

RESEARCH ARTICLE

Stochastic vagus nerve stimulation affects acute heart rate dynamics in rats

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Abstract

Vagus nerve stimulation (VNS) is an approved therapy for treatment of epilepsy and depression. While also shown to be promising in several preclinical and clinical studies to treat cardiovascular diseases, optimal therapeutic stimulation paradigms are still under investigation. Traditionally, parameters such as frequency, current, and duty cycle are used to adjust the efficacy of VNS therapy. This study explored the effect of novel stochastic VNS (S-VNS) on acute heart rate (HR) dynamics. The effect of S-VNS was evaluated in Sprague Dawley rats by comparing the acute HR and HR variability (HRV) responses to standard, periodic VNS (P-VNS) across different frequencies (FREQs, 10–30 Hz). Our results demonstrate that both S-VNS and P-VNS produced negative chronotropic effects in a FREQ-dependent manner with S-VNS inducing a significantly smaller drop in HR at 10 Hz and 20 Hz compared to P-VNS ($p < 0.05$). S-VNS demonstrated a FREQ-dependent drop in the SD1/SD2 ratio, a measure of HRV, which was absent in P-VNS, suggesting that S-VNS may acutely modulate the nonlinear relationship between short- and long-term HRV. In conclusion, S-VNS is a novel stimulation procedure that may provide different physiological outcomes from standard P-VNS, as indicated by our analysis of HR dynamics. Our study provides a rationale for further detailed investigations into the therapeutic potential of S-VNS as a novel neuromodulation technique.

Introduction

In recent years, there has been an emergence of interest in using neuromodulation techniques to enhance or suppress activity of the nervous system for the treatment of various pathological conditions, including but not limited to systemic inflammation, obesity, spinal cord injuries, and neurological disorders. One such therapy is vagus nerve stimulation (VNS), which was approved by the FDA as a clinical therapy for the treatment of refractory epilepsy in 1997 and for the treatment of medication-resistant depression in 2005 [1]. As a result, VNS has become one of several non-pharmacological treatment options to control epileptic seizures worldwide.

The benefits of using VNS to treat cardiovascular diseases, including cardiac arrhythmias, heart failure (HF), hypertension, and myocardial ischemia, have been extensively explored in

research materials (vagus nerve electrodes), but did not have any additional role in the study design, data collection and analysis, and decision to publish. The specific roles of these authors are articulated in the 'Author Contributions' section.

Competing interests: Drs. Imad Libbus, Bruce H. KenKnight, and Elena G Tolkacheva are employees of LivaNova, PLC (formerly Cyberonics, Inc.). We would like to report this work was completed with resources provided by LivaNova, PLC. This does not alter our adherence to PLOS ONE policies on sharing data and materials. The other authors (SWL, KK, and EMA) have no competing interests that might have influenced the performance or presentation of the work described in this manuscript. There are no patents, products in development, or marketed products to declare.

past decades, with several studies suggesting the therapy to be both safe and potentially effective as a long-term cardiac therapy [2–5]. However, randomized clinical trials evaluating VNS in chronic HF patients have shown discrepant findings [6]. While the INOVATE-HF and NECTAR-HF trials failed to demonstrate significant improvements, both ANTHEM-HF and the subsequent extension study (ENCORE) yielded favorable and encouraging results including reduced HF symptoms and improved ventricular function [5,7–9]. In addition to different inclusion and exclusion criteria of HF patients, the different choice of stimulation parameters is another attributable factor for these mixed results.

Traditionally, the therapeutic intensity of VNS is selected to demonstrate engagement of the autonomic nervous system (typically via observed changes in heart rate (HR)) [10]. The stimulation parameters used to treat cardiac diseases, however, are highly variable depending on different models and studies [2,4,11,12]. Hence, there is an ongoing search for the optimal settings that will maximize the benefits of VNS. This can include the traditional approach of varying the stimulation frequency, current/voltage, and pulse width [12,13], as well as developing novel stimulation paradigms, such as the recently developed "microburst" VNS [10]. Typically, chronic periodic VNS (P-VNS) is applied, and its stimulation profile consists of administering periodic, repetitive electrical pulses with the level of stimulation (i.e. intensity) determined by a selected frequency, pulse duration, pulse amplitude, and duty cycle which all can be adjusted via radiofrequency communication link using a proprietary programming computer. In this study, we propose and evaluate an innovative stimulation technique: stochastic VNS (S-VNS). The version of S-VNS used in this study delivers electrical impulses to the nerve according to a Gaussian distribution of stimulation frequencies, instead of a constant frequency used in standard, P-VNS.

It is known that the cardiac beat-to-beat time intervals vary stochastically due to the inherent variability in HR (HRV). This variability can be attributed to many physiological factors including the influence of circadian rhythms, respiratory rhythms, temperature regulations, and others [14]. In fact, it has been extensively reported that both increased HR and loss of HRV are associated with increased mortality in HF patients and are candidate markers for patients at risk of sudden cardiac death [15,16]. An abundance of literature supports that acute P-VNS effectively decreases HR and improves HRV via the extensive innervation of the vagus nerve into the sinoatrial (SA) and atrioventricular (AV) nodes [17,18]. Even though the exact mechanisms responsible for the coupling between VNS and HR and HRV remains unclear, it is postulated that it is mediated through muscarinic receptor activation via the neurotransmitter acetylcholine (ACh) released at parasympathetic nerve terminals [19]. However, while P-VNS may effectively modulate HR dynamics, there may exist a 'physiological compensation mechanism' in which the vagus nerve and the central-peripheral nervous system can adapt to the periodicity of the stimulation, a phenomena which can be similar to pharmacological tolerance [20–22]. We propose that S-VNS might help to avoid this potential adaptation, and therefore, the goal of our study is to evaluate the influence of acute S-VNS therapy on instantaneous HR and HRV by using *in-vivo* rat hearts, and compare it to the standard P-VNS.

Materials and methods

Sprague-Dawley rats (n = 8, 250–400 g, Charles River Laboratories, Wilmington, MA) were used in this study. All experiments were performed in accordance with the guidelines set forth by the National Institutes of Health Guide for the Care and Use of Laboratory Animals and were approved by the University of Minnesota Institutional Animal Care and Use Committee (IACUC).

Vagus nerve bipolar cuff electrode implantation

Rats were anesthetized with isoflurane (5% for induction and 1.5% for maintenance). After hair shaving and skin cleaning, aseptic technique was used to make a ventral midline incision in the neck, and the skin and muscles were retracted. After identifying and isolating the right cervical vagus nerve and common carotid artery bundle, a custom 1.5 mm diameter helical lead bipolar cuff electrode (Cyberonics, USA) was implanted around the bundle. The right vagus nerve was chosen for this study because it primarily innervates the atria and the sino-atrial node, and thus, right-side stimulation may induce significant change in cardiac rhythm [23]. The electrode was then connected to a battery-driven, constant current stimulator (Stimulus Isolator, model A385, World Precision Instruments, USA). At the end of the study, the anesthetized rats were euthanized by thoracotomy followed by heart explantation.

VNS protocols and study design

Both P-VNS and S-VNS were continuously delivered for 2 minutes at a pulse duration of 500 μ sec and an output current of 1.0 mA [2–4]. The same number of stimuli was used for both P-VNS and S-VNS.

P-VNS was delivered at different frequencies (FREQ) of 10, 20, or 30 Hz that were kept constant throughout the 2 minutes of stimulation.

S-VNS was used with stochasticity (STOCH) of 10% or 20%. Three values of mean FREQ (i.e. 10, 20, and 30 Hz) were used similar to P-VNS, and STOCH (using a Gaussian distribution) was incorporated as:

$$\text{FREQ}_{\text{STOCH}} = \frac{\text{FREQ}}{1 \pm [\text{FREQ} * \delta(\text{STOCH})]}$$

where $\delta(\text{STOCH})$ is a random number with a mean of zero and a standard deviation of STOCH/FREQ. A custom LabView program was used to generate $\delta(\text{STOCH})$ for each pulse.

The detailed study design is shown in [Fig 1](#). Briefly, the vagus nerve was first stimulated at FREQ = 20 Hz, starting with P-VNS (P-VNS #1) followed by S-VNS with STOCH = 20% after a 1-minute stabilization interval. After completion, a similar run was conducted for a FREQ = 30 Hz which was followed by a run at FREQ = 10 Hz. After a 10-minute stabilization interval, both P-VNS (P-VNS #2) and S-VNS with STOCH = 10% runs were conducted in the FREQ order of 20, 30 and 10 Hz. Each protocol for a certain FREQ consisted of the following stimulation:

1. 2 minutes of baseline recording before VNS (**PRE**)
2. 2 minutes of continuous VNS (**ON**)
3. 2 minutes of recovery after VNS (**POST**)

Data analysis for in-vivo ECG recordings

Electrocardiogram (ECG) was recorded from each anesthetized rat (IX-ECG-12, iWorx, USA). ECG electrodes were placed subcutaneously into the limbs. For both P-VNS and S-VNS, **PRE**, **ON**, and **POST** ECG data were recorded for 2 minutes. The *in-vivo* ECG recordings were used to quantify changes in the HR and HRV throughout the study using Kubios HRV 2.0 software [24]. Noisy data segments and ectopic beats were excluded and steady-state data after adjustment of the HR to acute VNS were used for the analysis. Hence, all runs of **PRE**, **ON**, and **POST** for all animals consisted of ~1-minute and 40 seconds interval for analysis.

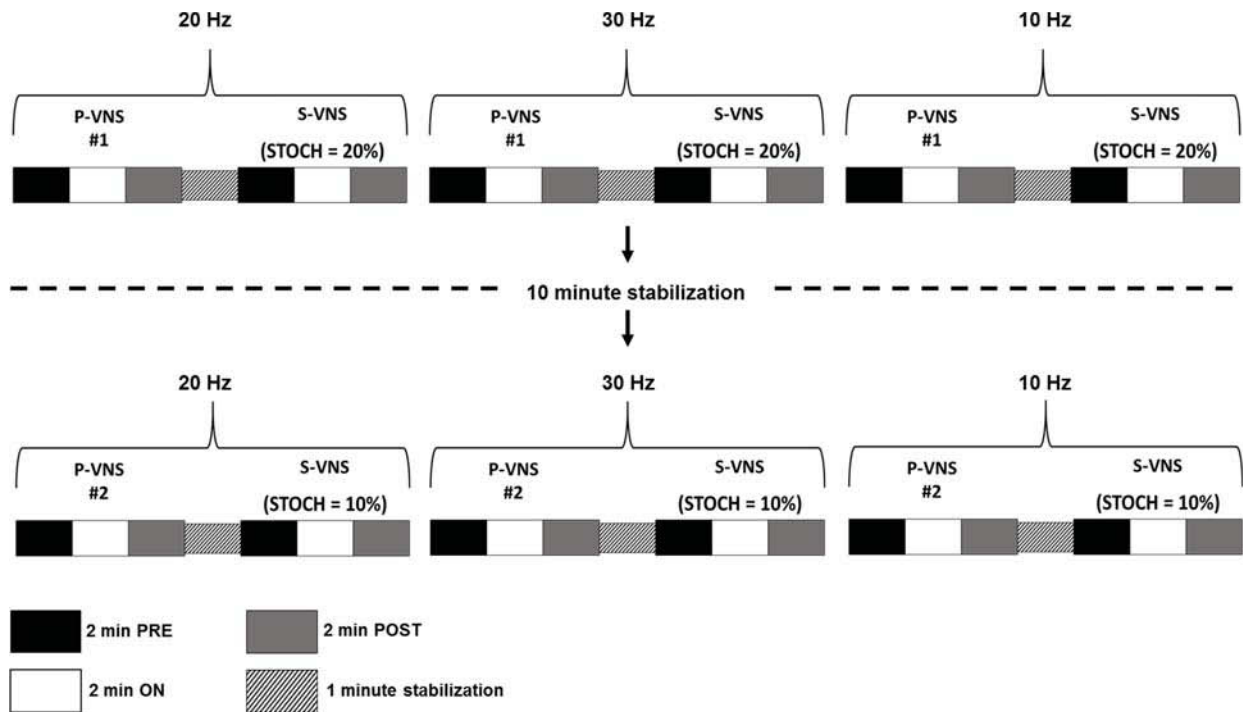


Fig 1. Detailed schematic of the experimental VNS protocol. Standard P-VNS or S-VNS with different degree of stochasticity (STOCH, 10% and 20%) was administered across different frequencies (FREQ, 20, 30, and 10 Hz) with stabilization times between protocols and conditions. P-VNS, periodic vagus nerve stimulation; S-VNS, stochastic vagus nerve stimulation; STOCH, stochasticity; PRE, baseline recording; ON, stimulation; VNS, vagus nerve stimulation.

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To account for variations in baseline HR, the chronotropic effect of VNS was determined by calculating a relative change in HR, ΔHR_{oxy} , as the following:

$$\Delta HR_{ON} = \left(\frac{HR_{ON} - HR_{PRE}}{HR} \right) * 100\%$$

where HR_{ON} and HR_{PRE} are the mean HR during the VNS ON and PRE periods, respectively.

After the termination of VNS, we assessed the recovery of HR to HR_{PPE} baseline value

Therefore, the HR post-VNS ratio (HR_{POST_Ratio}) was calculated as

$$HR_{POST\ Ratio} = \left| \frac{\Delta HR_{POST}}{\Delta HR_{ON}} \right|$$

and

$$\Delta HR_{POST} = \left(\frac{HR_{POST} - HR_{ON}}{HR_{ON}} \right) * 100\%$$

where HR_{POST} is the mean HR during the VNS **POST** period. A $HR_{POST\ Ratio} = 1$ means that the HR_{POST} returned to the baseline HR_{PRE} value. Whereas, if $HR_{POST\ Ratio} < 1$ (or > 1), then the HR_{POST} was less (or greater) than the baseline HR_{PRE} value.

To quantify HRV, Poincaré plots were used to provide a two-dimensional representation of the correlation between successive RR intervals, as described previously [24]. Briefly, the spatial distribution of all points was fitted to an ellipse with the center coinciding with the average

RR interval. The two dispersions (perpendicular (SD1) and parallel (SD2) to the line of identity $y = x$) and their ratio, SD1/SD2, were then calculated. Previously, it was established that SD1 is a measure of the short-term HRV, while SD2 provides an assessment of both the short- and long-term HRV [24,25].

Statistical analysis

All data are presented as means \pm standard error. Statistical analyses were performed using Origin 9.1 software (Northampton, MA). Statistical comparisons were performed using a one-way ANOVA with repeated measures: 1) FREQ-dependent effect of P-VNS and S-VNS, 2) between S-VNS and P-VNS; 3) between the two P-VNS protocols, and 4) FREQ-dependent effect on SD1/SD2 for **PRE**, **ON**, and **POST**. A one sample *t*-test was performed to assess if the mean of the $HR_{POST\ Ratio}$ is equal to one. Values of $P < 0.05$ were considered to be statistically significant. If appropriate, post hoc comparisons with Tukey's test were performed, with *P*-values corrected for repeated comparisons.

Results

[Fig 2](#) shows typical examples of ECG traces during P-VNS (Panel A) and S-VNS for STOCH = 10% and 20% (Panels B and C, respectively). Note the presence of small spikes on the ECG traces during the VNS **ON** periods corresponding to VNS stimulation artifacts, and the VNS-evoked decrease in HR. Furthermore, note the periodic, repetitive pattern of the VNS artifacts during P-VNS, and the absence of periodicity of the artifacts during S-VNS.

Frequency-dependent chronotropic effects of P-VNS and S-VNS

The FREQ-dependent chronotropic effects of P-VNS and S-VNS are shown in [Fig 3](#). Note that the reported mean and SEM of the P-VNS protocols were averages of the P-VNS protocols (i.e., mean of the average of P-VNS #1 and P-VNS #2 protocols, $n = 8$). Relative changes in HR, ΔHR_{ON} , in response to VNS across FREQ ranging from 10 to 30 Hz are shown for P-VNS (Panel A) and S-VNS (Panels B, C). For both P-VNS and S-VNS (both STOCH = 10% and 20%), no significant reduction in HR was observed as FREQ was increased from 10 Hz to 20 Hz. However, further increase of FREQ to 30 Hz produced a significant drop in HR.

It has been shown in several species such as dogs [26] and rats [27] that VNS FREQ is linearly related to heart period, while the relation to HR is nonlinear. To account for this possibility, we also calculated the relative changes in heart period for both VNS protocols. [Fig 3](#) (Panels D-F) indicate that FREQ-dependent chronotropic effect for heart period is similar to the one for HR (compare to [Fig 3](#), Panels A-C).

Effects of STOCH on HR and HRV

The relative changes in HR, ΔHR_{ON} , in response to 10% (Panel A) and 20% S-VNS (Panel B), respectively, compared to its immediate preceding P-VNS at different FREQs are shown in [Fig 4](#). Overall, S-VNS had significantly smaller negative chronotropic effect relative to P-VNS at FREQ = 10 and 20 Hz. At 30 Hz, however, this reduced effect was eliminated as all the protocols exhibited similar relative drop in HR. Moreover, note that both P-VNS runs, P-VNS #1 and P-VNS #2, exhibited similar drop in HR at different FREQs, as indicated in Panel C, suggesting the stability of the experimental conditions over time. To investigate whether the order of application of P-VNS and S-VNS protocols contribute to the effects shown in Panels A and B, 20% S-VNS was compared to its subsequent P-VNS #2 at different FREQs. [Fig 4D](#) demonstrates similar effect in the relative change in HR responses when compared to Panels A and B,

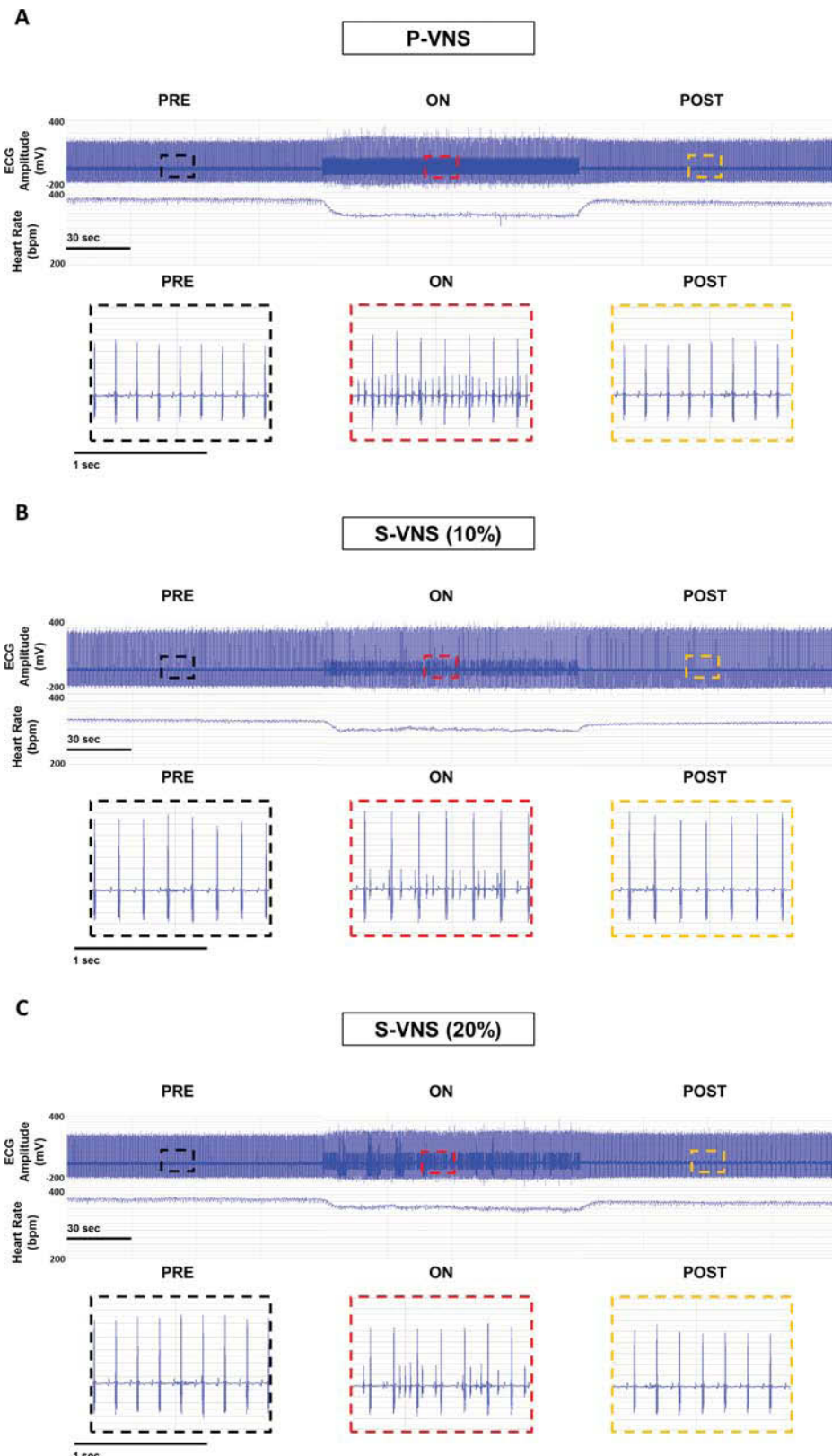


Fig 2. ECG recordings and corresponding HR responses. Representative segments of ECG recordings and corresponding heart rate (HR) response for an anesthetized rat for PRE, ON, and POST during (A) P-VNS, (B)

S-VNS (10%), and (C) S-VNS (20%) of the right cervical vagus nerve. Zoomed-in snapshots of **PRE** (black), **ON** (red), and **POST** (yellow) highlights VNS artifacts during stimulation. Here, VNS was continuously delivered at 20 Hz, 500 μ s pulse width, and 1.0 mA for 2 minutes.

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indicating that the effect is attributed to the S-VNS protocol itself, but not the different permutation of the two stimulation protocols.

Representative Poincaré plots of both P-VNS (Panel A) and S-VNS (Panel B) and their quantitative assessment, SD1/SD2 (Panel C) are shown in Fig 5. While our results suggest that S-VNS did not significantly affect SD1/SD2 when compared to P-VNS (as indicated in Fig 5C), we observed a FREQ-dependent effect. As shown in Fig 5C, for P-VNS protocol, the SD1/SD2 values were similar for all FREQs for **PRE**, **ON**, and **POST** periods. Similarly, no significant difference in SD1/SD2 values was observed for S-VNS for **PRE** and **POST** periods. However, for VNS **ON** period, S-VNS showed significant reduction in SD1/SD2 values between FREQ = 10 Hz and 30 Hz. In addition to Poincaré analysis, we also evaluated HRV as a ratio of the standard deviation of RR interval (SDRR) to mean RR interval (mean RR), as described previously [28]. From this approach, no significant difference was observed for all FREQs for **PRE**, **ON**, and **POST** periods for all VNS protocols (S1 Fig).

To evaluate the recovery of HR to baseline values after VNS application, $HR_{POST\ Ratio}$ was calculated across different FREQs. Our results from Panels B-D in Fig 6 indicate that for FREQ = 30 Hz, the HR could recover back to its **PRE** value in S-VNS protocols. However, for FREQ = 10 or 20 Hz, the $HR_{POST\ Ratio}$ was less than 1, reflecting a persistent reduction in HR after the end of the VNS cycle.

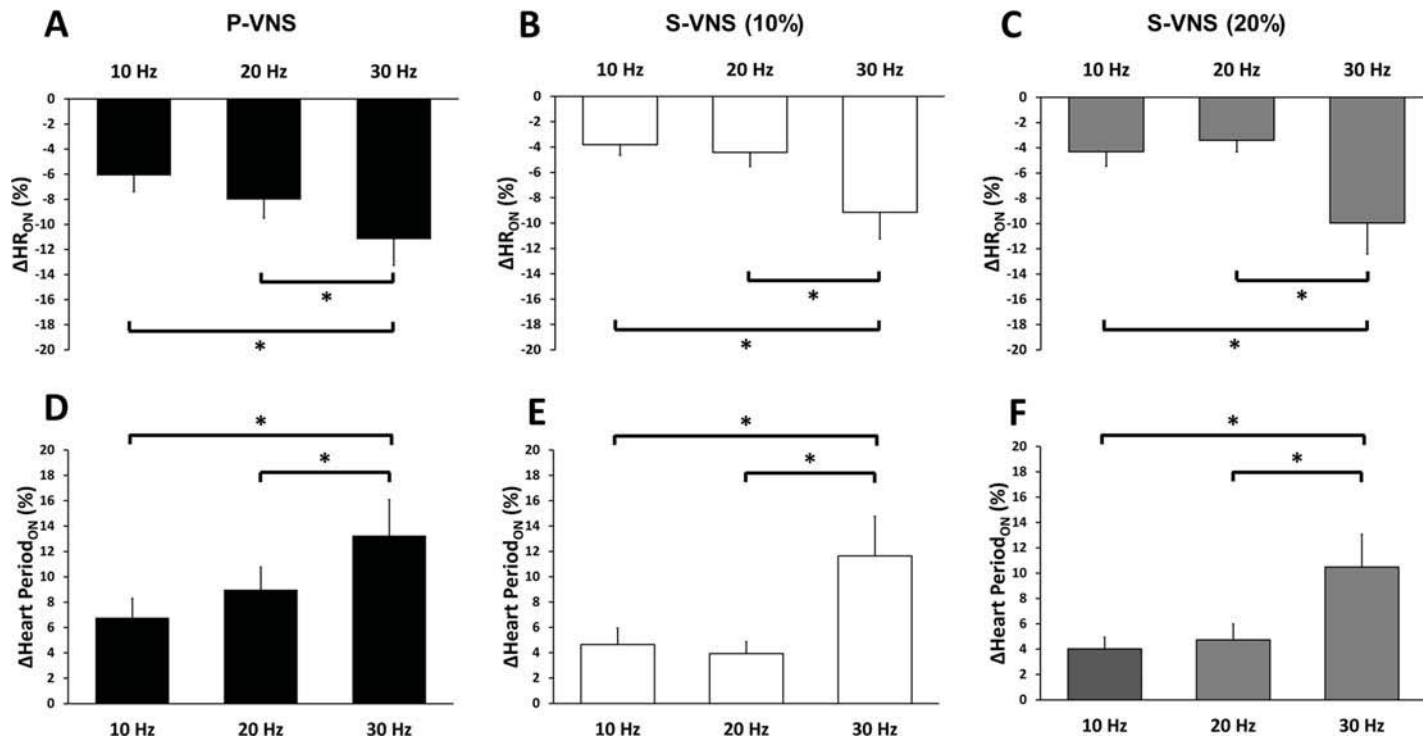


Fig 3. Effects of VNS FREQ on HR and heart period. Mean percent drop in HR for (A) P-VNS, (B) S-VNS (10%), and (C) S-VNS (20%) at different FREQ. Mean percent drop in Heart Period for (D) P-VNS, (E) S-VNS (10%), and (F) S-VNS (20%) at different FREQ. Note data reported here for P-VNS is the mean and SEM of the average of P-VNS #1 and P-VNS #2 protocols ($n = 8$). (* $p < 0.05$).

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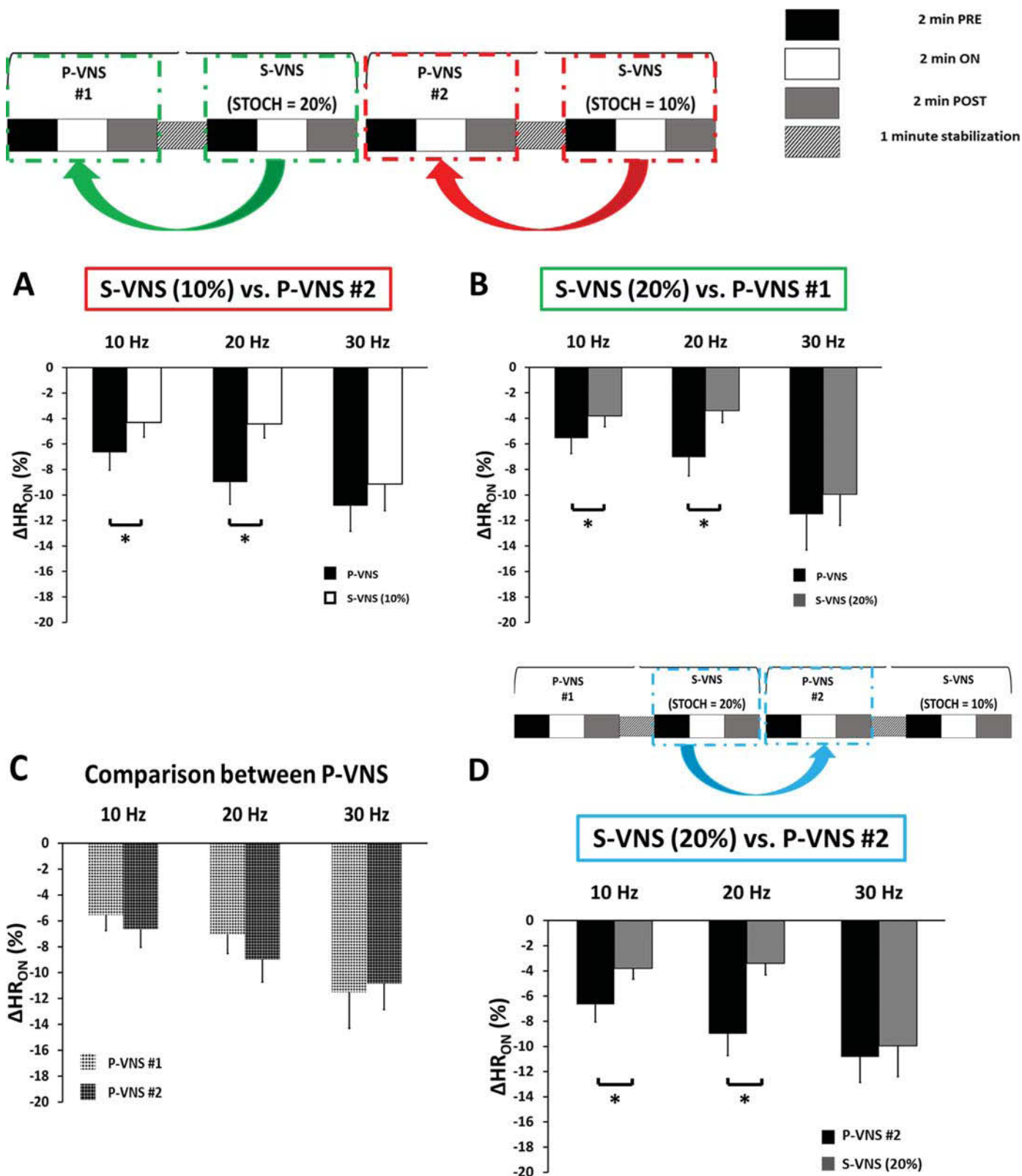


Fig 4. Effects of STOCH on the chronotropic effects of VNS. Mean percent drop in HR for (A) S-VNS (10%) and (B) S-VNS (20%) being compared to its immediate preceding P-VNS protocol at different FREQ. (C) Comparison of mean relative drop in HR between P-VNS protocols and (D) S-VNS (20%) and its subsequent P-VNS protocol at different FREQ. (*p < 0.05).

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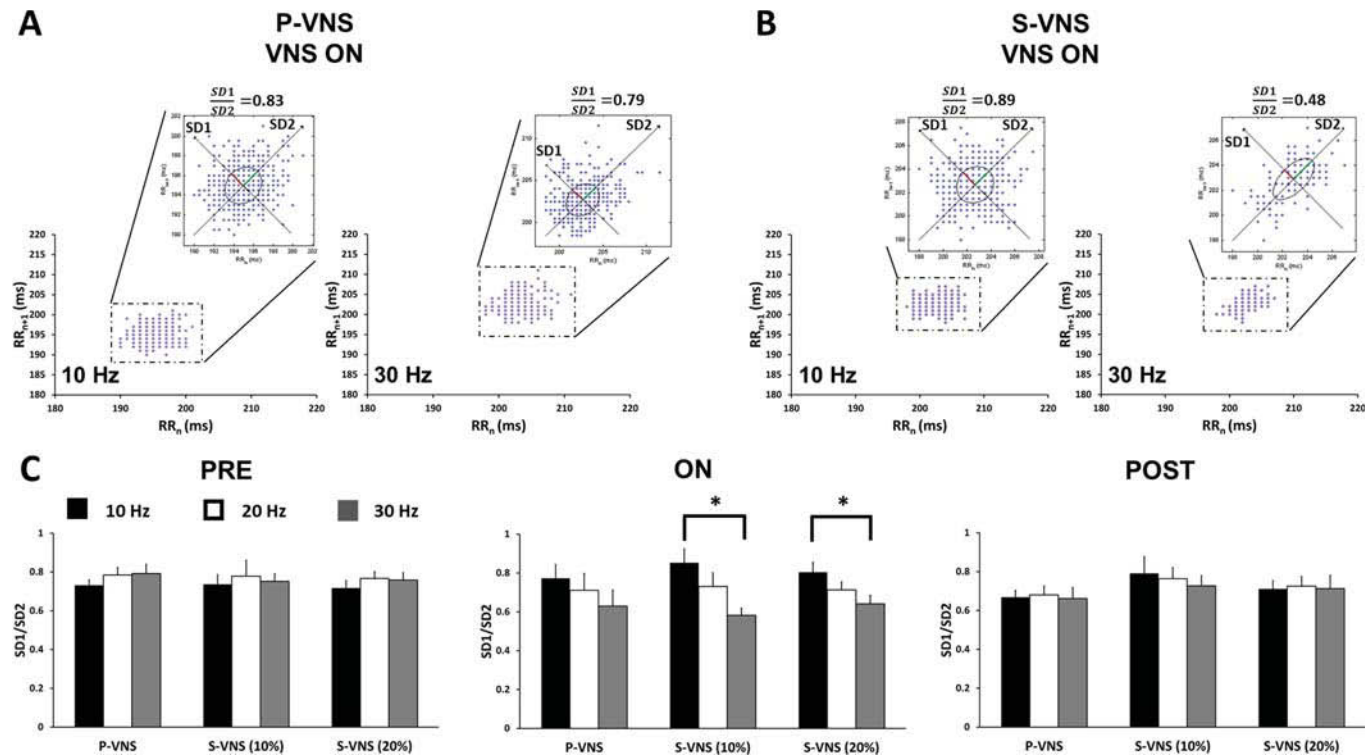


Fig 5. Effects of VNS on HRV using Poincaré analysis. Representative Poincaré plots of (A) P-VNS and (B) S-VNS (10%) during VNS stimulation (ON) at 10 Hz and 30 Hz demonstrating the elliptical fitting of the beat-distribution cloud and the standard deviation of short-term (SD1) and long-term (SD2) variability. (C) Mean SD1/SD2 ratio for PRE, ON and POST across different FREQ. (*p < 0.05).

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Discussion

The aim of the present study was to develop and test a new autonomic modulation stimulation protocol using stochastic stimulation (S-VNS), and compare it with traditional stimulation waveforms (P-VNS). Specifically, the evoked chronotropic response to P-VNS (delivered at a constant FREQ of 10, 20, or 30 Hz) were evaluated, and compared to S-VNS where stochastic variability was imposed in delivered FREQs with constraints for 10% or 20% around these three FREQs. The major findings of this study are as follows: (1) HR is reduced by both S-VNS and P-VNS in a FREQ-dependent manner. However, the negative chronotropic effect of S-VNS at 10 Hz and 20 Hz is significantly smaller than that of P-VNS and (2) S-VNS may acutely modulate the nonlinear relationship between short- and long-term HRV differently than P-VNS, as indicated by SD1/SD2 measurements. Overall, the results of our study suggest that acute S-VNS affects the dynamic changes in *in-vivo* rat hearts, specifically HR and HRV in a different manner, when compared to traditional P-VNS. Hence, our study further supports the notion that different modes of stimulation provides different outcomes as measured by means of change in HR. We also highly suggest further investigations into characterizing the effects and elucidating the mechanisms of S-VNS as a novel neuromodulation technique under different diseased conditions.

The need to optimize VNS parameters

It has been well-established that an important feature of the adverse pathophysiological changes seen in HF is autonomic imbalance, characterized by sympathetic hyperactivation and parasympathetic withdrawal [29]. VNS has emerged as a promising therapy to normalize

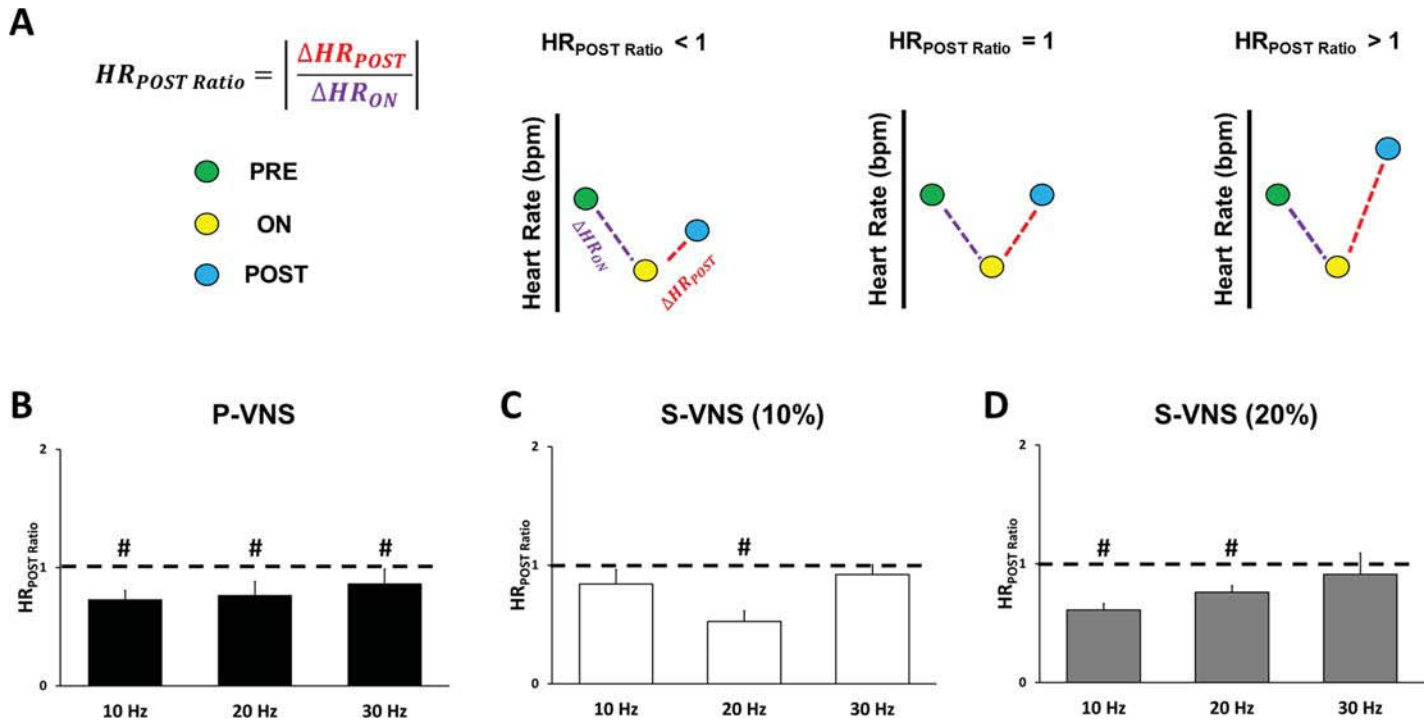


Fig 6. Effects of VNS FREQ on HR recovery. (A) Schematic representation of different scenarios of HR recovery based on the value of $HR_{POST\ Ratio}$. Mean $HR_{POST\ Ratio}$ values for (B) P-VNS, (C) S-VNS (10%), and (D) S-VNS (20%) at different FREQ. # $p < 0.05$ compared to a mean theoretical value of 1.

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parasympathetic activation of cardiac control and restores normal levels of autonomic regulatory function. In fact, several animal studies suggest VNS exhibits both anti-arrhythmic and anti-inflammatory effects and attenuates myocardial damage and cardiac myocyte apoptosis [2,4,11]. Based on the encouraging results of the preclinical studies, several clinical VNS studies have been conducted. Both ANTHEM-HF and the subsequent extension study (ENCORE) yielded favorable and encouraging results as patients who received VNS experienced significant improvements in left ventricular (LV) structure and function, decreased HF symptom expression and decreased T-wave alternans, a measure of electrical instability [5,7]. However, other randomized clinical trials, such as the INOVATE-HF and NECTAR-HF, failed to demonstrate significant improvements in clinical outcomes owing to imperfect inclusion/exclusion criteria and sub-therapeutic VNS dosing in significant (~30–50%) of the enrolled population [8,9].

Yet despite these and other studies, there still does not exist a universally accepted published prospective stimulation paradigm, which further highlights the importance and complexity of parameter optimization. Hence, to further optimize and improve VNS efficacy, this study introduced S-VNS, a novel technique that stimulates the vagus nerve stochastically since it is clinically well-established that healthy hearts have been associated with high HRV [14].

Effects of stochasticity on HR and HRV

Our study demonstrated a FREQ-dependent reduction in HR as a result of VNS, which was previously shown in several studies asserting that increasing the stimulation FREQ alters the rate of propagation of action potential through the vagus nerve fibers [13]. However, we also observed that the drop in HR is significantly smaller for S-VNS as compared to P-VNS, even

though mean FREQ was similar for both techniques. One of the possible reasons for this is that although the same number of stimuli were applied during both stimulation techniques, stimuli during S-VNS were distributed more randomly. In P-VNS, there was a consistent number of stimuli being delivered between RR intervals, as highlighted in [Fig 2](#).

To determine the residual functional effects occurring after cessation of VNS, we calculated $HR_{POST\ Ratio}$. Our results show that the post HR of both S-VNS and P-VNS at certain FREQs did not return to baseline. This could be attributed to the fact that we applied continuous stimulation which is much longer than the typical intermittent VNS (e.g. duty cycle of 10% or 20%). Therefore, while other studies report a recovery to baseline at end of their intermittent stimulation [7,13], continuous stimulation may require more than the collected 2 minutes **POST** for the HR to return to baseline. Another possible explanation is that the cardiac effects of VNS may persist beyond the end of stimulation, and exhibit a “memory effect,” a phenomenon similar to cardiac memory. Cardiac memory is an established property of paced cardiac myocytes to reflect the influence of pacing history and adapt and respond to novel stimuli. Several experimental studies and numerical simulations have shown that memory plays a role in modulating cardiac dynamics [30,31]. Indeed, in future studies, we will examine post-VNS recovery after a longer time delay.

In this study, we performed both linear and nonlinear approaches to evaluate the effects of VNS on HRV. Poincaré plots of RR intervals is a nonlinear approach used to monitor autonomic changes by means of SD1 and SD2. Physiologically, SD1 is an index of instantaneous recording of the beat-to-beat variability and reflects primarily the parasympathetic input to the heart, while SD2 is an index of long term variability which reflects the overall variability influenced by both the parasympathetic and sympathetic contributions [25,32]. Hence, their ratio (SD1/SD2) provides a measurement between both short term and long-term variability in the RR intervals. Libbus and colleagues recently reported that patients in the ANTHEM-HF trial who received chronic, right-sided intermittent P-VNS expressed an increase in the SD1/SD2 ratio, which suggested that right-sided VNS increases the balance in favor of parasympathetic dominance over sympathetic activity [32]. While our acute P-VNS did not induce any changes in SD1/SD2 ratio in rats, our experiments demonstrated that acute continuous S-VNS significantly decreased this ratio, specifically between 10 Hz and 30 Hz. We also observed no difference in SD1 and SD2 values with increase in stimulation FREQ across **PRE**, **ON**, and **POST** for P-VNS and S-VNS ([S2 Fig](#)). This data suggests that FREQ and STOCH not only play a role in the chronotropic effects of VNS, but also in the engagement of the autonomic nervous system via relative difference in ACh release between the two VNS techniques. Nonetheless, it is worth noting that brief changes in HRV, i.e. during VNS **ON** phase, may or may not have any significant effects on clinical outcomes, and will need to be further investigated. While Poincaré maps suggest a change in autonomic activity, our ratio of SDRR to mean RR saw no significant HRV difference induced by both S-VNS and P-VNS ([S1 Fig](#)). Indeed, there are several metrics to gauge autonomic activity which include time-domain (e.g. SDNN, RMSSD, . . .), frequency-domain (e.g. LF, HF, LF/HF, . . .), and nonlinear (Poincaré, sample entropy, detrended fluctuation analysis, . . .) methods [33]. Although previous studies have shown a possible correlation between the different measures [33,34], an exact equivalence has not been reported. It is possible that Poincaré plots may detect abnormalities that are not as easily detectable with traditional time-domain measures, given that each parameter has different sensitivity to noise and other experimental conditions [35]. Hence, further investigations to fully uncover and understand the physiological relevance of these autonomic assessment methods are warranted in order to determine the efficacy of the different metrics in accurately quantifying autonomic activity”.

Mechanistic insights into acute S-VNS

In the present study, we observed a smaller drop in HR and a reduction in SD1/SD2 ratio in S-VNS compared to P-VNS. From a physiological standpoint, this was a surprising finding since applying S-VNS does not necessarily correlate to stochastic release of ACh and stimulation of the muscarinic receptors of the cardiomyocytes. Rather, the efferent vagal fibers which are responsible for carrying descending information from the brain to the heart do not directly synapse with cardiomyocytes but rather with the intrinsic cardiac nervous system (ICNS), which acts as a buffer to register and send commands to the cardiomyocytes [13]. The ICNS is comprised of an intricate network of ganglia (and its neurons) that can independently act or communicate with its counterparts in the intrathoracic extracardiac, brain stem, spinal cord, and cortex to regulate intracardiac reflexes [13]. Although detailed studies will need to be performed to quantify the level of ACh release during the two stimulation protocols, we hypothesize that the observed minimal HR drop can be due to the fact that the temporal summation of ACh release is lower during S-VNS when compared to P-VNS protocol. This could be due to the fact that the instantaneous, effective FREQ of stimulation during S-VNS protocol is different (either less or more) from the corresponding FREQ of P-VNS protocol.

Phase response curves (PRC) detail how a system with its own intrinsic properties evolve with an external perturbation. The heart is an example of such dynamical systems that when given an external stimulus can cause either phase mismatch or entrainment. It has been well established that the chronotropic response to vagal stimulation is phase-dependent [36]. In fact, depending on the timing, duration, and magnitude of the vagal pulse, repetitive vagal stimulation has been shown to be able to regulate the SA cells to beat at rates that are different from its intrinsic cycle length [37]. Therefore, another possible explanation for our findings is that incorporating STOCH in our VNS may affect the level of entrainment between the vagal stimuli and the cycle length of the SA cells. Future additional studies should be conducted to understand how S-VNS may affect the intrinsic PRC.

There has been an emerging interest in using closed-loop neuro-stimulation to achieve targeted control of cardiac parameters [38]. In fact, significant developments have been made in this area based on state transition models where an algorithm optimally selects the VNS parameters to regulate instantaneous HR synchronously with cardiac cycles [39]. We hypothesize that incorporating STOCH in closed-loop VNS may potentially lead to improved therapeutic efficacy. Physiologically, the heart rhythm is not deterministic and is known to exhibit beat-to-beat variability (HRV) caused by numerous physiological factors including the influence of circadian rhythms, respiratory rhythms, temperature regulations, and others [40]. In fact, it has been associated that low HRV is a predictor for pathological cardiac conditions [16]. Hence, we hypothesize that by recording real-time changes in HRV and then integrating this “inherent stochastic feedback” into the stimulation of the VNS through STOCH may lead to a more physiological stimulation of the vagus and increase the efficacy of VNS to treat cardiac disease. This will require a fully functional, implantable, and chronic version of the S-VNS system that has real-time monitoring capabilities.

Limitations

One of the limitations of our study is that we only investigated the differences between P-VNS and S-VNS with respect to HR and HRV, and used a healthy rat model. Furthermore, we only evaluated the acute effects. Beneficial effects have been observed when chronic P-VNS was applied in both healthy and diseased conditions [3,4,7,11]; therefore, additional chronic studies in both healthy and diseased conditions need to be performed in order to evaluate the full potential of S-VNS.

Moreover, our results suggest that S-VNS may enable some flexibility in nerve response by its ability to vary the stimulation intervals and providing a 'pseudo-intermittent' stimulation. Therefore, even though the same mean FREQ in both P-VNS and S-VNS were delivered, S-VNS had a smaller drop in HR. This could make S-VNS suitable for treatment of diseased conditions such as hypertension, where a controlled drop in blood pressure with a minimal change in HR are desired. This study suggests that incorporating STOCH may provide room for titration of other stimulation parameters (i.e. duty cycle and output current). In fact, in terms of treatment of brain disorders, non-periodic stimulation is currently being studied in the preclinical setting with promising results. Examples of stochastic neuromodulation techniques include but not limited to stochastic galvanic vestibular stimulation and transcranial random noise stimulation [41,42]. Therefore, there is the potential for S-VNS to induce changes in physiological parameters other than HR and HRV, making it a plausible candidate to improve current VNS treatment for epilepsy and depression, but this will require more in-depth investigation.

Furthermore, it is well accepted that the cervical vagus nerve is composed of both myelinated A and B fibers and unmyelinated C fibers [43], and our study does not address the question as to which fibers are indeed stimulated. Zagon et al have demonstrated that providing higher stimulation amplitudes, i.e. 200 μ A, activates both unmyelinated and myelinated fibers in anesthetized rats [44]. Hence, this suggests that our stimulation current of 1 mA is sufficient to activate the whole thin vagal trunk of our rats. However, the discharge frequencies between afferent and efferent fibers may be different, and thus, future studies are needed to determine whether the mechanisms of both P-VNS and S-VNS occurs primarily through the afferent or efferent pathways.

Another limitation of our study is that the sequence testing was not randomized. However, we did not monotonically increase our FREQ from 10 Hz to 30 Hz. Instead the sequence order was 20 Hz, 30 Hz, and 10 Hz, but nonetheless, there may be the potential of order effects. Finally, the ECG recordings from rats were obtained with the use of general anesthesia, which is known to affect the HR of animals as compared to conscious animal ECG telemetry recordings. In fact, it has been reported that high-dose isoflurane can suppress the central cardiac parasympathetic activity in rats [45,46]. As a result, we ensured that the dosage of isoflurane administered was low (~1.5%) and remained identical across all rats throughout the duration of the study; therefore, it should not affect the interpretation of our results.

Due to the long duration of the experimental procedures, we only tested the effect of STOCH values of 10% and 20% in this present manuscript. Additional experiments will need to be performed to explore the full range of STOCH values. Furthermore, we also applied fixed values of current amplitude and pulse width. It is well accepted that VNS would elicit an immediate change in HR correlating with VNS current amplitude and pulse width [7]. Hence, while a different magnitude drop in HR value may be observed, our interpretation of the effects of incorporating STOCH should not be affected. In addition, while we only observed a difference of 2% HR drop between P-VNS and S-VNS which means extrapolating this to human may make these changes negligible, it is worth noting that the average HR of healthy rats is roughly 300–350 bpm which is about five times faster than the average resting HR of humans [46,47]. Therefore, we propose these exploratory findings to first be confirmed in larger animal studies under both control and diseased conditions before being tested in humans. Nevertheless, we plan to further evaluate the robustness and efficacy of our proposed S-VNS in chronic future studies and on non-anesthetized murine models.

Conclusions

S-VNS induces dynamical changes in *in-vivo* rat hearts and produces minimal changes in HR compared to P-VNS. Hence, our study provides evidence that stochastic and periodic modes

of stimulation induce different cardiac responses, and we suggest further detailed investigations into understanding the full therapeutic potential of S-VNS under different diseased conditions.

Supporting information

S1 Fig. Time domain HRV analysis. HRV was calculated as a ratio of the standard deviation of RR interval (SDRR) to mean RR interval (mean RR), as described previously [28]. From this approach, no significant difference was observed for all FREQs for **PRE**, **ON**, and **POST** periods for all VNS protocols.

(TIF)

S2 Fig. Values of SD1 and SD2. Mean (A) SD1 and (B) SD2 values for **PRE**, **ON**, and **POST** across different FREQs. No significant difference in SD1 and SD2 values were observed with increase in stimulation FREQ across **PRE**, **ON**, and **POST** for P-VNS and S-VNS.

(TIF)

S3 Fig. Quantitative analysis of 1-minute stabilization interval. In regards to whether 1-minute stabilization intervals were sufficient to avoid carryover effects on **PRE**, we performed the following quantitative analysis. The timeline of HR change during entire duration of an experiment is shown in Panel A for one rat. Vertical dashed lines represent the 1-minute stabilization intervals. The red squares indicate the HR before (yellow circle) and after (blue circle) 1-minute stabilization interval. As can be seen from Panel A, there are no carryover effects. To quantitatively show this, we have calculated the difference between these two HR values ($\Delta\text{HR} = \text{HR}_{\text{yellow}} - \text{HR}_{\text{blue}}$). Panel B represents ΔHR for the duration of our experiments, indicating very stable preparation with negligible $\Delta\text{HR} < 7 \text{ bpm}$ with respect to HR ranged from 300–450 bpm. We performed similar analysis for all our rats' data ($n = 8$).

(TIF)

S4 Fig. Quantitative analysis of steady state of the 'last' 120 seconds of recording data. In regards to whether the last 100 seconds of the 120 seconds were in steady state, we performed the following quantitative analysis: we calculated the difference between mean HR during the first and last 20 seconds (see green boxes in Panel B) of each rat's run in our study for all **PRE**, **ON**, and **POST** datasets for all S-VNS (STOCH = 10% and 20%) and P-VNS protocols. From this quantitative analysis we observed that there was minimal difference (<12 bpm) between the first and last 20 seconds. Mean difference between the mean HR during the first and last 20 seconds for **PRE**, **ON**, and **POST** are shown for C) 10 Hz, D) 20 Hz, and E) 30 Hz.

(TIF)

S5 Fig. Histogram distributions of P-VNS and S-VNS protocols. Representative histogram distributions of one representative rat for P-VNS and S-VNS protocols for **PRE**, **ON**, and **POST**, delivered at 20 Hz.

(TIF)

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