Obstructive sleep apnea, chronic obstructive pulmonary disease and hypertensive microvascular disease: a cross-sectional observational cohort study

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Hypertensive microvascular disease is associated with an increased risk of diastolic heart failure, vascular dementia and progressive renal impairment. This study examined whether individuals with obstructive sleep apnoea (OSA) had more retinal hypertensive microvascular disease than those with chronic obstructive pulmonary disease (COPD) and hospital controls. This was a single-centre, cross-sectional, observational study of participants recruited consecutively from a general respiratory clinic and a general medical clinic. OSA was diagnosed on overnight polysomnography study (apnoea-hypopnoea index ≥ 5), and controls with COPD had a forced expiratory volume/forced vital capacity (forced expiratory ratio) < 70%. Individuals with both OSA and COPD were excluded. Hospital controls had no COPD on respiratory function testing and no OSA on specialist physician questioning. Study participants completed a medical questionnaire, and underwent resting BP measurement, and retinal photography with a non-mydriatic camera. Images were deidentified and graded for microvascular retinopathy (Wong and Mitchell classification), and arteriole and venular calibre using a semiautomated method at a grading centre. Individuals with OSA (n = 79) demonstrated a trend to a higher mean arterial pressure than other hospital patients (n = 143) (89.2 ± 8.9 mmHg, p = 0.02), and more microvascular retinopathy (p < 0.001), and narrower retinal arterioles (134.2 ± 15.9 μm and 148.0 ± 16.2 μm respectively, p < 0.01). Microvascular retinopathy and arteriolar narrowing were still more common in OSA than hospital controls, after adjusting for age, BMI, mean arterial pressure, smoking history and dyslipidaemia (p < 0.01, p < 0.01, respectively). Individuals with OSA demonstrated a trend to a higher mean arterial pressure than those with COPD (n = 132, 93.2 ± 12.2 mmHg and 89.7 ± 12.8 mmHg respectively, p = 0.07), and more microvascular retinopathy (p = 0.0001) and narrower arterioles (134.2 ± 15.9 and 152.3 ± 16.8, p < 0.01). Individuals with OSA alone had more systemic microvascular disease than those with COPD alone or other hospital patients without OSA and COPD, despite being younger in age.

Obstructive sleep apnoea (OSA) affects 5–10% of middle-aged adults, and is even more common in the elderly. Its prevalence is increasing with the obesity epidemic. OSA is characterized by repeated episodes of partial or complete upper airways obstruction during sleep, due to relaxation of the tongue and airways muscles. This leads to snoring, and a reduction (‘hypopnoea’) or blockage (‘apnoea’) of airflow. Apneic episodes result in acute physiological stress including arterial desaturation, and surges of sympathetic activity with tachycardia and hypertension.

At least 50% of people with OSA have hypertension, and hypertension worsens with more severe disease. OSA predisposes not only to nocturnal hypertension but also sustained daytime hypertension. OSA increases cardiac risk and mortality, independent of the traditional risk factors. Type 2 diabetes is more common.
too because sleep-associated oxygen desaturation impairs glucose tolerance independent of obesity. OSA also results directly in coronary microvascular dysfunction and subclinical coronary artery disease partly through oxidative stress.

Chronic obstructive pulmonary disease (COPD) is another chronic respiratory disease but is characterized by inflammation in the airways and not fully reversible airflow obstruction. It affects 10% of the population over the age of 40, most of whom have been smokers. Individuals with COPD have a two- to three- times increased risk of cardiac disease, but smoking alone does not explain this association, and shared genetic risk factors, inflammatory mechanisms, oxidative stress and neurohumoral responses have all been suggested. COPD also results in microvascular disease. COPD overlaps with OSA and 10 -30% of individuals with COPD also have OSA. Any study of the consequences of OSA on the retinal microvasculature must exclude individuals who also have COPD, but this is rarely undertaken.

Small vessel disease in the retina reflects systemic small vessel disease including the risk of cardiac events and stroke. Small vessel changes are attributable not only to hypertension but also depend on the macrovascular risk factors, of age, gender, diabetes, smoking, family history and dyslipidemia. Features of small vessel disease include arteriovenous nicking, hemorrhage, exudates, and focal and generalized arteriolar narrowing (Keith-Wagener-Barker classification, modified by Wong and Mitchell). However calibre is also affected by systemic inflammation, for example, with diabetes obesity, and smoking.

The advantages of retinal small vessel examination over other methods of vascular assessment are that it is accessible, fast, inexpensive and highly reproducible. The primary outcome of this study was thus to compare the occurrence of microvascular retinopathy in individuals with OSA alone, in hospital patients without OSA or COPD, or in individuals with COPD without OSA. The secondary outcome was to compare the effect of hypertension in OSA, COPD and other hospital patients on retinal microvascular calibre. Other studies have examined retinal microvascular disease in OSA, but to date none has excluded individuals with both OSA and COPD; some used a self-administered questionnaire for diagnosis (‘sleep-disordered breathing’); one examined retinal photographs up to 3 years after the polysomnograms; one diagnosed hypertension on history and did not take current BP into account; and two did not consider diabetes or smoking when interpreting small vessel calibre.

Patients and methods

Study design. This was a single centre, cross-sectional, observational study of consecutive individuals with OSA recruited from the respiratory or general medical clinic of a metropolitan teaching hospital over a 6 month period. Hospital patients without OSA or COPD and those with COPD were recruited from the same clinics.

Recruitment, data capture and retinal photographs were coordinated in a single episode during a clinic visit. Individuals diagnosed with OSA within the previous year but without COPD on respiratory function testing were invited to participate. Exclusion criteria were COPD on respiratory function testing (to enable us to determine microvascular disease in OSA alone) or ungradable retinal images.

Hospital controls and patients with COPD were used as controls because their vascular risk factors and respiratory function were known.

Patients were assisted to complete a structured questionnaire, and then underwent BP measurement, and retinal photography. Retinal images were deidentified, and examined for microvascular retinopathy and vessel calibre by trained graders. The presence and severity of retinal microvascular abnormalities and calibre measurements were compared between subjects with OSA, hospital controls without OSA or COPD, and subjects with COPD. There were no changes to the study design after its commencement and no interim analysis.

The study was approved by the Northern Health Human Research Ethics Committee according to the Principles of the Declaration of Helsinki, and all participants provided signed, informed consent.

Participants. OSA was diagnosed on a polysomnography study (apnoea: hypopnea index ≥ 5) in an accredited diagnostic sleep laboratory where all studies were interpreted by a specialist sleep physician and conducted within the previous year. Testing included electroencephalography, electrooculography, electromyography, electrocardiography, pulse oximetry, oral and nasal airflow, thoracic and abdominal motion using standardized criteria (Compumedics software, Melbourne). Apnea was defined as cessation of airflow lasting ≥ 10 s, and hypopnea as a decrease in tidal volume with a ≥ 4% reduction in oxyhaemoglobin saturation. The apnoea: hypopnea index (AHI) was the average number of apneas plus hypopneas per hour of objectively measured sleep. OSA was classified as mild (apnoea-hypopnea index at least 5 but < 15), moderate (at least 15 but < 30) or severe (at least 30). Other OSA features that were recorded included the Epworth sleepiness scale score (by questionnaire), hypoxia duration and REM-predominant sleep.

Hospital controls were patients who had no previous diagnosis of COPD, asthma or bronchiectasis and had normal respiratory function tests within the past year. Individuals with COPD were recruited consecutively from a general respiratory clinic and had been diagnosed with an FER (ratio of FEV1 to forced vital capacity) of < 70%. Respiratory function tests were performed using a computerized spirometer (Sensor Medics Legacy 29D, Yorba Linda, USA) by a trained technician according to a standard protocol. Hospital controls and individuals with COPD were confirmed by a physician (JS) to have no clinical features of OSA, including excess daytime sleepiness, witnessed breathing interruptions, or awakenings due to gasping or choking. These are features consistent with a negative score on the STOP-Bang questionnaire.

Participants were assisted to complete a structured questionnaire for demographics (age, gender), and vascular risk factors (smoking, cigarette pack years, diabetes, hypertension, dyslipidemia), and laboratory test results (haemoglobin, lipids, estimated glomerular filtration rate) were obtained from their medical records. Hypertension was defined as a physician-made diagnosis, even if treated, and dyslipidemia with a cholesterol ≥ 5.0 mmol/l.
HDL ≤ 2.0 mmol/l, or statin use. Blood pressure was recorded after 5 min sitting using a Hg sphygmomanometer, and mean arterial pressure and pulse pressure were calculated from the systolic and diastolic BP.

Measurements. Retinal imaging was performed, and images graded for retinopathy. All participants underwent colour retinal photography using a non-mydriatic camera (KOWA 7, Japan; Canon CR5-45NM, Japan). Standard 45º images were taken of both eyes, with one view centred on the macula and another on the optic disc. All images were deidentified, and graded for microvascular retinopathy (Wong and Mitchell classification27 by an ophthalmologist and a trained observer. Microvascular retinopathy was classified as mild (arteriovenous nicking, focal arteriolar narrowing, silver-wiring or a decreased arteriovenous ratio), moderate (haemorrhage or exudates) or severe (papilloedema) 27. In all cases the grade of the more severely-affected eye was used in the assessment.

Retinal images were sent to the Centre for Eye Research Australia for measurement of the retinal arteriole and venular calibre by trained graders using a standardized protocol and Knudtson’s revision of the Parr–Hubbard formula39. Briefly, all vessels passing through a zone 0.5–1 disc diameter from the optic disc margin were examined using a semi-automated computer imaging program (University of Wisconsin, WI), and measures based on the 6 largest vessels were combined into the Central Retinal Artery and Vein Equivalents (CRAE and CRVE). This method was highly reproducible.

Statistical analysis. Demographic data were compared using Fisher’s exact test, chi square test or ANOVA. Possible determinants of calibre were examined using univariate analysis and then in multivariate linear regression models. Statistical analyses were performed using STATA version 11.2 software (STATACORP Inc, College Station Texas). A p value < 0.05 was considered significant and p < 0.10 a trend.

Results Seventy-nine subjects with OSA, 143 hospital controls, and 132 with COPD were studied (Table 1). Thirty-five further individuals with OSA (31%) were excluded because they also had COPD. Six (8%) and 13 (10%) subjects were excluded from the OSA and COPD cohorts respectively because their retinal images were ungradable.

Subjects with OSA. Fifty-six (71%) subjects with OSA had hypertension, with an overall mean arterial pressure of 93.2±12.2 mmHg (Table 1). Their mean BMI was 35.9±8.5 kg/m² and 25 (32%) had diabetes. Twenty (25%) had mild, 22 (28%) had moderate and 37 (47%) had severe OSA. Their mean Epworth Sleepiness Scale was 10.4±5.3 (2–24), 16 (20%) had an Epworth Sleepiness Scale ≥ 15, and their mean hypoxia duration was 8.7±17.9 s. Fifty-three (67%) were prescribed CPAP but it was unclear how many adhered to treatment adequately.

| Clinical characteristics                      | OSA (n = 79) | Hospital controls (n = 143) | OSA versus hospital controls 95% CI, p value | COPD (n = 132) | OSA versus COPD, 95% CI, p value |
|-----------------------------------------------|--------------|----------------------------|---------------------------------------------|----------------|---------------------------------|
| Age (mean, SD, years)                         | 62.5, 12     | 69.3, 7.7                  | 4.2–9.41, < 0.01                            | 69.9, 9.7      | 4.2–10.1, < 0.01                |
| Gender (male)                                 | 44 (56%)     | 75 (52%)                   | 1.14, 0.63–2.06, 0.64                       | 66 (50%)       | 1.26, 0.7–2.3, 0.43            |
| BMI (mean, SD, kg/m²)                         | 35.0, 8.5    | 26.5, 6                    | − 11.30 to − 7.41, < 0.01                   | 27.1, 6.6      | − 10.8 to − 6.6, 0.01          |
| Smoking history                               | 42 (53%)     | 77 (54%)                   | 1.03, 0.57–1.87, 0.92                       | 126 (96%)      | 0.06, 0.02–0.2, 0.01           |
| Pack years (mean, SD)                         | 31.6, 19.5   | 33.2, 25.4                 | − 7.97 to 10.73, 0.77                       | 46.3, 26.4     | 5.5–23.8, 0.01                 |
| Hypertension diagnosis                        | 56 (71%)     | 75 (52%)                   | 2.21, 1.19–4.17, 0.01                       | 81 (61%)       | 1.53, 0.8–2.94, 0.16           |
| Mean arterial pressure (mean, mmHg)           | 93.2, 12.2   | 89.2, 8.9                  | 0.65–7.2, 0.02                              | 89.7, 12.8     | − 0.32 to 7.28, 0.07           |
| Mean pulse pressure (mean, mmHg)              | 54.5, 13.5   | 55.2, 13.4                 | − 89.82 to 69.42, p = 0.74                  | 60.1, 16.9     | − 10.36 to 6.84, 0.02          |
| Diabetes                                      | 25 (32%)     | 34 (24%)                   | 1.48, 0.77–2.85, 0.21                       | 32 (24%)       | 1.45, 0.74–2.81, 0.24          |
| Dyslipidemia                                  | 43 (54%)     | 43 (30%)                   | 2.78, 1.51–5.10, 0.01                       | 55 (42%)       | 1.67 (0.92–3.05), 0.07         |

Table 1. Baseline demographic and clinical characteristics of subjects with OSA, hospital controls and subjects with COPD. Significant values are in bold.
Almost all subjects with OSA (97%) had a retinal microvascular retinopathy with mild (60, 76%) or moderate (17, 21%) features (Table 1, Figs. 1, 2). Moderate microvascular retinopathy was not associated with age, smoking history, diabetes, hypertension, dyslipidemia or BMI (p all > 0.05). Moderate retinopathy was also not associated with OSA features (severity, p = 1.00), increased Epworth Sleepiness Scale score (p = 0.56), longer hypoxia duration (p = 0.49) or more REM-predominant sleep (p = 1.00).

Subjects with OSA had a mean retinal arterial calibre of 134.2 ± 15.9 µm and mean retinal venular calibre of 197.3 ± 24.0 µm. Those with OSA and a moderate microvascular retinopathy had narrower arterioles and venules than those without a retinopathy (136.0 ± 14.6 µm and 165.0 ± 9.7 µm respectively, 95% CI 9.0–54.3 µm, p < 0.01; and 198.8 ± 13.5 µm and 250.0 ± 52.2 µm, 95% CI 18.3–86.3, p < 0.01) (Table 2).

When subjects with OSA alone were examined according to quartiles of their retinal arteriole calibre, there were no differences between the quartiles in OSA severity measured by apnoea: hypopnoea index (p = 1.00), Epworth Sleepiness Scale (p = 0.38), hypoxia duration (p = 0.24), or REM-predominant sleep (p = 0.50). These observations suggest that the retinal arteriole narrowing did not worsen directly with any of the commonly-used measures of OSA severity.

When subjects with OSA alone were examined according to quartiles of their retinal venular calibre, there were again no differences between the quartiles in OSA severity measured by apnoea; hypopnoea index (p = 1.00), Epworth Sleepiness Scale (p = 0.52), hypoxia duration (p = 0.89), or REM predominant sleep (p = 0.07). Again these observations suggest that the retinal venular narrowing did not worsen directly with any of the commonly-used measures of OSA severity.

Subjects with OSA compared with hospital controls. Subjects with OSA were younger (p < 0.01), had a higher BMI (p < 0.01), more dyslipidemia (p = 0.01), more hypertension (p = 0.01) and a higher mean arterial pressure (p = 0.02) than hospital controls (Table 1, Fig. 2). Their likelihood of smoking, diabetes or a higher pulse pressure were not different.
Subjects with OSA had more microvascular disease than other hospital patients (p < 0.001). They had more mild retinopathy (p < 0.001), and there was a trend for more moderate retinopathy (p = 0.08). Microvascular retinopathy was still more common in subjects with OSA than hospital controls after adjusting for age, BMI, hypertension (p = 0.04) and dyslipidemia (p = 0.03).

Subjects with OSA had a smaller mean arteriole calibre than other hospital patients (134.2 ± 15.9 µm and 148.0 ± 16.2 µm, respectively, mean difference = 31.2 to −12.3 µm, p < 0.01), but venular calibre was not different (197.3 ± 24.0 µm and 213.9 ± 26.2 µm, 95% CI = 44.9 to -19.1 µm, p = 0.23).

When features in subjects with OSA plus hospital controls were compared by quartile of arteriolar calibre, a smaller calibre was associated with hypertension (p = 0.04), lower BMI (p = 0.02), non-smoking status (p = 0.06), and smaller venular calibre (p < 0.01) (Table 3). For the corresponding quartiles in venular calibre, smaller vessels correlated with increasing age (p = 0.01), fewer cigarette pack years (p = 0.07) and smaller arterioles (p < 0.01) (Table 4).

Microvascular retinopathy and narrowed arterioles and venules were still more common in subjects with OSA than other hospital patients after adjusting for possible confounders (p < 0.01, p < 0.01 and p < 0.01) (Table 5).

Subjects with OSA had more microvascular disease than other hospital patients (p < 0.001). They had more mild retinopathy (p < 0.001), and there was a trend for more moderate retinopathy compared with none. CRAE central retinal artery equivalent, CRVE central retinal vein equivalent. Significant values are in bold.

### Table 2. Correlation of Microvascular retinopathy with OSA features and retinal arteriolar and venular calibre.

| Feature | Retinopathy | p value |
|---------|-------------|---------|
| OSA (n = 79) | None (n = 2) | Mild (n = 61) | Moderate (n = 13) |
| Mild (n = 20) | 0 | 14 | 3 |
| Moderate (n = 22) | 1 | 18 | 3 |
| Severe (n = 37) | 1 | 29 | 7 |
| CRAE (mean, SD) | 165.0, 9.7 | 132.7, 16.4 | 136.0, 14.6 |
| CRVE (mean, SD) | 250.0, 52.2 | 197.7, 25.2 | 198.8, 13.5 |

### Table 3. Clinical features and retinal arteriole calibre in OSA plus hospital controls by quartile. Smaller retinal arteriole calibre was associated with a higher BMI (p = 0.02), a hypertension diagnosis (p = 0.04) and a smaller venular calibre (p < 0.01), and there was a trend with less smoking history (p = 0.06). In the patients with OSA, all OSA was associated with a smaller arteriole calibre (p < 0.001). Moderate and severe OSA were associated with a smaller arteriole calibre (p = 0.04, p < 0.001 respectively). Retinal arteriole calibre was not associated with mean pulse pressure (p = 0.60). Significant values are in bold.

| Retinal arteriole calibre | Clinical features in OSA and controls | 90–129 µm (n = 48) | 130–140 µm (n = 49) | 141–155 µm (n = 49) | 156–190 µm (n = 48) | OR, 95%CI, p value |
|--------------------------|--------------------------------------|--------------------|--------------------|--------------------|--------------------|--------------------|
| Age (mean, SD, years) | 66.9, 9.4 | 62.7, 11.1 | 69.1, 8.4 | 66.7, 8.6 | 0.93 |
| Gender (male) | 31 (65%) | 26 (53%) | 21 (43%) | 22 (46%) | 0.10 |
| BMI (mean, SD, kg/m²) | 31.1, 6.6 | 32.5, 9.4 | 28.2, 6.8 | 27.5, 7.6 | 0.02 |
| Smoking history | 21 (44%) | 26 (53%) | 28 (57%) | 31 (65%) | 0.06 |
| Pack years (mean, SD) | 32.5, 21.5 | 34.2, 17.8 | 27.9, 21.3 | 35.8, 32 | 0.7 |
| Hypertension history | 33 (69%) | 26 (53%) | 29 (59%) | 22 (46%) | 0.04 |
| Mean arterial pressure (mean, SD, mmHg) | 92.7, 13.2 | 91.8, 11.4 | 90.7, 10 | 89, 8 | 0.17 |
| Dyslipidemia | 16 (35%) | 23 (47%) | 20 (41%) | 15 (31%) | 1.00 |
| Diabetes | 11 (23%) | 16 (33%) | 12 (25%) | 13 (27%) | 0.80 |
| Retinal venular calibre (mean, SD, µm) | 189.6, 18.4 | 202, 20 | 211.2, 25.1 | 229, 23.8 | < 0.01 |

| OSA patients only | n = 34 | n = 29 | n = 10 | n = 6 | < 0.001 |
|-------------------|-------|-------|-------|------|--------|
| Mild (n = 20) | 7 (9%) | 8 (10%) | 3 (4%) | 2 (3%) | 0.12 |
| Moderate (n = 22) | 9 (11%) | 8 (10%) | 4 (5%) | 1 (1%) | 0.04 |
| Severe (n = 37) | 18 (23%) | 13 (16%) | 3 (4%) | 3 (4%) | < 0.001 |
| Mean pulse pressure (mean, SD, mmHg) | 58.4, 13 | 56.1, 13 | 52.5, 17 | 51.0, 13 | 0.60 |
Subjects with OSA compared with COPD. Subjects with OSA were younger, had a higher BMI, and
Subjects with OSA had more microvascular retinopathy (p < 0.001) than those with COPD alone (Table 1). Microvascular retinopathy was still more common after adjusting for possible confounders (OR 6.67, 95% CI 1.27–34.82, p = 0.02) (Table 5).

Subjects with OSA also had narrower arterioles (134.3 ± 16.7 µm and 152.3 ± 16.8 µm, respectively, mean difference, − 18.0 µm, − 12.9 to -23.1 µm, p < 0.01) and venules than those with COPD (199.5 ± 25.2 µm and 224.8 ± 28.1 µm, mean difference, − 25.3, − 17.1 to − 33.5 µm, p < 0.01). Retinal arteriole and venular calibre were still less after adjusting for potential confounders (Table 5).

Discussion

This study found more systemic small vessel disease in individuals with OSA than in other hospital patients. It also found more small vessel disease in OSA after patients with COPD were excluded than in those with COPD alone. Nearly all subjects with OSA in this study had a microvascular retinopathy and narrowed arterioles despite being younger than the other cohorts. The increase in retinal small vessel disease in OSA is consistent with the increased risk of systemic microvascular disease associated with diastolic heart failure, vascular dementia and progressive renal failure.

Individuals with OSA had more microvascular retinopathy and narrower arterioles than hospital controls and the COPD cohort. Poorly-controlled hypertension may be a contributor to the small vessel disease seen in OSA. Diagnosed hypertension was more common in OSA than hospital controls. The mean arterial pressure was higher in OSA than the hospital controls and those with COPD. However pulse pressure which corresponds to arterial stiffness was not higher in OSA than in the other cohorts.

These results suggest a disparity between diagnosed hypertension, clinic blood pressure readings and retinal microvascular changes. Possible explanations include that the blood pressure was measured only once in this study during a clinic visit and that patients were more likely to take their antihypertensive medication on the day of a medical appointment. Secondly, previous reports of an association between OSA and hypertension did not exclude individuals with COPD which will have distorted the results. Thirdly, smoking in COPD increases both venular and arteriole calibre and complicates the measurement of vessel calibre. Finally and importantly, hypertension control in OSA is difficult to assess since hypertension is mainly nocturnal.

Repetitive apnoeas in OSA at night increase the sympathetic drive and trigger fluctuations in blood pressure and heart rate. They also suppress nitric oxide production. The mechanism of vascular disease may be through the release of free radicals, and reduced vasodilation. Endothelial dysfunction is a key event that precedes atherosclerosis and represents a pathogenic link with cardiovascular disease. In addition the hypertension in OSA is now correlated more directly with the damage from intermittent hypoxemia and ischemic reperfusion injury rather than the sustained hypoxemia seen in COPD.

This study also did not demonstrate worse microvascular retinopathy or a consistently narrower calibre with more ‘severe’ OSA as assessed by an apnoea: hypopnea index > 30. Possible explanations include that the cohort included many individuals with milder OSA, the sleep studies were performed prior to treatment, and that CPAP was commonly prescribed. In diseases other than OSA, improved blood pressure control reverses small vessel abnormalities, and in OSA, CPAP treatment reverses the microvascular dysfunction.

OSA coexists with COPD in up to 30% of individuals but this study minimised the risk of overlap by specifically questioning all participants for features of OSA and testing them for COPD. This study is, we believe, the first to examine microvascular disease in OSA independent of COPD. Arterioles and venules are usually larger in COPD because of vascular remodelling and intimal and medial thickening from the accumulation of inflammatory cells and fibroblasts. The increased calibre demonstrated here in OSA compared with hospital controls or those with OSA reflects this exaggeration.

The strengths of this study were its high recruitment rate; the completeness of the data; the near contemporaneous nature of the sleep studies and retinal imaging; the robustness of the retinal microvascular assessments; and the use of multiple measures of hypertension. The control cohorts of hospital patients and subjects with COPD were chosen because their medical comorbidities were well-characterised. Some other studies have not even considered the diagnosis of hypertension, whereas we included a previous diagnosis of hypertension, and even treated hypertension since treatment is often inadequate. The measurement of hypertension in OSA is further complicated by its nocturnal nature. We also examined the effect of mean arterial pressure, which weights systolic and diastolic BP, and considered the effect of the blood pressure in the different cohorts.

The study’s major limitations were its cross-sectional and single centre nature, and the exclusion of OSA by clinical questioning rather than with sleep studies. It was difficult to evaluate an effect of CPAP since few patients appeared strictly adherent to treatment. However other studies have demonstrated a beneficial effect of 3–12 months treatment with CPAP on small vessel disease.

The number of participants recruited was typical of studies examining retinal microvascular calibre in OSA. The proportion of patients with OSA excluded clinically with COPD or with ungradable retinal images approximated previously reported frequencies. Diabetes is a common comorbidity in OSA occurring in 24–86% of many cohorts, and OSA exacerbates diabetic retinopathy where this is present. Our analysis corrected for diabetes but did not exclude this group so that the cohort was representative of all patients found in a respiratory clinic. Indeed the proportion of individuals with diabetes was not different in the cohorts with OSA (32%), COPD (24%) or the controls (24%).

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This study suggests that subjects with OSA have an increased risk of small vessel disease that is greater than is found in COPD or in other hospital patients. These comparisons indicate where physician time and hospital resources should be directed in terms of minimising the risks of systemic small vessel disease.

Data availability
All deidentified data used and analysed in the current study is available from the corresponding author on reasonable request.

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References
1. Kripke, D. F. et al. Prevalence of sleep-disordered breathing in ages 40–64 years: A population-based survey. *Sleep* **20**(1), 65–76 (1997).
2. Olson, L. G., King, M. T., Hensley, M. J. & Saunders, N. A. A community study of snoring and sleep-disordered breathing. Health outcomes. *Am. J. Respir. Crit. Care Med.* **152**(2), 717–720 (1995).
3. Young, T. et al. The occurrence of sleep-disordered breathing among middle-aged adults. *N. Engl. J. Med.* **328**(17), 1230–1235 (1993).
4. Stradling, J. R. & Crosby, J. H. Predictors and prevalence of obstructive sleep apnoea and snoring in 1001 middle aged men. *Thorax* **46**(2), 85–90 (1991).
5. Sleep-related breathing disorders in adults: Recommendations for syndrome definition and measurement techniques in clinical research. The Report of an American Academy of Sleep Medicine Task Force. *Sleep* **22**(5), 667–689 (1999).
6. Somers, V. K., Dyken, M. E., Clary, M. P. & Abboud, F. M. Sympathetic neural mechanisms in obstructive sleep apnea. *J. Clin. Invest.* **96**(4), 1897–1904 (1995).
7. Silverberg, D. S., Oksenberg, A. & Iaina, A. Sleep-related breathing disorders as a major cause of essential hypertension: Fact or fiction?. *Curr. Opin. Nephrol. Hypertens.* **7**(4), 353–357 (1998).
8. Phillips, C. L. & O’Driscoll, D. M. Hypertension and obstructive sleep apnea. *Nat. Sci. Sleep* **5**, 43–52 (2013).
9. Hla, K. M. et al. Electrocardiographically indicated cardiovascular disease in sleep-disordered breathing. *Sleep Breath* **12**(3), 251–258 (2008).
10. Gottlieb, D. J. et al. The occurrence of sleep-disordered breathing among middle-aged adults. *Circulation* **122**(4), 352–360 (2010).
11. Redline, S. et al. Obstructive sleep apnea-hypopnea and incident stroke: The sleep heart health study. *Am. J. Respir. Crit. Care Med.* **182**(2), 269–277 (2010).
12. Young, T. et al. Sleep disordered breathing and mortality: Eighty-year follow-up of the Wisconsin sleep cohort. *Sleep* **31**(8), 1071–1078 (2008).
13. Bradley, T. D. & Floras, J. S. Obstructive sleep apnoea and its cardiovascular consequences. *Lancet* **373**(9657), 82–93 (2009).
14. Polotsky, V. Y. et al. Intermittent hypoxia increases insulin resistance in genetically obese mice. *J. Physiol.* **552**(Pt 1), 253–264 (2003).
15. Ip, M. S. et al. Obstructive sleep apnea is independently associated with insulin resistance. *Am. J. Respir. Crit. Care Med.* **165**(5), 670–676 (2002).
16. Morra, S. & Roubille, F. Obstructive sleep apnoea: From respiratory events to coronary microvascular dysfunction. *Acta Cardiol.* **73**(4), 319–324 (2018).
17. Shpilsky, D. et al. Association of obstructive sleep apnea with microvascular endothelial dysfunction and subclinical coronary artery disease in a community-based population. *Vasc. Med.* **23**(4), 331–339 (2018).
18. Wang, N. et al. Relationship between obstructive sleep apnoea and coronary microcirculatory function among patients with cardiac syndrome X. *Coron. Artery Dis.* **25**, 35–39 (2013).
19. Kocak, H. E. et al. Does pediatric obstructive sleep apnea syndrome cause systemic microvascular dysfunction?. *J. Craniofac. Surg.* **29**(4), e381–e384 (2018).
20. Buchner, N. J. et al. Microvascular endothelial dysfunction in obstructive sleep apnea is caused by oxidative stress and improved by continuous positive airway pressure therapy. *Respiration* **82**(5), 409–417 (2011).
21. Halbert, R. J. et al. Global burden of COPD: Systematic review and meta-analysis. *Eur. Respir. J.* **28**(3), 523–532 (2006).
22. Chew, S. K. et al. Hypertensive/microvascular disease and COPD: A case control study. *Kidney Blood Press Res.* **31**(1), 29–39 (2016).
23. McNicholas, W. T. COPD–OSA overlap syndrome: evolving evidence regarding epidemiology, clinical consequences, and management. *Chest* **152**(6), 1318–1326 (2017).
24. Duncan, B. B., Wong, T. Y., Tyloleer, H. A., Davis, C. E. & Fuchs, F. D. Hypertensive retinopathy and incident coronary heart disease in high risk men. *Br. J. Ophthalmol.* **86**(9), 1002–1006 (2002).
25. Wong, T. Y. et al. Retinal arteriolar narrowing and risk of coronary heart disease in men and women. The Atherosclerosis Risk in Communities Study. *JAMA* **287**(9), 1153–1159 (2002).
26. Wong, T. Y. et al. Retinal microvascular abnormalities and blood pressure in older people: The Cardiovascular Health Study. *Br. J. Ophthalmol.* **86**(9), 1007–1013 (2002).
27. Wong, T. Y. & Mitchell, P. Hypertensive retinopathy. *N. Engl. J. Med.* **351**(22), 2310–2317 (2004).
28. Wong, T. Y. et al. Retinal vascular caliber, cardiovascular risk factors, and inflammation: The multi-ethnic study of atherosclerosis (MESA). *Invest. Ophthalmol. Vis. Sci.* **47**(6), 2341–2350 (2006).
29. Wang, J. J. et al. Retinal vessel diameters and obesity: A population-based study in older persons. *Obesity (Silver Spring)* **14**(2), 206–214 (2006).
30. Kifley, A. et al. Long-term effects of smoking on retinal microvascular caliber. *Am. J. Epidemiol.* **166**(11), 1288–1297 (2007).
31. Boland, L. L. et al. Sleep-disordered breathing is not associated with the presence of retinal microvascular abnormalities: The Sleep Heart Health Study. *Sleep* **27**(3), 467–473 (2004).
32. Fraser, C. L. et al. A prospective photographic study of the ocular fundus in obstructive sleep apnea. *J. Neuroophthalmol.* **33**(3), 241–246 (2013).
33. Shankar, A. et al. Sleep-disordered breathing and retinal microvascular diameter. *Atherosclerosis* **226**(1), 124–128 (2013).
34. Chew, M. et al. Sleep apnea and retinal signs in cardiovascular disease: The Multi-Ethnic Study of Atherosclerosis. *Sleep Breath.* **20**(1), 15–23 (2016).
35. Ferris, B. G. Epidemiology standardization project (American Thoracic Society). *Am. Rev. Respir. Dis.* **116**(6 Pt 2), 1–120 (1978).
36. ATS statement—Snowbird workshop on standardization of spirometry. *Am. Rev. Respir. Dis.* **119**(5), 831–838 (1979).
37. Epstein, L. J. et al. Clinical guideline for the evaluation, management and long-term care of obstructive sleep apnea in adults. *J. Clin. Sleep Med.* **5**(3), 263–276 (2009).
38. Chung, F. et al. STOP questionnaire: A tool to screen patients for obstructive sleep apnea. *Anesthesiology* **108**(5), 812–821 (2008).
39. Knudtson, M. D. et al. Revised formulas for summarizing retinal vessel diameters. Curr. Eye Res. 27(3), 143–149 (2003).
40. Nagle, M. P. et al. Retinal microvascular dysfunction in heart failure. Eur. Heart J. 39(1), 47–56 (2018).
41. Lesage, S. R. et al. Retinal microvascular abnormalities and cognitive decline: The ARIC 14-year follow-up study. Neurology 73(11), 862–868 (2009).
42. Ooi, Q. L. et al. The microvasculature in chronic kidney disease. Clin. J. Am. Soc. Nephrol. 6(8), 1872–1878 (2011).
43. Saed, S. et al. Characteristics of hypertension and arterial stiffness in obstructive sleep apnea: A Scandinavian experience from a prospective study of 6408 normotensive and hypertensive patients. J. Clin. Hypertens. (Greenwich). 24(4), 385–394 (2022).
44. Narkiewicz, K. M. N., Cogliati, C., van de Borne, P. J. H., Dyken, M. E. & Somers, V. K. Altered cardiovascular variability in obstructive sleep apnea. Am. Heart Assoc. 98, 1071–1077 (1998).
45. Iij, H. & Pg, D. Nitric Oxide (NO) and obstructive sleep apnea (OSA). Sleep Breathing. 7(2), 53–61 (2003).
46. Trzepizur, W. G. F. et al. Microvascular endothelial function in obstructive sleep apnea: Impact of continuous positive airway pressure and mandibular advancement. Sleep Med. 10, 746–752 (2009).
47. Atkeson, A. & Jelic, S. Mechanisms of endothelial dysfunction in obstructive sleep apnea. Vasc. Health Risk Manag. 4(6), 1327–1335 (2008).
48. Sawyer, A. M. et al. A systematic review of CPAP adherence across age groups: Clinical and empiric insights for developing CPAP adherence interventions. Sleep Med. Rev. 15(6), 343–356 (2011).
49. Hughes, A. D. et al. Effect of antihypertensive treatment on retinal microvascular changes in hypertension. J. Hypertens. 26(8), 1703–1707 (2008).
50. Lin, G.M., Redline, S., Klein, R., Colangelo, L.A., Cotch, M.E., Wong, T.Y., et al. Sex-specific association of obstructive sleep apnea with retinal microvascular signs: The multi-ethnic study of atherosclerosis. J. Am. Heart Assoc. 5(7) (2016).
51. Lattimore, J. L., Walcox, I., Skilton, M., Langenfeld, M. & Celermaier, D. S. Treatment of obstructive sleep apnoea leads to improved microvascular endothelial function in the systemic circulation. Thorax 61(6), 491–495 (2006).
52. Harris, B. et al. The association of systemic microvascular changes with lung function and lung density: A cross-sectional study. PLoS ONE 7(12), e05224 (2012).
53. Krakenburge, A. R. et al. Enhanced expression of fibroblast growth factors and receptor FGFR-1 during vascular remodeling in chronic obstructive pulmonary disease. Am. J. Respir. Cell Mol. Biol. 27, 517–525 (2002).
54. Arikawa, T. et al. Microvascular endothelial function in patients with obstructive sleep apnoea syndrome. Vasc. Failure 2(1), 53–58 (2018).
55. Nguyen, P. K., Katikireddy, C. K., McConnell, M. V., Kushida, C. & Yang, P. C. Nasal continuous positive airway pressure improves myocardial perfusion reserve and endothelial-dependent vasodilation in patients with obstructive sleep apnea. J. Cardiovasc. Magn. Reson. 12, 50 (2010).
56. Adderley, N. J. et al. Obstructive sleep apnea, a risk factor for cardiovascular and microvascular disease in patients with type 2 diabetes: Findings from a population-based cohort study. Diabetes Care 43(8), 1868–1877 (2020).
57. Nannapaneni, S., Rama, K. & Surani, S. Effect of obstructive sleep apnea on type 2 diabetes mellitus: A comprehensive literature review. World J. Diabetes. 4(6), 238–244 (2013).

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Author contributions
S.C. recruited the study participants, took the retinal photographs, performed the statistical analysis and wrote the first draft; D.C. supervised the retinal photography and grading of the microvascular retinopathy; A.H. supervised the statistical analysis; P.H. supervised participant recruitment; L.H. undertook the retinal microvascular calibre measurements; J.S. oversaw the project, and prepared the final manuscript. All authors have approved the final manuscript.

Competing interests
The authors declare no competing interests.

Additional information
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