Abstract—In the ICU, fluid therapy is conventional strategy for the patient in shock. However, only half of ICU patients have well-responses to fluid therapy, and fluid loading in non-responsive patient delays definitive therapy. Prediction of fluid responsiveness (FR) has become intense topic in clinic. Most of conventional FR prediction method based on time domain analysis, and it is limited ability to indicate FR. This study proposed a method which predicts FR based on frequency domain analysis, named instantaneous pulse rate variability (iPRV). iPRV provides a new indication in very high frequency (VHF) range (0.4-0.8Hz) of spectrum for peripheral responses. Twenty six healthy subjects participated this study and photoplethysmography signal was recorded in supine baseline, during head-up tilt (HUT), and passive leg raising (PLR), which induces variation of venous return and helps for quantitative assessment of FR individually. The result showed the spectral power of VHF decreased during HUT (573.96±756.36 ms² in baseline; 348.00±434.92 ms² in HUT) and increased during PLR (573.96±756.36 ms² in baseline; 718.92±973.70 ms² in PLR), which present the compensated regulation of venous return and FR. This study provides an effective indicator for assessing FR in frequency domain and has potential to be a reliable system in ICU. Based on these for FR in patients is important in clinical. It is important to develop the reliable prediction method.

Conventional methods usually predict FR by using time domain analysis, such as impedance cardiography (ICG) and pleth variability index (PVI). ICG is a non-invasive method to detect electrical and impedance changes in the thorax by using dual sensors. The electrical and impedance changes are used to calculate hemodynamic parameters for evaluating fluid response. The reliability of hemodynamic parameters in ICG is based on placed position of dual sensors, which needs to operate by professional paramedic. PVI is a time domain analysis for predicting FR in mechanically ventilated patients. It adopts pulse oximeter waveform and then calculates dynamic change in perfusion index (PI) during respiratory cycle. Nevertheless, time domain analysis effects by motion artifact easily. For avoiding inaccuracy in time domain analysis by motion artifact, this study proposed a reliable method which predict FR by frequency domain analysis.

Heart rate variability (HRV) can be measured by interbeat intervals (RRi) on electrocardiogram (ECG), which provides time domain and frequency domain analysis to assess autonomic nervous system (ANS). Especially, frequency domain analysis is used to adopt fast Fourier transform (FFT) for spectral analysis. Furthermore, spectral is divided into low-frequency (LF) range (0.04-0.15Hz) to present sympathetic nervous system activities, and high-frequency (HF) range (0.15-0.4Hz) to present parasympathetic nervous system activities mainly. However, HRV studies are restricted by the feasibility and the reproducibility with inconvenient measurement [1]. Therefore, pulse rate variability (PRV) was proposed as a substitute measurement of HRV. PRV uses pulse wave, which collected from photoplethysmography (PPG), to...
replace ECG recording in HRV and has been examined as a surrogate of HRV during non-stationary conditions in previous study [2]. Moreover, the arterial pulse wave is regulated from complex physiological controls which make PRV provide much more information than HRV.

However, frequency domain analysis of HRV and PRV are both limited by the timescale of RRi and pulse wave time intervals. Since timescale limitation, the indication of high frequency range in spectral analysis was restricted by time resolution. For breaking timescale limitation, a novel adaptive method, named instantaneous pulse rate variability (iPRV), was proposed [3]. It adopted the frequency range extension method based on ensemble empirical mode decomposition (EEMD) [4] and instantaneous period (iPR) projecting technique [5] help for PRV spectral analysis. Therefore, iPRV provides a new indication, named very high frequency (VHF) range (0.4-0.8Hz) for the neural regulatory estimation and peripheral responses [6]. The literature has proposed that VHF of HRV is as a novel index of left ventricular function evaluation [7], which further indicates the cardiac function, venous return, and FR. However, the variation and interpretation of VHF of iPRV still needs further exploration and examination. There is a common clinical experiment, named passive leg raising (PLR), which induces the increase of venous return and helps for the quantitative assessment of FR [8]. Previous study has examined that iPRV is reliable by using PPG during non-stationary condition, such as head-up tilt (HUT) [9]. This study performed the clinical experiment, known as HUT and PLR, for the further exploration and examination. Since several studies revealed that the VHF index of HRV is a reliable evaluation of cardiovascular diseases [7], VHF of iPRV has potential to indicate more physiological responses.

The aim of this study is to 1) explore the potential indication of VHF of iPRV during HUT and PLR, and 2) interpret the physiological meaning of VHF of iPRV in different non-stationary conditions.

II. METHODS

A. Subjects and Data Collection

Twenty six healthy subjects (male: 14; age: 24±1), who had no history of cardiovascular disease, participated this study. All recruited subjects performed four trials in whole experiment. First, subjects were rest in supine position with 10-minute recording as baseline. Second, subjects were tilting up passively (HUT) on the automatic tilting table and kept in tilt-up position for 10 minutes. Then, subjects were back to the supine position with 5 minutes for recovering to baseline. Finally, subjects were raising leg passively (PLR) for 10 minutes. All measurements were performed in a quiet temperature controlled room and the experiment was approved by institutional review board of the hospital. The study was approved by institutional review board of Tungs’ Taichung Metro Harbor Hospital. Informed consent was obtained from all participants before the experiment.

The ECG signal was recorded by BEST-C-04056 (BioSenseTek Corp., Taiwan) and the PPG signal was recorded by Nonin 8500 (Nonin Medical Inc., Plymouth, MN) with a sampling frequency 200Hz.

B. Instantaneous Pulse Rate Variability

The algorithm of iPRV analysis shows at Fig. 1. At first, the blood pulse signal was extracted from PPG signal as the pulse wave component by sifting process in EEMD. Sifting process is an iteratively detrending operation which uses to compute finite set of components, named intrinsic mode functions (IMFs), from source non-stationary data. Moreover, before sifting process, EEMD provides noise-assisted method into original data for eliminating multiple characteristic problem in IMFs. After mixtures of added noise and source data, detrending operation contains several steps. First, local extrema of data $x(t)$ are identified by peak-valley detection. The upper envelope $U(t)$ and lower envelope $L(t)$ are generated by cubic spline interpolation according to the local maxima and local minima. The trend in current timescale is computed by calculating the mean of $U(t)$ and $L(t)$, as $M(t)$.

$$M(t) = (U(t) + L(t))/2$$

The new timescale $H(t)$ is representation after detrending operation by data $x(t)$ subtracting the trend.

$$H_k(t) = H_{k-1}(t) - M_k(t), k \geq 1$$

Where $H_0(t) = x(t)$. After $k$ times detrending operation, if the trend of $H_k(t)$ satisfies the criterion as the steady constant trend, then the components $H_k(t)$ were extracted from $x(t)$ as IMF. After $n$ sifting process, $x(t)$ was decomposed into $n$ IMFs, $IMF_1(t)$~$IMF_n(t)$, and one residue $r(t)$.

$$x(t) = \sum_{i=1}^{n} IMF_i(t) + r(t)$$

Since IMFs were decomposed from different mixtures, the ensemble IMFs are computed by averaging each corresponding IMF. However, the resolution of timescale still limit spectral analysis. For breakthrough timescale limitation, iPRV adopts iPR of blood pulse signal for proposing variation by using normalized direct quadrature (NDQ) [5]. NDQ contains several steps. First, the amplitude frequency modulation (FM) signal $F(t)$ of $IMF_{main}$ is eliminated by iteratively normalization. Then, the empirical frequency modulation (FM) signal $F(t)$ of $IMF_{main}$ is assumed to be cosine function, and its quadrature $\sin \Phi(t)$ can be computed directly.

$$\sin \Phi(t) = \sqrt{1 - F^2(t)}$$

The instantaneous phase $\Phi(t)$ is calculated by taking arctangent of FM signal and its quadrature, then the iPR in instant is obtained from inverse of the derivative of instantaneous phase.

$$\Phi(t) = \tan^{-1}(\sqrt{1 - F^2(t)}/F(t))$$

Finally, fast Fourier transform was performed as the spectral analysis in each frequency band of IP. The spectral power of LF, HF and VHF were calculated by spectral integration as the clinical indicators. The spectral analysis programs in this study was developed by using commercial software platform (LabVIEW version 2013, National Instruments Corp., Austin, USA).
C. Time Domain Sequential Analysis

This study used a correlation analysis to time domain sequential for ensuring similarity between iPRV and HRV during non-stationary conditions. Correlation analysis has several steps as follows (Fig. 2). First, low frequency band of iPR was filtered by low-pass filter in order to receive instantaneous pulse rate (iPR_{LF}) series for comparing with interpolated time series of RRi. Then, cross correlation \( R(d) \) and mean square error between iPR_{LF} and RRi were calculated to measure the similarity between iPRV and HRV as time domain sequential analysis.

\[
R(d) = \frac{\sum_{n=0}^{N}[\bar{f}_{iPR}(t) - \bar{f}_{iPR}](\bar{g}_{RRi}(t) - \bar{g}_{RRi})}{\sqrt{\sum_{n=0}^{N}[\bar{f}_{iPR}(t) - \bar{f}_{iPR}]^2 \cdot \sum_{n=0}^{N}[\bar{g}_{RRi}(t) - \bar{g}_{RRi}]^2}}
\]  

(6)

Where \( N \) presents data length number of iPR_{LF} and RRi. \( d \) is the time shift. \( \bar{f}_{iPR} \) and \( \bar{g}_{RRi} \) are the means of the corresponding series.

D. Statistic Analysis

Variation of spectral power in each frequency band were compared between different analysis method and different condition using paired-sample t test for significant difference. P value less than 0.05 was considered statistically significant. Results of spectral power are reported as mean ± standard deviation. Statistical analysis was performed using commercial statistics software.

III. RESULTS

A. Time Domain Sequential Analysis Between Baseline and Non-stationary States

The comparison of time series between iPR_{LF} and RRi was presented in Fig. 3 in one of the participants as an example. The iPR_{LF} and RRi were similar fluctuation in different conditions sequentially in time domain.

B. Spectral Analysis of Different Conditions in HRV and iPRV

The results of the spectral analysis in different conditions were summarized in Table II. In HRV spectrum, the power of LF increased both in HUT and PLR. The power of HF decreased in HUT. The power of VHF in each condition was small in HRV spectrum. In iPRV spectrum, the power of LF increased both in HUT and PLR, which were the same variation as HRV. The power of HF decreased in HUT but increased in PLR. The power of VHF decreased significantly during HUT and increased during PLR. The illustration of the iPRV spectrum were summarized in Fig. 4 in one subject for example. The results of all participants’ iPRV spectrum were similar with subtle change of the freqency peaks’ locations. In LF band, there is a spectral peak around 0.1Hz in each experiment. In HF band, the peak is around 0.3 Hz in each experiment. In VHF band, there is a peak around 0.7 to 0.8 Hz when subject during supine and PLR position.
Fig. 3. The comparison of fluctuation of time series during different conditions. (a) RRi from ECG, (b) iPRV from PPG

Table II. The Result of HRV and iPRV Spectrum

<table>
<thead>
<tr>
<th></th>
<th>Baseline</th>
<th>Head-up tilt (HUT)</th>
<th>Passive leg raising (PLR)</th>
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<tbody>
<tr>
<td>HRV</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LF</td>
<td>229.73±249.77</td>
<td>303.92±531.35</td>
<td>268.54±412.64</td>
</tr>
<tr>
<td>HF</td>
<td>228.23±192.75</td>
<td>203.23±184.64</td>
<td>226.58±183.88</td>
</tr>
<tr>
<td>VHF</td>
<td>22.38±27.75</td>
<td>16.00±10.87</td>
<td>28.27±30.34</td>
</tr>
<tr>
<td>iPRV</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LF</td>
<td>324.58±393.67</td>
<td>401.32±771.51</td>
<td>427.50±691.18</td>
</tr>
<tr>
<td>HF</td>
<td>474.23±430.27</td>
<td>363.00±284.71</td>
<td>515.50±448.09</td>
</tr>
<tr>
<td>VHF</td>
<td>573.96±756.36</td>
<td>348.00±434.92*</td>
<td>718.92±973.70</td>
</tr>
</tbody>
</table>

The form is (mean ± standard deviation); * means p<0.05 compared with HRV. LF denotes low frequency band (0.04-0.15Hz); HF denotes high frequency band (0.15-0.4Hz); VHF denotes very high frequency band (0.4-0.8Hz).

IV. DISCUSSION

The component represents pulse wave was extracted from PPG signal by using EEMD. As a result, NDQ is a reliable technique to transform pulse wave component into iPR, which indicates the regulation of ANS in spectral analysis [3]. The reliability of iPRV in frequency domain had been examined in the literature [9]. In addition, the information from iPR is more than from RRi by using frequency extension method to break limitation of time resolution in RRi. For the verification of the reliability of iPR in time domain, this study performed time sequential comparison of iPRV and RRi for further examination. The result illustrated that the time series of iPR is similar with RRi and contains much more intrinsic components with high frequency component, which provides much more physiological information for the assessment. Moreover, iPRV has high positive correlation with HRV in time domain. It has the potential usefulness as an indicator for cardiovascular circulation assessment.

Previous study had examined iPRV spectrum assessed the new indicator (VHF) to show more physiological information [9]. Though some literatures investigated that VHF of HRV is a reliable evaluation of left ventricular function [7], VHF of iPRV still needs further examination. It had been examined that VHF contains parasympathetic activities and peripheral responses, which are influenced by venous return and cardiac function. The influences of respiration on VHF were examined by paced respiration study [6]. The mechanism of the VHF indication needs more exploration.
This study applied the clinical experiment, known as HUT and PLR, for the further examination. HUT and PLR served as the simple clinical experiments for cardiovascular circulation evaluation. HUT causes temporarily and induces sympathetic activation. On the other hand, the sympathetic activities increased during HUT, and the power of LF in HRV also increased, which quantitatively assessed the sympathetic activation. In another experiment, PLR causes the increase of blood volume in upper body and then causes the increase of venous return. The cardiac function was increased temporarily and induces the peripheral FR. The PLR induces sympathetic activation and the power of LF both in HRV and iPRV also increased. The power in VHF in iPRV increased when venous return increased during PLR. However, the power of HF in iPRV during PLR had different variation with HRV. This is probably due to effect of respiratory frequency. Even respiratory influence, the iPRV spectrum still demonstrated similar variation along with HRV spectrum in LF and HF during HUT and PLR. Besides, the results showed that VHF has potential to indicate the relevant change of venous return and monitor the FR. It is reliable to observe FR in frequency domain analysis during different status of venous return. Predict FR also can be simple and intuitive by data acquisition from PPG and using method of iPRV to analyze.

Though iPRV analysis provided more information of venous return and FR in VHF, it has some limitations. First, it needed to combine with clinical trial (PLR) for observing variation, and it is improper as a real-time application. Second, the data acquisition depends on PPG sensor, which is sensitive and easy to be influenced by body movement and unstable measurement.

However, arterial blood pressure signal can substitute as another source signal for iPRV analysis, but the relevant measurement instrument is expensive and is not simple for usage. Third, the iPRV analysis based on the EEMD method which needed to set appropriate parameters and the results are mainly influenced by the parameters setting, which is different while the source signal and the signal properties, such as sampling rate, are different. In addition to these limitations, VHF in iPRV is reliable indicator for FR. The indicator would be examined on non-responders to verify the effectiveness and quantify level of variation of VHF, such as the threshold for the diagnostic reference that evaluate patient who has well FR or not.

V. CONCLUSION

This study provides an effective indicator for assessing FR in frequency domain. It has potential to be a reliable system for ICU which avoid delaying definitive therapy or additional damage to patient. In the future, this indicator needs to implement on the patients for further exploration.

ACKNOWLEDGMENT

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