Combination of obstructive sleep apnoea and insomnia treated by continuous positive airway pressure with the SensAwake pressure relief technology to assist sleep: a randomised cross-over trial protocol

Jean Louis Pepin,1 Frédéric Gagnadoux,2 Alison Foote,1 Rachel Vicars,3 Bhavi Ogra,3 Véronique Viot-Blanc,4 Meriem Benmerad,1 Marie-Pia D’Ortho,5 Renaud Tamisier1

ABSTRACT

Introduction Obstructive sleep apnoea (OSA) is a common sleep breathing disorder affecting up to 17% of the middle-aged population. Continuous positive airway pressure (CPAP) is the primary treatment for patients with OSA, but acceptance and adherence to therapy is suboptimal in specific subgroups particularly those with insomnia or poor sleep quality (40%–80% of patients with OSA). Pressure intolerance, particularly during periods of wakefulness, inhibiting sleep onset or return to sleep, is one reason for poor CPAP adherence. AutoCPAPs continually monitor airflow changes and only increase the pressure when the upper airway requires it. Reducing the pressure during wakefulness-sleep transition and wakefulness-after-sleep-onset (WASO) may improve therapy comfort and potentially adherence without compromising therapy efficacy. We hypothesise that SensAwake, a pressure relief function that reduces CPAP pressure on the transition from sleep to wakefulness and on WASO, may improve objective sleep quality.

Methods and analysis This is a multicentre, randomised double-blind crossover clinical trial on patients with both OSA and insomnia. Insomnia is defined as Insomnia Severity Index >15 at screening. Baseline data, including actigraphy, are collected for 1 week before randomisation (1:1) to either conventional AutoCPAP or AutoCPAP with SensAwake for 4 weeks. After an evaluation visit, patients are switched to the other treatment arm for a further 4 weeks. Allowing for 20% dropout, 48 patients are required. If applicable, repeated measures analysis of variance will be used to assess differences in WASO measured by actigraphy (primary outcome), other actigraphy measures, AutoCPAP compliance, subjective questionnaire scores (Epworth Sleepiness Scale, Pittsburgh Sleep Quality Index, Short-Form 12 Health Survey) and 24 hours blood pressure (secondary outcomes).

Ethics and dissemination The protocol was approved by the regional Ethics Committee (CPP Sud-Est–V, IRB N°6705) on 9 December 2015, is registered on ClinicalTrials.gov (NCT02721329) and started in June 2016 with expected publication of primary outcome results in 2018.

Trial registration number NCT0272132; Pre-results.

Strengthen and limitations of this study

► This multicentre randomised crossover trial could potentially lead to an alternative to conventional Autocontinuous positive airway pressure (CPAP) therapy in a well-defined common obstructive sleep apnoea (OSA) having a high rate of CPAP non-compliance.
► While the sample size is just sufficient to analyse objective sleep indices (primary outcome) in a crossover design, it is too small to inform on 24 hours blood pressure and patient-centred outcomes (secondary outcomes).
► The wrist actimeter used has not been previously validated for the measurement of wake-after-sleep-onset in this specific patient phenotype associating OSA and insomnia.
► Only one night of blood pressure monitoring is done in each arm of the study. Longer Ambulatory Blood Pressure Monitoring (ABPM) would be desirable, but wearing yet another device may bias the study.
► Despite having several study sites the sample size is rather small, one limitation being the difficulties in recruitment of willing patients naïve to CPAP therapy; restricting the relevance of any secondary analyses such as those on blood pressure.
airway pressure (CPAP) is the primary treatment for patients with OSA. Despite the effectiveness of CPAP in abolishing upper airway obstruction, acceptance of and adherence to therapy is often suboptimal particularly in specific phenotypes, including the combination of OSA and insomnia or insomnia symptoms.

Pressure intolerance is one possible reason for this lack of adherence. Conventional CPAP generally delivers higher pressure than necessary for much of the night as the needed CPAP pressure is selected based on one night’s titration (in a sleep laboratory) or during several nights at home and pressure requirements can vary considerably with sleeping posture, sleep stage and environmental influences such as alcohol and sedative use. AutoCPAPs address this problem by continually monitoring airflow changes and only increasing the pressure when the upper airway requires it. Research suggests that AutoCPAP and only increasing the pressure when the upper airway requires it. Research suggests that AutoCPAP generally delivers an overall lower mean treatment pressure than conventional CPAP. Despite this, there is limited evidence to suggest that AutoCPAP therapy can considerably improve CPAP adherence and acceptance in an unselected population, but this might be different in patients with OSA with concurrent insomnia.

Conceptually, a patient’s awareness of pressure occurs only during wakefulness. Thus, reducing the pressure during wakefulness may improve therapy comfort and potentially adherence without compromising therapy efficacy. SensAwake (Fisher & Paykel Healthcare, Auckland, New Zealand) is a pressure relief technology that accurately detects irregularity in the flow signal indicative of the transition from sleep to wake. When the transition from sleep to wake is detected, the device promptly reduces the pressure to help facilitate a return to sleep. CPAP/AutoCPAP with SensAwake has been used in the general OSA population and has been shown to provide the same treatment efficacy at a lower overall pressure as CPAP/AutoCPAP without SensAwake and patients have judged it to be more comfortable and preferred it to CPAP without SensAwake.

The prevalence of insomnia symptoms in patients with OSA is estimated to be 40%–80%, and existence of insomnia has been shown to negatively affect CPAP compliance in some studies. It is proposed that insomnia patients are preoccupied with external factors that may be perceived as a threat to sleep, which results in a higher wake-after-sleep-onset (WASO), the amount of time a person spends awake from when they first fall asleep to when they do not attempt to go back to sleep. Besides waking it takes account of difficulty in getting back to sleep, which may be further exacerbated by the presence of CPAP. It is therefore hypothesised that the pressure relief that SensAwake provides during wakefulness may be of a greater benefit to patients with OSA and insomnia if it can facilitate the return to sleep. There is no known published data on the use of SensAwake in the OSA/insomnia population.

Primary research objective
The primary objective is to compare the at-home objective sleep quality (WASO) when using AutoCPAP with SensAwake versus AutoCPAP without SensAwake in patients with a diagnosis of OSA and insomnia.

Secondary research objectives
The secondary objectives are to compare AutoCPAP compliance, other measures of objective sleep quality (total sleep time (TST), sleep onset latency (SOL) and sleep efficiency (SE)), daytime sleepiness, subjective sleep quality, insomnia, quality of life and 24 hours blood pressure.

METHODS AND ANALYSIS
Study design
This is multicentre prospective 1:1 randomised, double-blind, crossover trial.

Patient entry and screening for insomnia
Outpatients diagnosed with OSA by polysomnography at one of the participating tertiary hospital sleep centres (Grenoble, Angers and Bichat and Lariboisière hospitals in Paris, France) between November 2016 and October 2017, eligible for CPAP treatment under local requirements (Apnoea Hypoxia Index >30 with no more than 20% central respiratory events) are asked to answer an Insomnia Severity Index (ISI) questionnaire to screen for insomnia. If they meet the study inclusion/non-inclusion criteria, they are asked by the sleep physician for their written informed consent (available as a Supplementary file 1) and are enrolled into the study.

Materials
Nighttime actigraphy, the gold standard for measuring objective sleep quality in the home, is recorded using a wGT3X-BT wrist monitor from ActiGraph, Pensacola, Florida, USA. This is a standard actigraphy device that uses an adapted version of the Cole-Kripke Algorithm. The AutoCPAP device is the ICON+Auto from Fisher & Paykel Healthcare. This has an integrated heated humidification system and is intended for use on adult patients for the treatment of OSA at home or in a sleep laboratory. The ICON+ treats OSA by delivering a continuous flow of air at a pressure prescribed by the physician to maintain the airway open. In AutoCPAP mode, the device auto-adjusts the therapeutic pressure between a set minimum and maximum in response to respiratory events (apnoea, hypopnoea and flow limitation). SensAwake responsive pressure relief technology is a comfort feature that is available in the ICON+Auto device. It functions by detecting wakefulness using the flow signal and promptly reduces the pressure to a more comfortable level to allow the patient to return to sleep. The ICON+ records and reports industry standard metrics such as adherence, leaks and treatment efficacy data.
The choice of CPAP mask is left to the patient, physician, and/or home care provider and is the same as for usual care with CPAP.

Twenty-four hours ambulatory blood pressure monitors are those normally used by the centres and are fitted and data collected by qualified clinical research assistants (CRA) blinded to the study arms during weeks 1, 5 and 9 of the study.

Answers to the validated French versions of the self-reported questionnaires: ISI, Epworth Sleepiness Scale (ESS), Pittsburgh Sleep Quality Index (PSQI) and Short-Form 12 Health Survey (SF-12) will be analysed as recommended by the authors of the questionnaires.

Randomisation
Participants are block randomised via a secure electronic website to receive AutoCPAP either with or without SensAwake.

- SensAwake ‘off’ arm: SensAwake function off; OR
- SensAwake ‘on’ arm: SensAwake function on. When wakefulness is detected, SensAwake will automatically drop the pressure to the set SensAwake pressure. In the ICON+Auto Device, the minimum pressure is also the SensAwake pressure. The SensAwake pressure is the pressure that the device will drop to during wakefulness. So if the patient is experiencing a pressure of 12 cm H₂O, and their SensAwake pressure is 4 cm H₂O, then it will drop from 12 cm H₂O to 4 cm H₂O. The default SensAwake pressure is set to 4 cm H₂O, however, for patients with higher therapeutic pressures, 4 cm may be too low and result in discomfort. Therefore, the SensAwake pressure can be increased to 6 cm H₂O to account for this.

The allocation list was computer generated by a statistician independent from the study investigators.

Treatment
Participants receive training in use of the AutoCPAP device as per usual care. Usual care is standardised across the participating centres. Participants receive in-home therapy for 4 weeks. During the last week of arm one, blood pressure is monitored over one 24-hour period, whereas the actimeter is worn during the 4 weeks of each treatment arm.

Crossover
Participants return to the centre for the crossover visit. Full AutoCPAP data (recorded by the device) and actigraphy data are downloaded. They again complete the questionnaires: ISI, ESS, PSQI and SF-12, and are issued with a new sleep diary. The patient’s AutoCPAP device settings are switched over to the opposite treatment arm by the site coordinator without showing them to or discussing them with the participant or disclosing the settings to the physician. Participants receive at-home therapy for a further 4 weeks. During the last week of arm two, blood pressure is monitored over one 24-hour period.

End of study
Participants attend the study centre where full AutoCPAP data (entire folder on the device’s USB) and actigraphy data are downloaded; they hand in their sleep diaries and again complete the ISI, ESS, PSQI and SF-12 questionnaires. If patients prefer the AutoCPAP with SensAwake, they will be able to continue to use this feature after the conclusion of the study. Participants may obtain a summary of trial results after these have been submitted for publication.

Withdrawal and stopping criteria
Patients have the right to withdraw from the study at any time. In addition, the investigators may withdraw a patient at any time for the following reasons: protocol violation,
serious illness or adverse event. In the event of a serious adverse event, unblinding may be done through the site coordinator.

**Statistics**

**Sample size**
The sample size was calculated based on an assumption of a WASO of 58 min±SD\(^4\) and allowing for the cross-over nature of the study. It assumes that the SD of the difference between the two treatments is approximated by the SD derived from WASO single time assessments. The largest estimate of the WASO between individual SD was used: 46 min in the study by Natale et al.\(^4\) On this basis, a sample size of 40 completers of both treatments is required (2 sequence groups of 20/group each) to detect a difference of 15 min or more as statistically significant (two-tailed alpha=0.05) with 80% power. In a crossover study, it is advised to over-recruit to allow for dropouts, so the minimum sample size was set at 48 (24 per group) to allow for 20% dropout, with 12 patients per centre.

**Statistical analysis**
All consenting and enrolled patients will be included in the intention-to-treat analysis. Withdrawal and non-adherence to treatment are outcome measures, thus data on any withdrawn or non-adherent patients will be included. A complete description of the study population will be presented with continuous variables expressed as median and IQR, and categorical variables as frequencies and percentages (see table 1 for all outcome measures by study visit or period).

Each 4-week treatment arm includes a first-week ‘washout’ period, where data will not be analysed. Repeated measures analysis of variance (ANOVA) will be used to assess differences between the two treatments for actigraphy measures (WASO, TST, SOL, SE), treatment compliance, subjective questionnaire results (ISI, ESS, PSQI and SF-12) and 24 hours blood pressure measurements (minima, maxima, mean values of the systolic, diastolic and mean arterial pressures and dipping profile). If requirements for parametric repeated measures ANOVA are not met then a non-parametric Wilcoxon signed-rank test will be used. The analysis will include time-related factors (WASO, TST, SOL) and treatment as within-subject factors and treatment sequence as a between-subjects factor. The interaction between treatment and treatment sequence will be tested to ensure there are no carry-over effects influencing the comparison of the treatments. In the case of missing data for the primary outcome, it will be derived from the available WASO data at both post-treatment times (V2, V3). For both post-treatment times, missing data will be imputed from baseline outcome measures and any postbaseline assessments. Statistical data analysis will be performed using IBM SPSS Statistics V.22 and tested with a significance level of 0.05, by an independent statistician.

### Table 1 Outcome measurements at study visits

| Variable | Measurement | Measurement points* |
|----------|-------------|---------------------|
| **Primary end point** | | |
| Objective sleep quality | Actigraphy | Days 1–7 (baseline) (downloaded at V1) During week 4 of each treatment arm (days 29–35 and days 56–63; downloaded at V2 and V3, respectively) |
| ► Wake after sleep onset (WASO) | | |
| **Secondary end points** | | |
| Objective sleep quality | Actigraphy | Days 1–7 (baseline) (downloaded at V1) During week 4 of each treatment arm (days 29–35 and days 56–63; downloaded at V2 and V3, respectively) |
| ► Total sleep time | | |
| ► Sleep-onset latency | | |
| ► Sleep efficiency | | |
| Treatment compliance | CPAP (data recorded by AutoCPAP) downloads | Weeks 2–4 of each treatment arm (days 15–35 and days 28–35; downloaded at V2 and V3, respectively) |
| Subjective sleep quality, insomnia, daytime sleepiness and quality of life: | Self-reporting questionnaires: PSQI, ISI, ESS, SF-12 | Baseline (V1) on day 7 Crossover visit (V2) on day 35 End of study (V3) on day 63 |
| ► Sleep quality | | |
| ► Insomnia severity | | |
| ► Daytime sleepiness | | |
| ► Quality of life | | |
| 24 hours blood pressure | 24 hours ambulatory blood pressure monitor | 24 hours during days 1–7 (baseline) 24 hours during week 4 of treatment arm 1 (between day 29 and day 35) 24 hours during week 4 Treatment arm 2 (between day 56 and day 63) |

*All±2 days.
PSQI, Pittsburgh Sleep Quality Index; ISI, Insomnia Severity Index; ESS, Epworth Sleepiness Scale; SF-12, Short Form 12 Health Survey; V, study visit.
treatment, but insomnia can reduce tolerance and adherence of patients to CPAP therapy. The addition of SensAwake to AutoCPAP may improve the comfort of AutoCPAP therapy, and therefore may increase sleep quality and duration, improve a patient’s adherence to CPAP therapy and improve quality of life. One strength of our study is to combine objective assessments of sleep indices and patient-centred outcomes. As the addition of insomnia to OSA increases the prevalence of hypertension, the valuation of blood pressure by means of 24 hours ambulatory blood pressure monitoring adds another strength to the study. We will also investigate whether conjointly improving sleep apnoea and sleep duration/quality will allow a better control of nocturnal blood pressure.

The results of this study will help both OSA and insomnia specialists in their decision as whether to prescribe AutoCPAP with or without SensAwake for CPAP treatment of patients with OSA who also have insomnia. If positive, this study will be a step forward for personalised therapy in the frequent subgroup of OSA plus insomnia.

**Author affiliations**

1. Hypoxia-pathophysiology Laboratory - INSERM U1042, Grenoble Alps University Hospital, Université Grenoble Alpes, Grenoble, France
2. Department of Pneumology, INSERM URM 1063, Angers University Hospital, Bretagne Loire University I, Angers, France
3. Fisher & Paykel Healthcare Limited, Auckland, New Zealand
4. Sleep Clinic - Lariboisière Hospital, Assistance Publique—Hôpitaux de Paris, Paris, Ile-de-France, France
5. Department of Physiology and Functional Exploration, Bichat Hospital, Denis Diderot University, Paris, France

**Acknowledgements** The authors would like to thank Drs Marie Destors, Pascaline Priou, Wojtek Trzepizur, Helene Benzaquen and Ruben Wanono for their advice and participation, Marie Peeters for trial management and Rebecca Thomson for help in revising the manuscript.

**Contributors** JLP, FG, RV, BO, MPO, VVB and RT designed the study and wrote the study protocol. AF and JLP wrote the article based on the study protocol. FG, MPO, RV and BO critically revised the manuscript. MB calculated the sample size and wrote the study statistical analysis plan. JLP, FG, MPO, VVB and RT are currently including patients in the study. All authors approved the submitted manuscript.

**Funding** This study is funded by unrestricted grant from Fisher & Paykel Healthcare Limited (Auckland, New Zealand).

**Competing interests** RV and BO are employed by Fisher & Paykel Healthcare Limited.

**Patient consent** Study protocol: the inclusion of patients is ongoing. Written informed consent is an inclusion criterion.

**Ethics approval** French CPP Sud-Est V, IRB N°6705.

**Provenance and peer review** Not commissioned; externally peer reviewed.

**Open Access** This is an Open Access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/

© Article author(s) (or their employer(s) unless otherwise stated in the text of the article) 2017. All rights reserved. No commercial use is permitted unless otherwise expressly granted.
REFERENCES

1. Lévy P, Kohler M, McNicholas WT, et al. Obstructive sleep apnoea syndrome. *Nat Rev Dis Primers* 2015;1:15015.

2. Peppard PE, Young T, Barnet JH, et al. Increased prevalence of sleep-disordered breathing in adults. *Am J Epidemiol* 2013;177:1006–14.

3. Sullivan CE, Issa FG, Berthon-Jones M, et al. Reversal of obstructive sleep apnoea by continuous positive airway pressure applied through the nares. *Lancet* 1981;1:862–5.

4. Gay P, Weaver T, Loubé D, et al. Evaluation of positive airway pressure treatment for sleep related breathing disorders in adults. *Sleep* 2006;29:381–401.

5. Weaver TE, Kribs NB, Pack AI, et al. Night-to-night variability in CPAP use over the first three months of treatment. *Sleep* 1997;20:278–83.

6. Kribs NB, Pack AI, Kline LR, et al. Objective measurement of patterns of nasal CPAP use by patients with obstructive sleep apnea. *Am Rev Respir Dis* 1993;147:887–95.

7. Luyster FS, Buysse DJ, Strollo PJ. Comorbid insomnia and obstructive sleep apnea: challenges for clinical practice and research. *J Clin Sleep Med* 2010;6:196–204.

8. Berry RB, Parish JM, Harts KE. The use of auto-titrating continuous positive airway pressure for treatment of adult obstructive sleep apnea. *Sleep* 2002;25:148–73.

9. Oksenberg A, Silverberg DS, Arons E, et al. Prospective randomized comparison of impedance-controlled auto-continuous positive airway pressure with constant CPAP. *Sleep Med* 2004;9:563–4.

10. Olsenberg A, Arons E, Silverberg DS, et al. The effect of nasal CPAP for obstructive sleep apnea/hypopnea syndrome on hypoxemia, respiratory embarrassment, sleep movements, body mass index, and age. *Chest* 1999;116:1000–6.

11. Ayas NT, Patel SR, Malhotra A, et al. Auto-titrating versus standard continuous positive airway pressure for the treatment of obstructive sleep apnea results of a meta-analysis. *Sleep* 2004;27:249–53.

12. Bubbeheini K, Yen FC, Lucas EA, et al. A sleep laboratory evaluation of an automatic positive airway pressure system for treatment of obstructive sleep apnea. *Sleep* 1998;21:485–91.

13. Boudewyns A, Griller-Lanoir V, Villetinen MJ, et al. Two months follow up of auto-CPAP treatment in patients with obstructive sleep apnoea. *Thorax* 1999;54:147–9.

14. d’Ortho MP. Auto-titrating continuous positive airway pressure for treating adult patients with sleep apnea syndrome. *Curr Opin Pulm Med* 2004;10:495–9.

15. d’Ortho MP, Griller-Lanoir V, Levy P, et al. Constant vs. automatic continuous positive airway pressure therapy: home evaluation. *Chest* 2000;118:1010–7.

16. Randerath WJ, Galetke W, David M, et al. Auto-titrating versus standard-controlled auto-continuous positive airway pressure (APAP/LOT) with constant CPAP. *Sleep Med* 2001;115:24.

17. Randerath WJ, Galetke W, David M, et al. Prospective randomized comparison of impedance-controlled auto-continuous positive airway pressure (APAP/LOT) with constant CPAP. *Sleep Med* 2001;2:115:24.

18. Teschner H, Wessendorf TE, Farhat AA, et al. Two months auto-adjusting versus conventional nCPAP for obstructive sleep apnoea syndrome. *Eur Respir J* 2000;15:990–5.

19. To KW, Chan WC, Cho KL, et al. A randomized cross-over study of auto-continuous positive airway pressure versus fixed-continuous positive airway pressure in patients with obstructive sleep apnoea. *Respirology* 2008;13:79–86.

20. Ayappa I, Norman RG, Whiting D, et al. Irregular respiration as a marker of wakefulness during titration of CPAP. *Sleep* 2013;36:99–104.

21. Dungan GC, Marshall NS, Hoyos CM, et al. A randomized crossover trial of the effect of a novel method of pressure control (SensAwake) in automatic continuous positive airway pressure therapy to treat sleep disordered breathing. *J Clin Sleep Med* 2011;7:261–7.

22. Glidewell RN, Renn BN, Roby E, et al. Predictors and patterns of insomnia symptoms in OSA before and after PAP therapy. *Sleep* 2014;15:899–905.

23. Lévy P, Kohler M, McNicholas WT, et al. The sleep supine position has a major effect on optimal nasal continuous positive airway pressure treatment for sleep related breathing disorders in adults. *Sleep* 2002;25:148–73.

24. Lévy P, Kohler M, McNicholas WT, et al. The sleep supine position has a major effect on optimal nasal continuous positive airway pressure treatment for sleep related breathing disorders in adults. *Sleep* 2002;25:148–73.