STUDY PROTOCOL Open Access

Multiple N-of-1 trials to investigate hypoxia therapy in Parkinson's disease: study rationale and protocol

Jules M. Janssen Daalen¹, Marjan J. Meinders^{1,2}, Federica Giardina³, Kit C. B. Roes³, Bas C. Stunnenberg^{4,5}, Soania Mathur⁶, Philip N. Ainslie⁷, Dick H. J. Thijssen⁸ and Bastiaan R. Bloem^{1*}

Abstract

Background: Parkinson's disease (PD) is a neurodegenerative disease, for which no disease-modifying therapies exist. Preclinical and clinical evidence suggest that hypoxia-based therapy might have short- and long-term benefits in PD. We present the contours of the first study to assess the safety, feasibility and physiological and symptomatic impact of hypoxia-based therapy in individuals with PD.

Methods/Design: In 20 individuals with PD, we will investigate the safety, tolerability and short-term symptomatic efficacy of continuous and intermittent hypoxia using individual, double-blind, randomized placebo-controlled N-of-1 trials. This design allows for dose finding and for including more individualized outcomes, as each individual serves as its own control. A wide range of exploratory outcomes is deployed, including the Movement Disorders Society Unified Parkinson's Disease Rating scale (MDS-UPDRS) part III, Timed Up & Go Test, Mini Balance Evaluation Systems (MiniBES) test and wrist accelerometry. Also, self-reported impression of overall symptoms, motor and non-motor symptoms and urge to take dopaminergic medication will be assessed on a 10-point Likert scale. As part of a hypothesis-generating part of the study, we also deploy several exploratory outcomes to probe possible underlying mechanisms of action, including cortisol, erythropoietin and platelet-derived growth factor β. Efficacy will be assessed primarily by a Bayesian analysis.

Discussion: This evaluation of hypoxia therapy could provide insight in novel pathways that may be pursued for PD treatment. This trial also serves as a proof of concept for deploying an N-of-1 design and for including individualized outcomes in PD research, as a basis for personalized treatment approaches.

Trial registration: ClinicalTrials.gov Identifier: NCT05214287 (registered January 28, 2022).

Keywords: Parkinson's disease, Hypoxia, Treatment, Disease-modifying, Mitochondrial dysfunction, Clinical trial

Background

Parkinson's disease (PD) currently affects 10 million people worldwide and its prevalence is projected to rise exponentially in the coming decades [1]. Several symptomatic treatments are available, the mainstay of which has been levodopa for over half a century. Many patients continue to experience significant disability, despite deployment of all available management strategies. Therefore, additional treatment modalities are needed.

Anecdotal evidence from individuals with PD suggests that ascending to high-altitude areas occasionally improves motor symptoms of PD, in a subacute

Full list of author information is available at the end of the article $% \left(1\right) =\left(1\right) \left(1\right)$



© The Author(s) 2022. **Open Access** This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativeccommons.org/licenses/by/4.0/. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated in a credit line to the data.

^{*}Correspondence: Bas.Bloem@radboudumc.nl

¹ Center of Expertise for Parkinson & Movement Disorders; Nijmegen, the Netherlands, Department of Neurology, Donders Institute for Brain, Cognition and Behavior, Radboud University Medical Center, Nijmegen, The Netherlands

way. These findings were recently confirmed in a survey that we conducted among individuals with PD who had recently been on vacation (Janssen Daalen et al., manuscript submitted). We hypothesize that the positive effect of altitude on PD symptoms results from the decreased arterial oxygen tension at high altitude, which serves as an acute bodily stimulus for multisystem adaptations that potentially have protective effects on cellular homeostasis and survival [2, 3]. Therefore, altitude simulation has been the topic of research for potential therapeutic application in a variety of diseases.

Preclinical studies have suggested that hypoxia provokes release of survival-enhancing neurotransmitters. Specifically, the short-term clinical effects of hypoxia therapy appear to be related to augmented dopamine release from the substantia nigra [4–9]. Hypoxia therapy may improve parkinsonian symptoms via stabilization of hypoxia inducible factor 1 (HIF-1) and its downstream pathway, which in turn activates tyrosine hydroxylase (TH), the main rate-limiting enzyme in the production of dopamine [10, 11]. Several studies have demonstrated that HIF-1 stabilization leads to an increase in TH production, and consequently a rise in cellular dopamine content [8-13]. In addition, hypoxia protocols have a different influence on sympathetic nervous system activity, which regulates the body's stress response [12, 14-16]. For example, long-term hypoxia increases noradrenalin-adrenalin ratio [12], which might ameliorate nondopaminergic symptoms [17]. Taken together, these converging observations in animals and humans, provide a rationale that explains the potential positive effects of hypoxia on PD symptoms.

In addition to the short-term effects mentioned above, other studies also suggest that repeated exposure to hypoxia induces an evolutionary well-conserved adaptive mechanism. This adaptive response involves improves cellular energy metabolism as impaired by mitochondrial dysfunction, inhibits oxidative stress and induces adaptive plasticity, suggesting that in addition to the acute symptomatic effects, hypoxia might also exert long-term neuroprotective effects [18-20]. The concept behind these neuroprotective effects is the phenomenon of hypoxic preconditioning (HPC): induction of a sub-toxic hypoxic stimulus to improve the (systemic) tolerance of cells and tissues to subsequent more severe toxic stimuli. Although there is debate regarding the most potent hypoxia treatment regimen, clinical and preclinical evidence suggests that these effects are more pronounced when applied using a regime of intermittent hypoxia therapy (IHT) as compared to continuous hypoxia, meaning that hypoxia is present for short periods (i.e., minutes), interspersed with short periods of normoxic recovery. To date, hypoxia therapy, mostly IHT, has been used in a variety of populations, including fragile ones such as individuals with spinal cord injury, COPD, cardiac morbidity and multimorbidity, without any significant side effects [21–30]. However, the safety, feasibility and efficacy of hypoxia-based therapy have not been systematically investigated yet in individuals with PD. In this exploratory trial, we will assess the potential of hypoxia-based therapy in PD by assessing the physiological response to hypoxia, while also measuring the short-term symptomatic effects. To assess both continuous and intermittent hypoxia treatment regimens, this trial will deploy a double-blind, randomized placebocontrolled N-of-1 design, which allows for testing all selected hypoxia protocols in all participants [31].

Study objectives

Primary objectives

- (i) toevaluate the safety and feasibility of both intermittent and continuous hypoxiatherapy in individuals with PD under well-controlled circumstances.
- (ii) to explore the responsiveness of acute symptomatic outcome measures ofintermittent and continuous hypoxia therapy in individuals with PD underwell-controlled circumstances.

Secondary objectives

- (iii) to assess the acute symptomatic effects on selected subjective and standardized symptom scales.
- (iv) as a hypothesis-generating addition, to explore the potential mechanisms of (intermittent) hypoxia therapy on PD.

Hypothesis

We hypothesize that short-term hypoxia-based protocols are safe to apply in individuals with PD without cardiorespiratory comorbidity, and that IHT has short-term effects on dopamine, noradrenalin and stress-responsive symptoms in PD.

Methods/Design

Multiple N-of-1 trials

In this study, we will conduct multiple series of randomized, double-blind and placebo-controlled N-of-1 trials (also known as single participant cross-over trials). This design includes the testing of multiple hypoxia protocols in every participant, and it allows for the analysis of treatment effects in the individual participant (in addition to group effects) because the participant serves as his or her own control. Lastly, this design can result in a higher power when considering small populations as compared to other traditional designs, making N-of-1 trials appropriate for studying rare diseases and personalized treatment [32, 33]. N-of-1 trials are especially suitable to investigate treatments in chronic, symptomatic conditions, where period effects (i.e. changes in disease state) and carry-over effects (i.e. lingering treatment effects) are limited. Given the slowly progressive nature of PD with relative stable symptoms, several N-of-1 trials have already been successfully performed to study symptomatic treatments in PD [34, 35].

Study population

Twenty individuals with a diagnosis of PD (established by a neurologist according to the international Movement Disorder Society criteria) and Hoehn & Yahr stages between 1.5 and 3 will be included. Higher Hoehn & Yahr stages will be excluded for two reasons: firstly because of the significantly greater damage to dopaminergic pathways in advanced PD, which might limit the identification of nigrostriatal-mediated effects of hypoxia-based interventions; and secondly because of the high burden of testing participants in the OFF state in this more severely affected population. We aim to enrich our study population with individuals that have experienced (subjective) positive symptomatic effects at high altitude. In addition, we aim to include at least five individuals without prior experience with the positive effects of high altitude on their symptoms. This enrichment approach ensures that participants are more likely to be responders and allows us to validate our approach to identify each individual's optimal protocol for clinical benefit. We believe that the selection bias introduced by our approach is justifiable because of the study's explorative nature. In addition, PD is a very heterogeneous disease, for which a one-size-fitsall treatment approach is unlikely to be successful. Therefore, we wish to investigate for which individuals this therapy could be beneficial and the proposed approach is likely to be most promising. Main exclusion criteria relate to cardiorespiratory comorbidity and unstable PD medication. All inclusion and exclusion criteria can be found in Table 1.

Sample size

Because this is the first study in which the clinical effects of hypoxia therapy will be measured in PD and because of the study's exploratory nature, a formal sample size calculation cannot be performed. A previously published power and sample size simulation study aggregated N-of-1 trials with multiple cycles of intervention and placebo per participant, and found that under certain assumptions N-of-1 trials needed only one-third of the sample size of an RCT to reach a similar power and type I error [36]. This was confirmed in a recent aggregated N-of-1 trial that investigated the effectiveness of the drug mexiletine in myotonia, in which the aggregated N-of-1 trial showed comparable treatments effects on a personalized Likert scale (i.e., the same outcome measure as we propose) with inclusion of 11 participants vs 57 participants in a traditional cross-over RCT [32]. Our proposed sample size of 20 was motivated by this work, in combination with the sample size consensus for feasibility studies of 20–30 [37].

The observed effect sizes and standard deviations of the different hypoxia protocols that are found to be clinically meaningful at the participant group level in this study (N=20) will be used to simulate a power and sample size calculation for a future follow-up study that will be powered for efficacy using the data generated in this study.

Recruitment, screening and inclusion

Individuals with PD will be recruited via the web-based ParkinsonNEXT recruitment website (www.parkinsonn ext.nl) and, if necessary, from the outpatient registry of our university medical center – all patients on this

Table 1 Inclusion and exclusion criteria

Inclusion criteria	Exclusion criteria	
Clinical diagnosis of Parkinson's disease by a movement-disorder specialized neurologist	Individuals with diseases leading to restrictive and obstructive pulmonary diseases, pulmonary diff deficits, apnea and cardiac output deficits, such as pulmonary fibrosis, COPD, sleep apnea or excess alcoholic intake, and congestive heart failure respectively	
Hoehn and Yahr staging 1.5 to 3	Arterial blood gas abnormalities at screening day	
Age > 18 years	Individuals with shortness of breath or other airway or breathing-related inconvenience related to dopaminergic medication will be excluded	
Participant can provide informed consent	Inability to comply to intervention in off-medication condition (for example due to extreme discomfort, distress or severe head tremor due to being OFF, i.e. without dopaminergic medication)	
	Individuals with unstable dopaminergic medication dose (changes in the last month)	
	Individuals likely to start dopaminergic treatment in the next month, also judged by their treating neurologist	
	Individuals with active deep brain stimulation	
	Individuals unable to provide informed consent	

list have already consented to be contacted for research purposes. After a first telephone contact with the coordinating investigator, potential participants will receive an email with detailed information about the study, including an overview of necessary time investment and the risks of participation, as well as the informed consent form. Specifically, we will ask participants with a positive altitude experience to share this project in their own networks of hikers or mountaineers.

Individuals will have a consultation by telephone conducted by the coordinating investigator to answer questions about the research and the informed consent. If the participant is still interested, screening questionnaires will be sent before the physical screening day. The coordinating investigator will contact the participant by phone to check if potential exclusion criteria are met, to prevent any unnecessary participant visits. This process will be registered in a prescreening list. In this way, a transparent overview of study pre-selection will be available. The informed consent will be completed before initiation of the screening day. The participant has the right to withdraw consent at any moment during the study period. Drop-outs will be replaced.

Intervention

We will study the response to multiple exposures of different pre-selected hypoxia protocols per participant, as at present there is no conclusive evidence for the effect of different hypoxia stimuli. This will optimize the chances for any changes in outcomes being identified, and allowing the possibility of defining the optimal dose of hypoxia as a therapeutic intervention individually. The selection of interventions was based on multiple studies that investigated and reviewed the hypoxia regimes that are currently perceived as being most effective [38–43]. In this N-of-1 trial design, every participant will receive two sets of five different conditions, with one being the placebo and the other four variants of an active intervention, consisting of either intermittent or continuous hypoxia at FiO2 0.127 (~4000 m) and 0.163 (~2000 m). All interventions are administered twice to enhance intra-individual discriminative power towards a sufficiently low or high probability (see Statistical Analysis). As it is shown that hypoxic preconditioning effects may linger for up to four days [44], a wash-out of at least five days between sessions will be built-in. Once a week, one of the following five conditions will be administered following the order resulting from a Latin square randomization (see Statistical Analysis):

Active interventions:

- Continuous hypoxia for 45 min, at ~ 2000 m (16.3% O₂)
- Continuous hypoxia for 45 min, at ~4000 m (12.7% O₂)
- Intermittent hypoxia with 5×5 -min at ~ 2000 m (16.3% O_2), interspersed with 5-min normoxic recovery
- Intermittent hypoxia with 5×5 -min at ~ 4000 m (12.7% O_2), interspersed with 5-min normoxic recovery
 - Placebo intervention:
- Continuous normoxia for 45 min (20.9% O₂)

The intervention will be performed at Radboud university medical center in Nijmegen, the Netherlands. A commercially available hypoxicator will be used to deliver the hypoxic bout (B-cat High Altitude, the Netherlands). The hypoxicator (as an oxygen concentrator) is ISO 13485:2016 certified and uses a process called pressure swing adsorption to filter ambient air and to extract oxygen from that air. The principle is widely used in oxygen concentrators, but is inversely applied in the case of hypoxicators. Instead of oxygen-rich air, the resultant hypoxic gas is administered to the patient [45]. The hypoxicator is connected via medical-grade tubes to two large (>50L) reservoir air bags that buffer the hypoxic gas mixture, which optimizes stability of the delivered FiO2. Intermittent hypoxia and the placebo situation is administered by switching a three-way Hans Rudolph® valve to either room air or the hypoxic circuitry. The intraday and between-day reliability of the hypoxicator were tested on multiple days before application in participants. Intraday variability was low (average range 0.2 percent points around desired FiO2) and between-day variability was within 0.1 percent points of desired FiO2. Seven days before the start of the first intervention, a one week at-home registration of self-reported outcomes of symptom severity will be conducted by a short daily morning survey. This assures a solid baseline of symptom severity that will be used in the Bayesian analysis (see Statistical Analysis).

Study procedures

Screening

Before inclusion, potential participants will come to the hospital for a dedicated screening day, during which extensive safety screening will be conducted. Firstly, this will consist of standard pulmonary function testing (PFT) using spirometry to assess the presence of any previously unknown restrictive or obstructive pulmonary diseases. Second, a carbon monoxide diffusion test for any unknown diffusion deficits is conducted to exclude any unknown comorbidity that might pose a health risk

during the study. Finally, peak cough flow and mean inspiratory pressure (MIP) are assessed, as well as subjective respiratory problems via a selection of questions that predicted PD-related respiratory problems (Supplementary Table 1) in a recent study [Van de Wetering et al., submitted]. All participants will also undergo an electrocardiogram (ECG) to screen for any cardiac abnormalities such as dysrhythmias or signs of ischemia.

Subsequently, participants will be blindly exposed to stepwise decreasing levels of FiO2 until an FiO2 of 0.127 is administered fully. This will be performed in four steps. Before every step (step duration of approximately 5 min), arterial blood gas (ABG) and vital parameters are evaluated and compared to the pre-determined stop criteria (Supplementary Table 2). Vital parameters including peripheral oxygen saturation will be collected every 5 min and will be correlated to the ABG results to evaluate whether these vital parameters reflect the ABG results. If one or more of the stop criteria are met and the breathed-in air does not yet contain an FiO2 of 0.127, the intervention will be stopped. Moreover, the participant will then be excluded from further interventions and will be replaced.

Pilot phase

To ensure optimal safety, a pilot study of the first two participants selected for this study protocol will be deployed before conducting this study protocol at full pace. During the screening procedure, all individuals will be exposed to gradually decreasing FiO2's until an FiO2 of 0.127 (~4000 m altitude) is reached. In these two individuals, the most intense intervention (continuous hypoxia at FiO2 0.127) will be administered for 45 min, to evaluate whether exhaustion in individuals with PD will limit the maximum safe duration as clinical experience is limited. In the remaining participants, the 4000 m intervention will be administered until ABG parameters have stabilized. When no serious adverse events occur, the protocol will be continued. During this pilot phase, the same stop criteria will be adopted as during the interventions.

Intervention

After completion of the screening day, participants will visit the hospital on a weekly basis, for 10 consecutive weeks. Treatments and in-hospital assessments will be conducted in the practically-defined OFF phase and will therefore commence in the morning. The treatment session in the OFF state is preceded by a baseline clinical assessment and is immediately followed by a post-intervention clinical assessment and an assessment after 30 min (Table 3). The intervention is administered in a hospital by an experienced lab technician, who also continuously monitors participants during and after the

intervention. During the intervention, the inhaled FiO2 is measured continuously using the COSMED® metabolic system (*Quark CPET* metabolic cart for cardiopulmonary testing, COSMED Srl, The Metabolic Company, Italy), which also measures peripheral oxygen saturation, blood pressure, respiratory rate and tidal volume. In addition, the subjective participant experience regarding dizziness, discomfort and stress is registered by the lab technician during the intervention. In addition, to maximize safety of the participants during the intervention, clear stopping criteria are defined. If these are met, the lab technician will halt the intervention ('Stopping Criteria,' Supplementary Materials).

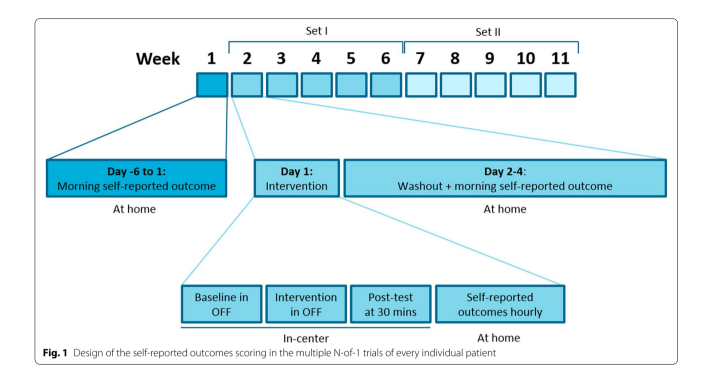
Randomization and blinding

Randomization

Subjects will be equally divided in 5 groups with different interventions order according to a Latin square design (5 periods/interventions). Allocation to intervention order will be randomized by a computer-generated randomization scheme. Interventions will be randomized at study start in two sets. As depicted in Fig. 1, set I will be randomized for the first five interventions and set II for the second five interventions. This reduces the risk of placebo effects in Set II potentially provoked by any positive effects that may have been experienced in Set I. This balanced randomization scheme minimizes bias introduces by the resulting order effect and period effect. Carry-over effects are minimized by ascertaining a sufficient washout period between every treatment.

Blinding

The intervention is double-blind for the participant and assessor, but not for the lab technician (because of safety and monitoring purposes). Ventilation during the administration is expected to increase by ~20%, mostly due to increased tidal volume and less often due to respiratory rate [46]. Risk of unblinding due to these limited ventilatory effects is unlikely and has not been reported in previous hypoxic experiments [28, 46-48]. Additionally, complaints of discomfort or shortness of breath are sometimes reported, regardless of the administered FiO2, indicating that this will not hamper blinding [23-25]. Changes in respiratory parameters will be available for the lab technician during the experiments, but will not be disclosed to the assessor. The assessor will only enter the lab after the intervention has been fully completed and will not have access to the vital parameters and blood parameters of the participant from which it might be possible to infer the assigned intervention. This problem is also mitigated by adding the placebo condition. We will assess



the success of the blinding procedure by asking the participant in what sequence the different treatments were administered after the first and second treatment set.

Outcomes and measurements Primary outcomes

The main primary outcomes of this study are the safety and feasibility of IHT under well-controlled circumstances in a lab-based setting. Safety outcomes will be measured by the lab technician during the intervention and by the reporting of adverse events. The feasibility questionnaire (Supplementary Table 3) is designed together with a

patient researcher. The questionnaire's content is based on the main categories from a widely used feasibility framework [49]. Statements in every category (e.g. acceptability, expectancy) were subsequently inspired by previously published healthcare feasibility questionnaires. Primary outcomes are summarized in Table 2.

Secondary outcomes Self-reported outcomes

In this exploratory study, we will use a battery of outcomes to determine which symptoms or signs might respond to hypoxia. In accordance with accepted N-of-1

Table 2 Primary outcomes

Primary outcome variable	Measurement frequency		
Nature and number of adverse events	Every 10 min during intervention, up to one hour post-intervention, one time next morning post-intervention		
Self-reported dizziness, discomfort and stress on a ten-point scale	Every 10 min up to one hour post-intervention, one time next morning post-intervention		
Number of serious or irreversible adverse events	Every 10 min, up to one hour post-intervention, one time next morning post-intervention		
	Measured continuously for safety, recorded:		
- Blood pressure systolic-diastolic	Baseline and every 5 min—> 30 min post-intervention		
- Heartrate	Baseline and every 5 min—> 30 min post-intervention		
- Respiratory rate	Baseline and every 5 min—> 30 min post-intervention		
- Oxygen saturation	Baseline and every 5 min—> 30 min post-intervention		
Feasibility questionnaire (total score and subscores) (in <i>Supplementary Materials</i>)	After 1 st , 5 th and 10 th intervention		

trial design recommendations and previously conducted trials [32, 50, 51], we have chosen to predefine personalized (self-reported) outcomes for each individual, scored on a 10-point Likert scale. Contrary to standardized PD scales, these outcomes better reflect important effects for the individual, and are potentially more sensitive to subtle treatment effects, reducing the risk of type II errors. Therefore, we included three participant-rated outcomes that assess on a 10-point Likert scale (allowing half points) general symptom impression, the urge to take a next dose of dopaminergic medication and the effect on one specific symptom that participants can choose themselves, based on goal attainment scale principles (Table 3). This symptom must fulfill the following criteria: it must fluctuate in severity throughout the day or between days, changes in its severity should be swiftly apparent and, if applicable, it is a symptom that previously improved at high altitude. To ensure a baseline score on these outcomes, participants will report these on a daily basis in the morning (in OFF) during the baseline period, i.e. 7 days preceding the start of the intervention period. On intervention days, participants will report these self-reported outcomes pre-intervention, directly post-intervention, 30 min post-intervention and from that moment on, another five times, on an hourly basis. In addition, these will be measured once every morning (i.e. in OFF) for the next three mornings after the intervention.

The measurement scheme of all outcomes is displayed in Supplementary Fig. 1. Patients will report all selfreported outcomes digitally.

Assessor-rated outcomes

Assessor-rated outcomes are summarized in Table 3. These consist of gold-standard general motor tests (Movement Disorders Society Universal Parkinson's Disease Rating Scale part III, or MDS-UPDRS part III) and non-motor tests (symptoms from the MDS Non-motor symptoms scale, or NMSS), supplemented with specific tests for bradykinesia (Purdue pegboard test [52]), gait

Table 3 Secondary outcomes

Self-reported outcome variable	Researcher	Participant	Moment of measurement (in addition to baseline)	Measurement unit
General symptom impression		X	Directly after intervention and post- intervention 30 min, 1 h, 2 h, 3 h, 4 h and 5 h, and once every morning on the next three days	10-point Likert scale allowing half points
Urge for dopaminergic medication on the usual moments of intake		X	Directly after intervention and post- intervention 30 min, 1 h, 2 h, 3 h, 4 h and 5 h, and once every morning on the next three days	10-point Likert scale allowing half points
Participant-selected symptom (goal attainment scale)		X	Directly after intervention and post- intervention 30 min, 1 h, 2 h, 3 h, 4 h and 5 h, and once every morning on the next three days	10-point Likert scale allowing half points
Motor symptom severity (self-selected, symptom that improved most during high altitude exposure, if applicable)		X	Directly after intervention and post- intervention 30 min, 1 h, 2 h, 3 h, 4 h and 5 h, and once every morning on the next three days	10-point Likert scale allowing half points
Motor symptoms MDS-UPDRS part III	Χ		30 min post-intervention	Total score and subscores
Hand tremor Accelerometry/gyroscope	Χ		Directly post-intervention, 30 min post-intervention	Amplitude
Rapid alternating movements Accelerometry/gyroscope	Χ		30 min post-intervention	Amplitude
Bradykinesia Modified Perdue pegboard test	Χ		30 min post-intervention	Number of pins
Gait Timed Up & Go Test	Χ		30 min post-intervention	Steps and seconds
Balance MiniBESTest	Χ		30 min post-intervention	Total score and subscores
Non-motor symptoms MDS Non-motor Scale symptoms	Χ		30 min post-intervention	10-point Likert scale (half points)
Stress Heartrate variability	X		During intervention, directly after intervention, 30 min post-intervention	RR interval

(Timed Up & Go Test [53]), balance (Mini-BESTest [54]) and quantified versions of the UPDRS items on tremor and pronation-supination (performed using accelerometry). These secondary outcomes are measured 30 min post-intervention, after which hypoxic preconditioning is believed to have its initial first window effects peak. After 30 min, the acute effects of hypoxia therapy peak, as shown in people with a cervical spinal cord injury [55], with the second window only occurring approximately 3 h post-intervention.

Various selected outcomes, including the UPDRS part III, are sensitive to the Hawthorne effect: individuals with PD try to perform as good as possible due to the awareness of being observed. To mitigate such effects, the placebo condition is added, for which the multiple N-of-1 design is also particularly suited [31, 36, 56].

Baseline characteristics that will be collected include age, gender, H&Y, quality of life (Parkinson's disease questionnaire 39 [57]) and other medication. Potential effect modifiers include levodopa-equivalent dose (LED), sleep quality (4-point ordinal scale) and physical activity (International Physical Activity Questionnaire – Short Form, IPAQ-SF [58]) and these will be measured during every pre-intervention phase.

Mechanistic markers

Lastly, three markers will be measured in serum, which serve as a hypothesis-generating addition to our study. These measures may provide insight into potential pathways involved in the (individual) responses to hypoxia in our study. These measures include platelet-derived growth factor receptor β (PDGFR β), cortisol and erythropoietin (EPO).

PDGFR β is a pericyte-shedded marker in response to hypoxia [59–62] and is associated with blood–brain barrier (BBB) permeability [60, 61, 63–65]. Disruption of the BBB is a central process involved in PD pathophysiology [66] and therefore, acute effects of hypoxia therapy on PDGFR β would give insight in the influence that hypoxia might have on BBB integrity in the long term [65].

Cortisol is a marker of physiological and mental stress and is hypothesized to rise during systemic hypoxic challenges [67]. As stress is one of the main determinants of variations in symptom severity, we investigate whether cortisol release might be associated with the short-term symptomatic efficacy of hypoxia-based interventions. This will give insight in whether hypoxia has beneficial influence on the stress system, or that the physiological stress results in fatigue and mental stress, thus worsening PD symptoms in the short-term. Cortisol is measured twice pre-intervention and three times post-intervention, because of its concentration changing with circadian rhythm.

EPO is a protein that is primarily shedded peripherally by the kidney. Although it is employed as a marker of hypoxic dose, attention for its role in neuroprotection has risen in recent years [68]. In preclinical PD models, it prevents neurotoxicity and preserves neuronal functioning [69, 70].

All outcome measures are summarized in further detail in Supplementary Fig. 1.

Statistical analysis

All data will be collected using direct entry in CastorEDC, a widely used electronic data capture system for clinical data.

Analysis of primary study parameters

Safety outcomes will be analyzed using descriptive statistics of total number and percentages of adverse events, number of serious reversible or irreversible adverse events and vital parameters. Feasibility outcomes will be analyzed using descriptive statistics of feasibility outcomes including the feasibility questionnaire sum score and domain-specific scores.

Secondary study parameters Bayesian analysis

We will use a Bayesian model for the analysis of individual and aggregated N-of-1 trial results. This model allows for a direct estimation of the posterior probability that a treatment results in a clinically beneficial effect [32, 71–73]. The treatment effects resulting from the four different hypoxia protocols will be estimated per level of interest (i.e. the individual and group level). Every secondary outcome scored using the 10-point Likert scale will be modeled assuming a normal distribution centered around the patient's true mean and variance for each protocol. If the posterior probability of reaching a 0.75point difference between the secondary outcome after treatment and at baseline is greater than 80%, treatments are considered effective. Treatments are considered ineffective if the same posterior probability is less than 20%. Every treatment day has its own baseline (pre-intervention tests are performed) in order to further reduce period effects. The minimal clinically important difference for the secondary outcome MDS-UPDRS part III and Mini-BESTest is 4. One additional analysis will be performed including baseline characteristics and effect modifiers (see Assessor-rated outcomes) as covariates. Bayesian analyses will be performed using *JAGS* version 3.4.1 [74], run from *R* [75] using the package *rjags* [76].

Frequentist analysis

For comparison with the Bayesian analysis we will also perform an explorative analysis of the secondary outcomes. For those that are normally distributed, we will use dependent t tests to calculate mean treatment effects, significance levels, and confidence intervals on group level. P values are 2-sided, and P<0.05 will be the threshold for significance for all tests. Analyses will be performed using R [75].

Monitoring and registration

Quality assurance and monitoring

The study will be monitored regarding the health, safety and rights of participants, protocol adherence and quality of data and data reporting during this trial at study initiation, twice during the study (on-site) and once at study completion. On-site monitoring visits (performed by Radboudumc or designee) will assess the progress of the study, study procedures, used study materials and identify any concerns that result from review of the subject Informed Consent documentation, study records, collected data and study management documents. The study monitor will also ensure the Investigator adheres to all applicable regulations.

Data safety monitoring board

A data safety monitoring board (DSMB) will be established, which will consist of a PD-specialized neurologist, an anesthesiologist, a pulmonologist, a biostatistician and a patient representative. A first interim analysis of the safety outcomes will be performed after the pilot phase of the first two subjects that are halfway through the treatment protocol (which means every treatment is already administered once), to provide the DSMB with the latest data on adverse events and recruitment. The second interim analysis after 7 participants have completed their protocol will give insight in primary as well as secondary outcomes.

Discussion

In this study, we propose multiple N-of-1 trials to investigate the merits of hypoxia-based interventions as a new symptomatic therapy in persons with PD. For the first time, this design offers a unique opportunity to test for the first time the safety, feasibility and short-term efficacy of various interventions in this unexplored therapeutic area in PD. Therefore, a wide variety of participant-reported and assessor-rated outcome measures will be deployed.

Hypoxia-based therapy has been applied extensively in research in a wide spectrum of healthy participants and individuals with medical conditions, and both short- and long-term effects have been investigated. Examples of previously studied treatment goals in various populations include rehabilitation in spinal cord injury (SCI) [55, 77–81], cardiorespiratory control in

type I and II diabetes [82, 83], endurance and exercise tolerance and performance in healthy and geriatric individuals [24, 25, 27, 29, 84, 85], cognitive performance in geriatric and elderly individuals [22, 26, 28, 86], cardiovascular risk factors in obese individuals [87], reducing acute mountain sickness [88], and training of respiratory dysfunction [42, 48, 89–91]. However, clinical parameters or symptomatic efficacy of hypoxiabased therapy have thus far never been studied in PD, even though the aforementioned underlying working mechanisms of hypoxia would make PD an attractive disorder to study. One earlier brief report investigated the effects of (unspecified) IHT on the hypoxic ventilatory response in PD and found markedly reduced hypoxic ventilatory response, indicating a suboptimal response in breathing frequency to hypoxic challenges [48]. Because there is such limited previous experience with delivering hypoxia to persons with PD, several theoretical concerns must be addressed in this study proposed here. First, because respiratory abnormalities can already be observed in relatively early stages of PD [92], the safety and feasibility profile of different hypoxia protocols must be established first; this is one of the goals of the present study. Second, the shortterm as well as long-term effects should be investigated separately, as different mechanisms might be involved [Janssen Daalen et al., manuscript in preparation]. With regard to short-term effects, we might counter both beneficial and harmful effects (such as stress, increased oxidative stress), and the hypoxic dose for which this balance is optimal remains to be established in PD.

This exploratory study will provide the first insights into the potential of hypoxia-based therapy in PD. Additionally, our study might yield hypothesis-generating insights regarding its underlying working mechanisms. At the same time, the findings might also improve our understanding of the mechanisms of respiratory involvement in PD and on motor and non-motor symptom variability, that can be derived from the weekly administered neurological test battery. The findings in this study might partly be extrapolated to other neurodegenerative diseases, such as Alzheimer's disease [40] or mitochondrial diseases [20]. Although the mechanisms of hypoxia-based interventions remain to be fully elucidated, we believe this rationale warrants the first well-controlled randomized trial of hypoxia-based interventions in PD.

Abbreviations

PD: Parkinson's disease; HIF-1: Hypoxia-inducible factor 1; TH: Tyrosine hydroxylase; HPC: Hypoxic preconditioning; MiniBES: Mini Balance Evaluation System; MDS-UPDRS: Movement Disorders Society Unified Parkinson's Disease Rating scale; IHT: Intermittent hypoxia therapy; SCI: Spinal cord injury; DSMB: Data safety monitoring board; PFT: Pulmonary function testing; MIP: Mean inspiratory pressure; ECG: Electrocardiogram; ABG: Arterial blood ga; LED:

Levodopa-equivalent dose; IPAQ-SF: International Physical Activity Questionnaire – Short Form; PDGFR β : Platelet-derived growth factor receptor β ; EPO: Erythropoietin; BBB: Blood–brain barrier; MJFF: Michael J. Fox Foundation for Parkinson's Research.

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s12883-022-02770-7.

Additional file 1: Supplementary Table 1. Orientating questionnaire for respiratory problems in PD [Van de Wetering et al., submitted]. Supplementary Table 2. Stop criteria. Supplementary Table 3. Feasibility questionnaire, scored on 10-point Likert scale. Supplementary Figure 1. Summary of outcome measurements

Additional file 2: SPIRIT checklist for interventional trials

Acknowledgements

The authors thank Hans M. Groenewoud, PhD (Radboud University Medical Center) for his methodological and statistical input throughout the conceptual phase of this study, and Jeroen van Hees, PhD and Matthijs Kox, PhD (Radboud university medical center) for their contributions relating to monitoring of safety and respiratory parameters. We thank expert by experience mr. Gerben Kestens for his input on study methodology and logistics.

Authors' contributions

MJM, DHJT and BRB obtained funding for the study. JMJD, MJM, FG, KCBR, BCS, SM, PNA, DHJT and BRB contributed to study design. BRB is primary investigator, JMJD is coordinating investigator and responsible for outcome assessment and data collection. JMJD, MJM, DHJ and BRB are responsible for general study execution. JMJD, MJM, FG, KCBR and BCS will contribute to data analysis. JMJD, MJM and DHJT wrote the first draft of the manuscript. All authors have read, critically revised and approved the final version of the manuscript.

Funding

This study is fully supported by the Michael J. Fox Foundation (MJFF, Therapeutic Pipelines Program, funding number MJFF-019201). The study funder will be consulted and informed during the study, but is not involved in study design, data collection, analysis or interpretation, or in writing or publication matters. The Centre of Expertise for Parkinson & Movement Disorders was supported by a Centre of Excellence grant by the Parkinson Foundation.

Availability of data and material

Anonymized data will be shared with The Michael J. Fox Foundation for Parkinson's Research (the study funder). This data may be kept for storage at a central repository either hosted by The Michael J. Fox Foundation, its collaborators, or consultants and will be kept indefinitely. Anonymized data will be made publicly available by the Foundation for the intended use of research in Parkinson's disease as well as other biomedical research studies that may not be related to Parkinson's disease.

Declarations

Ethics approval and consent to participate

The study will be performed according to the guidelines stated in the 'Declaration of Helsinki' from 2000 and 'the Medical Research Involving Human Subjects Act (WMO)' of the Civil Code of the Netherlands. This study has been reviewed and approved by the Medical Research Ethics Committee East Netherlands, The Netherlands, (reference number NL.77891.091.22) and has been registered at clinicaltrials.gov (ClinicalTrials.gov Identifier: NCT05214287). Participants receive verbal and written information about the study by the coordinating investigator and written informed consent will be obtained before screening. Participants are covered by trial insurance issued specifically for this study by Radboud University Medical Center.

Consent for publication

Not applicable.

Competing interests

Bastiaan R. Bloem has received honoraria from serving on the scientific advisory board for Abbvie, Biogen, UCB, and Walk with Path; has received fees for speaking at conferences from AbbVie, Zambon, Roche, GE Healthcare, and Bial; and has received research support from The Netherlands Organisation for Scientific Research, the Michael J. Fox Foundation, UCB, Abbvie, the Stichting Parkinson Fonds, the Hersenstichting Nederland, the Parkinson Foundation, Verily Life Sciences, Horizon 2020, the Topsector Life Sciences and Health, and the Parkinson Vereniging. The other authors have nothing to disclose.

Author details

¹Center of Expertise for Parkinson & Movement Disorders; Nijmegen, the Netherlands, Department of Neurology, Donders Institute for Brain, Cognition and Behavior, Radboud University Medical Center, Nijmegen, The Netherlands. ²IQ Healthcare, Radboud Institute for Health Sciences, Radboud University Medical Center, Nijmegen, The Netherlands. ³Department of Health Evidence, Radboud Institute for Health Sciences, Radboud University Medical Center, Section Biostatistics, Nijmegen, The Netherlands. ⁴Department of Neurology, Donders Institute for Brain, Cognition and Behavior, Radboud University Medical Center, Nijmegen, The Netherlands. ⁵Department of Neurology, Rijnstate Hospital, Arnhem, Netherlands. ⁶UnshakeableMD, Oshawa, ON, Canada. ⁷Center for Heart, Lung and Vascular Health, School of Health and Exercise Sciences, University of British Columbia, Kelowna, Canada. ⁸Department of Physiology, Radboud University Medical Center, Nijmegen, The Netherlands.

Received: 23 May 2022 Accepted: 24 June 2022 Published online: 14 July 2022

References

- Dorsey ER, Bloem BR. The Parkinson Pandemic-A Call to Action. JAMA Neurol. 2018;75(1):9–10.
- Prabhakar NR, Semenza GL. Adaptive and maladaptive cardiorespiratory responses to continuous and intermittent hypoxia mediated by hypoxiainducible factors 1 and 2. Physiol Rev. 2012;92(3):967–1003.
- Semenza GL. Hypoxia-inducible factor 1 and cardiovascular disease. Annu Rev Physiol. 2014;76:39–56.
- Broderick PA, Gibson GE. Dopamine and Serotonin in Rat Striatum during Invivo Hypoxic-Hypoxia. Metab Brain Dis. 1989;4(2):143–53.
- Barath AS, Rusheen AE, Rojas Cabrera JM, Price JB, Owen RL, Shin H, Jang DP, Blaha CD, Lee KH, Oh Y. Hypoxia-Associated Changes in Striatal Tonic Dopamine Release: Real-Time in vivo Measurements With a Novel Voltammetry Technique. Front Neurosci. 2020;14:869.
- Orset C, Parrot S, Sauvinet V, Cottet-Emard JM, Berod A, Pequignot JM, Denoroy L. Dopamine transporters are involved in the onset of hypoxiainduced dopamine efflux in striatum as revealed by in vivo microdialysis. Neurochem Int. 2005;46(8):623–33.
- Akiyama Y, Koshimura K, Ohue T, Lee K, Miwa S, Yamagata S, Kikuchi H. Effects of hypoxia on the activity of the dopaminergic neuron system in the rat striatum as studied by in vivo brain microdialysis. J Neurochem. 1991;57(3):997–1002.
- 8. Witten L, Sager T, Thirstrup K, Johansen JL, Larsen DB, Montezinho LP, Mørk A. HIF prolyl hydroxylase inhibition augments dopamine release in the rat brain in vivo. J Neurosci Res. 2009;87(7):1686–94.
- Johansen JL, Sager TN, Lotharius J, Witten L, Mørk A, Egebjerg J, Thirstrup K. HIF prolyl hydroxylase inhibition increases cell viability and potentiates dopamine release in dopaminergic cells. J Neurochem. 2010;115(1):209–19.
- 10. Yuan G, Nanduri J, Bhasker CR, Semenza GL, Prabhakar NR. Ca2+/calmodulin kinase-dependent activation of hypoxia inducible factor 1 transcriptional activity in cells subjected to intermittent hypoxia. J Biol Chem. 2005;280(6):4321–8.
- Belikova MV, Kolesnikova EE, Serebrovskaya TV: Intermittent hypoxia and experimental Parkinson's disease. In: Intermittent hypoxia and human diseases. edn. Edited by Xi L, Serebrovskaya T. London: Springer; 2012: 147–153.
- 12. Rostrup M. Catecholamines, hypoxia and high altitude. Acta Physiol Scand. 1998;162(3):389–99.
- 13. Schnell PO, Ignacak ML, Bauer AL, Striet JB, Paulding WR, Czyzyk-Krzeska MF. Regulation of tyrosine hydroxylase promoter activity by the von

- Hippel-Lindau tumor suppressor protein and hypoxia-inducible transcription factors. J Neurochem. 2003;85(2):483–91.
- Hughson RL, Yamamoto Y, McCullough RE, Sutton JR, Reeves JT. Sympathetic and parasympathetic indicators of heart rate control at altitude studied by spectral analysis. J Appl Physiol (1985). 1994;77(6):2537–42.
- Hansen J, Sander M. Sympathetic neural overactivity in healthy humans after prolonged exposure to hypobaric hypoxia. J Physiol. 2003;546(Pt 3):921–9.
- Rowell LB, Johnson DG, Chase PB, Comess KA, Seals DR. Hypoxemia raises muscle sympathetic activity but not norepinephrine in resting humans. J Appl Physiol (1985). 1989;66(4):1736–43.
- Espay AJ, LeWitt PA, Kaufmann H. Norepinephrine deficiency in Parkinson's disease: the case for noradrenergic enhancement. Mov Disord. 2014;29(14):1710–9.
- Burtscher J, Syed MMK, Lashuel HA, Millet GP. Hypoxia Conditioning as a Promising Therapeutic Target in Parkinson's Disease? Mov Disord. 2021;36(4):857–61.
- Zhang Z, Yan J, Chang Y, Yan SSD, Shi H. Hypoxia Inducible Factor-1 as a Target for Neurodegenerative Diseases. Curr Med Chem. 2011;18(28):4335–43.
- Jain IH, Zazzeron L, Goli R, Alexa K, Schatzman-Bone S, Dhillon H, Goldberger O, Peng J, Shalem O, Sanjana NE, et al. Hypoxia as a therapy for mitochondrial disease. Science. 2016;352(6281):54–61.
- Vose AK, Welch JF, Nair J, Dale EA, Fox EJ, Muir GD, Trumbower RD, Mitchell GS. Therapeutic acute intermittent hypoxia: A translational roadmap for spinal cord injury and neuromuscular disease. Exp Neurol. 2022;347: 113891.
- Serebrovska ZO, Serebrovska TV, Kholin VA, Tumanovska LV, Shysh AM, Pashevin DA, Goncharov SV, Stroy D, Grib ON, Shatylo VB, et al. Intermittent Hypoxia-Hyperoxia Training Improves Cognitive Function and Decreases Circulating Biomarkers of Alzheimer's Disease in Patients with Mild Cognitive Impairment: A Pilot Study. Int J Mol Sci. 2019;20(21):5405.
- Saeed O, Bhatia V, Formica P, Browne A, Aldrich TK, Shin JJ, Maybaum S. Improved exercise performance and skeletal muscle strength after simulated altitude exposure: a novel approach for patients with chronic heart failure. J Card Fail. 2012;18(5):387–91.
- Burtscher M, Pachinger O, Ehrenbourg I, Mitterbauer G, Faulhaber M, Puhringer R, Tkatchouk E. Intermittent hypoxia increases exercise tolerance in elderly men with and without coronary artery disease. Int J Cardiol. 2004;96(2):247–54.
- Burtscher M, Haider T, Domej W, Linser T, Gatterer H, Faulhaber M, Pocecco E, Ehrenburg I, Tkatchuk E, Koch R, et al. Intermittent hypoxia increases exercise tolerance in patients at risk for or with mild COPD. Respir Physiol Neurobiol. 2009;165(1):97–103.
- Bayer U, Likar R, Pinter G, Stettner H, Demschar S, Trummer B, Neuwersch S, Glazachev O, Burtscher M. Intermittent hypoxic-hyperoxic training on cognitive performance in geriatric patients. Alzheimers Dement (N Y). 2017;3(1):114–22.
- Bonetti DL, Hopkins WG. Sea-level exercise performance following adaptation to hypoxia: a meta-analysis. Sports Med. 2009;39(2):107–27.
- Serebrovskaya TV, Karaban IN, Kolesnikova EE, Mishunina TM, Swanson RJ, Beloshitsky PV, Ilyin VN, Krasuk AN, Safronova OS, Kuzminskaya LA. Geriatric men at altitude: hypoxic ventilatory sensitivity and blood dopamine changes. Respiration. 2000;67(3):253–60.
- Bayer U, Likar R, Pinter G, Stettner H, Demschar S, Trummer B, Neuwersch S, Glazachev O, Burtscher M. Effects of intermittent hypoxia-hyperoxia on mobility and perceived health in geriatric patients performing a multimodal training intervention: a randomized controlled trial. BMC Geriatr. 2019;19(1):167.
- Christiansen L, Chen B, Lei Y, Urbin MA, Richardson MSA, Oudega M, Sandhu M, Rymer WZ, Trumbower RD, Mitchell GS, et al. Acute intermittent hypoxia boosts spinal plasticity in humans with tetraplegia. Exp Neurol. 2021;335: 113483.
- 31. Lillie EO, Patay B, Diamant J, Issell B, Topol EJ, Schork NJ. The n-of-1 clinical trial: the ultimate strategy for individualizing medicine? Per Med. 2011;8(2):161–73.
- Stunnenberg BC, Raaphorst J, Groenewoud HM, Statland JM, Griggs RC, Woertman W, Stegeman DF, Timmermans J, Trivedi J, Matthews E, et al. Effect of Mexiletine on Muscle Stiffness in Patients With Nondystrophic Myotonia Evaluated Using Aggregated N-of-1 Trials. JAMA. 2018;320(22):2344–53.

- 33. Facey K, Granados A, Guyatt G, Kent A, Shah N, van der Wilt GJ, Wong-Rieger D. Generating health technology assessment evidence for rare diseases. Int J Technol Assess Health Care. 2014;30(4):416–22.
- 34. Rascol O, Ferreira J, Nègre-Pages L, Perez-Lloret S, Lacomblez L, Galitzky M, Lemarié JC, Corvol JC, Brotchie JM, Bossi L. A proof-of-concept, randomized, placebo-controlled, multiple cross-overs (n-of-1) study of naftazone in Parkinson's disease. Fundam Clin Pharmacol. 2012;26(4):557–64.
- 35. Ferreira JJ, Mestre T, Guedes LC, Coelho M, Rosa MM, Santos AT, Barra M, Sampaio C, Rascol O. Espresso Coffee for the Treatment of Somnolence in Parkinson's Disease: Results of n-of-1 Trials. Front Neurol. 2016;7:27.
- Blackston JW, Chapple AG, McGree JM, McDonald S, Nikles J. Comparison of Aggregated N-of-1 Trials with Parallel and Crossover Randomized Controlled Trials Using Simulation Studies. Healthcare (Basel). 2019;7(4):137.
- Billingham SA, Whitehead AL, Julious SA. An audit of sample sizes for pilot and feasibility trials being undertaken in the United Kingdom registered in the United Kingdom Clinical Research Network database. BMC Med Res Methodol. 2013;13:104.
- Soo J, Girard O, Ihsan M, Fairchild T. The Use of the SpO2 to FiO2 Ratio to Individualize the Hypoxic Dose in Sport Science, Exercise, and Health Settings. Front Physiol. 2020;11: 570472.
- Navarrete-Opazo A, Mitchell GS. Therapeutic potential of intermittent hypoxia: a matter of dose. Am J Physiol Regul Integr Comp Physiol. 2014;307(10):R1181-1197.
- 40. Mateika JH, El-Chami M, Shaheen D, Ivers B. Intermittent hypoxia: a low-risk research tool with therapeutic value in humans. J Appl Physiol (1985). 2015;118(5):520–32.
- Serebrovska TV, Serebrovska ZO, Egorov E. Fitness and therapeutic potential of intermittent hypoxia training: a matter of dose. Fiziol Zh. 2016;62(3):78–91.
- 42. Garcia N, Hopkins SR, Powell FL. Intermittent vs continuous hypoxia: effects on ventilation and erythropoiesis in humans. Wilderness Environ Med. 2000;11(3):172–9.
- Baker TL, Mitchell GS. Episodic but not continuous hypoxia elicits longterm facilitation of phrenic motor output in rats. J Physiol. 2000;529(Pt 1):215–9.
- 44. Hausenloy DJ, Yellon DM. The second window of preconditioning (SWOP) where are we now? Cardiovasc Drugs Ther. 2010;24(3):235–54.
- 45. Grande CA. Advances in Pressure Swing Adsorption for Gas Separation. ISRN Chemical Engineering. 2012;2012: 982934.
- Liu X, Xu D, Hall JR, Ross S, Chen S, Liu H, Mallet RT, Shi X. Enhanced cerebral perfusion during brief exposures to cyclic intermittent hypoxemia. J Appl Physiol (1985). 2017;123(6):1689–97.
- Loeppky JA, Icenogle M, Scotto P, Robergs R, Hinghofer-Szalkay H, Roach RC. Ventilation during simulated altitude, normobaric hypoxia and normoxic hypobaria. Respir Physiol. 1997;107(3):231–9.
- Serebrovskaya T, Kolesnikova E, Karaban I, Mishunina T, Mankovskaya I, Safronova O. Intermittent hypoxia (IH) improves hypoxic ventilatory sensitivity and blood dopamine (DA) in patients with Parkinson's disease (PD). Pathophysiology. 1998;5:234–234.
- Bowen DJ, Kreuter M, Spring B, Cofta-Woerpel L, Linnan L, Weiner D, Bakken S, Kaplan CP, Squiers L, Fabrizio C, et al. How we design feasibility studies. Am J Prev Med. 2009;36(5):452–7.
- Vohra S, Shamseer L, Sampson M, Bukutu C, Schmid CH, Tate R, Nikles J, Zucker DR, Kravitz R, Guyatt G, et al. CONSORT extension for reporting N-of-1 trials (CENT) 2015 Statement. BMJ. 2015;350: h1738.
- Porcino AJ, Shamseer L, Chan AW, Kravitz RL, Orkin A, Punja S, Ravaud P, Schmid CH, Vohra S. SPIRIT extension and elaboration for n-of-1 trials: SPENT 2019 checklist. BMJ. 2020;368: m122.
- Tiffin J, Asher EJ. The Purdue Pegboard: norms and studies of reliability and validity. J Appl Psychol. 1948;32(3):234–47.
- 53. Podsiadlo D, Richardson S. The timed "Up & Go": A test of basic functional mobility for frail elderly persons. J Am Geriatr Soc. 1991;39(2):142–8.
- 54. Franchignoni F, Horak F, Godi M, Nardone A, Giordano A. Using psychometric techniques to improve the Balance Evaluation System's Test: the mini-BESTest. Journal of rehabilitation medicine: official journal of the UEMS European Board of Physical and Rehabilitation Medicine. 2010;42(4):323.
- Sandhu MS, Perez MA, Oudega M, Mitchell GS, Rymer WZ. Efficacy and time course of acute intermittent hypoxia effects in the upper extremities of people with cervical spinal cord injury. Exp Neurol. 2021;342: 113722.

- Guyatt G, Sackett D, Taylor DW, Chong J, Roberts R, Pugsley S. Determining optimal therapy–randomized trials in individual patients. N Engl J Med. 1986;314(14):889–92.
- Jenkinson C, Fitzpatrick R, Peto V, Greenhall R, Hyman N. The Parkinson's Disease Questionnaire (PDQ-39): development and validation of a Parkinson's disease summary index score. Age Ageing. 1997;26(5):353–7.
- Lee PH, Macfarlane DJ, Lam TH, Stewart SM. Validity of the international physical activity questionnaire short form (IPAQ-SF): A systematic review. Int J Behav Nutr Phys Act. 2011;8(1):115.
- Zhang SXL, Gozal D, Sachleben LR, Rane M, Klein JB, Gozal E: Hypoxia induces an autocrine-paracrine survival pathway via platelet-derived growth factor (PDGF)-B/PDGF-beta receptor/phosphatidylinositol 3-kinase/Akt signaling in RN46A neuronal cells. Faseb J 2003, 17(10):1709-+.
- Halder SK, Milner R. Mild hypoxia triggers transient blood-brain barrier disruption: a fundamental protective role for microglia. Acta Neuropathol Commun. 2020;8(1):175.
- Al Ahmad A, Gassmann M, Ogunshola OO. Involvement of oxidative stress in hypoxia-induced blood-brain barrier breakdown. Microvasc Res. 2012;84(2):222–5.
- Tsunekawa S, Ohi Y, Ishii Y, Sasahara M, Haji A. Hypoxic ventilatory response in platelet-derived growth factor receptor-beta-knockout mice. J Pharmacol Sci. 2009;110(3):270–5.
- Shen J, Xu G, Zhu R, Yuan J, Ishii Y, Hamashima T, Matsushima T, Yamamoto S, Takatsuru Y, Nabekura J, et al. PDGFR-beta restores blood-brain barrier functions in a mouse model of focal cerebral ischemia. J Cereb Blood Flow Metab. 2019;39(8):1501–15.
- Sagare AP, Sweeney MD, Makshanoff J, Zlokovic BV. Shedding of soluble platelet-derived growth factor receptor-beta from human brain pericytes. Neurosci Lett. 2015;607:97–101.
- Gaceb A, Ozen I, Padel T, Barbariga M, Paul G. Pericytes secrete proregenerative molecules in response to platelet-derived growth factor-BB. J Cereb Blood Flow Metab. 2018;38(1):45–57.
- 66. Padel T, Ozen I, Boix J, Barbariga M, Gaceb A, Roth M, Paul G. Platelet-derived growth factor-BB has neurorestorative effects and modulates the pericyte response in a partial 6-hydroxydopamine lesion mouse model of Parkinson's disease. Neurobiol Dis. 2016;94:95–105.
- 67. Coste O, Beers P, Bogdan A, Charbuy H, Touitou Y. Hypoxic alteration of cortisol circadian rhythm in man after simulation of a long duration flight. Steroids. 2005;70:803–10.
- Rey F, Balsari A, Giallongo T, Ottolenghi S, Di Giulio AM, Samaja M, Carelli S. Erythropoietin as a Neuroprotective Molecule: An Overview of Its Therapeutic Potential in Neurodegenerative Diseases. ASN Neuro. 2019;11:1759091419871420.
- Signore AP, Weng Z, Hastings T, Van Laar AD, Liang Q, Lee YJ, Chen J. Erythropoietin protects against 6-hydroxydopamine-induced dopaminergic cell death. J Neurochem. 2006;96(2):428–43.
- Jang W, Kim HJ, Li H, Jo KD, Lee MK, Yang HO. The Neuroprotective Effect of Erythropoietin on Rotenone-Induced Neurotoxicity in SH-SY5Y Cells Through the Induction of Autophagy. Mol Neurobiol. 2016;53(6):3812–21.
- McGlothlin AE, Viele K. Bayesian Hierarchical Models. JAMA. 2018;320(22):2365–6.
- Stunnenberg BC, Woertman W, Raaphorst J, Statland JM, Griggs RC, Timmermans J, Saris CG, Schouwenberg BJ, Groenewoud HM, Stegeman DF, et al. Combined N-of-1 trials to investigate mexiletine in non-dystrophic myotonia using a Bayesian approach; study rationale and protocol. BMC Neurol. 2015;15:43.
- Kravitz R, Duan N, Eslick I, Gabler N, Kaplan H, Larson E: Design and implementation of N-of-1 trials: a user's guide. Agency for healthcare research and quality, US Department of Health and Human Services 2014.
- Plummer M: JAGS: A program for analysis of Bayesian graphical models using Gibbs sampling. In.
- 75. Team RC: R: A language and environment for statistical computing. 2013.
- 76. Plummer M, Stukalov A, Denwood M, Plummer MM: Package 'rjags'. update 2019, 1.
- Christiansen L, Urbin MA, Mitchell GS, Perez MA: Acute intermittent hypoxia enhances corticospinal synaptic plasticity in humans. Elife 2018, 7.
- Hayes HB, Jayaraman A, Herrmann M, Mitchell GS, Rymer WZ, Trumbower RD. Daily intermittent hypoxia enhances walking after chronic spinal cord injury: a randomized trial. Neurology. 2014;82(2):104–13.

- Trumbower RD, Hayes HB, Mitchell GS, Wolf SL, Stahl VA. Effects of acute intermittent hypoxia on hand use after spinal cord trauma: A preliminary study. Neurology. 2017;89(18):1904–7.
- Trumbower RD, Jayaraman A, Mitchell GS, Rymer WZ. Exposure to acute intermittent hypoxia augments somatic motor function in humans with incomplete spinal cord injury. Neurorehabil Neural Repair. 2012;26(2):163–72.
- 81. Vivodtzev I, Tan AQ, Hermann M, Jayaraman A, Stahl V, Rymer WZ, Mitchell GS, Hayes HB, Trumbower RD: Mild to moderate sleep apnea is linked to hypoxia-induced motor recovery after spinal cord injury. Am J Respir Crit Care Med 2020.
- 82. Duennwald T, Bernardi L, Gordin D, Sandelin A, Syreeni A, Fogarty C, Kyto JP, Gatterer H, Lehto M, Horkko S, et al. Effects of a single bout of interval hypoxia on cardiorespiratory control in patients with type 1 diabetes. Diabetes. 2013;62(12):4220–7.
- 83. Duennwald T, Gatterer H, Groop PH, Burtscher M, Bernardi L. Effects of a single bout of interval hypoxia on cardiorespiratory control and blood glucose in patients with type 2 diabetes. Diabetes Care. 2013;36(8):2183–9.
- Faulhaber M, Gatterer H, Haider T, Patterson C, Burtscher M. Intermittent hypoxia does not affect endurance performance at moderate altitude in well-trained athletes. J Sports Sci. 2010;28(5):513–9.
- 85. Gatterer H, Menz V, Burtscher M: Acute Moderate Hypoxia Reduces One-Legged Cycling Performance Despite Compensatory Increase in Peak Cardiac Output: A Pilot Study. Int J Environ Res Public Health 2021, 18(7).
- Bayer U, Glazachev OS, Likar R, Burtscher M, Kofler W, Pinter G, Stettner H, Demschar S, Trummer B, Neuwersch S. Adaptation to intermittent hypoxia-hyperoxia improves cognitive performance and exercise tolerance in elderly. Adv Gerontol. 2017;30(2):255–61.
- 87. Gatterer H, Haacke S, Burtscher M, Faulhaber M, Melmer A, Ebenbichler C, Strohl KP, Hogel J, Netzer NC. Normobaric Intermittent Hypoxia over 8 Months Does Not Reduce Body Weight and Metabolic Risk Factors—a Randomized, Single Blind, Placebo-Controlled Study in Normobaric Hypoxia and Normobaric Sham Hypoxia. Obes Facts. 2015;8(3):200–9.
- Wille M, Gatterer H, Mairer K, Philippe M, Schwarzenbacher H, Faulhaber M, Burtscher M. Short-term intermittent hypoxia reduces the severity of acute mountain sickness. Scand J Med Sci Sports. 2012;22(5):e79-85.
- Bernardi L, Passino C, Serebrovskaya Z, Serebrovskaya T, Appenzeller O. Respiratory and cardiovascular adaptations to progressive hypoxia; effect of interval hypoxic training. Eur Heart J. 2001;22(10):879–86.
- Brugniaux JV, Pialoux V, Foster GE, Duggan CT, Eliasziw M, Hanly PJ, Poulin MJ. Effects of intermittent hypoxia on erythropoietin, soluble erythropoietin receptor and ventilation in humans. Eur Respir J. 2011;37(4):880–7.
- 91. Kolesnikova EE, Serebrovskaya TV. Parkinson's disease and intermittent hypoxia training. Hauppauge: Nova Science Publishers, Inc; 2009.
- van de Wetering-van Dongen VA, Kalf JG, van der Wees PJ, Bloem BR, Nijkrake MJ. The Effects of Respiratory Training in Parkinson's Disease: A Systematic Review. J Parkinsons Dis. 2020;10(4):1315–33.

Publisher's Note

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.