Frequency dependence of hand-arm vibration on palmar sweating response
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Frequency dependence of hand-arm vibration on palmar sweating response

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Objectives This study attempted to elucidate the effects of hand-arm vibration frequency on palmar sweating response.

Methods Palmar sweating was measured before and during vibration exposure on the right palm of six healthy men. The left hand was exposed for 3 minutes to the following root mean square (rms) acceleration magnitudes and frequencies of vibration: 5 m/s² at 31.5 Hz, 10 m/s² at 63 Hz, 20 m/s² at 125 Hz, 40 m/s² at 250 Hz, and 50 m/s² at 315 Hz. According to international standard ISO 5349, these vibration levels generate the same frequency-weighted acceleration magnitude of 2.5 m/s² rms. A control condition consisted of grasping a handle without vibration. As the index of the activated central nervous system, plasma 3-methoxy-4-hydroxyphenylglycol (MHPG) was measured before and immediately after each vibration exposure.

Results Each condition of vibration induced a palmar sweating response. Among the six vibration conditions, vibration of 125 Hz and 63 Hz caused large palmar sweating responses compared with those of 315 Hz and the control condition. Plasma MHPG did not increase significantly after either vibration exposure.

Conclusions The palmar sweating response to vibration with the same frequency-weighted acceleration magnitude suggested dependency on frequency. The study suggests that the somatosympathetic reflex is associated with different palmar sweating responses.

Key terms acute vibration stress, palmar sweating, somatosympathetic reflex, vibration frequency.
against dry heat loss) varied from 0.7 to 0.8. Written informed consent was obtained from all the subjects before they started the experiment.

**Measurements**

Palmar sweating was measured by the ventilated capsule method (Hidrograph; AMU-100, Kands, Kariya, Japan). A capsule of 1 cm² in size was mounted on the right palm, which was not exposed to vibration. Dry nitrogen gas was pumped into the capsule at a rate of 0.3 l/min at one side, and the gas exited from the other side after being moistened by the sweat. The sweating volume was estimated from the change in humidity inside the capsule.

As an index of an activated central nervous system, including the hypothalamus, the plasma 3-methoxy-4-hydroxyphenylglycol (MHPG) concentration was assessed from 5 milliliters of venous blood (6).

**Experimental procedure**

A series of experiments was performed in a soundproof room with a constant temperature of 22°C and a relative humidity of 60%. So that adverse reactions to vibration would be excluded, each subject experienced all the vibration frequencies used in the study several times before the experiment was started.

After resting in the waiting room for 20 minutes, the subject entered the experimental room. A blood sample for MHPG was obtained from the right arm after a 30-minute period of acclimatization. The sampling was followed by the commencement of continuous recording of palmar sweating. After the stability of the palmar sweating was established, for at least 5 minutes, the subject was asked to grasp the handle of a vibration generator with his left hand for 3 minutes. Sinusoidal vibration was produced in the vertical direction by an electrodynamic vibrator (Akashi ASE-12, Yokohama, Japan). Visual feedback through a digital dynamometer allowed the subject to keep a constant grasp at a power of 49 N. Immediately after the vibration exposure, another blood sample was obtained.

The sinusoidal vibration applied to the subject included 5 m/s² rms (root-mean-square) at 31.5 Hz, 10 m/s² rms at 63 Hz, 20 m/s² rms at 125, 40 m/s² rms at 250 Hz, and 50 m/s² rms at 315 Hz. These frequencies and unweighted acceleration magnitudes generate the same frequency-weighted acceleration magnitude of 2.5 m/s² rms according to ISO standard 5349 (7). These vibration levels did not induce a significant palmar sweating response in the subject. One of these vibration conditions a day, including a control condition of grasping a handle without vibration, was applied to each subject in a randomized order.

The study was performed in November and December.

**Statistical methods**

The volume of palmar sweating and the concentration of plasma MHPG were analyzed quantitatively before and during vibration exposure for the six vibration conditions. The percentage increase in palmar sweating caused by vibration was also assessed for the six conditions. A repeated-measures analysis of variance (ANOVA) was used to test the hypothesis of no difference in the palmar sweating response across the six vibration conditions. A P-value of 0.05 was set as the limit of statistical significance.

**Results**

Figure 1 presents the mean values of palmar sweating throughout the six vibration conditions. The palmar sweating volume before the vibration exposure did not change significantly across the six vibration conditions. Acute exposure to any vibration stress, including a control condition, caused an increase in the palmar sweating response (P<0.0001). Although the repeated measures...
ANOVA did not indicate a significant difference among the six vibration conditions \( (P=0.50) \), palmar sweating during vibration frequencies of 63 Hz and 125 Hz was the largest with a mean value of 0.22 mg (cm\(^2\)/min) and 0.21 mg (cm\(^2\)/min), respectively, followed by 31.5 Hz with a value of 0.19 mg (cm\(^2\)/min).

Figure 2 shows the percentage increase in the palmar sweating volume as a result of vibration exposure during the six conditions. Vibration with frequencies of 125 Hz and 63 Hz increased the palmar sweating responses by 57.9% and 50.0%, respectively. Vibration at 250 Hz and 315 Hz increased palmar sweating as much as the control of grasping only.

Figure 3 indicates the plasma MHPG concentration before and immediately after the vibration exposure. The mean values of the MHPG plasma concentration in the six vibration conditions were all within the normal range (3–6 ng/ml) for healthy male subjects. The plasma MHPG concentration before the vibration exposure did not change significantly across the six vibration conditions or during any vibration exposure. Nor did vibration exposure cause a significant increase in the plasma MHPG concentration.

**Discussion**

According to international standard ISO 5349, frequency weighting is to be used for the assessment of all biological effects of hand-arm vibration (7). In this context, the vibration conditions used in our study can be considered to have the same frequency-weighted acceleration magnitude of 2.5 m/s\(^2\) rms. In our study, therefore, we evaluated the effects of different vibration frequencies with the same frequency-weighted acceleration magnitude on palmar sweating response.

Some studies have shown that the vascular response to vibration stress depends upon the vibration frequency (8, 9). In a study (4) in which the effect of vibration frequency on vascular response to vibration exposure was observed in the contralateral finger vibration frequency of 63–250 Hz provoked a significantly large response, and the central sympathetic vasomotor reflex elicited by the pacinian corpuscles was suggested to be a possible mechanism for the response. A similar phenomenon was observed for palmar sweating response.

We have earlier shown that vibration at 125 Hz with a magnitude of 50 m/s\(^2\) rms, unweighted (equivalent to 6.25 m/s\(^2\) rms weighted), induces palmar sweating response to vibration exposure in the contralateral side (5). In our present study, on the other hand, we observed that even a smaller magnitude of vibration produces a palmar sweating response.

With the same weighted acceleration magnitude, exposure to 125 Hz vibration produced the largest palmar sweating response, followed by 63 Hz and 31.5 Hz, although no significant change was obtained. Vibration at 315 Hz only increased palmar sweating as much as grasping. This finding suggests that the palmar sweating response to hand-arm vibration depends upon the vibration frequency.

Two mechanisms are known for vibration to induce palmar sweating, excitation of the higher center of the sympathetic nervous system (10) and the somatosympathetic reflex (11). As we previously observed (5), vibration intensity expressed as acceleration magnitude
seems to be associated with the former mechanism for palmar sweating response. The greater the vibration, the more discomfort produced in the subject, the result being excitation of the hypothalamus. With the same frequency-weighted acceleration magnitude, on the other hand, vibration frequency can be considered to affect the palmar sweating response through the somatosympathetic reflex. Pacinian corpuscles are vibration receptors activated in frequencies above 60 Hz, and they are the most sensitive at 125 Hz (12). This fact is compatible with our result that exposure to vibration at both 63 Hz and 125 Hz increased palmar sweating to the greatest degree. The higher the vibration frequency, the larger the input as total vibration stress to the body, the result being an increase in palmar sweating. As observed in our study, however, vibration at frequencies above 250 Hz produces total vibration stress that is too weak to produce palmar sweating. The palmar sweating response to vibration stress seems to depend on its frequency in a certain range, with which the optimal frequency range of pacinian corpuscles is consistent. Further study is needed with a larger number of subjects to provide more information on the frequency dependence of vibration on the palmar sweating response.

MHPG is known as a major metabolite of norepinephrine in the human brain (13). According to Maas et al (6), estimates of the concentration of MHPG in plasma could provide a possible reflection of the activity of noradrenergic neurons in the brain. In our present study, MHPG did not increase significantly across the six vibration exposures; this finding suggests that vibration with frequency-weighted acceleration of 2.5 m/s² rms did not activate the central noradrenergic neurons to such an extent that the subject felt discomfort. This finding indicates the possibility that the different palmar sweating responses to the six vibration frequencies used in this study are responsible for the somatosympathetic reflex rather than the mechanism related to the hypothalamus.

In conclusion, the palmar sweating response to vibration suggests dependency on vibration frequency. The different palmar sweating responses to vibration frequencies suggest a contribution by the somatosympathetic reflex via pacinian corpuscles.

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