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Effect of routine low-dose oxygen supplementation on death and disability in adults with acute stroke

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TITLE PAGE
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      Effect of routine low-dose oxygen supplementation on death and disability in
 3
      adults with acute stroke: the Stroke Oxygen Study randomized clinical trial
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      Revision 2 10<sup>th</sup> August 2017, corrected
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31 KEY POINTS

32

Question: Does routine prophylactic low-dose oxygen supplementation after acute stroke improve
 functional outcome?

35

- 36 Findings: In this randomized clinical trial 8003 patients with acute stroke were allocated within 24
- 37 hours of admission to 3 days of continuous oxygen, nocturnal oxygen, or control. After 3 months
- 38 there was no significant difference in death and disability for the combined oxygen groups
- 39 compared with control (odds ratio 0.97), or for continuous oxygen compared with nocturnal oxygen
- 40 (odds ratio 1.03).

41

42 Meaning: Routine low-dose oxygen did not improve outcomes in non-hypoxic patients after acute
43 stroke.

45 **ABSTRACT**

46

47 and often undetected. Oxygen supplementation could prevent hypoxia and secondary neurological 48 deterioration and thus has the potential to improve recovery. 49 **Objective:** To assess whether routine prophylactic low-dose oxygen therapy is superior to control 50 in reducing death and disability at 90 days and, if so, whether oxygen given at night only, when 51 hypoxia is most frequent, and oxygen administration is least likely to interfere with rehabilitation, is 52 more effective than continuous supplementation. 53 Design, setting, and participants: In this single-blind randomized clinical trial 8003 adults with 54 acute stroke were enrolled from 136 participating centers within 24 hours of hospital admission, if 55 they had no clear indications for, or contraindications to, oxygen treatment (first patient enrolled 24-56 Apr-2008, last follow-up 27-Jan-2015). 57 Interventions: Participants were randomized 1:1:1 to continuous oxygen for 72 hours (n=2668), 58 nocturnal (21:00-07:00) oxygen for three nights (n=2667), or control (oxygen only if clinically 59 indicated, n=2668). Oxygen was given via nasal tubes at 3L/min if baseline oxygen saturation was $\leq 93\%$ and at 2L/min if > 93%. 60 61 Main outcomes and measures: The primary outcome was the modified Rankin Scale (mRS) score 62 (a measure of disability ranging from 0=no symptoms to 6=death, minimum clinically important difference 1 point), assessed at 90 days by postal questionnaire (participant aware, assessor 63 64 blinded). The mRS was analyzed by ordinal logistic regression, which yields a 'common' odds ratio 65 (OR) for a change from one disability level to the next better (lower) level; OR > 1.00 indicates improvement. Significance was set at $p \le 0.05$ for the primary outcomes and ≤ 0.01 for all other 66 67 outcomes.

Importance: Hypoxia is common in the first few days after acute stroke, frequently intermittent,

68	Results: 8003 patients (4398 (55%) males, mean age 72 (SD13) years; median NIHSS 5; mean
69	baseline oxygen saturation 96.6%) were enrolled. The primary outcome was available in 7677
70	(96%) participants. The unadjusted odds ratio for a better outcome (calculated via ordinal logistic
71	regression) was 0.97 (95% CI 0.89–1.05), p=0.47 for oxygen versus control, and 1.03 (95% CI
72	0.93–1.13), p=0.61 for continuous versus nocturnal oxygen. No subgroup could be identified that
73	benefited from oxygen. There were 348 (13.0%), 294 (11.0%), and 322 (12.1%) participants with at
74	least one serious adverse event in the continuous, nocturnal, and control groups respectively. No
75	significant harms were identified.

Conclusions and relevance: Among non-hypoxic patients with acute stroke the prophylactic use of
low-dose oxygen supplementation did not reduce death or disability at 3 months. These findings do
not support low-dose oxygen in this setting.

79 Trial Registration: ISRCTN52416964

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83 INTRODUCTION

Hypoxia is common during the first days after an acute stroke, $\frac{1}{2}$ and associated with higher rates of 84 neurological deterioration,² death and institutionalization,³ and greater mortality.⁴ While cells in the 85 86 ischemic penumbra are only viable for a few hours, brain cells beyond the ischemic core and penumbra remain at risk of delayed cell death for several days owing to vasogenic edema, 87 88 inflammation, and programmed cell death, particularly if metabolic disturbances are compounded by hypoxia. $\frac{5,6,7}{2}$ Continuous monitoring is associated with better outcomes, ⁸/₈ but even in intensively 89 90 monitored patients, hypoxia is not always identified and treated. Adverse outcomes were observed 91 to be increased when only some desaturations <90% were treated with oxygen, and reduced when all were treated.^{$\underline{3}$} 92

93 Supplemental oxygen could improve outcomes by preventing hypoxia and secondary brain damage, but could also have adverse effects.⁹ These include vasoconstriction and pulmonary toxicity with 94 95 high concentrations,⁹ respiratory tract infection due to contamination of the giving set, the tubing 96 acting as an impediment to mobilization, stress, and the direct effects of oxygen on vascular tone and blood pressure.¹⁰ Three small trials of short-term (≤ 12 hours), high-flow (10-45 L/min) 97 therapeutic oxygen, aimed at generating supra-physiological blood oxygen levels, have not shown 98 improved outcomes. ^{11, 12, 13} A larger trial (n=550) using low-dose supplemental oxygen (3 L/min 99 for 24 hours) also showed no benefit,¹⁴ but early neurological recovery was improved in a study 100 giving low-dose oxygen over 72 hours.¹⁵ 101

102 The primary aim of the Stroke Oxygen Study (SO₂S) was to determine whether low-dose oxygen 103 therapy during the first 3 days after an acute stroke improves outcome compared to usual care 104 (oxygen only when needed). As oxygen may restrict mobility and interfere with daytime activities, 105 the secondary hypothesis was that oxygen given at night only, when hypoxia is most likely, is more 106 effective than continuous oxygen supplementation. 107

METHODS 108

109 Study design

This was a multi-center randomized clinical trial of oxygen supplementation with single-blind 110 outcome assessment. The protocol (see online supplement),¹⁶ statistical analysis plan (see online 111 supplement), $\frac{17}{17}$ and data collection forms $\frac{18}{18}$ are published. Fully informed written or witnessed oral 112 113 consent was given by the participants or, if they did not have capacity to consent, by a legal 114 representative. The protocol was approved by the North Staffordshire Research Ethics Committee 115 (06/Q2604/109).

116 **Participants**

117 Adults (\geq 18 years) with a clinical diagnosis of acute stroke (see eText 1 for definition) within 24 h 118 of hospital admission, who had no clinical indications for, or contraindications to, oxygen treatment, or any concomitant condition likely to limit life expectancy to <12 months were eligible. 119

120

Randomization and interventions

121	Participants were allocated 1:1:1 via central web-based minimized randomization ¹⁹ to one of three
122	groups: i) continuous oxygen supplementation, ii) nocturnal oxygen supplementation only, iii) no
123	routine oxygen (control). The factors for which imbalances were minimized were: the 'six simple
124	variables' (SSV) prognostic index for independent survival at 6 months ²⁰ (cut-offs ≤ 0.1 , >0.1 to
125	\leq 0.35, >0.35 to \leq 0.70, >0.70), oxygen treatment before randomization (yes, no, unknown), baseline
126	oxygen saturation on air (<95 , $\ge 95\%$), and time since stroke onset (defined as the last time well for
127	wake-up strokes) (≤ 3 , >3 to ≤ 6 , >6 to ≤ 12 , >12 to ≤ 24 , >24 hours). No blocking was used. Oxygen
128	was administered per nasal tubes either continuously (day and night) during the first 72 h after
129	randomization or overnight (21:00–07:00) for three nights. Oxygen was given at a flow rate of 3

130 L/min if baseline saturation was 93% or below, or at a flow rate of 2 L/min, if baseline saturation was greater than 93%. In the control group no routine oxygen supplementation was given. 131 132 Vital signs were observed at least 6-hourly, with any abnormal findings treated independently of 133 trial allocation. Patients requiring oxygen in the control group, or in the nocturnal oxygen group 134 during the day, or needing changes in oxygen dosage for clinical reasons, were given the 135 appropriate concentration of oxygen irrespective of treatment group. In addition, for 4144 patients recruited in the latter half of the study, spot checks of treatment adherence were undertaken at 136 137 midnight and 6 am.

138 **Outcomes and blinding**

139 Outcomes were assessed at one week by a member of the local research team and at 90 days via 140 postal questionnaire. Telephone interviews were conducted with non-responders, or to clarify unclear or missing answers. The primary outcome was the modified Rankin Scale $(mRS)^{\frac{21}{2}}$ score (a 141 142 measure of disability ranging from 0=no symptoms to 6=death; minimum clinically important 143 difference 1 point) at 90 days. Secondary outcomes were: number of participants with neurological improvement, defined as a \geq 4-point decrease on the National Institutes of Health Stroke Scale 144 (NIHSS)^{22, 23} between randomization and day 7, the highest and lowest oxygen saturations within 145 146 the first 72 h, and mortality at one week. Further secondary outcomes at 90 days were: mortality, number of participants alive and independent (mRS≤2), number of participants living at home, 147 Barthel Index (BI) activities of daily living (ADL) score, $\frac{24}{24}$ quality of life (EuroQol EQ5D-3L), $\frac{25}{25}$ 148 and Nottingham Extended Activities of Daily Living (NEADL) score.^{$\frac{26}{10}$} For the NIHSS and BI, 149 deaths were recorded as the worst outcome on the scale.²⁷ Participants, their doctors, and local 150 151 research staff who recorded the one-week outcomes were not blind to the study interventions. 90day assessments were undertaken by the SO₂S study office blind to treatment allocation. 152

153 Study size

154 The initial recruitment target was 6,000 participants, which was estimated to provide 90% power to 155 detect small (0.2 mRS-point, e.g. a one-point improvement in one in 5 participants) differences 156 between oxygen (continuous and night only groups combined) and no oxygen at p≤0.01, and 90% 157 power at $p \le 0.05$ to detect small differences between continuous oxygen and oxygen at night only. The study size was subsequently revised to 8,000 participants, using ordinal methods, $\frac{16,17}{10}$ without 158 159 knowledge of interim results, to increase the number of patients with severe strokes and thereby 160 provide greater power to investigate any differential effectiveness of oxygen versus control within 161 subgroups (defined by severity).

162 Statistical analysis

The trial was designed to answer two key questions: firstly, whether oxygen supplementation improves outcome (mRS at 90 days) and secondly, whether giving oxygen at night is more effective than giving it continuously. The main comparisons therefore were of the two combined oxygen groups (continuous and night-time only) versus control, and of continuous oxygen versus oxygen at night only. The statistical analysis plan describes the analysis methods in detail (see online supplement).¹⁷

169 The mRS was analysed by ordinal logistic regression, which yields a 'common' odds ratio (OR) for 170 a move from one level to the next better (lower) level with an odds ratio more than 1.00 indicating 171 an improvement. For this and other outcome variables, a primary unadjusted analysis and a 172 secondary covariate-adjusted analysis were performed. Adjusted analyses incorporated the 173 following covariates: age, sex, baseline NIHSS score, baseline oxygen saturation, and the SSV 174 prognostic index for 6-month independence (or for analysis of mortality, the SSV prognostic index for 30-day survival). Sensitivity analysis for the mRS used multiple imputation of missing values 175 176 (using a chained equations method, with 20 imputed datasets). Additional imputations were 177 performed to allow for the possibility that data were missing not at random and were either i) better

178 or ii) worse than expected; missing values were thereby replaced by either very good (i.e. lowest) or 179 very poor (i.e. highest) scores on the mRS, as appropriate (see eTable 3 in supplementary appendix). Subgroups, for the mRS only, were analysed by an interaction term, and were predefined 180 in the statistical analysis $plan^{\frac{17}{12}}$ (see figure 2 for details of subgroups). 181 182 For continuous outcomes, means and standard deviations (SD) or medians and interquartile ranges 183 (IQR) are reported, as appropriate. Unadjusted analyses used unrelated t-tests, with the mean difference between treatments and corresponding confidence interval (CI) reported. The adjusted 184 185 analysis used analysis of covariance, with the covariates specified earlier included in the analysis. 186 For dichotomous outcomes, percentages were compared across the treatment comparisons using a 187 chi-squared test (unadjusted analyses). Adjusted analyses of dichotomous outcomes used binary 188 logistic regression, with the covariates listed earlier; ORs and confidence intervals are reported. 189 All analyses were by intention to treat, i.e. according to the treatment group to which participants 190 were allocated, irrespective of treatment actually received. Statistical significance was set at $p \le 0.05$ 191 with 95% CIs for the primary outcome and at $p \le 0.01$ with 99% CIs for secondary outcomes. All 192 reported p-values are 2-sided. The main analysis was performed in SAS® software for Windows, version 9.4, SAS Institute Inc., Cary, USA. IBM SPSS for Windows version 22, Armonk, New 193 194 York, USA was used for sensitivity analyses.

195 Interim analyses of safety and effectiveness were reviewed annually by an independent Data

196 Monitoring and Safety Committee. No alpha-spending adjustments were made.

197

198 **RESULTS**

199 **Participants**

8003 participants from 136 collaborating centers in the UK were randomized and followed up
between April 24th 2008 and January 27th 2015 (Figure 1). Baseline demographic and clinical

202 characteristics, including stroke severity and oxygen saturation at randomization, were well 203 balanced in the three groups (Table 1). The mean age of participants was 72 (SD 13) years, 4398 204 (55%) were male, and 7332 (92%) could undertake activities of daily living independently before 205 the stroke. The mean/median NIHSS was 7/5 (SD 6/IQR 3-9). 1601 (20%) had been given oxygen 206 prior to randomization, in the ambulance or in hospital. Patients were enrolled at a median 207 20h:43min (IQR 11:59-25:32) after symptom onset. The mean oxygen saturation at randomization 208 was 96.6% (SD 1.7%). All participants had a clinical diagnosis of stroke at the time of enrolment. 209 The final diagnosis at 7 days was ischemic stroke in most cases (n=6555, 82%), 588 (7%) had a 210 primary intracerebral hemorrhage, and 294 (4%) were strokes without CT diagnosis. 168 (2%) were 211 given a final diagnosis of transient ischemic attack, and 292 (4%) were found to have other non-212 stroke diagnoses, with missing data in 106(1%).

213 6991 (87%) of participants gave fully informed consent and 1012 (13%) had consent given by a

relative, carer, or an independent legal representative (eTable 1). Of the participants who were

unable to consent themselves and were included by a representative, 6(0.1%) refused consent at the

216 1-week reassessment and 22 (2%) at the 90-day assessment, and were withdrawn.

217 Treatment adherence

Adherence was similar in the continuous and night-time oxygen groups, with 2158 (81%) and 2225 (83%), respectively, prescribed oxygen for the full course of treatment (eTable 2). 433 (16%)

220 participants in the continuous and 361 (14%) in the night-time group discontinued oxygen

221 prematurely. The most common reason for early discontinuation of oxygen was discharge from

hospital. In the control group, trial oxygen was recorded as being given in 33 (1.2%), with no

recording of whether oxygen was given in 406 (15%).

224 Effect on oxygenation

225 Oxygen treatment resulted in a significant increase of 0.8% and 0.9% in the highest and lowest 226 oxygen saturations, respectively, during the 72 h of the intervention period in the continuous 227 oxygen group compared to controls, and of 0.5% and 0.4% for the highest and lowest oxygen 228 saturations, respectively, in the nocturnal oxygen group compared to controls (p<0.001 for all 229 comparisons, Table 2). Significantly more participants in the combined oxygen groups required 230 oxygen for clinical reasons during the intervention period than in the control group: 9% (463) vs. 231 7% (176), p<0.001. Similarly, more participants in the continuous than nocturnal oxygen group 232 required oxygen: 10% (254) vs. 8% (209), p=0.03.

233 Main outcome

- The primary analysis demonstrated that oxygen supplementation did not significantly improve
- functional outcome at 90 days (Figure 2). The unadjusted OR for a better outcome (lower mRS)
- 236 was 0.97 (95% CI 0.89–1.05, p=0.47) for combined oxygen versus control, and 1.03 (95% CI 0.93–
- 237 1.13, p=0.61) for continuous oxygen versus nocturnal oxygen. Secondary analyses adjusted for age,
- sex, baseline NIHSS score, baseline oxygen saturation, and the SSV prognostic index yielded very
- similar results: OR=0.97 (95% CI 0.89–1.06, p=0.54) for the combined oxygen group versus
- control and OR=1.01 (95% CI 0.92–1.12, p=0.81) for continuous oxygen versus oxygen at night
- only. With similar numbers of missing responses in the continuous oxygen (n=101), nocturnal
- 242 oxygen (n=106), and control groups (n=119), findings were much the same in sensitivity analyses
- using multiple imputation or analyzing adherers only (eTable 3).

Subgroup analysis (figure 3) found no indication that treatment effectiveness differed in any of the
 predefined subgroups, even those where most benefit might be expected – such as patients with
 more severe strokes or those for whom oxygen supplementation was started early after onset of

stroke.

248 Secondary outcomes

Analyses of secondary outcomes also showed no benefit from oxygen (Table 2). Neurological

250 impairment at one week improved from baseline to the same degree in all three groups, with median

251 NIHSS scores of 2 (IQR 1–6) by one week. Oxygen treatment did not increase the number of

252 participants who were alive and independent, or back in their home, the ability to perform basic (BI)

or extended (NEADL) activities of daily living, or quality of life (EQ5D-3L) at 90 days. The results

remained unchanged after adjustment for baseline prognostic factors (eTable 4). Mortality (figure 4)

was similar in the oxygen (both groups combined) and control groups (hazard ratio [HR] =0.97

256 [99% CI 0.78–1.21], p=0.75), and for continuous oxygen versus oxygen at night only (HR=1.15

257 [99% CI 0.90–1.48], p=0.15).

258 Exploratory analyses

There was no evidence of increased stress levels (higher heart rates, higher blood pressure and need for sedation) in oxygen-treated participants than in controls, or that oxygen treatment was associated with more infections, with little differences in the highest temperature or the need for antibiotics (Table 2).

263 Safety outcomes

The number of serious adverse events by 90 days was similar in the combined oxygen and control groups, but lower in the nocturnal oxygen group when compared to continuous oxygen (Table 2 and eTable 5). No oxygen-related adverse events (respiratory depression, drying of mucous membranes) were reported.

269 **DISCUSSION**

270 The key finding of this trial is that routine prophylactic low-dose oxygen supplementation did not 271 improve outcome in patients with acute stroke who were not hypoxic at baseline, whether given 272 continuously for 72 hours or at night only. This applied to the primary 90-day functional outcome 273 and to all other tested outcomes, including early neurological recovery, mortality, disability, 274 independence in basic and extended activities of daily living, and quality of life. The results 275 remained unchanged in analyses adjusted for baseline prognostic factors, and in sensitivity analyses 276 using multiple imputation or analyzing adherers only. Subgroup analyses did not identify any 277 characteristics that would make a patient more likely to benefit from oxygen treatment. This 278 includes enrolment between 3-6 hours after onset, patients with a lower baseline oxygen saturation, 279 severe strokes, a reduced level of consciousness, and a history of heart failure or lung disease; i.e. 280 those characteristics for which benefit from oxygen was most anticipated. Because of the large 281 overall size of this trial, these patient subgroups were each sufficiently large for the lack of 282 observed benefit to be likely real and not a false negative.

In contrast to the much smaller SOS Pilot study, $\frac{15}{15}$ this trial showed no evidence of better early 283 284 neurological recovery with oxygen. Subgroup analysis of an earlier study of low-dose oxygen supplementation in acute stroke¹⁴ suggested that oxygen might adversely affect outcome in patients 285 286 with mild strokes, possibly through formation of toxic free radicals. A more recent study of shortburst high-flow oxygen (45L/min) was terminated early after enrolment of 85 patients because of 287 excess mortality in the actively treated group. $\frac{13}{13}$ Hyperoxia was independently associated with 288 mortality in a large retrospective cohort study of ventilated patients with stroke.²⁸ While suggestive 289 290 of potential harm, these findings could be due to confounding factors. This trial showed no 291 difference in mortality, functional outcomes and adverse events and therefore provides reassuring 292 evidence that low-dose oxygen supplementation is safe in patients with acute stroke.

As a large pragmatic trial, this study included patients with a clinical diagnosis of acute stroke,
without radiological confirmation. The sample therefore included ischemic and hemorrhagic
strokes, and participants who were later found to have mimics or transient ischemic attacks.

296 This trial was a large pragmatic study aimed at unselected patients with stroke. Over half of all 297 acute stroke services in the UK participated and wide inclusion criteria allowed enrolment of a 298 representative sample of ischemic and hemorrhagic patients with stroke across the whole range of 299 severity. Stroke severity was similar to that of the UK stroke population as a whole, with a median 300 NIHSS of 5 in this trial and 4 in the UK Sentinel Stroke National Audit Programme, which includes every stroke patient admitted to UK hospitals.³¹ The median NIHSS of 127,950 patients with acute 301 ischemic stroke in the US Get with the Guidelines Register^{32} was 5, as in this trial. A median 302 303 NIHSS of 5 at baseline was also recorded in a large Dutch study of antibiotic prophylaxis after stroke, with similarly wide inclusion criteria. $\frac{33}{2}$ 304

305 This study has several limitations. Minor benefits from oxygen treatment might have been masked 306 by poor compliance. However, this seems unlikely given the high statistical power to detect even 307 small improvements. Moreover, sensitivity analyses did not show better outcomes in the adherers-308 only group (eTable 3). Furthermore, this trial found significant increases in the oxygen saturations 309 in the treated groups compared to control. Patients with acute stroke are often restless and confused. 310 Ensuring full adherence would ideally require 1:1 nursing. However, this is not possible outside an 311 intensive care setting. The main outcome was assessed by postal questionnaire, supported by telephone interviews in non-responders. This method has been used successfully in large pragmatic 312 trials, $\frac{29,30}{29,30}$ but has been replaced by remote multiple-rater video-recorded interviews or in-person 313 314 interview and examination by an allocation-blinded rater using formal structured assessments in 315 several more recent studies. Low-dose oxygen supplementation may not be sufficient to prevent severe desaturations; both the SOS Pilot $\frac{15}{15}$ and this trial found no significant difference in severe 316

317 desaturations between the treatment and control groups. A small (n=46) non-randomized study 318 comparing high-flow oxygen treatment via mask with low-flow supplementation via nasal cannulae 319 showed a trend towards lower mortality with high flow. However, evidence from randomized trials of high-flow oxygen treatment in acute stroke $\frac{11,12,13}{12}$ does not show that higher doses of oxygen are 320 321 associated with better outcomes. Early administration of high-dose oxygen might help maintain the 322 viability of the ischemic penumbra and allow a broader time window for neuroprotection or thrombolysis. This question was not addressed in this trial of prophylactic oxygen, but will be 323 324 tested in the PROOF trial (http://www.safestroke.eu/proof-trial/).

The median time from stroke onset to randomization in this trial was 20h 43min. However, 101 participants were enrolled early (within 3 hours of symptom onset). Subgroup analysis (figure 3) showed a similar lack of effect for oxygen in the small subset of patients enrolled early as in those enrolled later, but was underpowered. Larger trials in the early time window would be needed to

329 definitely exclude a benefit.

330 Conclusions

331 Among non-hypoxic patients with acute stroke the prophylactic use of low-dose oxygen

332 supplementation did not reduce death or disability at 3 months. These findings do not support low-

dose oxygen in this setting.

334

335 **CONTRIBUTORS**

336 CR and RG designed the trial. TN, CR, JS, NI, and RG ran the trial and CR recruited patients.

Analyses were planned by NI, RG and JS and undertaken by JB and JS; PF reviewed the literature.

338 CR, TN, JS and RG drafted the report and revised it with advice from all writing committee

members.

340

341 **DECLARATION OF INTERESTS**

342 C. Roffe received lecture and travel fees from Air Liquide and is an independent member of the343 data safety and monitoring committee of the PROOF trial. There are no other competing interests.

344

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346 The funders had no role in design and conduct of the study; collection, management, analysis, and 347 interpretation of the data; preparation, review, or approval of the manuscript; and decision to submit 348 the manuscript for publication.

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393 N Saravanan, N Thomas (133); *Queen's Hospital, Burton J Birch, R Damant, B Mukherjee** (131);

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395 Wright (129); Wansbeck Hospital, Northumberland C Ashbrook-Raby, A Barkat, R Lakey, C

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397 Eastwood, M James*, S Keenan (113); Royal United Hospital, Bath J Avis, D Button, D Hope, B

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400 G Thomas (105); York Hospital, York J Coyle*, N Dyer, S Howard, M Keeling, S Williamson

401 (105); University Hospital of North Durham, Durham E Brown, S Bruce, B Esisi*, R Hayman, E

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407	N Gautam, C Maity*, R Miller, C Mustill, M Salehin*, A Walker (87); Kent & Canterbury
408	Hospital, Canterbury H Baht, I Burger*, L Cowie, T Irani, A Thomson (84); New Cross Hospital,
409	Wolverhampton P Bourke, K Fotherby*, D Morgan, K Preece (84); Northwick Park Hospital,
410	Harrow L Burgess, D Cohen*, M Mpelembue (83); Barnsley District General Hospital, Barnsley
411	M Albazzaz*, R Bassi, C Dennis, K Hawley, S Johnson-Holland (82); Blackpool Victoria Hospital,
412	Blackpool H Goddard, J Howard, C Jeffs, J Mcilmoyle*, A Strain (82); North Tyneside General
413	Hospital, North Shields J Dickson, K Mitchelson, C Price*, V Riddell, A Smith (79); Eastbourne
414	District General Hospital, Eastbourne C Athulathmudali*, E Barbon (76); Warrington Hospital,
415	Warrington K Bunworth, L Connell, G Delaney-Sagar, K Mahawish*, O Otaiku*, H Whittle (75);
416	Princess Royal Hospital, Haywards Heath R Campbell*, A Nyarko (71); City Hospitals,
417	Sunderland S Crawford, C Gray*, D Gulliver, R Lakey, N Majmudar*, S Rutter (69); William
418	Harvey Hospital, Ashford L Cowie, D Hargroves*, T Webb (69); Stepping Hill Hospital, Stockport
419	A Brown, H Cochrane, S Krishnamoorthy*, J McConniffe (66); The James Cook University
420	Hospital, Middlesborough D Broughton*, K Chapman, L Dixon, A Surendran (66); Northampton
421	General Hospital (Acute), Northampton M Blake*, F Faola, A Kannan, P Lai, B Vincent (59);
422	Leicester General Hospital, Leicester M Dickens, D Eveson, S Khan, R Marsh, A Mistri*,(57);
423	Rotherham District General Hospital, Rotherham J Harris, J Howe, K McNulty, J Okwera* (56); St
424	Peter's Hospital, Chertsey R Nari*, E Young (56); Macclesfield District General Hospital,
425	Macclesfield A Barry, B Menezes, M Sein*, H Rooney, L Wilkinson (55); Manor Hospital, Walsall
426	S Hurdowar, K Javaid*, K Preece (54); Bradford Royal Infirmary, Bradford R Bellfield, B
427	Hairsine, L Johnston, C Patterson*, S Williamson (53); Luton & Dunstable Hospital, Luton F
428	Justin, S Sethuraman*, L Tate (50); Royal Blackburn Hospital, Blackburn A Bell, M Goorah, N
429	Goorah*, A Sangster (50); University College Hospital, London N Bhupathiraju, L Latter, P
430	Rayson, R Simister*, R Uday Erande (50); Addenbrooke's Hospital, Cambridge N Butler, D Day, E
431	Jumilla, J Mitchell, E Warburton* (48); Queen Alexandra Hospital, Portsmouth T Dobson, C

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457	S Ragab* (24	(4): Frenchav	Hospital.	Bristol N Baldwin*	. S Hierons.	H Skuse.	L Whelan ((22):

- 458 Princess Alexandra Hospital, Harlow L Brown, M Burton, A Daniel, S Hameed*, S Mansoor*
- 459 (22); West Suffolk Hospital, Bury St Edmunds A Azim*, M Krasinska, J White (22); The Ulster
- 460 Hospital, Dundonald M Power*, B Wroath (21); Watford General Hospital, Watford D Collas*, S
- 461 Sundayi, E Walker (21); Southampton General Hospital, Southampton M Brown, G Durward*, V
- 462 Pressly, B Watkins, N Weir*, D Whittaker (20); Craigavon Area Hospital, Portadown C Douglas,
- 463 M McCormick*, M McParland (19); Royal Lancaster Infirmary, Lancaster C Culmsee, P Kumar*
- 464 (18); Basildon Hospital, Basildon M Bondoc, B Hadebe, R Rangasami*, I Udeozor, U
- 465 Umansankar* (17); Birmingham City Hospital, Sandwell F Kinney, S Hurdowar, S Ispoglou*, S
- 466 Kausar* (17); City Hospital, Nottingham P Cox, A Ferguson, D Havard, F Shelton, A Shetty* (16);
- 467 Antrim Area Hospital, Antrim C Edwards, C McGoldrick, A Thompson, D Vahidassr* (15);
- 468 Pinderfields General Hospital, Wakefield G Bateman, P Datta*, A Needle (15); Royal Albert
- 469 Edward Infirmary, Wigan P Farren, S Herath* (15); Good Hope Hospital, Sutton Coldfield I
- 470 Memon*, S Montgomery (13); Hereford County Hospital, Hereford S Black, S Holloman, C
- 471 Jenkins*, F Price (13); South Tyneside District General Hospital, South Shields M Duffy, J
- 472 Graham, J Scott (13); Broomfield Hospital, Chelmsford A Lyle, F Mcneela, K Swan, J Topliffe, V
- 473 Umachandran* (12); Wythenshawe Hospital, Wythenshawe B Charles, E Gamble*, S Mawn (11);
- 474 Warwick Hospital, Warwick M Dean, B Thanvi* (10); Ipswich Hospital, Ipswich M Chowdhury*, J
- 475 Ngeh, S Stoddart (9); Kettering General Hospital, Kettering K Ayes*, J Kessell (9); Nevill Hall
- 476 Hospital, Abergavenny B Richard*, E Scott (9); Princess Royal University Hospital, Orpington L
- 477 Ajayo, E Khoromana, E Parvathaneni, B Piechowski-Jozwiak*, L Sztriha* (9); Scarborough
- 478 General Hospital, Scarborough L Brown, K Deighton, E Elnour, J Paterson*, E Temlett (9); Hull
- 479 Royal Infirmary, Hull A Abdul-Hamid*, J Cook, K Mitchelson (8); King's Mill Hospital, Sutton-in-
- 480 Ashfield M Cooper*, I Wynter (8); The Royal London Hospital, London P Gompertz*, O Redjep, J
- 481 Richards, R Uday Erande (8); Trafford General Hospital, Manchester S Anwar*, A Ingram, S

- 482 McGovern, S Musgrave*, L Tew (8); Altnagelvin Area Hospital, Londonderry J Corrigan*, C
- 483 Diver-Hall, M Doherty, M McCarron* (7); Darent Valley Hosptial, Dartford P Aghoram*, T
- 484 Daniel, S Hussein, S Lord (7); *Royal Berkshire Hospital, Reading* N Mannava, A van Wyk* (6);
- 485 Arrowe Park Hospital, Wirral J Barrett*, R Davies*, A Dodd, D Lowe*, P Weir (5); Basingstoke
- 486 and North Hampshire Hospital, Basingstoke D Dellafera, E Giallombardo* (5); Lincoln County
- 487 Hospital, Lincoln S Arif, R Brown, S Leach* (5); Hexham General Hospital, Hexham C Price*, V
- 488 Riddell (4); Manchester Royal Infirmary, Manchester J Akyea-Mensah, J Simpson* (4); Salisbury
- 489 *District Hospital, Salisbury* T Black*, C Clarke, M Skelton (4); *Croydon University Hospital,*
- 490 Croydon J Coleman, E Lawrence* (3); Russells Hall Hospital, Dudley A Banerjee*, A Boyal, A
- 491 Gregory (3); Worthing Hospital, Worthing S Ivatts*, M Metiu (3); Bedford Hospital, Bedford A
- 492 Elmarimi*, S Hunter (2); James Paget Hospital, Great Yarmouth H Benton, M Girling, P Harrison*,
- 493 H Nutt, S Mazhar Zaidi*, C Whitehouse (2); St Richard's Hospital, Chichester G Blackman, S
- 494 Ivatts* (2); Erne Hospital, Fermanagh M Doherty, J Kelly* (1); University Hospital Lewisham,
- 495 Lewisham M Patel* (1); Bronglais General Hospital, Aberystwyth P Jones* (0); Hillingdon
- 496 Hospital, Hillingdon A Parry* (0); Kingston Hospital, Kingston upon Thames L Choy* (0);
- 497 Morriston Hospital, Morriston (0); North Middlesex Hospital, Enfield T Adesina, A David, R
- 498 Luder* (0); Staffordshire District General Hospital, Stafford A Oke* (0); St Helier Hospital,
- 499 Carshalton V Jones*, P O'Mahony, C Orefo (0); Whipps Cross University Hospital, London R
- 500 Simister* (0).

501 **REFERENCES**

- 502 1. Roffe C, Sills S, Halim M, et al. Unexpected nocturnal hypoxia in patients with acute stroke.
 503 *Stroke*. 2003; **34**: 2641–5.
- 2. Rocco A, Pasquini M, Cecconi E, et al. Monitoring after the acute stage of stroke: a prospective
 study. *Stroke*. 2007; 38: 1225–8.
- 3. Bravata DM, Wells CK, Lo AC, et al. Processes of care associated with acute stroke outcomes. *Arch Intern Med.* 2010; **170**: 804-10.
- 4. Rowat AM, Dennis MS, Wardlaw JM. Hypoxaemia in acute stroke is frequent and worsens
 outcome. *Cerebrovasc Dis.* 2006; 21: 166–72.
- 5. Heiss WD. The ischemic penumbra: how does tissue injury evolve? *Ann N Y Acad Sci*.
 2012;1268:24-34.
- 512 6. Alawneh JA, Jones PS, Mikkelsen IK, et al. Infarction of 'non-core-non-penumbral' tissue after
- 513 stroke: multivariate modelling of clinical impact. *Brain*. 2011;**134**:1765-76.
- 514 7. Dreier JP. The role of spreading depression, spreading depolarization and spreading ischemia in
 515 neurological disease. *Nat Med.* 2011;**17**:439-47.
- 516 8. Ciccone A, Celani MG, Chiaramonte R, Rossi C, Righetti E. Continuous versus intermittent
- 517 physiological monitoring for acute stroke. *Cochrane Database Syst Rev.* 2013; 5: CD008444.
- 518 9. O'Driscoll BR, Howard L, Earis J, et al. BTS Guideline for oxygen use in adults in healthcare
- 519 and emergency settings. *Thorax.* 2017;**72**(suppl 1):i1-i90.
- 520 10. Floyd TF, Clark JM, Gelfand R, et al. Independent cerebral vasoconstrictive effects of
- 521 hyperoxia and accompanying arterial hypocapnia at 1 ATA. *J Appl Physiol.* 2003; **95**: 2453–61.

- 522 11. Padma MV, Bhasin A, Bhatia R, et al. Normobaric oxygen therapy in acute ischemic stroke: A
 523 pilot study in Indian patients. *Ann Indian Acad Neurol.* 2010; 13: 284–8.
- 524 12. Singhal AB, Benner T, Roccatagliata L, et al. A pilot study of normobaric oxygen therapy in
 525 acute ischemic stroke. *Stroke*. 2005; **36**: 797–802.
- 526 13. Singhal AB. Normobaric oxygen therapy in acute ischemic stroke trial ClinicalTrials.gov.
- 527 https://clinicaltrials.gov/ct2/show/NCT000414726 (accessed Sept 14, 2015).
- 528 14. Rønning OM, Guldvog B. Should stroke victims routinely receive supplemental oxygen? A
- 529 quasi-randomized controlled trial. *Stroke*. 1999; **30**: 2033–7.
- 530 15. Roffe C, Ali K, Warusevitane A, et al. The SOS pilot study: A RCT of routine oxygen
- supplementation early after acute stroke—effect on recovery of neurological function at one week. *PLoS ONE*. 2011; 6: e19113.
- 16. Roffe C, Nevatte T, Crome P, et al. The Stroke Oxygen Study (SO₂S) a multi-center study to
 assess whether routine oxygen treatment in the first 72 hours after a stroke improves long-term
 outcome: study protocol for a randomized controlled trial. *Trials*. 2014; 15: 99.
- 536 17. Sim J, Gray R, Nevatte T, Howman A, Ives N, Roffe C. Statistical analysis plan for the Stroke
 537 Oxygen Study (SO₂S): a multi-center randomized controlled trial to assess whether routine oxygen
 538 supplementation in the first 72 hours after a stroke improves long-term outcome. *Trials*. 2014; 15:
 539 229.
- 540 18. Stroke Oxygen Study. http://www.so2s.co.uk/ (accessed Jul 14, 2016).
- 541 19. Pocock SJ, Simon R. Sequential Treatment Assignment with Balancing for Prognostic Factors
- 542 in the Controlled Clinical Trial. *Biometrics. International Biometric Society.* 1975;**31**: 103–115.

543	20. Counsell C, Dennis M, McDowall M, Warlow C. Predicting outcome after acute and subacute
544	stroke: development and validation of new prognostic models. Stroke. 2002; 33: 1041-7.
545	21. van Swieten JC, Koudstaal PJ, Visser MC, Schouten HJ, van Gijn J. Interobserver agreement
546	for the assessment of handicap in stroke patients. <i>Stroke</i> . 1988; 19 : 604–7.
547	22. Brott T, Adams HP, Olinger CP, et al. Measurements of acute cerebral infarction: a clinical
548	examination scale. <i>Stroke</i> . 1989; 20 : 864–70.
549	23. Wityk RJ, Pessin MS, Kaplan RF, Caplan LR. Serial assessment of acute stroke using the NIH
550	Stroke Scale. <i>Stroke</i> .1994; 25 : 362–5.
551	24. Collin C, Wade DT, Davies S, Horne V. The Barthel ADL Index: a reliability study. Int Disabil
552	<i>Stud.</i> 1988; 10 : 61–3.
553	25. EuroQol Group. EuroQol – a new facility for the measurement of health-related quality of life.
554	<i>Health Policy</i> . 1990; 16 : 199–208.
555	26. Nouri FM, Lincoln NB. An extended activities of daily living scale for stroke patients. Clin
556	<i>Rehabil.</i> 1987; 1 : 301–5.
557	27. Ali M, Jüttler E, Lees KR, Hacke W, Diedler J, for the VISTA and DESTINY Investigators.
558	Patient outcomes in historical comparators compared with randomised-controlled trials. Int J
559	<i>Stroke</i> . 2010; 5 : 10–5.

- 560 28. Rincon F, Kang J, Maltenfort M, et al. Association between hyperoxia and mortality after
- 561 stroke: a multicenter cohort study. *Crit Care Med.* 2014; **42**: 387–96.

562	29. Dennis MS, Lewis SC, Warlow C; FOOD Trial Collaboration. Effect of timing and method of
563	enteral tube feeding for dysphagic stroke patients (FOOD): a multicentre randomised controlled
564	trial. Lancet 2005;26;365:764-72.
565	30. IST-3 collaborative group. The benefits and harms of intravenous thrombolysis with
566	recombinant tissue plasminogen activator within 6 h of acute ischaemic stroke (the third
567	international stroke trial [IST-3]): a randomised controlled trial. Lancet 2012;379:2352-63.

568 31. Smith CJ, Bray BD, Hoffman A, et al. Can a novel clinical risk score improve pneumonia
569 prediction in acute stroke care? A UK multicenter cohort study. *J Am Heart Assoc.* 2015; 4:
570 e001307.

571 32. Fonarow GC, Pan W, Saver JL, et al. Comparison of 30-day mortality models for profiling
572 hospital performance in acute ischemic stroke with vs without adjustment for stroke severity.
573 JAMA. 2012; 308: 257–64.

33. Westendorp WF, Vermeij JD, Zock E, et al. The Preventive Antibiotics in Stroke Study
(PASS): a pragmatic randomised open-label masked endpoint clinical trial. *Lancet*. 2015; 385:
1519–26.

577

578 33. López-Cancio E, Salvat M, Cerdà N, et al. Phone and Video-Based Modalities of Central
579 Blinded Adjudication of Modified Rankin Scores in an Endovascular Stroke Trial. *Stroke*.
580 2015;46:3405-10.

582 **TABLES**

583

584 **Table 1 Baseline characteristics**

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	Continuous oxygen n=2668	Nocturnal oxygen n=2667	Control n=2668
Demographic characteristics			1
Age; (years); mean (SD) ^a	72 (13)	72 (13)	72 (13)
Male sex; <i>n</i> (%)	1466 (55)	1466 (55)	1466 (55)
Prognostic factors			
Living alone before the stroke; n (%) ^a	861 (32)	857 (32)	907 (34)
Independent in basic ADLs before the stroke; n (%) ^a	2451 (92)	2431 (91)	2450 (92)
Normal verbal response; n (%) ^a	2190 (82)	2207 (83)	2196 (82)
Able to lift both arms; n (%) ^a	1998 (75)	2022 (76)	1996 (75)
Able to walk; $n(\%)^{a}$	660 (25)	704 (26)	677 (25)
Probability of 30-day survival; median $(IQR)^{\frac{20}{2}}$	0.92 (0.86-0.95)	0.92 (0.86-0.95)	0.92 (0.86-0.95)
Alive and independent at 6 m; <i>probability</i> median (IQR) $a \frac{20}{2}$	0.44 (0.12-0.71)	0.42 (0.12-0.71)	0.42 (0.12-0.71)
Blood glucose; mg/dl mean (SD)	127 (46)	126 (43)	128 (45)
Concomitant medical problems			
Ischemic heart disease; <i>n</i> (%)	573 (21)	515 (19)	514 (19)
Heart failure; <i>n</i> (%)	224 (8)	217 (8)	216 (8)
Atrial fibrillation; <i>n</i> (%)	638 (24)	673 (25)	684 (26)
Chronic obstructive pulmonary disease/asthma; n (%)	253 (9)	242 (9)	245 (9)
Other chronic lung problem; n (%)	29 (1)	24 (1)	19 (1)
Details of the qualifying event			
Time since symptom onset; hh:mm median (IQR) ^a	20:44 (11:53-25:33)	20:32 (12:05-25:31)	20:45 (11:57-25:31)
Diagnosis:			
Transient ischemic attack; $n (\%)^{b}$	52 (1.9)	50 (1.9)	66 (2.5)
Ischemic stroke; $n(\%)^{b}$	2187 (82.0)	2165 (81.1)	2203 (82.6)
Intracerebral hemorrhage; $n(\%)^{b}$	185 (6.9)	207 (7.8)	196 (7.3)
Stroke without imaging diagnosis; $n(\%)^{b}$	104 (3.9)	106 (4.0)	84 (3.1)
Not a stroke; $n (\%)^{b}$	101 (3.8)	98 (3.7)	93 (3.5)
Missing; $n (\%)^{b}$	39 (1.5)	41 (1.5)	26 (1.0)
Glasgow Coma Scale score (3–15); median (IQR) [range]	15 (15–15) [4-15]	15 (15–15) [5-15]	15 (15–15) [3-15]
Thrombolysed; $n (\%)^{b}$	447 (17)	410 (15)	447 (17)
NIHSS score (0–42); median (IQR)	5 (3–9)	5 (3-9)	5 (3-9)
Oxygenation			
Oxygen given prior to randomization; $n (\%)^{a}$	531 (20)	531 (20)	539 (20)
Oxygen saturation on room air; % mean (SD) ^a	96.6 (1.7)	96.6 (1.6)	96.7 (1.7)

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587 Data in this table were collected before randomization with the exception of items marked with ^b 588 which were recorded on day 7. Minimization variables are indicated by ^a. Activities of daily living 589 (ADLs). The probability of being alive and independent was calculated using the 'six simple 590 variables (SSV) prognostic index for independent survival at 6 months (m).²⁰ 'Normal verbal 591 response' was taken from the verbal sub-item of the Glasgow Coma Scale. See online supplement 592 eText 1 for definitions for diagnosis. The Glasgow Coma Scale score ranges from 3 (deep coma) to 593 15 (alert and oriented). The National Institutes for Heath Stroke Scale ranges from 0 (no deficit) to 42 (most severe deficit). Blood glucose was converted from mm/L to mg/dl by multiplying by afactor of 18.

Table 2 Secondary, exploratory and safety outcomes

5	9	8

	N=8003 (n)	Continuous oxygen n=2668	Nocturnal oxygen n=2667	Control n=2668	Comparison 1: Combined oxygen vs. control OR, MD, MdD or RR (99%CI); p-value	Comparison 2: Continuous vs. nocturnal OR, MD, MdD or RR (99%CI); p-value
Secondary outcomes at 72 hours						
Highest oxygen saturation (%) ^a	7860	99.1 (99.1-99.2)	98.8 (98.7-98.9)	98.3 (98.2-98.3)	MD 0.69 (0.61-0.77); p<0.0001 ^d	MD 0.32 (0.22-0.41); p<0.0001 ^d
Lowest oxygen saturation (%) ^a	7860	95.0 (94.9-95.1)	94.5 (94.4-94.6)	94.1 (94.0-94.2)	MD 0.62 (0.48-0.76); p<0.0001 ^d	MD 0.48 (0.32-0.63); p<0.0001 ^d
Oxygen saturation <90% ^b	7860	39 (1.5%)	30 (1.1%)	74 (2.8%)	OR 0.46 (0.30-0.71); p<0.0001 ^e	OR 1.30 (0.69-2.44); p=0.28 ^e
Oxygen saturation <95% ^b	7860	861 (32.9%)	1119 (42.9%)	1354 (51.5%)	OR 0.57 (0.51-0.65); p<0.0001 ^e	OR 0.65 (0.56-0.76); p=<0.0001 ^e
Need for additional oxygen ^b	7809	254 (9.8%)	209 (8.1%)	176 (6.7%)	OR 1.36 (1.07-1.73); p=0.0008 ^e	OR 1.23 (0.96-1.59); p=0.03 ^e
Secondary outcomes at 7 days						
National Institutes of Health Stroke Scale ^c	7778	2 (2–3)	2 (2–3)	2 (2–3)	MdD 0 (0–0); p=0.56 ^f	MdD 0 (0–0); p=0.95 ^f
Neurological improvement ^b	7778	1016 (39.2%)	1029 (39.7%)	1037 (39.9%)	OR 0.98 (0.86-1.11); p=0.68 ^e	OR 0.98 (0.85-1.13); p=0.71 ^e
Death by 7 days ^b	7959	50 (1.9%)	35 (1.3%)	45 (1.7%)	OR 0.95 (0.59-1.53); p=0.78 ^e	OR 1.43 (0.81-2.54); p=0.11 ^e
Secondary outcomes at 90 days						
Death by 90 days (mRS=6) ^b	7677	257 (10.0%)	236 (9.2%)	246 (9.7%)	OR 1.00 (0.81-1.23); p=0.96 ^e	OR 1.10 (0.86-1.40); p=0.3 ^e
Alive and independent $(mRS \le 2)^b$	7677	1325 (51.6%)	1316 (51.4%)	1337 (52.5%)	OR 0.96 (0.85-1.09); p=0.43 ^e	OR 1.01 (0.87-1.17); p=0.87 ^e
Living at home ^b	6859	1961 (85.8%)	1947 (84.8%)	1947 (85.4%)	OR 0.99 (0.82-1.20); p=0.91 ^e	OR 1.08 (0.87-1.34); p=0.35 ^e
Barthel ADL index [0 (worst)-100 (best)] ^a	6549	70.2 (68.2-72.2)	71.1 (69.1-73.1)	70.9 (68.9-72.8)	$ \begin{array}{c} \text{MD -0.18 (-2.60-2.24);} \\ \text{p=}0.85^{\text{d}} \end{array} $	MD -0.86 (-3.65-1.93); p=0.43 ^d
Nottingham Extended ADL [0 (worst)–21 (best)] ^a	7528	9.66 (9.29-10.02)	9.54 (9.17-9.90)	9.77 (9.40-10.14)	$ \begin{array}{c} \text{MD -0.17 (-0.62-0.28);} \\ \text{p=}0.32^{\text{d}} \end{array} $	$ \begin{array}{c} \text{MD } 0.12 \ (-0.40-0.64); \\ \text{p}=0.55^{\text{d}} \end{array} $

Table 2 continued	N=8003 (n)	Continuous oxygen n=2668	Nocturnal oxygen n=2667	Control n=2668	Comparison 1: Combined oxygen vs. control OR, MD, MdD or RR (99%CI); p-value	Comparison 2: Continuous vs. nocturnal OR, MD, MdD or RR (99%CI); p-value
Quality of life (EQ5D-3L) [- 0.59 (worst)-1 (best)] ^a	7248	0.50 (0.48-0.51)	0.50 (0.48-0.51)	0.49 (0.48-0.51)	MD 0.004 (-0.02-0.03) p=0.71 ^d	MD 0.003 (-0.03-0.03) p=0.78 ^d
Quality of life (VAS) [0 (worst)-100 (best)] ^a	6675	55.4 (53.8-67.1)	55.7 (54.1-57.3)	55.5 (53.8-57.1)	MD 0.10 (-1.93-2.12); p=0.90 ^d	MD -0.24 (-2.57-2.09) p=0.79 ^d
Exploratory outcomes						
Highest heart rate within 72 hours (beats per minute) ^a	7859	87.2 (86.3-88.0)	88.0 (87.2-88.8)	87.7 (86.9-88.4)	MD -0.07 (-1.06-0.92)	MD -0.83 (-2.01-0.35)
Highest systolic BP within 72 hours (mm Hg) ^a	7864	162.4 (161.2-163.7)	162.8 (161.5-164.0)	164.6 (163.3- 165.8)	MD -1.96 (-3.48-(-0.44))	MD -0.35 (-2.11-1.41)
Highest diastolic BP within 72 hours (mm Hg) ^a	7861	89.5 (88.7-90.2)	90.2 (89.4-90.0)	90.9 (90.1-91.7)	MD -1.10 (-2.06- (-0.15))	MD -0.72 (-1.82-0.37)
Highest temperature within 7days (Celsius) ^a	7877	37.1 (37.1-37.2)	37.2 (37.1-37.2)	37.1 (37.1-37.2)	MD 0.01 (-0.03-0.04)	MD -0.01 (-0.05-0.03)
Antibiotics given within 7 days ^b	7916	400 (15.2%)	393 (14.9%)	403 (15.2%)	OR 0.99 (0.83-1.17)	OR 1.02 (0.84-1.24)
Sedatives given within 7 days ^b	7916	140 (5.3%)	161 (6.1%)	154 (5.8%)	OR 0.98 (0.76-1.28)	OR 0.86 (0.63-1.17)
Sleep as good as before the stroke ^b	6584	1407 (64%)	1436 (65%)	1419 (65%)	OR 0.98 (0.85-1.13)	OR 0.96 (0.82-1.13)
No significant speech problems ^b	6716	1957 (88%)	1957 (87%)	1939 (87%)	OR 1.09 (0.89-1.32)	OR 1.06 (0.84-1.34)
Memory as good as before the stroke ^b	6646	981 (44%)	1000 (45%)	971 (44%)	OR 1.02 (0.89-1.16)	OR 0.97 (0.83-1.13)
Safety						
Serious adverse events (SAEs) ^a	8003	0.16 (0.14-0.18)	0.13 (0.11-0.16)	0.16 (0.13-0.18)	RR 0.94 (0.78-1.13); p=0.37 ^g	RR 1.19 (0.96-1.47); p=0.03 ^g
Participants with least one SAE ^b	8003	348 (13.0%)	294 (11.0%)	322 (12.1%)	OR 1.00 (0.83-1.20); p=0.96 ^e	OR 1.21 (0.97-1.51); p=0.02 ^e

601 Data are given as means and 99% confidence intervals,^a numbers and percentages,^b or medians and 99% confidence intervals.^c Mean differences 602 (MD) are reported for means, median differences (MdD) for medians, odds ratios (OR) for frequencies, and rate ratios (RR) for count data. ORs 603 604 < 1 indicate that the outcome is less likely with oxygen than with control (reference category) in comparison 1 and less likely with continuous oxygen than with nocturnal oxygen (reference category) in comparison 2. Significance testing was by unrelated t-test^d chi-squared test,^e 605 Wilcoxon rank sum test,^f or negative binomial regression.^g The highest and lowest oxygen saturations were the highest/lowest record of oxygen 606 saturation on the participant's observation chart during the 72 hours after randomization. Neurological improvement is a decrease of 4 or more or 607 608 to zero on the National Institutes of Health Stroke Scale (NIHSS). Death by 90 days is a modified Rankin Scale (mRS) score of 6. Alive and independent is a modified Rankin Scale score of 2 or less. Activities of daily living (ADL), quality of life (EQ5D-3L) and visual analogue scale 609 (VAS). As outlined in the statistical analysis plan, $\frac{17}{17}$ significance tests were not conducted on the exploratory data and the outcomes suggested 610 611 by patients and carers. HR: heart rate. 612

613 FIGURE LEGENDS

614

615 Figure 1 Trial profile

616 This figure shows participant enrollment, withdrawals, and follow up. Data on the number of

- 617 patients screened are not available.
- 618

619 Figure 2 Main outcome: modified Rankin Scale at 90 days

620

621 From the ordinal regression analysis the unadjusted odds ratio (OR) for a better outcome 622 (lower mRS) was 0.97 (95% CI 0.89–1.05, p=0.47) for combined oxygen versus control, and 1.03 (95% CI 0.93–1.13, p=0.61) for continuous oxygen versus nightly oxygen. Modified 623 624 Rankin Scale (mRS): 0 = no symptoms, 1 = few symptoms, but able to carry out all previous 625 activities and duties, 2 = unable to carry out all previous activities, but able to look after own 626 affairs without assistance, 3 = needs some help with looking after own affairs, but able to 627 walk without assistance, 4 = unable to walk without assistance and unable to attend to own 628 bodily needs without assistance, but I does not need constant care and attention, 5 = major 629 symptoms (bedridden and incontinent, needs constant attention day and night), 6 = death.

630 Figure 3 Subgroup analyses: mRS at 90 days oxygen versus control

631

632 Subgroup analyses are depicted as a forest plot; p-values relate to the test for interaction. The 633 x-axis depicts the 'common' odds ratio (OR) for a better outcome over all 7 levels of the modified Rankin Scale score (mRS). It is derived from ordinal logistic regression. ORs > 1 634 635 indicate that a good outcome (low mRS) is more likely with oxygen than with control 636 (reference category). n is the total number of participants in that subgroup category. The size 637 of the markers reflects the total sample size in the subgroup concerned, with larger markers equating to more precise estimates. The subgroup thresholds for oxygen concentration at 638 639 randomization were revised from the prespecified thresholds as the analysis did not converge using the prespecified values. NIHSS: National Institutes of Health Stroke Scale; TIA: 640 641 transient ischemic attack; SSV: 'six simple variables' risk score; COPD: chronic obstructive 642 pulmonary disease; CCF: congestive cardiac failure; GCS: Glasgow Coma Scale.

643 Figure 4 Mortality up to 90 days

644

This figure shows the probability of death in the control group (grey dashed line) and the combined oxygen group (black line) in the top panel. The bottom panel shows the probability

- of death in the nocturnal oxygen group (dashed grey line) and the probability of death in the
- 648 continuous oxygen group (black line). The cut off for mortality for this figure was 90 days.
- This is different from the 90-day mortality reported in table 2 and Figure 2, where responses
- 650 were accepted up to 6 months if 3-month outcomes were not returned. Median and [range] of
- duration of follow-up was 90 days [0-90] in each treatment group.