalence and incidence of diabetes with lower availability of healthy foods or physical-activity-promoting resources, but research on these mechanistic pathways is lacking in Spain and southern Europe in general. In particular, the association of contextual socioeconomic status and unhealthy food environments has not been thoroughly replicated in Europe and may actually follow a different gradient. We have previously shown that neighbourhoods in Madrid with improving socioeconomic status indicators have an increased proportion of supermarkets and decreased proportion of fruit and vegetable stores, a contextual change undesired by neighbours and perceived as not conducive to better diets. We have also previously shown that walkability may follow an inverse social gradient in Madrid (worse walkability in higher NSES areas), but that this association may not hold in gentrifying areas. In summary, understanding the mechanisms (and therefore potential intervention targets) linking NSES to diabetes may require studies that take into consideration changes in both the exposure and the outcome side.

WHO has identified social determinants as underlying many of the health inequities observed within countries, and resulting strategies to ameliorate social determinants through a system change are under way in countries including Spain. For diabetes, an unhealthy diet, lack of physical activity, and subsequent obesity are some of the main modifiable risk factors that are adversely impacted by social determinants. Understanding the contextual contributors to the social patterning of diabetes we have described in this study can offer opportunities for prevention through structural changes. Nonetheless, these strategies need not be restricted to macro-level changes. Globally, intensive lifestyle diabetes prevention programmes present an evidence-based opportunity that is not reliant on environmental structural change. Diabetes prevention programmes using this model have proven effective in reducing diabetes incidence in persons in lower income communities in the USA. There is also initial evidence that patient diabetes self-management...
programmes focused on barriers to care and social determinants can improve diabetes self-management skills, health behaviours and HbA1c in low-income patients and communities. For reference, our results regarding the 2-year incidence of diabetes in high socioeconomic status as compared with low socioeconomic status areas (HR=0.80 and 0.69 in men and women, respectively) have an association with reduced diabetes incidence similar to a 1.2 kg and 2.1 kg reduction in body weight in the DPP trial. Focusing diabetes prevention efforts in lower NSEs areas may help in ameliorating health inequities. Our study provides a framework to identify areas that may require more intensive efforts by linking diabetes outcomes with readily measurable NSEs.

CONCLUSION

To conclude, our study is the first to show a social gradient in diabetes burden by contextual measures of socioeconomic status in southern Europe. The use of universal EHR of an entire population improves representability and statistical power, providing a rich representation of population health patterns. Future studies should provide targets for intervention to address this population health inequity.

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Data sharing statement Neighborhood SES indicators are available online as detailed in the appendix. Health data was obtained from the primary care system and cannot be shared due to privacy concerns.

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REFERENCES


