Pattern of respiratory-induced changes in fingertip blood volume measured by light transmission

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ABSTRACT

Respiratory-induced fluctuations in heart rate and arterial blood pressure have been intensively investigated, but there is little information on the effect of respiration on peripheral blood volume. In the current study, blood volume changes in the finger, obtained by light transmission measurements, were measured during regular breathing (6 s periods) and long breathing (12 s periods). Respiratory chest-circumference changes were simultaneously measured in order to associate the pattern of tissue blood volume change with the respiratory cycle. Sixteen subjects were studied, and in fourteen finger blood volume increased during inspiration and decreased during expiration in the long-breathing periods. In all 14 subjects the start of blood volume decrease was significantly delayed from the start of expiration by mean ± SD 1.00 ± 0.65 s (p < 0.001, range 0 - 2.3 s). The start of blood volume increase was significantly delayed from the end of expiration by 3.45 ± 1.76 s (p < 0.005). In eight, finger blood volume started to increase more than 2 s before the start of inspiration. For the 6 s breathing period, blood volume decreased during inspiration in five examinations, and increased in seven. The increase in peripheral blood volume during inspiration could be attributed to the higher abdominal pressure during inspiration, and to the decrease in sympathetic activity during inspiration and the subsequent vasodilatation. The decrease in peripheral blood volume during inspiration is probably due to the negative thoracic pressure during inspiration and its mechanical effect on thoracic vessels.

Keywords: Light Absorption; Deep Breathing; Sympathetic Nervous system; Tissue Blood Volume

1. INTRODUCTION

Respiratory-induced fluctuations in heart rate—respiratory sinus arrhythmia—and similar fluctuations in arterial blood pressure have been intensively investigated, but in only a few studies the pattern of the change during inspiration and during expiration was investigated. Heart rate increases and systolic blood pressure decreases during inspiration [1-3] but there is strong variability in the relationship between heart rate or systolic blood pressure and the respiratory changes in chest circumference. The origin of these respiratory-induced fluctuations is not decisively known. They have been attributed to fluctuations in central autonomic activity caused by spontaneous oscillations in respiratory center activity or by respiratory-induced mechanical effects on the aortic baroreceptors and the pulmonary stretch receptors [2-4]. The direct mechanical effect of respiratory-induced thoracic pressure changes on arterial blood pressure and on central veins may also be significant [5-7].

Respiratory-induced fluctuations have also been found in the peripheral circulatory system. Deep inspiration reduces skin blood flow, measured by skin temperature [8,9] and by laser Doppler flowmetry [9-12], and this reduction has been attributed to higher sympathetic activity. Peripheral blood volume, as measured by light transmission through the tissue, has been shown to fluctuate at the respiratory rate [13-18], as it does with each heart beat, and so induces the photoplethysmographic (PPG) signal.

Two possible mechanisms have been suggested for the origin of tissue blood fluctuations with respiration: mechanical influence of the negative thoracic pressure during inspiration on the arteries and veins in the thorax [14-17] and respiratory changes in sympathetic activity [15-18]. The effect of respiration on sympathetic activity has been demonstrated by several studies [19-21], which showed higher muscle sympathetic nerve activity.
(MSNA) during expiration and very low MSNA at end-inspiration, when lung volume is maximal. Since sympathetic activity generally constricts skin blood vessels, the decrease of sympathetic activity during inspiration is expected to increase fingertip blood volume.

The mechanical effect of respiration on the tissue blood volume also depends on the relative contributions of abdominal respiration and thoracic respiration. The negative thoracic pressure during inspiration decreases blood pressure in the arteries and veins in the thorax, and consequently decreases peripheral tissue blood volume during inspiration. The higher abdominal pressure during inspiration increases blood pressure in the arteries and veins in the abdomen and consequently increases peripheral tissue blood volume during inspiration [22,23]. In particular, finger blood volume is expected to increase due to the higher blood pressure in the abdominal aorta, resulting in higher blood flow from the heart to the upper part of the body.

The respiratory changes in the peripheral blood volume can therefore exhibit different patterns. In the current study we measured the temporal relationship between changes in finger blood volume and the respiratory phase.

2. MATERIALS AND METHODS

2.1. Subjects and Examination

Sixteen non-smoker male subjects aged 21 - 63 years, with no known cardiovascular or neurological disease were studied. During the examination the subjects sat with their right hand comfortably laid on the table, at about heart level. A PPG probe was attached to the right index finger for the measurement of light transmission, while the finger tissue, and an optic-fiber sensor for the measurement of the respiratory chest-circumference changes (see later) was applied around their chest. Fingertip skin temperature was measured before the start of the examination by an alcohol thermometer held by the index finger and the two adjacent fingers. Room temperature was 21°C - 24°C; fingertip temperature was 29°C - 35°C.

After a rest period of five min the subjects were asked to breathe three series of 5 regular and 5 long respiration periods, where inspiration and expiration time were determined by a light point moving on a computer screen in the form of triangular waves. Regular respiration consisted of inspiration of 2 s and expiration of 2 s followed by 2 s of no-breath, and long breathing consisted of inspiration of 4 s and expiration of 3 s followed by 5 s of no-breath. It should be noted that in our study there were no constraints on the tidal volume, and it was allowed to change between long and regular breathing.

2.2. The Light Transmission and the Chest Circumference Sensors

The PPG probe consisted of an infrared light-source and photodetector of a pulse-oximeter probe (Oxisensor N25, Nelcor), mounted on the same plane (reflection PPG probe). The probe was attached to the index finger of the right hand of the subject. A low-pass filter (0 - 40 Hz) reduced high frequency noise. The signal was inverted so that a higher signal level corresponded with a higher blood volume. The signals were sampled at a rate of 500 Hz (16 bit) and digitally stored for offline processing.

In order to obtain the relationship between the light transmission changes and time of inspiration and expiration, we used an optic-fiber sensor previously developed by us [24] for the measurement of respiratory-induced changes in chest-circumference. The sensor is based on the dependence of light transmission through a bent optical fiber on its radius of curvature, and on the change of the latter when chest circumference changes. Some light rays, which are totally reflected by the core-cladding surface when the fiber is straight or slightly bent, may escape through the cladding when the fiber bending is higher, if the angle to the surface normal becomes lower than the critical angle. The details of the sensor were described elsewhere [24].

3. RESULTS

In 14 out of the 16 long-breathing period examinations (12 s each, including 5 s pause), tissue blood volume increased during inspiration and decreased during expiration. Figure 1 shows the tissue blood-volume and chest-circumference change as a function of time for one of these examinations. In this examination, finger blood volume started to decrease about two s after the start of expiration, while the increase of finger blood volume started before the start of inspiration, and continued to increase during inspiration. In two examinations, the pattern of blood volume change during long breathing was not in direct relationship with chest-circumference change. In one of these, an inverse pattern was found for the long breathing periods: the tissue blood volume in the finger decreased during inspiration and increased during expiration. In the other, the finger blood volume displayed double blood volume pulses for each breath.

Figure 2 shows two examples from the 14, showing the pattern of direct relationship between tissue blood volume and chest-circumference change for the long-breathing period examinations. In all those examinations the finger blood volume started to decrease after the start of expiration (range 0 - 2.3 s). In ten cases finger blood volume started to increase before the start of inspiration, and continued to increase during inspiration, indicating that the blood volume increase was also related to the
end of the previous phase of expiration. In 8, the changes started more than 2 s before inspiration, and in 2, just before. To allow the relationship between tissue blood volume and chest-circumference change to be seen more clearly, the pulses in tissue blood volume were eliminated by depicting only the diastolic (minimal) blood volume for each pulse, and Figure 3 shows the results for the same examinations as in Figure 2.

The time of the increase and decrease of the blood volume pulse relative to respiratory phase was characterized by two parameters: the time-difference TDI from the end of expiration to the start of blood volume increase, and the time-difference TDD from the start of expiration to the start of blood volume decrease, as shown in Figure 1. The time of the start of expiration was taken as the time of the maximum of the chest-circumference change curve; the time of the end of expiration was taken as the time of the end of the breathing instruction triangle (BG in Figure 2).

Figure 1. The changes in tissue blood volume (TBV) and chest-circumference (CC) as a function of time. TDI is the time difference between the end of decrease of CC and the start of increase of TBV; TDD is the time difference between the start of decrease of CC and the start of decrease of TBV.

Figure 2. Two examinations showing, for long breathing periods, a direct relationship between tissue blood volume (TBV) increase (upper curve) and chest circumference (CC) increase (middle curve). TBV changes during regular breathing show an inverse relationship (a) and a direct relationship (b) with chest circumference increase. Upward direction indicates increase in TBV and chest circumference. The lower curve in each examination (denoted by BG—breathing graph) shows the triangular waves displayed on the computer screen for breathing instruction.

Figure 3. The two examinations of Figure 2, showing the changes in the minimal (diastolic) blood volume (upper curve) and chest circumference increase (lower curve).
Figure 4 presents the detailed results for the mean values of TDI and TDD for each of the 14 subject examinations. These mean values were obtained by averaging, for each subject, the values for all the long-breathing pulses, not including the first in each group. The increase in blood volume was significantly delayed from the end of expiration by $\text{TDI} = 3.45 \pm 1.76 \text{ s}$ (mean $\pm$ SD, $p < 0.005$) and the decrease in blood volume was significantly delayed from the start of expiration by $\text{TDD} = 1.00 \pm 0.65 \text{ s}$ ($p < 0.001$).

Figures 2 and 3 also present results of blood volume measurements with regular breathing of the 6 s period. A pattern of direct relationship between tissue blood volume change and chest-circumference change (i.e. increase of tissue blood volume during inspiration and decrease during expiration) was found in some subjects as shown in Figure 2(b) for one examination, while for other subjects it was an inverse relationship (as in Figure 2(a)). For the 14 examinations, seven showed a direct relationship, five an inverse relationship and in two the relationship was not clearly defined. No quantitative results could be derived from the examinations of regular breathing due to the low signal-to-noise ratio in the blood volume measurement.

4. DISCUSSION

In the current study we measured the blood volume changes in the finger, as obtained by light transmission measurements, during regular breathing (6 s periods) and during long breathing (12 s periods). The examinations were performed simultaneously with measurement of chest-circumference change, in order to see whether inspiration was associated with an increase or decrease in tissue blood volume. The main finding of the study was that in all except two long-breathing examinations of finger blood volume changes, tissue blood volume increased during inspiration and decreased during expiration (direct relationship). In only one examination did tissue blood volume decrease during inspiration and increase during expiration (inverse relationship). This inverse relationship was also found in some regular-breathing examinations, even though the long-breathing examinations in these subjects were associated with a direct relationship pattern.

An important finding was the relative timing of blood volume change. The start of the decrease in finger blood volume followed the start of expiration, suggesting that the former was induced by the latter. On the other hand the start of the increase in finger blood volume preceded the start of inspiration in the majority of the examinations. Though the increase in finger blood volume related to inspiration, it is likely to have been initiated by the end of previous expiration.

As presented in the Background Section, two possible mechanisms have been suggested as the origin of the tissue blood volume fluctuations with respiration: mechanical influence of the thoracic pressure on the thoracic blood vessels [14-17] and sympathetic activity oscillations [15-18]. The sympathetic activity has been shown to decrease during inspiration [19-21], which can explain the increase in fingertip blood volume during inspiration, found in almost all of our long-breathing results. Though those studies on sympathetic activity (measured by MSNA) were performed with spontaneous breathing, with a respiration frequency which was even higher than our regular breathing, it seems reasonable to suggest that sympathetic activity is similarly modulated by the respiratory pattern of our long-breathing examinations.

In former publications the mechanical effect of breathing on the peripheral blood volume was attributed to the negative internal thoracic pressure during inspiration which decreases blood pressure in the thoracic arteries and increases blood volume in the thoracic veins. These two effects are expected to decrease peripheral tissue blood volume during inspiration. However, inspiration is generally also accompanied with increased abdominal pressure, which increases the blood pressure in the abdominal arteries and veins and diverts blood to the peripheral tissue [22,23]. This effect increases the blood volume in the peripheral blood vessels during inspiration and can explain the increase in tissue blood volume dur-
ing inspiration and its decrease during expiration (direct relationship) in long breathing. In addition to the effect of sympathetic activity decrease during inspiration, the respiratory change in tissue blood volume is probably affected by the two opposing effects of thoracic and abdominal breathing on the peripheral tissue blood volume.

5. CONCLUSIONS

While in most long-breathing examinations finger blood volume increased when chest-circumference increased, an inverse relationship between the two parameters was found in one long-breathing examination and also in several examinations of regular breathing. It is likely that different mechanisms are involved in the effect of respiration on peripheral blood volume, probably including lower thoracic pressure, higher abdominal pressure and lower sympathetic activity during inspiration.

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REFERENCES


