Diabetic nephropathy (DN) is the most common cause of end-stage renal disease (ESRD) in many countries (1) and has significant impact on patients and health care systems (2). DN progresses slowly, starting with microalbuminuria, which progresses into overt proteinuria in 20–40% of patients, and 20% of patients will have progressed to ESRD within 20 years after onset of overt proteinuria (1). The speed of DN progression is variable and largely dependent on blood pressure (BP), obesity, metabolic control, and other factors such as male sex and ethnicity (3,4).

The pathogenesis of DN is thought to be similar to other microvascular complications in which hyperglycemia and hypertension are thought to be fundamental to its development, as they promote increased oxidative and nitrosative stress (3). In addition, hemodynamic changes occur as a result of the activation of the renin-angiotensin-aldosterone (RAAS) and endothelin systems, resulting in increased systemic and intraglomerular pressure, causing hyperfiltration and albuminuria (3). Despite attempts to improve metabolic control and RAAS inhibition, DN remains very common, and many patients develop ESRD requiring renal replacement therapy. Hence, better understanding of the pathogenesis of DN is needed in order to develop more effective treatments.

Several reports, including our own, have shown a high prevalence of obstructive sleep apnea (OSA) in patients with type 2 diabetes (5,6). Patients with OSA and type 2 diabetes are at increased risk of diabetic peripheral neuropathy (6). Furthermore, OSA is associated with increased oxidative and nitrosative stress as well as impaired microvascular regulation in patients with type 2 diabetes (6). Hence, it is plausible that OSA complicating type 2 diabetes could facilitate the development and progression of microvascular complications including DN.

The primary aims of this study were to assess the relationship between OSA and DN and to assess the impact of OSA on the estimated glomerular filtration (eGFR) decline in patients with type 2 diabetes. Secondary aims were to assess the impact of OSA on the development of albuminuria and to explore the mechanisms by which OSA and DN could be linked.

**OBJECTIVE**—Diabetic nephropathy (DN) is a leading cause of end-stage renal disease (ESRD). Obstructive sleep apnea (OSA) is common in type 2 diabetes and increases oxidative stress. Hence, OSA could promote the development and progression of DN.

**RESEARCH DESIGN AND METHODS**—This was a cohort study in adults with type 2 diabetes. Patients with known OSA or ESRD were excluded. DN was defined as the presence of albuminuria or an estimated glomerular filtration rate (eGFR) <60 mL/min/1.73 m². DN progression was based on eGFR measurements. OSA was defined as apnea hypopnea index (AHI) ≥5 events/h. Serum nitrotyrosine abundance (a marker of nitrosative stress) was measured by ELISA.

**RESULTS**—A total of 224 patients were included. OSA and DN prevalence was 64.3 and 40.2, respectively. DN prevalence was higher in patients with OSA (OSA+) compared with those without OSA (OSA−) (49.3% vs. 23.8%, P < 0.001). After adjustment, OSA (odds ratio 2.64 [95% CI 1.13–6.16], P = 0.02) remained independently associated with DN. After an average follow-up of 2.5 (0.7) years, eGFR decline was greater in OSA+ compared with OSA− patients (median −6.8% [interquartile range −16.1 to 2.2] vs. −1.6% [−7.7 to 5.3%], P = 0.002). After adjusting, both baseline OSA (B = −3.8, P = 0.044) and AHI (B = −4.6, P = 0.02) remained independent predictors of study-end eGFR. Baseline serum nitrotyrosine abundance (B = −0.24, P = 0.015) was an independent predictor of study-end eGFR after adjustment.

**CONCLUSIONS**—OSA is independently associated with DN in type 2 diabetes. eGFR declined faster in patients with OSA. Nitrosative stress may provide a pathogenetic link between OSA and DN. Interventional studies assessing the impact of OSA treatment on DN are needed.
between 2009 and 2010 and were followed until the end of 2012. Patients with respiratory disease including previously diagnosed OSA or ESRD receiving renal replacement therapy were excluded. Patients were recruited consecutively from the diabetes clinics of two U.K. hospitals. Patients were approached consecutively in the waiting area by the investigator or a research nurse without any prior knowledge of their medical condition. Consent was obtained and ethnicity determined in accordance with the U.K. decennial census by the study participants. The project was approved by the Warwickshire Research Ethics Committee (REC no. 08/H1211/145).

OSA assessment
OSA was assessed by a single overnight home-based cardiorespiratory sleep study using a portable multichannel device (Alice PDX; Philips Respironics) and scored in accordance with the American Academy of Sleep Medicine guidelines (7). An apnea hypopnea index (AHI) ≥5 events/h was consistent with the diagnosis of OSA (8). Sleep studies with <4 h of adequate recordings were repeated and excluded if the quality remained poor. OSA severity was assessed based on the AHI categories (≤5 events/h no OSA, 5 to <15 mild OSA, 15 to ≤30 moderate OSA, and ≥30 severe OSA). AHI (as a continuous variable), the time spent with oxygen saturations <80%, and the nadir oxygen saturations during sleep. Patients diagnosed with OSA were referred to the sleep clinic and treated as per routine clinical care. Patients with no or mild OSA were not offered continuous positive airway pressure (CPAP) treatment, while all patients with moderate to severe OSA were offered CPAP. Patients with moderate to severe OSA were hence divided into those compliant with CPAP (average usage ≥4 h/night for 70% of days) and those not compliant (including patients who declined treatment) (9). Data regarding CPAP compliance were downloaded directly from the CPAP equipment.

DN assessment
DN was assessed using eGFR, calculated using the four-variable Modification of Diet in Renal Disease equation (10) and the urinary albumin-to-creatinine ratio (ACR) of a single early-morning urine measurement. Microalbuminuria was defined as ACR >3.4 mg/mmol, and macroalbuminuria was defined as ≥30 mg/mmol (11-13). Urine samples with evidence of urinary tract infection were repeated when free from infection. DN was defined as the presence of albuminuria (micro or macro) or an eGFR <60 mL/min/1.73 m² (14). ACR and eGFR were measured at baseline and study end. Study-end measurements were taken during patients visits to the follow-up appointments of the diabetes clinic. eGFR measurements during acute illness or after imaging that used contrast were excluded.

Nitrosative stress assessment
To explore possible mechanisms that might explain the relationship between OSA, DN, and progression of eGFR, we assessed serum nitrotyrosine levels. The rationale for the selection of serum nitrotyrosine was based upon prior reports of an association between nitrosative stress and DN in animal studies (15,16) as well as a single report that identified high levels of nitrotyrosine in the kidneys of patients with diabetes (17). All study participants were approached, and serum 3-nitrotyrosine was assessed in duplicate in all subjects who agreed using commercially available ELISA (Oxiselct; Cell Biolabs, San Diego, CA). The measurements had good precision (<10% difference between duplicate samples) and high reproducibility (correlation coefficient between repeated measurements = 0.9 and a mean [SEM] coefficient of variance of 10.9% [2.4%]). All assessments in the study (renal function, nitrosative stress, and biochemical profiles) were conducted blind to OSA status.

Outcome measures and analyses
Data analysis was performed using SPSS 21.0 software (SPSS, Chicago, IL). Data are presented as mean (SD) or median (interquartile range [IQR]). Independent continuous variables were compared using the Student t test or the Mann-Whitney test. Categorical variables were compared using the χ² test. Correlations between continuous variables were performed using the Pearson or Spearman tests. Differences between independent groups were assessed by ANOVA.

DN
For assessment of whether OSA status, OSA severity, or hypoxemia measures are independently associated with DN, multiple logistic regression (forced entry) was used. Multiple linear regression (forced entry) was used to assess independent associations of continuous variables. Proportion of variation explained by models (R²) plus statistical significance and effect size of OSA, OSA severity, and hypoxemia measures are reported. Variables included in both the logistic and linear regression models were based on known outcome-related risk factors or variables that differed between patients with and without OSA.

eGFR progression
For assessment of the impact of OSA on eGFR progression, only patients who had eGFR levels measured at baseline and study end were used. Baseline differences were accounted for by using linear regression with eGFR at study end and eGFR change from baseline as the outcome measures and OSA, baseline eGFR, and other confounders as the predictors. A repeat analysis assuming that baseline eGFR had not changed in those with missing eGFR at study end was performed. For assessment of progression of albuminuria, only patients with normal ACR at baseline were included and logistic regression was used.

Residuals and collinearity were considered in assessing fit of models to data. Sequentially removing variables involved in multicollinearity had limited impact on models estimates for the main exposure. Hence, final models presented include variables based on the known outcome-related risk factors, possible confounders, or variables that differed between patients with and without OSA, regardless of the presence of collinearity. Removing variables in order to minimize collinearity had little impact on the regression models.

Matched-groups analysis
For further exploration of the impact of baseline differences on the associations observed, a subgroup of 71 patients with and 59 without OSA were group matched for key DN risk factors. A P value < 0.05 was considered significant in all statistical testing.

RESULTS—A total of 266 patients consented for the project; data regarding DN (both eGFR and albuminuria) at baseline were available in 224 patients. Baseline and study-end eGFR measurements were available in 196 patients. Please see Fig. 1 for details.

Baseline data
The prevalence of OSA was 64.3% (144 of 224), with 38.4% (86 of 224) mild and 25.9% (58 of 224) moderate to severe. The prevalence of DN was 40.2% (90 of 224),
with 33.0% (74 of 224) of patients exhibiting albuminuria, and 10.3% (23 of 224) was macroalbuminuria. The eGFR was 90, 60–89, 30–59, 15–29, and 15 mL/min/1.73 m² in 45.5% (102 of 224), 37.9% (85 of 224), 15.2% (32 of 224), 1.3% (3 of 224), and 0% (0 of 224) of patients, respectively.

Baseline patient characteristics are summarized in Table 1. As expected, patients with OSA were older, heavier, and had a longer duration of diabetes and higher systolic BP, which required more antihypertensive medications. Patients with and without OSA had similar glycaemic control and total cholesterol, as well as similar use of RAAS inhibitors.

DN is more common in patients with OSA. Patients with OSA had a higher prevalence of DN, serum creatinine, albuminuria, and macroalbuminuria and lower eGFR levels regardless of ethnicity (Table 2). OSA is independently associated with DN: multivariable analysis. In multiple logistic regression, OSA was independently associated with DN despite adjustments for potentially confounding variables (Table 3). Other independent associations with DN included age, diabetes duration, HbA1c, and triglycerides. The relationship between OSA severity and hypoxemia measures is summarized in the Supplementary Data.

**Consort diagram for study participants. COPD, chronic obstructive pulmonary disease.**

**Longitudinal analysis**

As indicated in Fig. 1, at study end eGFR data were available on 196 (124 [63.3%] with and 72 [36.7%] without OSA) patients and albuminuria data on 163 (105 [64.4%] with and 58 [35.6%] without OSA) patients. The follow-up analysis was based only on patients in whom both baseline and follow-up data were available, which comprised a total of 196 patients for eGFR progression, 163 for albuminuria progression, and 169 for DN progression. The average length of follow-up was 2.5 (0.7) years. There was no difference in the follow-up duration between patients with and without OSA (2.5 [0.6] vs. 2.4 [0.7], P = 0.51). Comparing patients who had study end point data with those who were lost to follow-up showed that the two groups were largely comparable.

OSA and the AHI are associated with a greater decline in eGFR. eGFR declined from baseline to the study end in patients with OSA (81.7 [27.8] vs. 75.2 [28.8] mL/min/1.73 m², P < 0.001) and without OSA (90.9 [23.2] vs. 89.2 [21.9] mL/min/1.73 m², P = 0.1), but eGFR change was only significant in patients with OSA. The change in eGFR over the follow-up period was larger in the OSA group (−5.0 [−14 to 1.8] vs. −1.5 [−8.5 to 4.0] mL/min/1.73 m², P = 0.006). Calculating the change in eGFR as a percentage of baseline eGFR showed a greater decline in the OSA group (−6.8% [−16.1 to 2.2%] vs. −1.6% [−7.7 to 5.3], P = 0.002) in a stepwise manner (−1.4% [−7.7 to 5.2%] vs. −5.3% [−16.5 to 2.7%] vs. −8.7% [−16.1 to 2.0], P = 0.003, for no OSA vs. mild vs. moderate to severe OSA) (Fig. 2). Rapid eGFR decline, defined as 4% decline of eGFR per year (18), was more common in patients with OSA compared with those without OSA (39.5% (n = 49) vs. 20.8% (n = 15), P = 0.007, respectively).

In a linear regression model, baseline eGFR (R² = 0.84, adjusted R² = 0.84) was a predictor of study-end eGFR (B = 0.94, P < 0.001). After addition of OSA, age at diagnosis, diabetes duration, ethnicity, sex, BMI, mean arterial pressure, antihypertensive agent use, HbA1c, insulin use, oral antidiabetic agent use, total cholesterol, triglycerides, lipid-lowering therapy, antiplatelet use, and smoking to the model (R² = 0.86, adjusted R² = 0.85), OSA remained an independent predictor of study-end eGFR (B = −3.8, P = 0.04). The only other independent predictors were diabetes duration and baseline eGFR. After we replaced BMI with waist circumference (R² = 0.86, adjusted
Table 1—Participant characteristics in relation to OSA status

|                | OSA− | OSA+ | P    |
|----------------|------|------|------|
| n              | 80   | 144  |      |
| Male           |      |      |      |
| White Europeans|      |      |      |
| Age (years)    | 54.8 | 58.7 | 0.02 |
| Diabetes duration (years) | 9.5 | 11 | 0.03 * |
| BMI (kg/m²)    | 31.6 | 35.4 | 0.18 |
| Waist circumference (cm) | 106.8 | 116.9 | <0.001 |
| Waist-to-hip ratio | 0.98 | 1.0 | 0.18 |
| Neck circumference (cm) | 38.7 | 42.8 | <0.001 *
| Systolic BP (mmHg) | 125.8 | 132.7 | 0.004 |
| Diastolic BP (mmHg) | 77.0 | 70.5 | 0.29 |
| Mean arterial pressure | 93.3 | 96.6 | 0.03 |
| Total cholesterol (mmol/L) | 8.1 | 8.3 | 0.33 |
| Triglycerides (mmol/L) | 1.5 | 1.8 | 0.03 * |
| HDL (mmol/L)    | 1.2  | 1.3  | 0.03 * |
| Epworth sleepiness score | 5.0 | 8.0 | 0.003 * |
| Smoking (current or ex-smoker) | 31.38 | 57.39 | 0.09 |
| Alcohol (drinks alcohol) | 12 | 49 | 0.002 |
| Oral antidiabetes treatment | 78 | 130 | 0.044 |
| Insulin         | 33   | 85   | 0.01 |
| Insulin dose (units) | 60 | 81.5 | 0.007 * |
| GLP-1 analogs   | 5    | 20   | 0.08 |
| RAAS inhibitors | 53   | 108  | 0.16 |
| Duretics        | 18   | 64   | 0.001 |
| Antihypertensive agents | 59 | 125 | 0.02 |
| Number of antihypertensive agents used | 1 | 2 | <0.001 * |
| Lipid-lowering treatment | 69 | 119 | 0.48 |
| Anti-platelets  | 49   | 104  | 0.091 |
| NSAID use       | 2    | 3    | 1    |
| Stroke or TIA   | 5    | 17   | 0.18 |
| Ischemic heart disease | 14 | 32 | 0.40 |

Data presented as median (IQR) or mean (SD) unless otherwise indicated. Categorical variables are presented as n (% of OSA status). GLP-1, glucagon-like peptide-1; NSAID, nonsteroidal anti-inflammatory drugs; TIA, transient ischemic attack. Analysis performed using the χ² test for categorical variables, the independent t test for normally distributed variables, and the Mann-Whitney U test* for non-normally distributed variables. *P value was calculated assuming unequal variance.

R² = 0.85), OSA remained independently associated with study-end eGFR (B = -4.2, P = 0.03).

Replacing OSA with AHI as a marker of OSA severity in the above model demonstrated that baseline AHI was an independent predictor of study-end eGFR (B = -4.6, P = 0.02). Nocturnal hypoxemia measures were not independent predictors of study-end eGFR after adjustment.

Further analysis was performed by replacing the missing study-end eGFR measurements with the baseline eGFR measurements (giving a sample size of 234) and showed that baseline OSA (R² = 0.88, adjusted R² = 0.87) (B = -3.89, P = 0.02) and AHI (B = -3.45, P = 0.04) remained independent predictors of study-end eGFR.

The use of eGFR change instead of study-end eGFR as the outcome measure in the above-described regression models confirmed that OSA was an independent predictor of eGFR decline (Supplementary Data). For the impact of OSA on the development of albuminuria, see the Supplementary Data.

**Matched group analysis.** We aimed to minimize the clinical differences between patients with and without OSA by re-examining our findings in a subgroup of patients who were group matched for key diabetes risk factors (Supplementary Table 1). Despite matching for important baseline differences, as predicted by the statistical adjustments, DN prevalence remained significantly higher in patients with compared with those without OSA (46.5 vs. 28.8%, P = 0.039). The change in eGFR in absolute terms (median -5.0 [IQR -14.0 to 1.0] vs. -2.0 [-7.0 to 5.0] mL/min/1.73 m², P = 0.02) and as percent of baseline eGFR (−5% [−20 to 2%] vs. −2% [−8 to 6%], P = 0.016) was greater in the OSA group compared with those without OSA.

**CPAP treatment and the longitudinal analysis.** Patients who were CPAP compliant had a slower eGFR decline than noncompliant patients (Supplementary Data).

**Serum nitrotyrosine: an independent predictor of study-end eGFR.** Serum nitrotyrosine levels were available from 87 patients who had measurements of both baseline and study-end eGFR. There were no significant differences between patients who agreed and those who declined to provide serum samples (6). Serum nitrotyrosine levels were higher in patients with DN at baseline (n = 38) compared with those without (n = 45) (median 23.3 [IQR 17.7–34.8] vs. 20.8 [12.0–29.6] nmol/L, P = 0.04). There was also a modest correlation between serum nitrotyrosine abundance and baseline eGFR (r = −0.24, P = 0.03) as well as study-end eGFR (r = −0.26, P = 0.02). Serum nitrotyrosine correlated with the eGFR change when presented as percentage of baseline eGFR (r = −0.36, P = 0.001). Serum nitrotyrosine was nonsignificantly higher in patients with OSA compared with those without (22.8 [17.2–33.6] vs. 15.5 [11.8–26.5] nmol/L, P = 0.058).

For assessment of the relationship between serum nitrotyrosine and eGFR progression, multiple linear regression was used with study-end eGFR as the outcome measure and baseline eGFR and serum nitrotyrosine as the predictors (R²=0.84, adjusted R²=0.84). With use of this model, serum nitrotyrosine was an independent predictor of study-end eGFR (B = -0.25, P = 0.001). After addition of age at diabetes diagnosis, diabetes duration, sex, ethnicity, HbA1c, total cholesterol, triglycerides, mean arterial pressure, BMI, smoking, insulin use, antihypertensive use, and lipid-lowering therapy to the model (R² = 0.87, adjusted R² = 0.85), serum nitrotyrosine remained an independent predictor of study-end eGFR (B = -0.20, P = 0.02). Other independent predictors included baseline eGFR and diabetes duration. Interestingly, adding OSA to the same linear regression model (R² = 0.88, adjusted R² = 0.85) showed that serum nitrotyrosine remained an independent predictor of study-end eGFR (B = -0.20, P = 0.017), but OSA was not (B = -3.7, P = 0.23).
CONCLUSIONS—Our aim was to assess the relationship between OSA and DN and to study the impact of OSA on DN progression in patients with type 2 diabetes. We demonstrated a robust association between OSA and DN, which persisted after adjustment for a wide range of demographic and clinically relevant confounders. Importantly, baseline OSA status and AHI were independent predictors of future eGFR and eGFR change over an average of 2.5 years of follow-up. Serum nitrotyrosine abundance emerged as an independent predictor of study-end eGFR as well as the change of eGFR over the time course of the study despite adjustment for multiple confounders, including OSA. We have also identified a high prevalence of undiagnosed OSA in patients with type 2 diabetes.

The study sample was drawn from patients attending secondary care units for diabetes in the U.K. The participant characteristics were similar to those reported previously from a different region in the U.K. (19), suggesting that the current study sample was representative of the wider type 2 diabetic population in secondary care. However, whether our findings are applicable to patients typically managed in primary care and those with a shorter duration of diabetes remains to be examined. OSA prevalence in our sample is also consistent with other reports (5,20). Similarly, the prevalence of DN and albuminuria is consistent with that reported by others in patients with type 2 diabetes (12). It is important to note that patients with ESRD were excluded from our study. Patients with ESRD are well known to have a high prevalence of OSA (30–80%) (21); hence, our results are novel and important, as they apply to patients at an earlier stage of renal dysfunction.

As expected, demographic and metabolic factors differed between patients with and without OSA. Nevertheless, although these differences contributed to the observed relationship between OSA and DN, OSA remained independently associated with DN even after adjustment. Furthermore, OSA remained an independent predictor of study-end eGFR and eGFR change over the follow-up period after adjustment for a wide range of confounders. However, much of the variation of study-end eGFR is explained by baseline eGFR, which is consistent with previous studies (22). The matched group analysis also supported the findings of the multivariable analysis.

This is the first study to assess the impact of OSA on DN in patients with type 2 diabetes. Recently, some researchers postulated that sleep disorders (including OSA) might be a modifiable risk factor for the development and progression of chronic kidney disease, but to date there are little data to support such a link (23). The available evidence for a relationship between OSA and chronic kidney disease is generally conflicting and obtained from cross-sectional studies not specifically targeting patients with diabetes (24–31). In a study of 505 men, there was no association between eGFR and OSA after adjustment (24,25). In a cross-sectional study of 91 obese adults, there was no association between albuminuria and OSA, but AHI was independently associated with serum creatinine after adjustment (26). In another study of 496 adults, AHI was independently associated with albuminuria after adjustment.

Table 2—Relationship between OSA and DN in patients with type 2 diabetes in the total study population and in ethnicity subgroups

|                          | OSA+ | OSA− | P     |
|--------------------------|------|------|-------|
| Total cohort             | n = 80 | n = 144 | <0.001 |
| DN                       | 19 (23.8%) | 71 (49.3%) | |
| Albuminuria              | 16 (20.0%) | 58 (40.3%) | 0.002 |
| Macroalbuminuria         | 4 (5.0%) | 19 (13.2%) | 0.05  |
| Serum creatinine (µmol/L)| 74.4 (23.4) | 90.9 (36.8) | <0.001 |
| eGFR (mL/min/1.73 m²)    | 92.9 (25.1) | 82.2 (27.6) | 0.005 |
| eGFR <60 mL/min/1.73 m²  | 5 (76.3%) | 32 (22.2%) | 0.002 |

White Europeans  

|                          | n = 30 | n = 90 | P     |
|--------------------------|------|------|-------|
| DN                       | 5 (16.7%) | 42 (46.7%) | 0.004 |
| Albuminuria              | 4 (13.3%) | 32 (35.6%) | 0.02  |
| Macroalbuminuria         | 0 (0%) | 10 (11.1%) | 0.07  |
| Serum creatinine (µmol/L)| 72.6 (26.8) | 93.3 (36.6) | 0.005 |
| eGFR (mL/min/1.73 m²)    | 89.9 (28.6) | 80.0 (27.3) | 0.09  |
| eGFR <60 mL/min/1.73 m²  | 3 (10.0%) | 23 (25.6%) | 0.07  |

South Asians  

|                          | n = 50 | n = 54 | P     |
|--------------------------|------|------|-------|
| DN                       | 14 (28.0%) | 29 (53.7%) | 0.008 |
| Albuminuria              | 12 (24.0%) | 26 (48.1%) | 0.01  |
| Macroalbuminuria         | 4 (8.0%) | 9 (16.7%) | 0.2   |
| Serum creatinine (µmol/L)| 75.5 (21.3) | 86.9 (37.0) | 0.058 |
| eGFR (mL/min/1.73 m²)    | 94.7 (22.8) | 85.9 (27.9) | 0.08  |
| eGFR <60 mL/min/1.73 m²  | 2 (4.0%) | 9 (16.7%) | 0.04  |

Data are mean (SD) or n (%) unless otherwise indicated.

Table 3—Assessing the impact of possible confounders on the association between OSA and DN using logistic regression models

|                  | Nagelkerke R² | OR  | 95% CI          | P     |
|------------------|---------------|-----|-----------------|-------|
| Unadjusted       | 0.09          | 3.12| 1.70–5.75       | <0.001|
| Model 1          | 0.26          | 3.21| 1.59–6.47       | 0.001 |
| Model 2          | 0.43          | 2.38| 1.07–5.31       | 0.03  |
| Model 3          | 0.46          | 2.64| 1.13–6.16       | 0.02  |

The odds ratios (ORs) reported are the odds for having DN in OSA+ compared with OSA− patients. All patients with DN data (n = 224) were included in all models. Model 1 is adjusted for sex, ethnicity, age, and diabetes duration. Model 2 is adjusted for model 1, BMI, mean arterial pressure, HbA₁c, triglycerides, treatment with insulin, glucagon-like peptide-1 analogs, and antihypertensives. Model 3 is adjusted for model 2 plus total cholesterol, HDL, lipid-lowering treatment, aspirin, oral antidiabetes agents, alcohol (units/week), and smoking (current or ex-smoking vs. none). All medication use was inserted into the model as yes vs. no variables. Antihypertensives were inserted into the model individually and include RAAS inhibitors, diuretics, α-blockers, β-blockers, and calcium antagonists. Model 2 is the main model that is produced based on the main confounders, the model best, and minimizing multicollinearity. Model 3 is a fully adjusted model regardless of multicollinearity. Replacing BMI with waist circumference (or inserting them both together in the model) does not change the results significantly. The ORs are for OSA vs. no OSA.
(27), while other studies failed to identify such a relationship (28,29). For assessment of the issue of hypertension as a possible confounder for the effects of OSA on renal function, a cross-sectional study of 62 untreated hypertensive patients with OSA matched to 70 hypertensive patients without OSA found that albuminuria was 57% greater in patients with OSA versus those without and that albuminuria correlated with AHI and OSA severity (30).

Interestingly, our data demonstrated that even mild OSA (based on AHI) is associated with DN and worsening eGFR prospectively. We postulate that this may reflect the impact of intermittent hypoxemia, which is exaggerated in the presence of vulnerable tissue damaged because of chronic hyperglycemia. This finding is important for future interventional studies; as current guidelines do not recommend treating patients with mild OSA.

Although our data demonstrate an independent association between OSA and DN and that OSA is an independent predictor of worsening eGFR, OSA was not a predictor of the development of DN or albuminuria. This could be explained by the relatively short duration of follow-up or the small sample size of the subgroup that did not have DN (and/or albuminuria) at baseline. Another possible explanation is that OSA may not necessarily result in the development of DN, but once DN is present, OSA might contribute to a rapid progression and a decline in renal function. Furthermore, eGFR decline can be the only manifestation of DN progression in patients with diabetes without any evidence of albuminuria in up to 30% of cases (14,32).

There are several mechanisms that might explain the association between OSA and DN and the impact of OSA on eGFR. We have previously shown that OSA is associated with increased oxidative and nitrosative stress and impaired microvascular and endothelial regulation in patients with type 2 diabetes (6). Studies in patients with OSA but without diabetes have shown that OSA and/or intermittent hypoxia are associated with increased advanced glycation end products (33), altered protein kinase C signaling (34), decreased endothelial nitric oxide synthase and increased endothelin-1 levels (35), hypercoagulability (increased plasminogen activator inhibitor-1) (36), and increased inflammation (37). The repetitive episodes of reoxygenation after hypoxemia in OSA patients simulate ischemia-reperfusion injury, which results Figure 2 — The longitudinal impact of OSA on eGFR in patients with type 2 diabetes presented as eGFR change from baseline (A) and percentage of baseline eGFR (B and C). Data presented as mean and SEM.
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in the generation of reactive oxygen species (37). We have previously shown that OSA, AHI, and nocturnal hypoxemia are independently associated with serum nitrotyrosine abundance after adjustment for confounders in patients with type 2 diabetes (6). Our results give possible mechanistic insights into the pathophysiology of DN, since serum nitrotyrosine abundance emerged as an independent predictor of study-end eGFR despite adjustment for many possible confounders, suggesting that nitrosative stress is important in the progression of eGFR regardless of the cause of nitrosative stress (whether it was OSA, hyperglycemia, or hypertension). Our data are consistent with reports indicating an association of nitrotyrosine with DN in rodent models (38) and cell culture systems (39) and the demonstration of increased nitrotyrosine staining in the renal proximal tubules and thin limb of the loop of Henle in patients with DN (17). Furthermore, amelioration of nitrosative stress has been associated with improvement in DN in diabetic rodent studies (15,16). Interestingly, serum nitrotyrosine was a predictor of study-end eGFR even when OSA was included in the model. It is therefore tempting to speculate that nitrosative stress could mediate the relationship between OSA and eGFR, but our sample size is too small to confidently draw such a conclusion and it seems likely that OSA might influence eGFR decline via multiple mechanisms such as those described above.

The impact of CPAP treatment was not in the scope of this study. Interestingly, in our cohort, in patients with OSA, the decline in eGFR was smaller in the CPAP-compliant group than those who were noncompliant with CPAP. Additionally, a larger proportion of patients in the noncompliant CPAP group developed albuminuria during follow-up. However, it is difficult to assess CPAP efficacy from observational data owing to the small numbers of patients who were compliant with CPAP treatment. Nonetheless, these data provide further justification for assessing the impact that CPAP can have on the development and progression of DN but also highlight the challenges of CPAP compliance in patient with type 2 diabetes.

Our study has several strengths and limitations. Use of home-based portable multichannel respiratory devices rather than inpatient overnight polysomnography may be considered a limitation, but this approach is well established and validated (40). We used a single random measurement of urine albumin to assess albuminuria rather than three measurements; again, this approach has been used previously by many other investigators (12,13), and recent data suggest that single urinary albumin measurements are accurate in predicting nephropathy and appropriate for epidemiological studies (13). The cross-sectional analysis of the relationship between OSA and DN makes it difficult to ascertain causality, but the longitudinal analysis of the impact of OSA on eGFR decline suggests a cause-effect relationship, although this needs to be tested in prospective interventional studies. The missing follow-up data, particularly for albuminuria, is a weakness and a potential source for bias; however, no differences in characteristics of participants versus patients lost to follow-up were observed. A strength of our study was the well-characterized population, with measurement of a wide range of demographic and clinical variables, which allowed adjustment for a wide range of potential confounders. Furthermore, the study included patients of both sexes and South Asian and white European ethnicity and was prospective, which allowed the assessment of the impact of OSA on eGFR decline. In addition, we focused on patients relatively early in the course of DN; this population would be a prime target for treatments that might slow the progression toward ESRD.

OSA is very common in patients with type 2 diabetes attending secondary care units in the U.K. Patients with OSA and type 2 diabetes are more likely to have DN compared with those with type 2 diabetes but without OSA despite adjustment for a wide range of confounders. Renal function (assessed by eGFR) declined faster in patients with OSA and type 2 diabetes compared with those with type 2 diabetes alone. The eGFR decline in patients with OSA was very clinically meaningful, as more patients in the OSA group had a rapid decline in eGFR (≥10% over 2.5 years), and indeed, 25% of patients with OSA exhibited an eGFR decline >16%. Nitrosative stress may provide an important pathogenic link between OSA, type 2 diabetes, and DN. This study could form the basis for interventional studies to examine the impact of OSA treatment on the development and progression of DN in patients with type 2 diabetes.

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A.A.T. conceived and designed the study, contributed to analysis and interpretation, wrote the first draft of the manuscript, and gave final approval of the manuscript. A.A. designed the study, reviewed the draft of the manuscript, and gave final approval of the manuscript. N.T.R. performed statistical analysis and interpretation, reviewed the draft of the manuscript, and gave final approval of the manuscript. S.B., K.D., Q.-A.A., M.K.P., and A.H.B. designed the study, reviewed the draft of the manuscript, and gave final approval of the manuscript. M.J.S. conceived and designed the study, contributed to interpretation, and gave final approval of the manuscript. A.A.T. is the guarantor of this work and, as such, had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

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