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Sleep Apnea and Fetal Growth Restriction (SAFER) study: protocol for a pragmatic randomised clinical trial of positive airway pressure as an antenatal therapy for fetal growth restriction in maternal obstructive sleep apnoea

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ABSTRACT

Introduction  Fetal growth restriction (FGR) is a major contributor to fetal and neonatal morbidity and mortality with intrauterine, neonatal and lifelong complications. This study explores maternal obstructive sleep apnoea (OSA) as a potentially modifiable risk factor for FGR. We hypothesise that, in pregnancies complicated by FGR, treating mothers who have OSA using positive airway pressure (PAP) will improve birth weight and neonatal outcomes.

Methods and analysis  The Sleep Apnea and Fetal Growth Restriction study is a prospective, block-randomised, single-blinded, multicentre, pragmatic controlled trial. We enrol pregnant women aged 18–50, between 22 and 31 weeks of gestation, with established FGR based on second trimester ultrasound, who do not have other prespecified known causes of FGR (such as congenital anomalies or intrauterine infection). In stage 1, participants are screened by questionnaire for OSA risk. If OSA risk is identified, participants proceed to stage 2, where they undergo home sleep apnoea testing. Participants are determined to have OSA if they have an apnoea-hypopnoea index (AHI) ≥5 (if the oxygen desaturation index (ODI) is also ≥5) or if they have an AHI ≥10 (even if the ODI is <5). These participants proceed to stage 3, where they are randomised to nightly treatment with PAP or no PAP (standard care control), which is maintained until delivery. The primary outcome is unadjusted birth weight; secondary outcomes include fetal growth velocity on ultrasound, enrolment-to-delivery interval, gestational age at delivery, birth weight corrected for gestational age, stillbirth, Apgar score, rate of admission to higher levels of care (neonatal intensive care unit or special care nursery) and length of neonatal stay. These outcomes are compared between PAP and control using intention-to-treat analysis.

Ethics and dissemination  This study has been approved by the Institutional Review Boards at Washington University in St Louis, Missouri; Hadassah Hebrew University Medical Center, Jerusalem; and the University of Rochester, New York. Recruitment began in Washington University in November 2019 but stopped from March to November 2020 due to COVID-19. Recruitment began in Hadassah Hebrew University in March 2021, and in the University of Rochester in May 2021. Dissemination plans include presentations at scientific conferences and scientific publications.

Trial registration number  NCT04084990.

Strengths and limitations of this study

► Sleep Apnea and Fetal Growth Restriction study is a multicentre, pragmatic trial conducted in a tertiary care setting.
► Studying patients with both fetal growth restriction and home sleep apnoea testing-diagnosed obstructive sleep apnoea (OSA) increases the potential for benefit from the intervention but limits generalisability.
► The intervention is positive airway pressure, a routine, safe and effective therapy for OSA in non-pregnant populations.
► While investigators and care providers are blinded to group allocation in stage 3, participants are unblinded.
► Positive findings will increase OSA awareness and treatment in pregnancy.

BACKGROUND

Fetal growth restriction (FGR) affects up to 10% of all pregnancies and is a major contributor to fetal and neonatal morbidity and mortality with intrauterine, neonatal and lifelong complications.1 2 FGR is second only to prematurity as a leading cause of perinatal morbidity and mortality. Both FGR and prematurity are independent risk factors for the development of cognitive delay, poor academic achievement and adult diseases.
such as obesity, type 2 diabetes mellitus, coronary artery disease and stroke.\(^3\) There are many potential causes of FGR, but in the absence of underlying genetic conditions, congenital anomalies or intrauterine infection, FGR is typically due to impaired uteroplacental perfusion.\(^2\)\(^4\) Current assessment is based on repeated ultrasound assessments of fetal growth, antenatal testing including non-stress test and/or biophysical profile (including electronic fetal heart rate monitoring) and umbilical artery Doppler velocimetry.\(^5\) Intervention is limited to antenatal steroid administration and interventional delivery when the risk of stillbirth is deemed too high to continue the pregnancy. There is no intervention currently available to improve uteroplacental blood flow and fetal growth in utero, so there is often no alternative to interventional delivery.\(^6\) When the pregnancy is remote from term, interventional delivery exposes an already compromised fetus to additional complications of prematurity, in particular to neonatal brain injury from intraventricular haemorrhage.\(^7\)\(^8\)

Pregnancy is associated with a higher incidence of sleep disordered breathing (SDB), a group of chronic conditions involving recurrent episodic partial or complete cessation of breathing throughout the night.\(^3\) Obstructive sleep apnoea (OSA) is an increasingly common form of SDB in both the general and pregnant population.\(^10\) OSA is characterised by complete (apnoea) or incomplete (hypopnoea) collapse of the upper airway during sleep leading to recurrent episodic cessation or limitation of normal breathing. This in turn leads to recurrent oxygen desaturation and hypercapnia, frequent night-time arousals and excessive daytime sleepiness.\(^11\)\(^12\)

OSA in pregnancy has been associated with poor maternal-fetal outcomes, including gestational hypertension, pre-eclampsia, gestational diabetes, FGR, low birth weight, preterm delivery and higher rates of neonatal intensive care unit (NICU) admission.\(^13\)\(^14\)\(^15\)\(^16\)\(^17\) Patients with OSA are more likely to have negative neonatal outcomes,\(^21\) and severe maternal morbidity and mortality,\(^17\) regardless of their obesity status. The relationship between severity of OSA in pregnancy and adverse outcomes is an ongoing area of research.\(^22\) Unfortunately, because few pregnant women are referred for polysomnography (PSG), it is likely that OSA and other sleep disorders are underdiagnosed,\(^23\) with the OSA-related symptoms of snoring, disrupted sleep and fatigue being frequently attributed to transient features of normal pregnancy.\(^24\)

Recurrent apnoeic and hypopnoeic episodes are associated with intermittent oxygen desaturation and hypercapnia. Recurrent hypoxia leads to oxidative stress, sympathetic activation and inflammation that may be harmful to both the mother and her fetus.\(^25\) Acute hypercapnia in pregnant mice and rats has been shown to cause acute placental hyperperfusion and acute fetal asphyxia.\(^26\) There is very little information about the effect of these episodes on the human fetus. One small observational study demonstrated fetal heart rate decelerations accompanying maternal oxyhaemoglobin desaturation,\(^27\) while another found no association.\(^28\)

Positive airway pressure (PAP) is a common and effective treatment for OSA which acts by mechanically splitting the upper airway with pressurised air to prevent collapse. PAP delivered at a single pressure (continuous PAP or CPAP) usually requires a sleep study to identify optimal settings; it also cannot respond to changes in upper airway function related to changes in habitus, body position, sleep stage or other factors. In contrast, auto-titrating PAP (aPAP) can detect upper airway resistance and respond by adjusting the delivered pressure (within a selected range). The delivered pressures from aPAP can therefore be lower than from CPAP, which makes aPAP more tolerable for most patients.\(^29\) We use aPAP as the intervention for all patients in the Sleep Apnea and Fetal Growth Restriction (SAFER) study. However, as much of the medical literature is still based on CPAP rather than aPAP, we use the generic term PAP throughout the remainder of this manuscript.

PAP is a proven low-risk therapeutic intervention for patients with OSA in the general, non-obstetric population. Multiple studies have shown that PAP use in patients with OSA reduces the incidence of death from cardiac-related complications including congestive heart failure, coronary artery disease, arrhythmia and stroke,\(^30\)\(^31\)\(^32\) and may improve outcomes related to diabetes.\(^33\)\(^34\) Small studies of PAP in pregnant women with severe pre-eclampsia and OSA diagnosed by PSG demonstrated improved maternal haemodynamic profiles and cardiac output\(^35\)\(^36\) although no obstetric or neonatal outcomes were measured.

The hypothesis of the SAFER study is that, in pregnancies complicated by FGR where the mothers have been diagnosed with OSA, maternal PAP therapy will improve intrauterine fetal growth and birth weight, increase randomisation-to-delivery interval and the gestational age at delivery and improve neonatal well-being. Ultimately, if PAP therapy is shown to improve intrauterine fetal growth in pregnancies with OSA and FGR, obstetric practice would be expected to change. Such a finding would lead to a clinical imperative to screen, diagnose and treat OSA in pregnancy, particularly in the presence of FGR.

**Methods**

**Research design overview**

The Human Research Protection Office at Washington University School of Medicine in St Louis, Missouri, the Research Subject Review Board at the University of Rochester, New York, and the Helsinki Committee for Ethics in Research in Human Subjects in Hadasah Hebrew University Medical Center, Jerusalem, Israel, all approved the study. The choice of these sites was determined by the location of the principal investigators involved in initiation of this study; however, all sites are academic, high-risk obstetric, tertiary referral centres with large numbers of pregnancies complicated by FGR. There is wide

Hincker A, et al. BMJ Open 2021;11:e049120. doi:10.1136/bmjopen-2021-049120
geographic and demographic diversity between these centres, which adds to the generalisability of the study. The SAFER study detailed in this protocol adheres to the Standard Protocol Items: Recommendations for Interventional Trials checklist, and lists the 24 items in the WHO Trial Registration Data Set. The protocol version is dated 5 February 2020. The SAFER study is a three-stage trial that assesses pregnant patients with diagnosed FGR. The overall flow of participants through the SAFER study is shown in figure 1.

We enrol women with pregnancies complicated by FGR between 22 and 31 weeks of gestation. Stage 1 is a brief telephone or in-person questionnaire to identify which of these pregnant women are at elevated risk for OSA; these women are then recruited into stage 2. Stage 2 is a prospective observational study using home sleep apnoea testing (HSAT) to confirm the diagnosis of OSA. Participants meeting OSA diagnostic criteria progress to stage 3. Stage 3 is a pragmatic randomised clinical trial of PAP as the intervention, versus a control group of standard care which does not include PAP. The primary outcome of stage 3 is birth weight; secondary outcomes include fetal growth velocity on ultrasound, birth weight corrected for gestational age, enrolment-to-delivery.
interval, gestational age at delivery and neonatal outcomes.

All three stages of this international, multicentre study will be conducted at three academic medical centres, which all serve a large and diverse population of high-risk obstetric patients. These centres are: Washington University School of Medicine in St Louis, Missouri (Barnes-Jewish Hospital); University of Rochester, New York (Strong Memorial Hospital); and Hadassah Hebrew University Medical Center, Jerusalem, Israel (Hadassah Hospital, Ein Karem and Hadassah Hospital, Mt Scopus). The demographic characteristics of these hospitals are summarised in table 1.

There is lack of clarity in current guidelines over the use of the term FGR. We use the American College of Obstetrics and Gynecology definition of FGR as estimated fetal weight below the 10th percentile for a population, and small for gestational age (SGA) as birth weight below the 10th percentile.

Patient and public involvement

In an unpublished pilot study completed prior to completion of this protocol, pregnant volunteers used HSAT as described in stage 2, and OSA-positive subjects used PAP as described in stage 3. The pilot study allowed investigators to assess the night-time tolerability of both HSAT and PAP for participants. Out of 57 patients who received HSAT, 52 had data adequate for analysis. Of these 52 patients, 10 met OSA criteria for PAP use (see below); however, one had an urgent delivery before PAP could be distributed. Of the remaining nine subjects, one declined to use PAP, one could not tolerate it and all the remaining seven met PAP adherence criteria (see below). Generally, adherence and comfort were good for both HSAT and PAP.

As part of the effort to design the study with a tolerable intervention and a meaningful outcome, two of our principal investigators met with community obstetricians, primary care physicians and with a non-profit organisation advocating for high-risk pregnant maternal health, in order to discuss the study and obtain feedback well in advance of its final iteration.

Study participants

Inclusion criteria

This study enrolls pregnant women, aged 18–50 years, who have been diagnosed with established FGR. FGR is defined as estimated fetal weight <10th percentile based on at least one routine second trimester ultrasound without a subsequent increase to >15th percentile on any ultrasounds prior to enrolment. If the ultrasound that identified FGR was performed prior to 21 completed weeks, a repeat scan after 22 weeks is required to confirm the diagnosis of FGR prior to enrolment. The lower limit of gestational age at enrolment to stage 1 is 22+0 weeks; the upper limit of gestational age at enrolment to stage 1 is an adequate gestational age to be able to complete stages 1 and 2 and, if appropriate, to receive stage 3 intervention by no later than 32+0 weeks.

Exclusion criteria

The following exclusion criteria apply at the time of enrolment: prespecified independent cause of FGR (congenital or genetic anomalies, suspected aneuploidy with two minor or one major markers, intrauterine infection or multiple gestation); active labour; a concrete decision already made to induce labour or perform caesarean delivery within 2 days; reverse end-diastolic flow in the umbilical artery (note that other abnormal Doppler flow velocities such as absent end-diastolic flow in the umbilical artery or uterine artery notching or increased pulsatility index are not exclusion criteria unless other exclusion criteria are present); pre-existing formal diagnosis of OSA; chronic pulmonary disease; haemoglobinopathies (including thalassemia major and sickle cell disease, but

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Table 1 Demographic characteristics of the medical centres in the SAFER study (data for 2019)

| Institution | Deliveries | CS rate (%) | US protocol used for estimated fetal weight | US fetal growth nomogram used for FGR | Birthweight nomogram used for SGA |
|-------------|------------|-------------|-------------------------------------------|-----------------------------------|-----------------------------------|
| University of Washington, Barnes Jewish Hospital, St Louis, Missouri | 3576 | 31.4 | Hadlock et al.56 | A US national reference for fetal growth.56 | Revised Fenton growth chart for preterm infants.57 |
| University of Rochester, Strong Hospital, Rochester, New York | 2859 | 34.9 | Hadlock et al.58 | Local nomogram from 115 000 births in the nine-county Finger Lakes Region, 2004–2013.59,60 (For Rochester birthweight nomogram see online supplemental file 2). | Local nomogram from 115 000 births in the nine-county Finger Lakes Region, 2004–2013.59,60 (For Rochester birthweight nomogram see online supplemental file 2). |
| Hadassah Hebrew University Medical Center, Jerusalem, Israel | 8050 Ein Karem campus | 17.1 | Hadlock et al.60 | International estimated fetal weight standards of the INTERGROWTH-21st Project.61 | Birthweight standards in the liveborn population in Israel.62 |
| | 5812 Mt Scopus campus | 16.8 | | | |

CS, caesarean section; FGR, fetal growth restriction; SAFER, Sleep Apnea and Fetal Growth Restriction; SGA, small for gestational age; US, ultrasound.
not sickle trait or thalassemia minor without clinical manifestations); maternal craniofacial anomalies which might impair the ability of the participant to use PAP appliances; lack of proficiency in English (Rochester or St Louis) or either Hebrew, Arabic or English (Jerusalem); inability to understand the consent; and unwillingness or inability to participate adequately in OSA screening (stage 1) or in HSAT (stage 2). Note that poor adherence to PAP (stage 3) is not an exclusion criterion. PAP usage is recorded but outcomes are assessed by intention to treat.

Stage 1: initial screening for OSA risk
Stage 1 of the trial is designed in accordance with the Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) statement (https://www.strobe-statement.org). Potential participants with FGR as defined above are identified through referral from the obstetric ultrasound service, directly from the obstetrician or by review of the electronic medical record. Those who meet inclusion and exclusion criteria are approached for consent either by telephone, in the outpatient clinics or as inpatients by a member of the research team. Participants undergo a brief screening questionnaire to assess for risk of OSA. Participants are deemed to be at risk for OSA if they report loud snoring, report witnessed gasping for air during sleep, have an Epworth score >10 or a Faccos score >75 (see figure 1).20 Participants with evidence of OSA risk are invited to proceed to stage 2.

Stage 2: HSAT and OSA screening tools
Participants identified as being at risk for OSA in stage 1 progress to stage 2. The primary assessment for confirming OSA diagnosis in this study is HSAT. Stage 2 of the trial is an observational study designed in accordance with the STROBE statement (https://www.strobe-statement.org/).

Prior to HSAT, participants are assessed for demographic data, medical comorbidities, body mass index, neck circumference and the presence of any craniofacial abnormalities. They also undergo a more detailed OSA assessment consisting of component parts of five standard OSA screening tools (STOP-BANG, Berlin Questionnaire, American Society of Anesthesiologists checklist, Flenmons Index and Epworth Sleepiness Scale). These screening tools for OSA have each been validated in the non-pregnant population; some but not all have been validated in pregnancy.20 28 38 39

HSAT is performed with a Food and Drug Administration-approved type III home sleep monitor (ResMed Apnea Link Air, ResMed, San Diego, California), which records the following: (1) air flow using nasal cannula and pressure transducer; (2) respiratory effort using elastic respiratory inductance plethysmography belts around the chest and abdomen; (3) ECG; and (4) pulse oximetry.40 A trained member of the study team shows each study participant how to apply the sensors and use the HSAT monitor and instructs her to wear the HSAT monitor for two consecutive nights. The HSAT monitor data are downloaded, and studies are reviewed and scored by a registered polysomnographic technologist per standard scoring protocols for apnoea-hypopnoea index (AHI) and oxygen desaturation index (ODI).41 AHI is defined as the number of apnoeas plus hypopnoeas on average each hour.

Hypopnoea is defined by the American Academy of Sleep Medicine recommended rule 1A; where a hypopnoea requires all of the following to be met: (1) the peak signal for respiratory excursions drops by ≥30% from pre-event baseline; (2) the duration of the ≥30% drop is ≥10 s; and (3) there is a ≥2% oxygen desaturation from pre-event baseline and/or the event is associated with an arousal.41 ODI is defined as the number of ≥3% oxygen desaturations per hour of sleep without the required drop in peak signal excursions.

HSAT generally underestimates AHI compared with PSG, as the entire study duration is used as the denominator when calculating AHI. These errors are most evident in mild OSA.42 To maximise sensitivity to detect OSA yet improve specificity in this pragmatic study, we add an ODI ≥5 criterion in cases where AHI is between 5 and 10. Accordingly, an HSAT that is positive for OSA is defined as either an AHI ≥5 with an ODI ≥5 or as an AHI ≥10 regardless of the ODI.

The second night of HSAT monitoring serves only as a backup in case the first night is inadequate to determine the AHI and ODI. Variables derived from HSAT include AHI, ODI, SaO₂ nadir and time with SaO₂ <90%. Investigators, treating physicians and participants are blinded to these results until after delivery. The only result that is unblinded prior to delivery is the binary ‘OSA positive’ versus ‘OSA negative’ HSAT result. Participants with an HSAT that is positive for OSA will proceed to stage 3. All participants from stage 2 are followed through the time of delivery and assessed for maternal and fetal outcomes after delivery (details below).

Stage 3: randomised trial of PAP therapy
Participants diagnosed with OSA in stage 2 proceed to stage 3, which is a randomised controlled trial (RCT) of PAP versus standard care control. The RCT is designed in accordance with the Consolidated Standards of Reporting Trials statement (http://www.consort-statement.org). The primary outcome is unadjusted birth weight.

Participants randomised to PAP are asked to use PAP whenever sleeping, from the time of randomisation until delivery. The PAP device is the ResMed AirSense 10 ‘AutoSet for Her’, set at the widest pressure range of 4–20 cmH₂O. The low end of the range is increased if the participant indicates discomfort from a sensation of insufficient air pressure. During set-up, participants select from full face mask, nasal mask or nasal pillow interfaces, depending on comfort and fit. Initial set-up is performed by a team member with clinical experience with PAP treatment. Participants can change their mask interfaces ad libitum. Adherence to PAP is continuously assessed remotely using a cloud-based monitoring system,
Airview (ResMed). In case of malfunction of this system, adherence is also recorded by a memory card in each PAP device. Adherence is monitored twice in the first week, then weekly until the end of the study. We have defined acceptable adherence to PAP as ≥4 hours of treatment on ≥70% of nights from the first night after PAP set-up to the night prior to delivery. This definition is based on the Centers for Medicare and Medicaid Services guideline. If a participant does not meet this benchmark based on the nights studied, she receives a call from a team member and is offered a troubleshooting visit to reassess mask fit, machine settings and other changes in equipment in order to try to improve adherence.

Recruitment, randomisation and blinding
Participant recruitment occurs simultaneously at all sites (Barnes-Jewish Hospital, Washington University in St Louis; Strong Memorial Hospital, University of Rochester; and Hadassah Hebrew University Medical Center, Jerusalem, Israel). Randomised allocation in stage 3 occurs only after participant enrolment to avoid selection bias. Randomised allocation is performed centrally by random number generator using block randomisation with 1:1 allocation to PAP versus non-PAP control, using randomised blocks of either two or four participants. Randomisation is stratified by study site; within each site, blocks are trichotomised by AHI into low (AHI 5–15), medium (AHI 16–30) or high (AHI >30) as defined above. Only the coordinating-centre study coordinator is unblinded to absolute AHI or AHI category. All sites produce a weekly report on the number of eligible participants, the number of participants approached and the number recruited. To aid data homogeneity, a central data entry system is used (Research Electronic Data Capture: REDCap). At each centre, the investigators have a secure password allowing access to the protected database. After randomised allocation, the participant is notified by her allocation by the site investigator; subjects in the PAP group are then instructed on PAP use and a mask fitting session is scheduled. Participants are instructed not to reveal their group allocation to the technicians performing follow-up ultrasounds or to the obstetricians making clinical management decisions.

Outcome measures and analysis

Primary outcome
The goal for stage 1 is to identify OSA risk as defined above for the purpose of recruiting participants into stage 2. The goal for stage 2 is to identify OSA based on HSAT criteria as defined above for the purpose of recruiting participants into stage 3. The goal for stage 3, and the primary outcome of the SAFER study, is the impact of the randomised treatment intervention (PAP vs standard care control) on unadjusted birth weight. Birth weight is an endpoint driven by multiple factors including intrauterine fetal growth velocity and gestational age at delivery. These factors may not be independent of each other, as frequently women with severe FGR have interventional delivery performed remote from term, either for fetal indications or for maternal indications (typically when FGR is accompanied by pre-eclampsia). The primary endpoint, unadjusted birth weight, will be corrected for gestational age and by appropriate racial and local nomograms in multivariate analysis.

Secondary outcomes
The secondary outcome of stage 1 is the percentage of patients with FGR who screen positive for OSA. The secondary outcome for stage 2 is the percentage of patients who test positive for OSA in HSAT. These outcomes will help determine the clinical feasibility and cost-effectiveness of this strategy in the general population. An additional secondary outcome of stage 2 is to assess the predictive value of each of the OSA screening tools for identifying OSA in pregnancy. Secondary outcomes of stage 3 are as follows: (1) obstetric outcomes: ultrasound fetal growth velocity, enrolment-to-delivery interval, gestational age at delivery, birth weight (as a Z-score) corrected for gestational age, using an algorithm to standardise gestational age, stillbirth; (2) neonatal outcomes: Apgar score, rate of admission to higher levels of care (NICU or special care nursery), length of neonatal stay.

Exploratory outcomes of stage 3 are as follows: (1) obstetric outcomes: mode of delivery, umbilical cord gases, umbilical artery blood flows; (2) maternal outcomes: postpartum depression Edinburgh Postnatal Depression Scale (EPDS) score, rate of hypertensive disorders of pregnancy, rate of gestational diabetes mellitus; (3) neonatal outcomes: postnatal hypoglycaemia (<40 mg/dL at any time) and/or requirement for intravenous glucose treatment based on American Academy of Pediatrics guidelines, rates of hypoxic ischaemic neonatal encephalopathy, neonatal death in first year of life.

Primary and secondary outcomes are compared between the two study groups in stage 3 (OSA-positive patients randomised to standard care with no PAP vs those randomised to PAP therapy). We use intention-to-treat analysis.

Planned subgroup analyses
We plan the following subgroup analyses:

PAP use
The effect of PAP on primary and secondary outcomes based not on intention to treat, but rather on PAP adherence (as described above). The clinical justification of this subgroup allocation is to assess whether a negative or a borderline primary outcome of SAFER may mask a true clinical effect of PAP when used with good adherence. This subgroup analysis will also accommodate the unlikely crossover of a patient allocated to the no-PAP control in stage 3, but who is subsequently referred by their primary physician or obstetrician for OSA work-up (not current standard of care) and who goes on to receive PAP.
Effect of untreated OSA

The effect of PAP therapy on primary and secondary outcomes with stratification by OSA severity (mild, AHI 5–15; moderate, AHI 15–30; severe >30). We hypothesise that maternal and fetal outcomes will be most improved following PAP therapy in patients with the most severe cases of OSA. The clinical justification of this subgroup allocation is to assess whether a negative or borderline primary outcome of SAFER may mask a true clinical effect of PAP when used in patients with more severe OSA; this may also aid in the determination of an optimal target population based on OSA severity for clinical use of PAP in FGR.

Planned exploratory subgroup analyses

PAP use

The effect of PAP usage as a continuous variable (mean hours of PAP use per night) on primary and secondary outcomes; the purpose of this exploratory subgroup analysis is to explore the possible dose-response effect of PAP.

OSA severity

The effect of PAP on primary and secondary outcomes with stratification by secondary measures from the HSAT: ODI, SpO₂ nadir and time with SpO₂ <90%; where the stratification into mild, moderate and severe will be determined after examining the distribution of data; the purpose of this exploratory subgroup analysis is to explore whether other measures of OSA severity, particularly oxygen desaturation, are useful to direct which patients may benefit from PAP therapy.

Effect of untreated OSA

Comparison between OSA-negative patients from stage 2 versus untreated OSA-positive patients (non-PAP control) from stage 3 (we anticipate this will be approximately a 3:1 ratio); the purpose of this exploratory subgroup analysis is to assess the deleterious effects of OSA on maternal and fetal outcomes in these patients.

Statistical analysis

Data are assessed for normality by visual inspection of the frequency plot (histogram) and Q-Q plot of each outcome variable and by the Kolmogorov-Smirnov test. Normally distributed data are presented as mean (SD); non-normally distributed data are presented as median (IQR). Our primary outcome (birth weight) (derived from stage 3) is assessed by two-tailed parametric t-test (normal distribution) or non-parametric Wilcoxon rank-sum test (non-normal distribution) as appropriate, using intention to treat as the grouping variable. The same tests are used for other continuous secondary outcome variables (gestational age at delivery; enrolment-to-delivery interval; birth weight corrected for gestational age; Apgar and umbilical cord gas). For categorical outcomes (higher level nursery admission; stillbirth; compliance with PAP), the \( \chi^2 \) test or Fisher’s exact test (if expected cells are small) is used as appropriate. Additionally, we use a mixed effects regression model to assess the effect of PAP on the longitudinal association between estimated fetal weight and Doppler ultrasound umbilical artery flow, while controlling for maternal, obstetric and fetal variables that may affect the primary outcome. Only the intercept will be specified as the random component to account for any omitted variable. By standard convention, statistical significance is based on a two-sided p value <0.05 to determine the significance of association. Statistical testing is currently planned to use SAS V.9.4 (SAS Institute).

Sample size calculation

The baseline birth weight of infants with FGR in the population from our primary study centre (Barnes-Jewish Hospital, St Louis, Missouri) is 2535±234 g (unpublished data). Based on these data, we calculated that 104 evaluable participants with OSA will need to be randomised to either PAP or control (stage 3) in order to have 90% power to detect a 150 g difference in birth weight in the PAP group compared with control. This is based on an alpha of 0.05, anticipated 5% loss to follow-up and a two-tailed t-test.

The risk of OSA in high-risk obstetric populations (such as FGR) is as high as 35%. As we are screening participants for OSA risk by a brief telephone or in-person questionnaire (stage 1) prior to HSAT testing (stage 2), we estimate that there will be an OSA-positive HSAT in 30%–50% of these screened participants. Consequently, we estimate that we will need to assess HSAT in 200–350 participants in stage 2 in order to achieve our sample size of 104 evaluable participants for stage 3. We estimate needing to screen some 500–1000 participants in stage 1 in order to identify these 200–350 participants for stage 3. Based on an estimated combined total of 20000 births annually with an estimated incidence of 15% of FGR at the above centres, we estimate 3000 pregnancies complicated by FGR each year. Accounting for participants who will not meet study inclusion criteria, or will decline to participate, the recruitment process is expected to last approximately 2 years.

Analysis of pragmatic elements of the SAFER study

The SAFER study is a pragmatic trial. The pragmatic elements of the study were quantified using the Pragmatic Explanatory Continuum Indicator Summary-2 tool (https://www.precis2.org). Data were obtained from the principal investigators in all study centres. According to six of the nine criteria, the SAFER study is largely a pragmatic study (figure 2). The study is particularly pragmatic for experimental intervention (a standard commercially available product), follow-up, relevance of clinical outcomes and analysis. It is more explanatory for patient selection/recruitment and organisational intervention. This is because OSA work-up is not currently a standard of care for FGR and patients only reach stage 3 if they have FGR with HSAT-verified OSA. In this way, the study targets patients most likely to experience benefits in fetal
growth from PAP, but the generalisability to other populations may be more limited.

**Strengths and limitations**

The SAFER study has important strengths. It is a multicentre study of a safe, routinely available intervention that examines an easily measured outcome with important short-term and long-term ramifications for the well-being of neonates and possibly for later adult life. SAFER is a pragmatic study and the intervention does not preclude any existing approaches for the management of these high-risk pregnancies. As PAP is worn only when sleeping, typically at home, it is relatively easy to keep the research team and clinical providers blinded to group allocation. As the PAP device used in this study records PAP compliance, we do not have to rely on compliance self-reporting which may overestimate nightly device use.

The SAFER study has several limitations. There is currently no definitive estimate for the impact of PAP on birth weight; hence, the effect measure used in the sample size estimation may be inaccurate, which may affect the study’s power in regard to our primary and secondary endpoints. It is likely that follow-up studies will be warranted, especially regarding whether or not the results from this specific target population can be generalised to other pregnant women with FGR, or pregnant women with OSA who may have pregnancy complications without FGR, for example, pre-eclampsia. It is the intention of the investigators to follow-up a positive result (that PAP increases intrauterine fetal growth) with a larger study to assess long-term neurological outcome.

Estimates of fetal weight were all based on the Hadlock formula using the standard biometric parameters (biparietal diameter, head circumference, abdominal circumference, femur length) [52, 53]. However, when converting fetal weight to a calculated percentile, this must reference a ‘population or customized standard’. As this is a pragmatic multicentre study, each study centre defines FGR and SGA by the population or customised standard used in their routine care (see table 1). This will lead to a slight difference between centres in FGR and SGA definition. However, fetal growth varies between different ethnic and geographical populations, so a single absolute imposed standard is of limited relevance clinically. The impact of this factor in stage 3 is reduced by stratification of randomised allocation by study centre.

Although investigators, treating physicians and ultrasound technicians are blinded to participant group allocation in this study, we do not blind participants. We are confident that our primary outcome and all of the secondary outcome measures (with the exception of the EPDS) can be assessed blindly, which minimises the risk of bias. We considered using sham PAP as a placebo control. While this could minimise bias in evaluating the effect of PAP, wearing an ineffective mask does increase the risk of discomfort and sleep disturbances for the control group, which potentially biases the results toward the active group. A recent study demonstrated that there was no difference in outcome between sham PAP and a device-free control; however, most subjects in the sham PAP group guessed correctly that they were receiving the placebo intervention.

Home sleep testing may underestimate the true presence of SDB for several reasons, most notably because it
uses the study duration as the denominator to calculate AHI. In addition, in this study, our protocol may be over-reliant on oxygen desaturation, as that feature may not be as prominent in a young female pregnant population.56

Hence, our methods may underestimate the true prevalence and severity of SDB, and preferentially select the study population with more severe SDB. This selected population with more severe SDB may have a greater likelihood of exhibiting a clinical effect with PAP treatment. However, in this pragmatic study, we need to use HSAT so that the intervention can be started within as narrow a time frame as possible.

This study uses intention-to-treat analysis to assess the efficacy of the intervention. The adherence to PAP is relatively low across all populations.51 54 Adherence is generally higher for aPAP devices, as used in this study, when compared with CPAP devices.51 Two recent large national database studies reported conflicting results regarding gender effects on PAP adherence—either lower in men51 or lower in women,54 although it is not known whether any of these women were pregnant in those studies. It is possible that pregnant women concerned about potentially optimising fetal and maternal outcomes would be more adherent to PAP. We performed a pilot study in preparation for this protocol (see the Patient and public involvement section) in which we observed PAP adherence in seven out of nine subjects tested. Our calculated power does not specifically accommodate non-adherence. Accordingly, we also describe a planned subgroup analysis in which PAP adherence is used rather than intention to treat. Ultimately, this pragmatic study can only assess whether administering a PAP device will improve outcomes in pregnancy. If PAP is not effective during pregnancy because women do not adhere to it, that raises a separate question that will require addressing separately.

Potential benefits, risks and alternatives

Benefits
All participants reaching stage 2 of the study will potentially benefit from being identified as at risk for OSA and being referred to their primary care physician for evaluation by a sleep physician after delivery.

Risks
The likelihood of adverse events in this study is low as the devices that will be used in the study (HSAT and aPAP) are ones that have been used in the general and obstetric population for years. In the unlikely event that serious side effects occur, these will be documented and reported to the human research protection office and to the study’s Data Safety Monitor.

Minimisation of risks to confidentiality
All participants are assigned a unique study ID number. The link to identifiers will be destroyed at the end of the study. Data will be stored under lock and key (office, file cabinet) and only the investigators and research team will have access. If data are published, there will be no link to identifiers. Study data are not entered into participants’ medical records. Data regarding PAP compliance are downloaded from ResMed’s secure, cloud-based patient management system to a secure server at the primary site (Washington University), where they are stored for the duration of the study.

Data from this study will be recorded using REDCap, a web-based, Health Insurance Portability and Accountability Act-compliant application. Access to the data is password protected. REDCap servers are housed in a secure data centre and information transmission is encrypted.

Adverse event reporting and safety monitoring
The research team continuously monitors the study for adverse events. All serious adverse events (SAEs) are reported to the Institutional Review Board (IRB) according to IRB stipulations. Additionally, an attending anaesthesiologist at Washington University who is not involved in the study serves as the Data Safety Monitor; given the small size and relatively low-risk nature of the protocol, an individual physician rather than a full Data Safety Monitoring Board is used. This monitor reviews all adverse events annually and reviews SAEs or unexpected adverse events as they occur.

Premature study termination
The only interim analysis to be performed in this study will be a single, blinded analysis performed by the Data Safety Monitor at the mid-way point, after 52 patients have completed stage 3. This interim analysis will assess differences between groups in birth weight, gestational age at delivery, intrauterine fetal death or reported major adverse events. The study code will not be broken, and other secondary endpoints will not be assessed. If there is a statistically significant difference between groups in one of these selected outcomes, then the Data Safety Monitor will break the code. If the intervention is associated with smaller birth weight, lower gestational age at delivery, more intrauterine fetal death or more reported major adverse events, then this will be brought to the attention of the investigators and the primary study site IRB, for consideration of possible premature termination of the study on the grounds of increased harm due to the intervention. There will be no premature termination of the study due to increased benefit of the intervention. Unless there is an increased harm of the intervention, the Data Safety Monitor will not give any information to the investigators regarding the interim analysis.

Ethics and dissemination
Informed consent for stage 1 may be by telephone, electronic or written consent. Informed consent for stages 2 and 3 is obtained by a study investigator using electronic or written consent (see online supplemental file 1). There are no additional risks associated with the screening study (stage 1) or the observational HSAT study (stage 2). There
is a potential ethical concern regarding randomisation of HSAT-positive participants (identified from stage 2 as having OSA) to either a PAP (intervention) or a no-PAP (control) group. However, this concern is minimised by the following arguments: (A) HSAT, or a formal sleep study, is not a standard care for investigating FGR, and the diagnosis of OSA is rarely made in pregnancy; hence, the diagnosis is only made as a result of this study,23 24, (B) PAP is not a standard care for sleep disturbance in pregnancy as many practitioners and participants rightly or wrongly assume that this will improve after delivery,24 while PAP is seen as a long-term therapeutic intervention. After delivery, participants with AH1 ≥5 will be given a standard IRB-approved letter along with their HSAT results. This information can be presented to their primary care physician for possible referral to a sleep specialist.

The trial steering committee is responsible for all major decisions regarding changes to the protocol. The committee communicates these changes to the IRB, the trial registry and appropriate parties. Data will be shared in keeping with the data sharing statement below. Dissemination plans include presentations at scientific conferences. The results of the SAFER trial will be published in a peer-reviewed journal. Dissemination of results to study participants and their family members will be available on request. Updates and results of the study will be available to the public at ClinicalTrials.gov.

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