Novel Clinimetric Toolset to Quantify the Stability of Blood Pressure and Its Application to Evaluate Cardiovascular Function After Spinal Cord Injury

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Profound dysfunction of the cardiovascular system occurs after spinal cord injury (SCI), which is a leading cause of mortality in this population. Most individuals with chronic SCI experience transient episodes of hypotensive and hypertensive blood pressure in response to daily life activities. There are currently limited tools available to evaluate the stability of blood pressure with respect to a reference range. The aim of this study was to develop a clinimetric toolset for accurately quantifying stability of the blood pressure measurements and taking into consideration the complex dynamics of blood pressure variability among individuals with SCI. The proposed toolset is based on distribution of the blood pressure data points within and outside of the clinically recommended range. This toolset consists of six outcome measures including 1) total deviation of the 90% of the blood pressure data points from the center of the target range (115 mmHg); 2) The area under the cumulative distribution curve starting from the percentage of blood pressure measurements within the range, and the percentage of values within symmetrically expanded boundary ranges, above and below the target range; 3) the slope of the cumulative distribution curve that is calculated by fitting an exponential cumulative distribution function and the natural logarithm of its rate parameter; 4) its x- and 5) y-axis intercepts; and 6) the fitting error. These outcome measures were validated using blood pressure measurements recorded during cardiovascular perturbation tests and prolonged monitoring period from individuals with chronic SCI and non-injured controls. The statistical analysis based on the effect size and intra-class correlation coefficient, demonstrated that the proposed outcome measures fulfill reliability, responsiveness and discrimination criteria. The novel methodology proposed in this study is reliable and effective for evaluating the stability of continuous blood pressure in individuals with chronic spinal cord injury.

Keywords: spinal cord injury, cardiovascular dysfunction, cardiovascular perturbation, spinal cord epidural stimulation, blood pressure stability
INTRODUCTION

Spinal cord injury (SCI) leads to life-long autonomic cardiovascular dysfunction for which there is no adequate treatment (Teasell et al., 2000; Wecht et al., 2000; Wecht et al., 2003; Karlsson, 2006; Krassioukov, 2009; Wecht et al., 2013; Wecht et al., 2017; Biering-Sorensen et al., 2018; Wecht and Bauman, 2018; Wecht et al., 2020a). For the estimated millions of people living with SCI globally (Singh et al., 2014; Kumar et al., 2018), altered autonomic cardiovascular regulation leads to chronic arterial blood pressure instability (Hubli et al., 2015; Katzelnick et al., 2019) with variation between blood pressure that can be exacerbated by orthostasis (e.g., position changes) and severe hypertension triggered by autonomic dysreflexia (Blackmer, 1997; Teasell et al., 2000; Wecht et al., 2003; Dolinak and Balraj, 2007; Krassioukov et al., 2009; Wecht et al., 2013; Zhu et al., 2013). Studies on individuals with chronic SCI have identified blood pressure instability as one of the determinants of cardiovascular morbidity and mortality, including a 4-fold increased stroke risk in the SCI population (Su and Miao, 2005; Jegede et al., 2010; Rothwell et al., 2010; Wu et al., 2012; Muntner et al., 2015; Piatt et al., 2016; Stevens et al., 2016; Parati et al., 2018; Wecht et al., 2018; Ditterline et al., 2020; Parati et al., 2020). Even in those without SCI, mounting evidence links blood pressure instability during orthostasis with poorer general physical and mental health (Wilkie et al., 1976; Wessely et al., 1990; Pilgrim et al., 1992; Rosengren et al., 1993; Barrett-Connor and Palinkas, 1994; Vasudev et al., 2010; Vasudev et al., 2011; Regan et al., 2013; Briggs et al., 2016; Briggs et al., 2018; Shanhag et al., 2018).

Recent studies have demonstrated spinal cord epidural stimulation (scES) can stabilize blood pressure toward a more normotensive range and alleviate symptoms of orthostatic hypotension when using cardiovascular-specific stimulation parameters (Harkema et al., 2018a; Aslan et al., 2018; Harkema et al., 2018b; West et al., 2018; Darrow et al., 2019). Acute use of cardiovascular scES in individuals with SCI that exhibit chronic hypotension can immediately increase blood pressure and maintain it within a target range of 110–120 mmHg (Harkema et al., 2018b). It is also able to acutely mitigate orthostatic stress and maintain blood pressure during a position change (Harkema et al., 2018c). It is suggested that long term use of scES may facilitate adaptive neuroplasticity to restore underlying autonomic nervous system defects: in individuals with SCI and profound cardiovascular dysregulation, use of daily cardiovascular-specific scES to stabilize blood pressure led to mitigation of orthostatic hypotension even without active scES, as well as restoration of cardiovascular autonomic reflexes (Harkema et al., 2018c; Legg Ditterline et al., 2021). Blood pressure stability generated by cardiovascular-specific scES is similar to that seen in non-injured individuals, and does not reflect the profound variability between blood pressure measurements typically observed in individuals with SCI.

Despite this, outcome measures of cardiovascular dysfunction that are both statistically and clinically relevant and can detect change in blood pressure and identify a relationship to a normative range are limited (Chen et al., 2010) and there is no comprehensive measure designed to accurately quantify the complex dynamic changes that occur during continuous blood pressure recordings in individuals experiencing cardiovascular dysfunction. In this study, we propose and validate a novel blood pressure stability toolset based on the cumulative distribution of data points within and outside of a normative range. This straightforward and intuitive method can comprehensively capture the complex and dynamic blood pressure variability that SCI population experience in their day-to-day life. The proposed framework provides a reliable toolset to accurately quantify effects of SCI on blood pressure instability, as well as provide a foundation for statistical comparison among SCI groups, e.g., in response to perturbation, or with and without interventions to stabilize blood pressure.

MATERIALS AND METHODS

We used systolic blood pressure data from individuals with SCI as well as non-injured participants to evaluate stability of blood pressure in response to perturbation and over a prolonged period of time. Data from a passive sit-up test was obtained from three research centers: Kentucky Spinal Cord Injury Research Center (KSCIRC), James J Peters VA Medical Center (JJP), and Kessler Foundation (KF). Systolic blood pressure data recorded during a 70° head-up tilt maneuver, over 24 h, and over 6 h was obtained at KSCIRC.

Cardiovascular Perturbation Tests

Sit-Up Test

Individuals lay supine for 5 min (at JJP and KF) or 15 min (at KSCIRC) and then were passively moved to the seated position with hips and knees at 90° angle (Figures 1A,B) The participants remained in seated position for 5 min (at JJP and KF) or 15 min (at KSCIRC) or as long as they could tolerate without showing symptoms of presyncope. More details about this procedure is reported previously (Wang et al., 2020). In this study, we included the individuals with SCI that were identified in the previous study (Wang et al., 2020) as having a large drop in systolic blood pressure after being moved to sitting position. In total, blood pressure data from 45 individuals with SCI and 48 non-injured participants (with no known cardiovascular dysfunction) were used for this test.

70° Tilt Test

Individuals lay supine for 5 min and once supported by restraints at the knees, hips, and chest the table would tilt upright to a 70° angle (Figures 1A,C); individuals would remain upright for 30 min or as long as tolerated without experiencing symptoms of pre-syncope. Blood pressure data of 34 individuals with SCI and nine non-injured participants (with no known cardiovascular dysfunction) were used for this test.

In both perturbation assessments, beat-by-beat systolic and diastolic blood pressure were obtained from finger plethysmography with intermittent brachial blood pressure measurements. Finger blood pressure measurements were
calibrated to brachial blood pressure measurements offline using a 2-point calibration method (Wang et al., 2020).

**Twenty Four-Hour Blood Pressure Monitoring**

In addition to the two blood pressure perturbation tests described above, we tested the outcome measures of the proposed toolset with systolic blood pressure data measured over 24 h to ensure the toolset could evaluate what individuals experience daily. Blood pressure of individuals with SCI and Non-injured individuals was monitored over 24-h using an automatic brachial recording device. Blood pressure data was recorded every 15 min during awake time and every 30 min during overnight sleeping; the schedule was determined in advance by each individual. All participants kept a diary of their daily routines including the time they slept at night and the time they woke up in the morning. Participants with too many missing data points were excluded. In total, the data from 22 individuals with SCI and 12 Non-injured participants (with no known cardiovascular dysfunction) were used for the analysis.

**Six-Hour Blood Pressure Recording With and Without scES Intervention**

Blood pressure data were recorded every 10 or 15 min over 6 h of awake time using an automatic brachial recording device from two groups of individuals with SCI: one group with implanted scES targeting cardiovascular regulation (n = 9) and a control group without scES (n = 15). We used the blood pressure stability outcome measures that we developed in this study to see if these measures can accurately describe the cardiovascular effects of scES on blood pressure stability in individuals with SCI.

All research participants provided written informed consent about the procedures and publications of findings of the study which was approved by Institutional Review Boards at University of Louisville, James J Peters VA Medical Center, and Kessler Foundation.

**Development of the Blood Pressure Stability Toolset**

In the following sub-sections, details of the components of the proposed blood pressure stability toolset are described. This method describes the distribution of systolic blood pressure data with respect to the 110–120 mmHg reference range. The selection of this range as the target does not imply that all individuals should have a systolic blood pressure within 110–120 mmHg. Rather, it is a physiologically meaningful reference range because the participants’ pre-injury systolic blood pressure is unknown to us (i.e., their own “healthy” systolic blood pressure) and 110–120 mmHg is within the “healthy” systolic blood pressure range established by the American College of Cardiology (Whelton et al., 2017).

**Total Deviation From Target**

The first outcome measure is defined as the overall amount of deviation of the range of 90% of the blood pressure data points
from the center of the target range (115 mmHg), called the Total deviation from target. We selected the data within the 5th (lower limit) and 95th (upper limit) percentiles, including 90% of all data points for analysis in order to remove effect of outliers. Total deviation from target is the sum of the deviation of measurements above 115 mmHg and deviation of data points below 115 mmHg (Figures 2A–C, Figures 3A,C). If all the blood pressure data points only fall on one side of the 115 mmHg line, Total deviation from target is calculated as the distance between 115 mmHg and the lower limit—if 90% of data points fall below 115 mmHg—or the upper limit—if 90% of data points fall above 115 mmHg only (Figure 3B). While this measure quantifies the distance of the farthest data points (i.e., those data points that are not considered outliers) from the center of the target range, it does not describe the distribution pattern of all the data points with respect to the target range.

Cumulative Distribution Curve
In order to quantify the distribution pattern of blood pressure data points with respect to the target range, we developed a new methodology based on the theory of cumulative distribution function (Montgomery and Runger, 2018). In this method, a cumulative distribution curve is built based on the percentage of systolic blood pressure measurements plotted against their corresponding ranges. The face-color of circle markers match the colors of the expanded ranges in D. The adjacent table shows the steps to calculate the area under the curve (AUC) from summation of trapezoidal areas between the curve and X-axis.

FIGURE 2 | Development of blood pressure stability toolset. (A) Schematics of brachial blood pressure recording and the table of recorded systolic blood pressure measurements over 6 h of awake time in an individual with chronic SCI. (B) Plot of 6 h of systolic blood pressure measurements during awake time. The limits of the shaded area in purple illustrate the 110–120 mmHg target range, while the horizontal green line illustrates the center of the blood pressure target range (115 mmHg). (C) Plot of 6 h of systolic blood pressure (same as B) with red horizontal dashed lines to indicate the 5th (lower limit) and 95th (upper limit) percentiles of systolic blood pressure measurements. Two red vertical curly brackets indicate the distance between the upper and lower limits and 115 mmHg line (dashed green), which give the deviation above target line and deviation below target line. The sum of these two deviations gives us the Total deviation from target. (D) Plot of 6 h of systolic blood pressure (same as B and C) with Y-axis limits expanded to include 40–230 mmHg. Horizontal gradient gray dashed lines mark each expansion of the upper and lower boundaries beyond the target range with blue–pink gradient colored backgrounds (Expansion Rate is set at 10 mmHg); the adjacent table shows the percentage of data points within each expanded range, colored to correspond to the related range on the plot. (E) Cumulative distribution curve: the percentages of systolic blood pressure measurements are plotted against their corresponding ranges. The face-color of circle markers match the colors of the expanded ranges in D. The adjacent table shows the steps to calculate the area under the curve (AUC) from summation of trapezoidal areas between the curve and X-axis. (F) Fitting exponential cumulative function (shown in red) to the cumulative distribution curve: λ is the rate parameter that shows the slope, X0 and Y0 show the intercepts with X and Y-axis, and the fitting error (E) which is the difference between the cumulative curve and the exponential curve; and (G) Values of six outcome measures.
starting with reference (110–120 mmHg), and then by expanding the upper and lower boundaries of this range by a given Expansion Rate (below); with each new expansion, the percentage of systolic blood pressure measurements within each range is calculated until the final range includes all possible systolic blood pressure values, i.e., 40–230 mmHg (Figures 2D,E). It should be noted that there is a floor-effect with this analysis: the lower range ends at 40 mmHg while the upper range expands to include the largest possible systolic blood pressure value (230 mmHg) to reflect the physiological limits of systolic blood pressure. The Expansion Rate is a parameter that defines the spatial resolution of the proposed method and it can be any values between 1 mmHg (highest spatial resolution) and 10 mmHg. For beat-to-beat blood pressure recordings where there are many data points, it is ideal to set the Expansion Rate to 1 mmHg so the method can capture all details about the distribution patterns of data points. For brachial blood pressure recordings (Figures 2A,B) where fewer data points are obtained, and especially when data points can be distributed widely throughout the range, the Expansion Rate can be set at 5 or 10 mmHg to achieve a smoother cumulative distribution curve. We have performed sensitivity analysis to test the effects of variable Expansion Rates on the outcome measures. This analysis and findings are described in the validation and results sections.

**Area Under the Curve**

From the cumulative distribution curve, we calculated the area under the curve (AUC): a percentage calculated as the sum of all trapezoidal areas contained between each pair of consecutive points on the curve and the x-axis (Figure 2E). To calculate the AUC, the x-axis is mapped linearly and equidistant between 0 and 1 based on the selected Expansion Rate. The calculation formula for AUC is as following:

\[
AUC = \sum_{i=1}^{N} \frac{1}{2} (x_{i+1} - x_i) (y_i + y_{i+1})
\]

Where N is the number of expansions; \(x_i\) and \(x_{i+1}\) represent values between 0 and 1 for consecutive expanded ranges and \(y_i\) and \(y_{i+1}\) represent consecutive percentages of the measurements within ranges \(x_i\) and \(x_{i+1}\). An AUC of 100% means all observed systolic blood pressure measurements are within the reference range (Wecht et al., 2020b). Values less than 100% but higher than 95% indicate that the measurements are distributed very close to the target range. Lower AUC values indicate that the blood pressure data points are farther away from the target.

**Exponential Cumulative Distribution Function**

In order to quantitatively describe the shape of the cumulative distribution curve, we fit an exponential cumulative distribution function (Figure 2F) to this curve with the following equation:
that are either inside the target range or distributed very close it. We used the exponential function for fitting to the cumulative curve because the fast-rising quality of this function can accurately model stable blood pressure data points that are either inside the target range or distributed very close it.

**Natural Logarithm of the Rate Parameter**

The rate parameter ($\lambda$) in the cumulative exponential function describes the slope of the curve, with higher $\lambda$ values representing steeper slope. Since the relationship between $\lambda$ and the slope of the exponential function is non-linear (i.e., small changes in $\lambda$ at lower values affect the slope more than greater changes in $\lambda$ at higher values) we are reporting the natural logarithm of $\lambda$ as an outcome measure to remove this nonlinearity. The natural logarithm of $\lambda$ can be any value between zero (shallowest slope) and 4 (steepest slope). Larger values of $\ln(\lambda)$ indicate the blood pressure data points have formed clustered distribution (either close to the target range or far away), whereas smaller values of $\ln(\lambda)$ indicate data points have more scattered distribution either on one side (hypertension or hypotension) or both sides (episodes of both hypertension and hypotension) of the target range.

**x- and y-Axis Intercepts**

The intercepts of the fitted exponential function on the x- and y-axis ($X_0$, $Y_0$), are included as outcome measures (Figure 2F). The $X_0$ and $Y_0$ are non-negative values that can both be 0, but both values cannot be greater than 0 at the same time. If $Y_0 = 0$ and $X_0 > 0$, none of the blood pressure measurements fall within the target range of 110–120 mmHg and the value of $X_0$ indicates the penultimate range without any blood pressure data points. If $Y_0 > 0$ and $X_0 = 0$, a percentage of data points ($Y_0$) fall within the target range. When $Y_0 = 100\%$, all recorded data points are within the target range. If $Y_0 = 0$ and $X_0 = 0$, it means that there are no data points within the target range, but the adjacent range includes a percentage of the measurements, indicating blood pressure observations are distributed close to the target range.

It should be noted that although the rate parameter $X_0$ and $Y_0$ are important outcomes to report individually, together, these three outcomes quantify the shape of the cumulative distribution curve and therefore for the statistical comparison, we combined them as $\frac{\ln(\lambda)}{4} + \frac{Y_0}{100} - X_0$.

Maximum value of $\ln(\lambda)$ is 4 and $Y_0$ is a percentage and therefore by dividing them to their maximum, we normalize them to a value between 0 and 1. $X_0$ is already between 0 and 1 and because of its inverse effect on blood pressure stability, $X_0$ is subtracted from the sum of normalized $\ln(\lambda)$ and $Y_0$. As mentioned earlier, $X_0$ and $Y_0$ cannot both have positive values and therefore they either enhance ($Y_0 > 0$ and $X_0 = 0$) or reduce ($Y_0 = 0$ and $X_0 > 0$) the effect of normalized $\ln(\lambda)$ in the combined formula.

**Fitting Error**

The Fitting error ($E$) is calculated as the mean value of absolute differences between the data points on the fitted exponential function and the cumulative distribution curve (Figure 2F). The smallest $E$ value is zero which means that the exponential function was a perfect fit to the cumulative distribution curve. Smaller non-zero $E$ values indicate that the blood pressure measurements follow an exponential distribution pattern which means that data points are clustered very closely to each other (Figures 3A,B). Larger $E$ values mean that the blood pressure measurements do not follow an exponential distribution pattern, i.e., data points are widely distributed on one or both sides of the target range with partial clusters (Figure 3C).

The combination of these six outcome measures can accurately describe various distribution patterns of systolic blood pressure recordings to quantify the differences between systolic blood pressure that is stable and within normative range (Figure 3A), stable but hypotensive (or hypertensive) systolic blood pressure (Figure 3B), and unstable systolic blood pressure with episodes of hypotension and hypertension (Figure 3C).

**Validation of the Stability Measure**

The proposed stability measures are validated based on the effect size (ES) to analyze discrimination (SCI vs NI) and responsiveness (supine vs sit/tilt for SCI), and based on the intra-class correlation coefficient (ICC) to evaluate the test-retest reliability (multiple assessments during screening in supine/awake for SCI).

**Evaluating Discrimination**

The discrimination refers to the ability of the measure to distinguish groups of individuals known to be different (ability to detect inter-group differences). In this case, it is important for this measure to distinguish between individuals who have cardiovascular dysfunction and those who do not. Discrimination is evaluated with effect size (Cohen, 1988; Lakens, 2013). The effect size is the standardized mean difference between the two groups. It quantifies the observed difference in terms of the pooled standard deviation. The effect size is classified as tiny (< 0.01), very small (0.01–0.2), small (0.2–0.5), medium (0.5–0.8), large (0.8–1.2), very large (1.2–2.0) and huge (> 2.0) (Cohen, 1988; Sawilowsky, 2009). An effect size of 0.5 or higher is considered relevant (Norman et al., 2003).

**Evaluating Responsiveness**

A measure is responsive if it can detect a change within individuals. During the sit-up and tilt maneuver, we want to know if the measure can detect changes in blood pressure when moving from one position (supine) to another (sit or tilt) or from awake to sleep during 24-h blood pressure monitoring. Responsiveness was also measured with the effect size of paired differences.

**Evaluating Reliability**

We have also measured the consistency of the outcome measures over time, i.e., the test-retest reliability. The test-retest reliability refers to the property of a measure to be statistically stable across time when the individuals do not experience any change. In this study, reliability (also called consistency) was evaluated using
intra-class correlation coefficient (ICC) and standard error of measurement (SEM) (Weir, 2005; Koo and Li, 2016). The ICC is calculated using mixed models. Let $\sigma_r^2$ be the variance of the random effect and $\sigma_e^2$ be the variance of the model error term. The ICC is calculated by the formula: $ICC = \frac{\sigma_r^2}{\sigma_r^2 + \sigma_e^2}$. Reliability has been classified as poor (0–0.25), fair (0.25–0.5), moderate (0.5–0.75), good (0.75–0.90) and excellent (> 0.90) (Portney and Watkins, 2008).

The SEM is also a clinically useful metric which allows practitioners to make inferences about individual changes in a test. SEM values are in the same units as the units of the variable being analyzed.

Figure 3 shows three examples of systolic blood pressure recordings with stable and two different unstable patterns and the corresponding outcome measures for each example.

**Sensitivity Analysis: Effects of Expansion Rate, Averaging and Down-Sampling on Proposed Outcome Measures**

In analysis of continuous blood pressure recordings obtained from the finger, it is routine to average the data points over a short period of time to remove clinically insignificant variabilities in the data that occur due to acquisition methodology. Down-sampling the data points is also sometimes used to remove excessive number of data points. The definition of the stability measures also involves the choice of the Expansion Rate parameter from the target range 110–120 mmHg. A sensitivity analysis was performed to evaluate whether the results obtained would change based on a different choice of these parameters.

**Data Averaging**

Analyzing systolic blood pressure data might use beat-to-beat data or averaging over a selected time interval. To evaluate any effect on the results found, multiple averaging schemas were performed (5, 10, 15, . . . , 60 s). Stability measures were calculated for each averaging and were compared to those obtained with no averaging (beat-to-beat data).

**Down-Sampling**

Down-sampling refers to a systematic choice of fewer points from the sample. The effect of picking every nth systolic blood pressure value on the obtained stability measure was evaluated by varying $n$ from 2 (every other data point) to 20 (every 20th data point) and calculating the corresponding stability measures and comparing them to what is obtained when every measurement it considered.

**Expansion Rate**

To calculate the area under the curve, boundaries are expanded symmetrically from the target range 110–120 mmHg with equal jumps and the percentage of values falling in that range are calculated. The effects that the choice of $R$ would have on the calculated outcome measures were evaluated by comparing the values found when $R = 2, 3, 4, \ldots, 10$ mmHg and comparing to $R = 1$ mmHg for beat-to-beat blood pressure recordings.

**Statistical Analysis**

Participants characteristics (demographics and injury details) were summarized using mean and standard deviation (SD) for continuous descriptors, and frequency count and percentage for categorical descriptors.

Discrimination was evaluated between groups known and statistically confirmed to have physiologically different responses, i.e., comparing non-injured vs individuals with SCI in the Sit Up Test, 70° tilt test, and 24 h BP Monitoring assessments. The measure used is effect size using Cohen’s d (for similar sample sizes) or Hedges’ g (for different sample sizes) formulas, calculated as the standardized difference between the mean values of the non-injured and SCI groups in sitting and tilt positions and during awake time from 24-h recordings divided by the Satterhwaite standard deviation. The systolic blood pressure data from Sit-up test and 70° Tilt test was used to evaluate responsiveness given the known and statistically confirmed changes that occur in individuals with SCI from supine to sitting or from supine to tilt. Paired changes, supine to sitting or tilt, were calculated for each individual and used to calculate the effect size (mean change divided by change standard deviation). The ICC value used to evaluate reliability was obtained from mixed models using data recorded in supine from SCI participants who have had two measurements during the screening phase without any changes to their day-to-day life and statistically confirmed no change in their cardiovascular function in between. These models included a random intercept for each participant. The variance of the residuals $\sigma_e^2$ and the variance of the random intercepts $\sigma_r^2$ were obtained and used in the formula of the ICC as the estimates for the variance of the error term and random effect, respectively. Statistical confirmation was evaluated with $p$-value from 2-sample t-tests or non-parametric Wilcoxon Rank Sum tests for discrimination and paired t-test or non-parametric Signed Rank test for responsiveness and test-retest reliability. The choice between t-tests or non-parametric equivalents was based on the normal distribution of the data tested using the Kolmogorov-Smirnov test.

The evaluation of the effects of averaging, down-sampling and Expansion Rate were performed using paired t-test of the stability measure values resulting from different scenarios described in Sensitivity analysis: Effects of expansion rate, averaging and downsampling on proposed outcome measures.

All tests were 2-sided with a significance level of 0.05. Data preprocessing and analyses were performed in MATLAB R2017b and SAS 9.4 (SAS Inc., Cary, NC).

**RESULTS**

**Participants Characteristics**

Sit up test: The combined Kessler, VA, and KSCIRC non-injured group of 48 individuals were 40 ± 13 years old and 67% were males. The combined SCI group ($n = 45$) was composed of 37 ± 11 years old individuals, 27 ± 12 years after injury, 84% males with 78% cervical injuries distributed across AIS A-D.
70° tilt test: The nine non-injured participants were 31 ± 11 years old at the time of experiments and 67% were males. The 24 SCI were 39 ± 11 years old, 75% males, 71% cervical injuries, 50% AIS A, 38% AIS B and 13% AIS C, and 11 ± 8 years after injury. A subset of SCI \( (n = 10, \text{all cervical injuries}) \) was used for test-retest reliability (80% male, 33 ± 13 years old, 7 ± 4 years post injury).

24-h blood pressure monitoring: Of the twelve non-injured individuals included in the 24-h blood pressure monitoring, 58% males and 27 ± 5 years old at the time of assessment. The individuals with SCI were divided into two groups: the screening group (SCI-G1: \( n = 13 \)) and the scES implanted group (SCI-G2: \( n = 9 \)); during the 24-h blood pressure monitoring assessment, stimulation remained OFF throughout the recording. The SCI-G1 were 54% males and 37 ± 14 years old at the time of assessment and 9 ± 6 years post injury. The SCI-G2 were 89% males and 31 ± 9 years old at the time of screening and 7 ± 4 years post injury. Only SCI-G1 data was used for discrimination and responsiveness analysis to avoid the interference of possible effects of scES on daily blood pressure in SCI-G2. A subgroup of SCI-G1 and all participants data in SCI-G2 were used for test-retest analysis \( (n = 15) \). The test-retest blood pressure recordings for SCI-G2 were performed with scES off and prior to scES-cardiovascular training, therefore no effects from the stimulation were expected.

6-h blood pressure monitoring for scES effects: Individuals with SCI without intervention for cardiovascular stability...
TABLE 1 | Validation characteristics of the proposed blood pressure stability outcome measures. Reliability, discrimination and responsiveness were evaluated. Responsiveness and discrimination were evaluated using effect size (ES), ES is classified as tiny (< 0.01), very small (0.01–0.2), small (0.2–0.5), medium (0.5–0.8), large (0.8–1.2), very large (1.2–2.0) and huge (> 2.0). An ES of 0.5 or larger has been classified as relevant. Reliability was measure with the intraclass correlation coefficient (ICC) and standard error of measurement (SEM). ICC is classified as poor (0–0.25), fair (0.25–0.5), moderate (0.5–0.75), good (0.75–0.90) and excellent (> 0.90). A measure with ICC of 0.5 is considered to be reliable. Estimate values above this 0.5 represent a discriminatory, responsive or reliable measure. SE: Standard Error.

| Measure                          | Position      | n  | Mean ± se | Evaluation measure | Classiﬁcation | Comment |
|---------------------------------|---------------|----|-----------|-------------------|----------------|---------|
| **TOTAL DEVIATION FROM 115 MMHG** |               |    |           |                   |                |         |
| Sit Up Test                     | SCI sitting position | 45 | 40 ± 1.88 | ES: 1.38 | Large effect | Discriminatory |
| 70-degree Tilt Maneuver         | SCI sitting position | 24 | 53 ± 4.70 | ES: 1.31 | Large effect | Discriminatory |
| 24Hr BP                         | SCI awake      | 13 | 52 ± 4.69 | ES: 1.18 | Large effect | Discriminatory |
| **Area Under the curve**        |               |    |           |                   |                |         |
| Sit Up Test                     | SCI sitting position | 48 | 94 ± 1.04 | ES: 1.18 | Large effect | Discriminatory |
| 70-degree Tilt Maneuver         | SCI sitting position | 45 | 83 ± 1.53 | ES: 1.29 | Large effect | Discriminatory |
| **DISCRIMINATION**             |               |    |           |                   |                |         |
| Sit Up Test                     | SCI sitting position | 48 | 1.03 ± 0.05 | ES: 1.10 | Large effect | Discriminatory |
| 70-degree Tilt Maneuver         | SCI sitting position | 45 | 0.68 ± 0.04 | ES: 1.08 | Large effect | Discriminatory |
| 24Hr BP                         | SCI awake      | 13 | 0.90 ± 0.07 | ES: 0.94 | Large effect | Discriminatory |
| **RESPONSIVENESS**             |               |    |           |                   |                |         |
| Sit Up Test                     | SCI supine position | 45 | 23 ± 1.79 | ES: 1.40 | Very large effect | Responsive |
| 70-degree Tilt Maneuver         | SCI supine position | 24 | 21 ± 2.03 | ES: 1.82 | Very large effect | Responsive |
| **TOTAL DEVIATION FROM 115 MMHG** |               |    |           |                   |                |         |
| Sit Up Test                     | SCI supine position | 45 | 40 ± 1.88 | ES: 1.82 | Very large effect | Responsive |
| 70-degree Tilt Maneuver         | SCI supine position | 24 | 53 ± 4.70 | ES: 1.08 | Large effect | Responsive |
| **Area Under the curve**        |               |    |           |                   |                |         |
| Sit Up Test                     | SCI supine position | 45 | 93 ± 1.12 | ES: 1.08 | Large effect | Responsive |
| 70-degree Tilt Maneuver         | SCI supine position | 24 | 92 ± 1.52 | ES: 1.07 | Large effect | Responsive |
| **EXPONENTIAL FITTING ERROR**   |               |    |           |                   |                |         |
| Sit Up Test                     | SCI supine position | 45 | 1.02 ± 0.06 | ES: 1.05 | Large effect | Responsive |
| 70-degree Tilt Maneuver         | SCI supine position | 24 | 0.64 ± 0.04 | ES: 0.81 | Large effect | Responsive |
| **TEST-RETEST RELIABILITY**     |               |    |           |                   |                |         |
| Sit Up Test                     | SCI Supine Day 1 | 8  | 18 ± 2.70 | ICC: 0.96 | Excellent | Reliable |
| 70-degree Tilt Maneuver         | SCI Supine Day 2 | 8  | 18 ± 2.43 | SEM: 1.14 | Good | Reliable |
| 24Hr BP                         | SCI Supine Day 1 | 10 | 15 ± 1.32 | ICC: 0.81 | Good | Reliable |
| 24Hr BP                         | SCI Supine Day 2 | 10 | 16 ± 1.75 | SEM: 2.1 | Good | Reliable |
| Monitoring                      | SCI awake Day 1 | 15 | 52 ± 3.37 | ICC: 0.64 | Moderate | Reliable |
| Monitoring                      | SCI awake Day 2 | 15 | 53 ± 3.30 | SEM: 7.99 | — | — |

(Continued on following page)
(n = 15) were 60% male, 36 ± 15 years old, 10 ± 9 years post injury; individuals with the scES implant targeting cardiovascular function (n = 9) were 78% male, 31 ± 6 years old, 7 ± 3 years post injury.

Validation of Blood Pressure Stability Outcome Measures

Results of the validation measures for the proposed blood pressure stability toolset, i.e., discrimination, responsiveness and test-retest analysis, are presented in Figures 4A–C and Table 1. The validation analysis demonstrates all proposed outcome measures for blood pressure stability are discriminatory, responsive, and test-retest reliable based on the ES for data recorded during sit-up test, 70° tilt test, and 24-h blood pressure monitoring. While all four outcome measures demonstrate the difference between non-injured and SCI (discrimination) and SCI in supine versus sitting positions (responsiveness), total deviation from 115 mmHg has the greatest ES values (statistically significant); this is expected, since this measure depends on the most (non-outlier) extreme blood pressure values that individuals experience during the perturbation tests and daily variability of their blood pressure. The AUC and integrated value for \( \ln(\lambda) \), \( Y_0 \), and \( X_0 \) demonstrate the difference in variability of systolic blood pressure with respect to the target range between individuals with SCI and non-injured individuals and they both show statistically significant differences between the groups for discrimination and responsiveness. The Fitting Error demonstrates how closely the cumulative distribution curve follow an exponential cumulative curve and shows statistically significant differences between all groups for discrimination and responsiveness except the NI vs SCI 24-h comparison. In test-retest analysis performed on repeated blood pressure recordings (Figure 4C), all outcome measures demonstrate no significant difference between the two perturbation tests or 24-h tests across individuals with chronic SCI based on ICC and p-values with the exception of Fitting Error for SCI 24-h data day 1 vs day 2.

Table 1 presents the details of discrimination, responsiveness and test-retest analysis with the classification of the evaluation measures.

Effects of scES Intervention on Cardiovascular Function

The effects of scES stimulation on cardiovascular function between two groups of individuals with chronic SCI, one group with and the other group without a scES intervention that targets cardiovascular regulation are depicted in Figure 4D. The results are illustrating all outcome measures proposed in this study can demonstrate the improvement in the blood pressure stability in the presence of scES and the differences between the two groups are statistically significant.

Sensitivity Analysis

Results of sensitivity analysis on the choice of averaging, down-sampling and expansion rate are reported in Table 2.
We found that averaging data has a significant effect on the resulting Stability Measure outcomes. Down-sampling did not have a significant effect on the AUC, total deviation from 115 mmHg, and SBP range containing 90% of the data for most choices. Down-sampling significantly altered the values obtained for the curve corresponding exponential function fitting error. The expansion rate had a significant effect on the AUC, and the fitting error but minor effect on the curve corresponding to exponential function parameter \( \frac{\ln(\lambda)}{4} + \frac{y_0}{100} - X_0 \).

**DISCUSSION**

The aim of this study was to develop a blood pressure stability toolset based on the cumulative distribution of data points around a normative blood pressure range and validate it with respect to discrimination, responsiveness, and reliability properties. We demonstrated that outcome measures introduced in this study, i.e., the area under the curve, the natural log of rate parameter of the fitted exponential curve with x-axis and y-axis intercepts, the fitting error, and the total deviation from 115 mmHg are effective at quantifying blood pressure instability and deviation from clinically recommended values and that they are reliable, responsive, and discriminatory. Each of the proposed outcome measures reveal different aspects of blood pressure variability and instability and could be used for classification and categorizing various types of cardiovascular dysfunction.

Traditionally, summary statistics, mainly mean and standard deviation or median, quartiles and extrema (minimum and maximum), have been used to measure blood pressure over time and response to treatments (Penzel, 1995; di Rienzo et al., 1983). The average provides the central tendency of the data over the recording period and can be highly discriminatory between two different recordings when the mean of the measurements changes. However, it fails to provide insights regarding how far the measurements are from a clinically accepted “normal” range and whether observed changes between two recordings indicate the data are trending closer or further from normal. Such insights are critical in order to determine the clinical relevance of the change in distribution. Additionally, mean and standard deviation are highly sensitive to extreme occurrences—one aberrant blood pressure value can highly impact the accurate quantification of the blood pressure recorded over time as it skews the mean and inflates the standard deviation. Furthermore, in order to understand the distribution of the blood pressure over time, the median and both quartiles with extrema must be included. This makes it difficult to evaluate a group of individuals or use it as a study outcome measure.

For these reasons, it is more desirable to compare the blood pressure measurements to a clinically valid normative range. One option is to use the percentage of measurements falling within a pre-specified range. This method will fail to account for values that inevitably fall outside the range, and given the variability inherent to cardiovascular function and the additional variability that occurs as a consequence of SCI, clinically relevant measurements that fall remarkably close to the boundaries will be ignored. This problem could be overcome by expanding the range, however, there is no consensus on how wide the range should be. The Total deviation from a target line introduced herein addresses the boundary issue and provides an overall view of how far the measurements fall from a target. It is also demonstrated to be highly discriminatory between different recordings.

Despite this, Total deviation alone does not comprehensively describe the distribution of data points with respect to the target. The proposed methodology based on cumulative distribution of blood pressure measurements (Figure 2), overcomes this limitation. It accounts for values falling within and outside the range, can include extreme measures without being significantly influenced by them, and it provides a comprehensive model of the distribution of blood pressure measurements within a recording. In order to quantitatively describe the cumulative distribution curve (Figure 2B), we proposed five outcome measures: the area under the curve, which is directly calculated from the cumulative distribution curve; the rate parameter (log) of the fitted exponential curve and its x and y intercepts (and the sum of their normalized values \( \frac{\ln(\lambda)}{4} + \frac{y_0}{100} - X_0 \)), as well as the fitting error. The analysis demonstrated all these proposed outcome measures are reliable, responsive, and discriminatory, and each describe a characteristic of blood pressure variability with respect to a normative range.

The sensitivity analysis demonstrated the commonly-used method of averaging blood pressure data points are detrimental to the blood pressure stability analysis: averaging essentially smooths the distribution and replaces actual recorded blood pressure values with artificial values calculated from smoothing. We recommend using all data points for the variability/stability analysis; if it is necessary to reduce the number of recordings, down-sampling is preferred as it has significantly less impact on the stability measure outcomes. Results presented in Table 2 demonstrate total deviation is more sensitive to down-sampling, followed by AUC; but outcome measures calculated from the exponential fitted curve \( \frac{\ln(\lambda)}{4} + \frac{y_0}{100} - X_0 \) are more robust and less affected by down-sampling. Similarly, AUC is more sensitive to the chosen expansion rate but \( \frac{\ln(\lambda)}{4} + \frac{y_0}{100} - X_0 \), and E are not significantly affected by this factor.

There are several considerations for which to account when using the proposed blood pressure stability toolset: this method is only sensitive to changes in blood pressure with respect to the target range, and it can accurately detect a clinically relevant improvement in the stability of the blood pressure. However, given two recordings—one of sustained hypertension and another of sustained hypotension equally distant from the target—outcome measures will not indicate a meaningful difference between them because, from the viewpoint of “blood pressure stability”, each recording represents far-from-stable blood pressure without improvement towards the target range. This method also does not account for the recording duration and it is recommended that when using these outcome measures, all measurements be of similar a duration. In applications where comparison between two recordings with different durations is unavoidable, the underlying presumption for these stability measures is the distribution pattern of data points in the shorter recording would have remained the same if it was recorded for

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**Mesbah et al. Blood Pressure Stability Toolset**

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| AVERAGING          | Mean (95%CI)          | Total deviation from 115 | Mean (95%CI) | Total deviation from 240 | Mean (95%CI) | Total deviation from 360 | Mean (95%CI) |
|--------------------|-----------------------|--------------------------|--------------|--------------------------|--------------|--------------------------|--------------|
| No averaging       | 80.94 (75.86, 86.01)  | 52.5 (43.02, 61.99)      | 1.26 (1.14, 1.39) | 5.43 (3.31, 7.56)       |
| Every 5 s          | 81.11 (76.08, 86.26)  | 50.18 (40.43, 59.93)     | 1.29 (1.16, 1.41) | 4.71 (3.04, 6.39)       |
| Every 10 s         | 81.28 (76.17, 86.39)  | 49.29 (39.77, 58.82)     | 1.29 (1.17, 1.42) | 4.49 (2.95, 6.03)       |
| Every 15 s         | 81.34 (76.22, 86.48)  | 47.7 (38.35, 57.04)      | <0.0001      | 4.27 (2.82, 5.71)       |
| Every 20 s         | 81.66 (76.59, 86.73)  | 46.29 (37.35, 55.23)     | <0.0001      | 4.21 (2.88, 5.54)       |
| Every 25 s         | 81.42 (76.22, 86.62)  | 46.15 (37.16, 55.15)     | <0.0001      | 3.93 (2.68, 5.18)       |
| Every 30 s         | 81.61 (76.48, 86.75)  | 45.69 (36.88, 54.74)     | <0.0001      | 3.82 (2.61, 5.03)       |
| Every 35 s         | 81.98 (76.67, 87.29)  | 45.21 (35.55, 54.86)     | <0.0001      | 3.8 (2.64, 4.98)        |
| Every 40 s         | 82.05 (76.82, 87.28)  | 43.97 (34.73, 53.22)     | <0.0001      | 3.75 (2.64, 4.86)       |
| Every 45 s         | 81.97 (76.61, 87.33)  | 44.43 (35.17, 53.69)     | <0.0001      | 3.57 (2.54, 4.6)        |
| Every 50 s         | 82.01 (76.64, 87.39)  | 43.51 (34.74, 52.29)     | <0.0001      | 3.56 (2.53, 4.58)       |
| Every 55 s         | 82.05 (76.76, 87.43)  | 42.67 (34.29, 51.06)     | <0.0001      | 3.57 (2.56, 4.59)       |
| Every 60 s         | 82.16 (76.89, 87.43)  | 41.5 (33.32, 49.69)      | <0.0001      | 3.2 (3.21, 4.1)         |
| Every point        | 80.94 (75.86, 86.01)  | 52.5 (43.02, 61.99)      | <0.0001      | 5.43 (3.31, 7.56)       |
| Every other point  | 80.96 (75.88, 86.03)  | 52.54 (42.99, 62.08)     | 1.26 (1.14, 1.39) | 5.32 (3.26, 7.37)       |
| Every 3rd point    | 80.93 (75.86, 86.01)  | 52.34 (42.89, 62.08)     | 0.968        | 1.26 (1.14, 1.39)       |
| Every 4th point    | 80.99 (75.93, 86.06)  | 52.49 (42.93, 62.05)     | 0.782        | 1.26 (1.13, 1.39)       |
| Every 5th point    | 80.96 (75.89, 86.03)  | 52.31 (42.76, 61.86)     | 0.9827       | 1.26 (1.14, 1.39)       |
| Every 6th point    | 80.97 (75.93, 86.01)  | 52.24 (42.59, 61.89)     | 0.9827       | 1.26 (1.14, 1.39)       |
| Every 7th point    | 80.93 (75.89, 85.96)  | 52.23 (42.9, 61.57)      | 0.6496       | 1.26 (1.14, 1.39)       |
| Every 8th point    | 80.97 (75.91, 86.04)  | 52.33 (42.69, 61.98)     | 0.7759       | 1.26 (1.14, 1.39)       |
| Every 9th point    | 80.88 (75.81, 85.95)  | 51.93 (42.45, 61.41)     | 0.3346       | 1.26 (1.14, 1.39)       |
| Every 10th point   | 80.9 (75.62, 85.98)   | 52.46 (42.81, 62.11)     | 0.9437       | 1.27 (1.23, 1.39)       |
| Every 11th point   | 80.97 (75.86, 86.09)  | 51.03 (41.79, 60.26)     | 0.0136       | 1.27 (1.23, 1.39)       |
| Every 12th point   | 80.97 (75.93, 86.09)  | 52.25 (42.76, 61.73)     | 0.6689       | 1.27 (1.14, 1.4)        |
| Every 13th point   | 80.74 (75.65, 85.83)  | 51.37 (42.51, 60.24)     | 0.0589       | 1.26 (1.13, 1.38)       |
| Every 14th point   | 80.93 (75.95, 85.9)   | 52.81 (43.38, 61.84)     | 0.8569       | 1.26 (1.13, 1.38)       |
| Every 15th point   | 80.79 (75.73, 85.84)  | 51.74 (42.38, 61.11)     | 0.2025       | 1.27 (1.13, 1.38)       |
| Every 16th point   | 80.83 (75.81, 85.84)  | 52.09 (42.18, 62.01)     | 0.4902       | 1.27 (1.15, 1.4)        |
| Every 17th point   | 80.62 (75.46, 85.79)  | 52.05 (43.15, 60.94)     | 0.4423       | 1.26 (1.13, 1.38)       |
| Every 18th point   | 80.71 (75.68, 85.73)  | 50.94 (41.93, 59.96)     | 0.0992       | 1.27 (1.14, 1.39)       |

(Continued on following page)
longer—i.e., increasing the observation time will not itself lead to improvements in stability of the data. This methodology is also not concerned with providing specific time points of stable or unstable blood pressure within a recording and only provides a measure of stability over the entire recording. These outcome measures can be used in conjunction with visualization of raw blood pressure measurements over time, the cumulative distribution curves and the mean values, especially when uncertainty arises.

The proposed clinimetric toolset for blood pressure stability is particularly valuable in the application of using epidural stimulation for improving cardiovascular function in individuals with chronic SCI. Given that the pre-injury blood pressure range in this population is often unknown, a clinically recommended range is used as a target. Individuals that are implanted with scES participate in experiments in which the examiners search for optimum stimulation parameters (electrode contact combinations, polarity, intensity, pulse width, and frequency) that maintain blood pressure within or close to the target range (Harkema et al., 2018b). By systematically changing stimulation parameters while measuring beat-by-beat blood pressure and heart rate, examiners can identify optimum stimulation parameters that lead to immediate increases in blood pressure from a chronically hypotensive state. Cardiovascular-specific scES can be used to maintain blood pressure within a target range in the research setting as well as in the community with the use of a pre-programmed brachial blood pressure device, allowing individuals to participate in their activities of daily life while their blood pressure is safely monitored. The proposed outcome measures (Total deviation, AUC and the combination of slope and x- and y-axis intercepts) would allow the researchers to quantitatively assess the changes in blood pressure recordings with scES and how close they get to the target range. In addition to measuring the overall deviation of data points from the target, the proposed outcome measures based on the expanding boundaries of the target range would also provide detailed assessment of the distribution patterns of the blood pressure measurements. This toolset provides a valuable framework for researchers and clinicians in the field to assess the effectiveness of scES quantitatively.

Utilizing the proposed method for expanding the boundaries of a target range and generating the cumulative distribution curve and related outcome measures can be expanded to other blood pressure stabilizing interventions and longer blood pressure monitoring periods (i.e., 48- or 96-h) which will be the focus of future studies. The proposed method also has implications beyond blood pressure analysis. With the rapid development of wearable sensors (Zhang et al., 2019; Moonen et al., 2020) and point of care tests (Huang et al., 2019; Liu et al., 2020; Xu et al., 2020) a swell the utilization of the methodology proposed in this study can be expanded into these other areas.

**CONCLUSION**

The blood pressure stability outcome measures proposed in this study are reliable, responsive, and discriminatory. They provide a solid ground for comprehensive and objective quantification of the

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**Table 2** (Continued) Effect of averaging, down-sampling and expansion rate on the proposed stability measure outcomes.

| EXPANSION-RATE | AUC | Total deviation from 115 | w(x, y) – x0 | Fitting error |
|----------------|-----|------------------------|--------------|--------------|
|                | Mean (95%CI) | p* | Mean (95%CI) | p* | Mean (95%CI) | p* |
| Every 19th point | 80.76 (75.78, 85.74) | 0.123 | 51.91 (43.17, 60.65) | 0.3212 | 1.26 (1.13, 1.39) | 0.883 |
| Expanding 1 mmHg | 80.87 (75.8, 85.94) | 0.554 | 51.18 (42.09, 60.27) | 0.0269 | 1.27 (1.14, 1.4) | 0.2349 |
| Expanding 2 mmHg | 80.9 (75.86, 86.01) | 0.9659 | — | — | 1.26 (1.12, 1.4) | 0.9039 |
| Expanding 3 mmHg | 80.58 (75.42, 85.74) | 0.0045 | — | — | 1.25 (1.11, 1.39) | 0.3698 |
| Expanding 4 mmHg | 80.56 (75.41, 85.72) | 0.0026 | — | — | 1.25 (1.11, 1.39) | 0.5422 |
| Expanding 5 mmHg | 80.89 (75.83, 85.95) | 0.7147 | — | — | 1.27 (1.13, 1.41) | 0.7722 |
| Expanding 6 mmHg | 79.4 (73.97, 84.83) | <0.0001 | — | — | 1.21 (1.07, 1.35) | 0.0007 |
| Expanding 7 mmHg | 79.96 (74.68, 85.22) | <0.0001 | — | — | 1.23 (1.09, 1.37) | 0.0643 |
| Expanding 8 mmHg | 79.73 (74.42, 85.04) | <0.0001 | — | — | 1.22 (1.08, 1.36) | 0.0104 |
| Expanding 9 mmHg | 78.73 (73.19, 84.27) | <0.0001 | — | — | 1.16 (1.02, 1.3) | <0.0001 |
| Expanding 10 mmHg | 80.74 (75.74, 85.75) | 0.1162 | — | — | 1.27 (1.13, 1.41) | 0.669 |

*Comparing averaging and down-sampling schemas with including every data point and comparing all >1 mmHg expansion rates with 1 mmHg expansion rate.
effects of SCI on cardiovascular function, and can evaluate the effectiveness of clinical interventions that target blood pressure stability in individuals with chronic SCI.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusion of this article will be made available by the authors, without undue reservation.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by Institutional Review Boards at University of Louisville, James J Peters VA Medical Center, and Kessler Foundation. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

SM contributed in conceptualization, methodology, software, original draft preparation, review and editing. BD contributed in investigation, resources, data curation, review and editing. SWang contributed in resources and data curation. SWu contributed in conceptualization and methodology. JWBeir contributed in formal analysis, review and editing. JWecht contributed in project administration, investigation and resources. GF contributed in project administration, investigation, resources and methodology. SH contributed in project administration, funding acquisition, supervision, investigation, resources, review and editing. BU contributed in methodology, formal analysis, validation, original draft preparation, review and editing.

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