Contribution of type 2 diabetes to all-cause mortality, cardiovascular disease incidence and cancer incidence in white Europeans and South Asians: findings from the UK Biobank population-based cohort study

Mirthe Muilwijk,1,2 Frederick Ho,3 Heather Waddell,1,2 Anne Sillars,2 Paul Welsh,2 Stamatina Iliodromiti,2 Rosemary Brown,2 Lyn Ferguson,2 Karien Stronks,1 Irene van Valkengoed,1 Jill P Pell,3 Stuart Robert Gray1,2 Jason Martin Regnald Gill,2 Naveed Sattar,2 Carlos Celis-Morales1,2,3,4

ABSTRACT

Objective To investigate whether the health implications of having type 2 diabetes (T2D) were different in South Asian compared with white European participants.

Research design and methods Prospective data from UK Biobank were used, and 457,935 participants of white European and 7102 of South Asian background were included. Cox proportional regression was performed to investigate the association between T2D and health outcome by ethnicity.

Results Over a mean of 7.0 years (IQR 6.3–7.6) of follow-up, 12,974 participants had died, and 30,547 and 27,159 developed cardiovascular disease (CVD) and cancer, respectively. South Asians had a higher risk for CVD mortality (HR: 1.42, 95% CI 1.07 to 1.89) and incidence (HR: 1.78, 95% CI 1.63 to 1.94), but a decreased risk for cancer mortality (HR: 0.59, 95% CI 0.41 to 0.85) and incidence (HR: 0.80, 95% CI 0.70 to 0.92) compared with white Europeans. Compared with individuals without T2D, both white Europeans and South Asians with T2D had a higher risk for all-cause mortality (1.59 (1.48 to 1.71) vs 2.83 (1.76 to 4.53)), CVD mortality (2.04 (1.82 to 2.28) vs 4.40 (2.37 to 8.16)) and CVD incidence (1.37 (1.31 to 1.44) vs 1.60 (1.31 to 1.95)), respectively. However, the magnitude of the risk was higher for South Asians than white Europeans.

Conclusions Although T2D was associated with a higher risk for all-cause mortality and CVD incidence and mortality, in both white Europeans and South Asians, the risk experienced by South Asians with T2D was higher than their white European counterparts.

INTRODUCTION

Type 2 diabetes (T2D) more than doubles the risk of cardiovascular disease (CVD) and all-cause mortality.1,2 South Asians typically develop T2D 5–10 years earlier than white Europeans, transition through the ‘pre-diabetes’ phase more rapidly, and once they develop diabetes disease progression is faster and microvascular complications develop more rapidly.3–5 Moreover, South Asians are at a twofold to sixfold increased risk of T2D compared with white Europeans.6,7 It might, therefore, be expected
that T2D contributes to all-cause mortality and CVD to a larger extent in South Asians than in white Europeans. If so, the potential impact of T2D prevention among South Asians may be larger than among white Europeans.

It is already known that South Asians, as well as being at increased risk of T2D, are at increased risk for CVD. But there is limited evidence as to whether the increased CVD risk among South Asians is due to the higher T2D prevalence and how all-cause mortality risk is influenced. There are few studies to date that have investigated mortality risk in South Asians and white Europeans with and without T2D. The results from these studies have been conflicting. A recent study showed that years of life lost due to T2D is higher among white Europeans than South Asians; this was substantiated by another study by Shah et al that showed all-cause mortality risk to be lower in South Asians than in white Europeans. In contrast to this, a study by Tillin et al showed that all-cause mortality risk is higher in South Asians compared with white Europeans with T2D, but the risk is similar between South Asians and white Europeans without T2D. These conflicting results may be due to differences in study design. The study by Shah et al did not adjust for important variables including smoking and adiposity measures. Furthermore, mortality risk may have changed over time due to improvements in healthcare; the study by Tillin et al was published recently, but the follow-up period started 20 years ago. The higher CVD mortality risk in South Asians may be balanced by a lower risk for cancer, as cancer and CVD together form the leading causes of death in high-income countries, resulting in no difference in all-cause mortality risk.

The aim of the current study was to compare the health risks associated with having T2D in South Asians and white Europeans. This was investigated for all-cause mortality, CVD and cancer mortality and incidence, both in terms of the increased risk associated with having T2D and the proportion of events attributable to T2D.

**METHODS**

**Population**

Between April 2007 and December 2010, the UK Biobank invited people aged 40–69 years old from the general population to participate in the study. From the initial invitations there was a 5.5% response rate, with 502,655 participants finally agreeing to participate. Participants attended one of 22 assessment centers across England, Wales and Scotland where they completed a touch-screen questionnaire, had physical measurements taken and provided biological samples, as described in detail later. Participants who indicated that they had been diagnosed with diabetes under the age of 30 were excluded to reduce the likelihood of including those with type 1 diabetes (1,563 participants (1,462 whites and 101 South Asians) were excluded on this basis), as shown previously. Participants were only included in the total sample if the data that were provided were complete. Therefore, after the exclusion of participants who did not meet the specified criteria, 457,935 white Europeans and 7,102 South Asians were considered in our analysis.

**Procedures**

Date of death was obtained from death certificates held by the National Health Service (NHS) Information Centre (England and Wales) and the NHS Central Register Scotland (Scotland). Date and cause of hospital admissions were identified via record linkage to Health Episode Statistics (England and Wales) and to the Scottish Morbidity Records (SMR01) (Scotland). Detailed information regarding the linkage procedure can be found at [http://www.ic.nhs.uk/services/medical-research-information-service](http://www.ic.nhs.uk/services/medical-research-information-service). At the time of analysis, mortality data were available up to 31 January 2016. Mortality analysis was therefore censored at these dates, or date of death if this occurred earlier. Hospital admission data were available until 31 March 2015, resulting in disease-specific analyses being censored at this date, or the date of hospital admission or death if this occurred earlier. CVD was defined as a hospital admission or death with International Classification of Diseases (ICD) 10th Revision code I05–I89.9 (myocardial infarction, heart failure, stroke, atrial fibrillation, cerebrovascular diseases and circulatory diseases). Cancer was defined as an ICD code of C0.0–C9.9, D3.7–9 or D4.0–8.

**Measurements**

Participants classified themselves into one of 16 ethnic groups consistent with the UK Office for National Statistics Census categories. This study was restricted to participants who classified themselves in one of the following groups: white European (British, Irish and other white European) or South Asian (Indian, Pakistani and Bangladeshi). Age was calculated from dates of birth and baseline assessment. Area-based socioeconomic status was derived from postcode of residence using the Townsend score, which is derived from census data on housing, employment, social class and car availability. Other sociodemographic information such as professional qualifications (college or university, A (advanced) or O (ordinary) levels, General Certificate of Secondary Education (GCSE); or equivalent levels) and income (<£18,000, £18,000–29,999, £30,000–51,999, £52,000–100,000 and >£100,000) was self-reported at baseline.

Dietary information was collected via a self-reported dietary questionnaire (Oxford WebQ). Participants were asked how many portions of specified foods they generally ate. Subjective sleep duration was obtained by asking ‘About how many hours sleep do you get in every 24 hours?’ from which the following ordinal categories were derived: short sleeper <7 hours/day, normal sleeper 7–9 hours/day or long sleeper >9 hours/day. Physical activity was measured via the International Physical Activity Questionnaire (IPAQ) short form, with...
participants reporting the frequency and duration of walking and moderate and vigorous activity undertaken in a typical week. Data were analyzed in accordance with the IPAQ scoring protocol, and total physical activity was computed as the sum of walking and moderate and vigorous activity, measured as metabolic equivalent-hours per week. Participants were excluded from the analyses if they recorded implausible values, defined as the sum of their total physical activity, sleeping time and total screen time exceeding 24 hours.

Height, body weight and waist circumference were measured by trained nurses during the baseline assessment. Body mass index (BMI) was calculated as (weight/height$^2$). Body fat percentage was measured using standardized bioimpedance protocols. Waist circumference was measured using standardized protocol. Further details of these measurements can be found in the UK Biobank online protocol (http://www.ukbiobank.ac.uk) and elsewhere.

T2D was based on self-reported physician-diagnosed diabetes. Medical history (physician diagnosis of cancer, CVD, diabetes and hypertension) and medication history relating to diabetes, dyslipidemia and hypertension were collected from the self-completed baseline assessment questionnaire.

Statistical analyses

Participants’ baseline characteristics by T2D status and ethnicity were presented as mean and SD for continuous variables and percentages and numbers of observations for categorical variables.

To investigate the association between T2D and health outcomes of interest (all-cause mortality and CVD and cancer incidence and mortality), Cox proportional hazard models with years of follow-up as the time-scale were performed and results presented as HR and their 95% CI. The proportional hazards assumption was checked by tests using the Schoenfeld residuals. Participants who self-reported having CVD (n=16 430 Europeans; n=367 South Asians) at baseline were excluded from the analyses when the outcome of interest was CVD. Similarly, when cancer was the outcome of interest, participants with a diagnosis of cancer (n=37 377 Europeans; n=249 South Asians) at baseline were excluded.

To investigate whether the hazard for health outcomes differs between white Europeans and South Asians, Cox regression analyses with white Europeans as the reference group were performed. To investigate whether the association between T2D and health outcomes differed by ethnicity, analyses were first carried out in each ethnic group separately, using individuals without T2D as the reference group. Following this it was tested whether the effect of T2D on health outcomes differed by ethnicity, by the addition of a multiplicative interaction term between T2D and ethnicity to the model.

Incremental adjustment for confounders were used; the models included age, sex, deprivation index, income and professional qualifications (model 0), and additionally smoking, red meat intake, processed meat intake, oily fish intake, fruit and vegetable intake, the frequency of alcohol intake, sedentary time, total physical activity, sleep duration and BMI (model 1). This was in accordance with a conceptual model developed with software from DAGitty. The directed acyclic graph (DAG) is shown at dagitty.net/mvWb2B and includes considered covariates (online supplementary figure 1).

Additionally, sensitivity analyses, where BMI was replaced by waist circumference and body fat percentage, as well as a mutually adjusted model to account for a potential fat distribution confounding effect, were conducted. Another sensitivity analysis was performed to assess if mortality risk between ethnic groups was similar when participants with T2D at baseline were excluded from all analyses.

Population attributable fractions (PAF) were calculated to estimate the proportions of mortality and morbidity outcomes attributed to T2D among white Europeans and South Asians. The potential impact fractions stratified by ethnicity were also determined for a 10%, 25%, 50% and 100% reduction in T2D prevalence. All analyses were performed using STATA V.15 statistical software and p values <0.05 were considered statistically significant.

RESULTS

Baseline characteristics

At baseline, 4.4% (n=20 068) of the included white Europeans and 16.4% (n=1166) of the included South Asians had T2D (table 1). South Asian participants, both with and without T2D, were somewhat younger than white European participants. Although South Asians lived more often in deprived areas than white Europeans, a higher percentage of South Asians had a college or university degree compared with white Europeans. A higher percentage of white Europeans reported current or former smoking. Red and processed meat intake, as well as oily fish and alcohol intake, was higher in white Europeans than in South Asians. Fruit and vegetable intake, on the other hand, was higher among South Asians. Waist circumference was higher in South Asian participants without T2D, compared with white European participants without T2D. However, waist circumference was higher in white European than South Asian participants when T2D was present.

Mortality, CVD and cancer risk by ethnicity

During a mean follow-up of 7.0 years (IQR 6.3–7.6), a total of 12974 deaths occurred, 30346 participants developed CVD and 27159 developed cancer (online supplementary table S1). South Asians and white Europeans had similar rates of all-cause mortality (figure 1 and online supplementary table S1). However, in the minimally adjusted model (model 0), the hazards for CVD mortality and incidence were higher in South Asians than in white Europeans (HR 1.73 (95% CI 1.31 to 2.29), p<0.0001 and HR 1.93 (95% CI 1.78 to 2.11), p<0.0001, respectively). The hazards for
| Table 1 | Baseline characteristics |
|---------|--------------------------|
|         | White European without T2D | South Asian without T2D | White European with T2D | South Asian with T2D |
| Total participants | 437867 | 5936 | 20068 | 1166 |
| Age (years) | 56.6 (8.0) | 52.5 (8.4) | 60.2 (6.8) | 57.4 (7.9) |
| Sex (% male) | 44.6 (195 302) | 50.1 (2975) | 61.3 (12 310) | 62.5 (1166) |
| Deprivation index | | | | |
| Lower (least deprived) | 35.3 (154 382) | 17.1 (1015) | 28.4 (5691) | 13.4 (156) |
| Middle | 34.3 (149 976) | 27.3 (1618) | 32.1 (6440) | 27.4 (320) |
| Higher (most deprived) | 30.5 (133 509) | 55.6 (3303) | 39.6 (7937) | 59.2 (690) |
| Income (£) | | | | |
| <18000 | 21.4 (81 024) | 28.1 (1280) | 36.2 (6189) | 44.0 (376) |
| 18000–51 999 | 52.9 (197 060) | 45.5 (2069) | 48.9 (8356) | 39.9 (341) |
| >52 000 | 26.7 (101 231) | 26.4 (1201) | 14.9 (2546) | 16.1 (138) |
| Professional qualifications | | | | |
| CSEs | 6.2 (23 671) | 7.1 (363) | 5.5 (927) | 7.6 (76) |
| O-levels | 24.7 (94 803) | 18.4 (943) | 23.9 (4043) | 19.3 (194) |
| A-levels | 13.1 (50 154) | 9.9 (507) | 11.4 (1934) | 8.1 (81) |
| College/university degree | 37.4 (143 468) | 48.0 (2466) | 26.7 (4521) | 38.5 (386) |
| None of the above | 16.7 (371 714) | 16.7 (855) | 32.5 (5499) | 26.5 (266) |
| Smoking | | | | |
| Never | 54.7 (239 392) | 80.1 (4755) | 42.2 (8477) | 73.2 (853) |
| Previous | 35.0 (153 400) | 11.4 (677) | 47.0 (9432) | 17.1 (199) |
| Current | 10.3 (45 075) | 8.5 (504) | 10.8 (2159) | 9.8 (114) |
| Red meat intake (portions/week) | 1.94 (1.38) | 1.26 (1.48) | 2.19 (1.56) | 1.34 (1.66) |
| Processed meat intake (portions/week) | 1.87 (1.05) | 1.18 (1.17) | 2.10 (1.03) | 2.10 (1.03) |
| Oily fish intake (portions/week) | 1.64 (0.92) | 1.17 (1.04) | 1.66 (0.96) | 1.19 (1.11) |
| Fruit and vegetable intake (g/day) | 325.9 (187.2) | 397.3 (279.9) | 343.4 (194.9) | 436.3 (356.5) |
| Alcohol intake (% of TE) | 2.84 (1.47) | 4.55 (1.67) | 3.38 (1.62) | 4.82 (1.58) |
| Sedentary time (hours/day) | 5.00 (2.26) | 4.80 (2.73) | 5.87 (2.61) | 5.07 (2.94) |
| Physical activity (MET/day) | 388.0 (540.7) | 305.3 (441.9) | 317.9 (490.3) | 247.2 (393.0) |
| Physical activity recommendations | | | | |
| Inactive | 44.9 (196 496) | 53.5 (3175) | 55.9 (11 209) | 60.4 (704) |
| Active | 55.1 (241 371) | 46.5 (2761) | 44.1 (8859) | 39.6 (462) |
| Sleep duration (hours) | 7.17 (1.06) | 7.04 (1.17) | 7.24 (1.32) | 7.17 (1.36) |
| Height (cm) | 168.6 (9.2) | 163.8 (9.3) | 169.6 (9.3) | 164.3 (9.2) |
| Weight (kg) | 77.5 (15.5) | 72.4 (13.6) | 91.2 (18.6) | 77.9 (15.0) |
| BMI (kg/m²) | 27.2 (4.6) | 26.9 (4.3) | 31.7 (5.9) | 28.8 (5.0) |
| Waist circumference (cm) | 89.5 (13.1) | 90.1 (11.6) | 103.5 (14.4) | 97.7 (11.7) |
| Body fat (%) | 31.2 (8.5) | 31.7 (8.1) | 34.1 (8.6) | 32.7 (8.1) |
| Cancer diagnosed | | | | |
| No | 92.1 (402 429) | 97.0 (5736) | 90.6 (18 129) | 96.4 (1117) |
| Yes | 7.9 (34 338) | 3.0 (180) | 9.4 (1883) | 3.6 (42) |
| CVD diagnosed | | | | |
| No | 96.9 (432 802) | 96.9 (5704) | 88.3 (17 703) | 89.2 (1031) |
| Heart attack | 1.9 (8499) | 2.3 (133) | 8.7 (1747) | 8.3 (96) |
| Stroke | 1.1 (4940) | 0.9 (50) | 2.9 (589) | 2.5 (29) |

Data are mean and SD for continuous variables and percentages and numbers for categorical variables. A, advanced; BMI, body mass index; CSE, Certificate of Secondary Education; CVD, cardiovascular disease; MET, metabolic equivalent; O, ordinary; T2D, type 2 diabetes; TE, total energy.
cancer mortality and incidence were lower in South Asians than in white Europeans (HR 0.56 (95% CI 0.39 to 0.80), p=0.002 and HR 0.77 (95% CI 0.67 to 0.89), p<0.0001), respectively. Adjustment for T2D (model 2) attenuated the increased risk of CVD mortality among South Asians compared with white Europeans, and was fully attenuated in the sensitivity analyses which excluded participants with T2D (model 3). Adjustment or exclusion of participants with T2D did not change the observed higher risk for CVD or lower risk for cancer incidence among South Asians compared with white Europeans. However, the lower risk for cancer mortality observed in South Asians was attenuated when participants with T2D were excluded (figure 1 and online supplementary table S1). The adjustment for different adiposity markers (BMI, waist circumference, body fat) did not change the direction and magnitude of the associations, as presented in online supplementary table S3.

**Mortality, CVD and cancer risk by T2D and ethnicity**

The risk of all-cause mortality, CVD mortality, CVD incidence and cancer mortality was, in both white Europeans and South Asians, higher in those with T2D compared with those without T2D (figure 2 and online supplementary table S2). Although there were no significant interactions between T2D and ethnicity for health outcomes (p>0.05), the associations were somewhat stronger among South Asians with T2D, compared with white Europeans with T2D, for all-cause mortality and CVD mortality and incidence. Compared with participants without T2D, the hazard for CVD mortality was 1.99 (95% CI 1.79 to 2.21) in white Europeans with T2D, whereas South Asians with T2D had a hazard of 4.26 (95% CI 2.32 to 7.80). Similar results were observed for all-cause mortality (HR: 1.62 (95% CI 1.53 to 1.72) vs 2.46 (95% CI 1.67 to 3.63)) and CVD incidence (HR: 1.36 (95% CI 1.31 to 1.43) vs 1.62 (95% CI 1.33 to 1.97)) in people with T2D of white European and South Asian ethnicsities, respectively (figure 2 and online supplementary table S2). Further adjustment for waist circumference or body fat percentage did not...
change the direction and magnitude of the associations, as presented in online supplementary table S4.

Potential impact of T2D prevention
Modeling the impact of a reduction in T2D prevalence showed that T2D contributed to all-cause mortality, CVD mortality and CVD incidence in both South Asians and white Europeans (table 2). However, due to its high prevalence among South Asians, T2D contributed to a greater extent to all-cause mortality, CVD mortality and CVD incidence in South Asians compared with white Europeans. For example, a reduction in T2D prevalence of 50% resulted in an estimated 1.97% (95% CI 1.43% to 2.45%) and 22.29% (95% CI 6.33% to 35.57%) reduction in the CVD mortality in white Europeans and South Asians, respectively.

DISCUSSION
The results of this study confirm the increased risk of all-cause mortality and CVD mortality and incidence in participants with T2D. Although the findings of this study did not show a significant interaction of T2D and ethnicity with health outcomes, the risks associated with having T2D were greater in South Asians compared with white Europeans for all-cause and CVD mortality and incidence. T2D appears to be a stronger risk factor for all-cause mortality, CVD and cancer mortality and incidence among South Asians (Relative Risk (RR) 1.63–4.30) than white Europeans (RR 1.27–1.89). This was further compounded by the higher T2D prevalence in South Asians, rendering the potential impact of T2D reduction being greater in South Asians compared with white Europeans.

Both white European and South Asian participants had a relatively similar risk of all-cause mortality. This is thought to be because the higher risk of mortality due to CVD among South Asians is counterbalanced by the lower cancer risk. Previous studies have not reached a consensus on this matter. Shah et al.10 suggested that the mortality risk in people with T2D was lower among South Asians.
Asians when compared with white Europeans. However, the Shah et al. study included participants who had recently been diagnosed with T2D. This was in contrast to the current study, where we included participants who had already been diagnosed with T2D at baseline. Therefore, participants within the current study had a longer duration of T2D. Additionally, the study by Shah et al. did not adjust for smoking or adiposity levels, with smoking rates generally much lower (as is also the case in this study) among South Asians compared with white Europeans. Study settings may also have led to the contrasting results, with the current study conducted in the UK and the study by Shah et al. conducted in Canada. Differences including those related to healthcare may have led to the observed study differences.

The results of the current study agree with the findings from a previous study conducted in the UK by Tillin et al., which suggested that the risk of CVD is higher in South Asians, relative to white Europeans, with T2D but not those without T2D. Although results from the study of Tillin et al. were published recently, follow-up started between 1988 and 1991. It was speculated therefore that over time mortality risk may have changed due to an improvement in CVD management. In addition, in past decades greater attention has been paid to the differences in disease risk between ethnicities within healthcare, for example, in diabetes screening. It is, however, interesting to note that, according to the current study, these changes have not affected the higher CVD and all-cause mortality risk among South Asians compared with white Europeans.

In accordance with previous studies, this study reported differences in CVD and cancer risk between white European and South Asian participants. In the future, it might be important to consider that when the high CVD risk among South Asians is better managed in these populations, this may impact on cancer risk. The higher risk for CVD mortality may partly result in the lower cancer mortality among South Asians, but if CVD mortality was lowered in South Asians then the risk of developing cancer due to a higher life expectancy may increase. On the other hand, South Asians may have a high prevalence of genetic polymorphisms that are protective of cancer, suggesting that the low cancer risk may actually be conserved in the case that the interplay with environmental risk factors is not changed.

Adjustment for T2D only partially explained the increased risk of CVD mortality among South Asians compared with white Europeans for CVD mortality. However, sensitivity analyses were used to exclude participants with T2D, which resulted in the full attenuation of the higher risk for CVD mortality in South Asians. However, it cannot be discarded that this may be attributable to our limited power to detect an association when T2D cases were excluded. In contrast to this finding, the excess CVD incidence risk in South Asians, compared with white Europeans, was not altered after the adjustment for T2D or exclusion of T2D cases from the analyses, resulting in a slight attenuation of the CVD risk.

Although this study provides evidence that T2D may not fully explain the higher CVD risk experienced by South Asians, further investigation is needed to underpin the risk factors that could play a key role in the increased T2D and CVD risk in South Asians. Especially since T2D and CVD share common denominators that may explain both the increased risk T2D and CVD risk, preventive measures for T2D may also impact on CVD risk independent of its development through T2D. Unfortunately, the underlying causes for the higher susceptibility to both CVD and T2D among South Asians compared with other populations remain unclear despite many hypotheses, mainly related to socioeconomic, genetic, developmental and lifestyle factors, having been investigated. However, preventive measures could still target known risk factors that impact T2D risk among both South Asians and white Europeans, such as dietary intake and physical activity.

Our PAF analyses highlight that the contribution of T2D to the CVD incidence and mortality in the general population is larger among South Asians than among white Europeans due to the higher prevalence of T2D in South Asians than in white Europeans. A reduction in the prevalence of T2D had the largest potential impact among South Asians, and therefore we suggest that preventive measures to prevent or delay the onset of T2D may have the largest impact within the South Asian population.

Limitations

This study is not exempt from limitations. First, the diagnosis of T2D within the current study was based on self-reporting of physician-diagnosed T2D. This may have biased our results as some participants may not recall their diagnosis accurately, at all, or may not want to report it. This bias may be greater if self-reporting was more accurate in one ethnic group compared with others. Furthermore, screening rates for T2D may be different among ethnic groups due to ethnic differences in the prevalence of T2D. However, previous studies using UK Biobank have shown that the use of self-reported diabetes is a valid method and can distinguish between participants with and without diabetes. Moreover, studies on the validity of self-reported physician-diagnosed diabetes have shown both excellent positive and negative predictive value.

In addition, misclassification of the type of diabetes was limited by excluding participants with diabetes diagnosed under the age of 30 years. Second, analyses were adjusted for confounders in accordance with the DAG. It can, however, not be excluded that analyses were overadjusted for confounders. This may have partly led to filtering out the direct effect of T2D on mortality. Third, participants included in the UK Biobank study may not be a representative sample of the British population due to ‘healthy volunteer’ selection bias. Therefore, future studies are needed to confirm the findings in representative populations of the UK but also in other countries where South
Asians represent a greater proportion of the population. Finally, South Asian populations consist of a very heterogeneous group of people with a wide variety of population characteristics. Heterogeneity exists both in genetic etiology and in lifestyle factors depending on ancestral background. Since diabetes is a common factor affecting all South Asian populations, the different underlying populations were studied together, but in future studies subpopulation-specific figures may be calculated.

CONCLUSION
In conclusion, South Asians have a higher risk of CVD mortality and incidence but a lower risk of cancer mortality and incidence compared with white Europeans. Although T2D was associated with a higher risk for all-cause mortality and CVD incidence and mortality, in both white Europeans and South Asians, the risk experienced by South Asians with T2D was higher than their white European counterparts.

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Patient consent for publication Not required.

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Data availability statement Data may be obtained from a third party and are not publicly available.

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ORCID iDs
Mirithe Muliwijk http://orcid.org/0000-0002-3686-5116

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