Cold-induced vasoconstriction at forearm and hand skin sites: the effect of age

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Cold-induced vasoconstriction at forearm and hand skin sites: the effect of age

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Abstract During mild cold exposure, elderly are at risk of hypothermia. In humans, glabrous skin at the hands is well adapted as a heat exchanger. Evidence exists that elderly show equal vasoconstriction due to local cooling at the ventral forearm, yet no age effects on vasoconstriction at hand skin have been studied. Here, we tested the hypotheses that at hand sites (a) elderly show equal vasoconstriction due to local cooling and (b) elderly show reduced response to noradrenergic stimuli. Skin perfusion and mean arterial pressure were measured in 16 young adults (Y: 18–28 years) and 16 elderly (E: 68–78 years). To study the effect of local vasoconstriction mechanisms local sympathetic nerve terminals were blocked by bretylium (BR). Baseline local skin temperature was clamped at 33°C. Next, local temperature was reduced to 24°C. After 15 min of local cooling, noradrenalin (NA) was administered to study the effect of neural vasoconstriction mechanisms. No significant age effect was observed in vasoconstriction due to local cooling at BR sites. After NA, vasoconstriction at the forearm showed a significant age effect; however, no significant age effect was found at the hand sites. [Change in CVC (% from baseline): Forearm Y: −76 ± 3 vs. E: −60 ± 5 (P < 0.01), dorsal hand Y: −74 ± 4 vs. E: −72 ± 4 (n.s.), ventral hand Y: −80 ± 7 vs. E: −70 ± 11 (n.s.)]. In conclusion, in contrast to results from the ventral forearm, elderly did not show a blunted response to local cooling and noradrenalin at hand skin sites. This indicates that at hand skin the noradrenergic mechanism of vasoconstriction is maintained with age.

Keywords Thermoregulation · Aging · Glabrous skin

Introduction

With advancing age, thermoregulatory defense mechanisms may be impaired. Recent studies indicate that even during mild cold exposure elderly are predisposed to risk of hypothermia (DeGroot and Kenney 2007). Skin blood flow is a major effector of heat loss in humans, as it transports heat from the body core to the skin. DeGroot and Kenney (2007) report a reduced vasoconstrictor response in elderly subjects compared with young adults. However, their biophysical analysis could not explain why elderly failed to defend their core temperature. In his editorial response van Someren proposed (Van Someren 2007) that there might be an underestimation in calculations of the actual heat loss, because the regions rich in arterio venous anastomoses (AVA) are generally not taken into account. AVAs are predominantly found in glabrous skin such as palms, fingers, soles, and forehead (Bergersen et al. 1997; Donadio et al. 2006; Gorgas et al. 1977; Lenasi and Strucl 2004). Together with their relatively large surface to volume ratio and high sensitivity to thermal changes (Defrin et al. 2009), these locations are ideally adapted as heat exchangers with the environment (Grahn and Craig Heller 2004; Grahn et al. 2009; Van Someren 2007). Hence, a blunted perfusion response in elderly in these skin areas might explain why
elderly fail to maintain their core temperature when exposed to cold (Gilbert et al. 1999). Clothing provides insulative protection to most of the body surface. Therefore, the initial defense to cold might be related to local cooling of exposed areas, which in fact are mainly glabrous skin sites (forehead and hands). Age effects on mechanisms of mediating local cold-induced vasoconstriction are well documented for non-glabrous human forearm skin, yet no information is available for glabrous (hand) skin.

Neurally mediated vasoconstriction works primarily through sympathetic adrenergic axon reflexes and alpha-2-adrenergic receptors (Hodges et al. 2007; Kellogg 2006; Pergola et al. 1993). However, in vitro research does not support findings of increased cold-related axonal release of sympathetic adrenergic agents (Johnson 2007). Therefore, non-neural mechanisms are likely involved. Non-neural mechanisms include the rho-kinase pathway and removal of nitric oxide (Hodges et al. 2006; Thompson-Torgerson et al. 2007a). Experimental evidence indicates that the early phase of vasoconstriction due to local cooling is mainly dependent on neural regulation and that late phase vasoconstriction relies more on non-neural mechanisms (Pergola et al. 1993).

From literature it is known that during local cooling, young adults and elderly show a similar level of vasoconstriction at the ventral forearm (Thompson et al. 2005a; Thompson-Torgerson et al. 2007b). Interestingly, the actual mechanisms by which the vasoconstriction is mediated are different. Elderly show a reduced response to noradrenergic agents (Thompson et al. 2005a), yet this is compensated by an increased vasoconstriction through non-adrenergic mechanisms (Thompson-Torgerson et al. 2007b). It is unclear whether young and elderly show different vasoconstriction due to local cooling at other skin sites than the forearm, such as glabrous and non-glabrous skin of the hand. Many studies describe that innervation and topology of cutaneous microcirculation are not homogenous (Donadio et al. 2006; Johnson et al. 1995; Kellogg 2006; Krogstad et al. 1995; Wilson et al. 2005; Yamazaki and Sone 2006). For instance, glabrous skin (palms, fingers, soles, lips) is innervated mostly by sympathetic noradrenergic nerves and is rich in AVAs (Alvarez et al. 2006; Charkoudian 2003; Donadio et al. 2006; Kellogg 2006; Krogstad et al. 1995). Studies by Ekenvall et al. have shown that vascular responses to local cooling and noradrenaline in human finger skin (glabrous) are primarily mediated through alpha-2-adrenergic receptors (Ekenvall et al. 1988; Lindblad et al. 1990). In contrast to glabrous skin, non-glabrous skin has no AVAs and is innervated by both noradrenergic and cholinergic nerves (Kellogg 2006).

In this study, we are interested in age-related differences in perfusion in response to cooling at the hand sites. No data on age effects on vasoconstriction due to local cooling at hand skin and mechanisms that mediate it are available in literature. Therefore, we tested the hypotheses that at hand sites (a) elderly show equal vasoconstriction due to local cooling and (b) elderly show reduced perfusion response to neuronal (non-adrenergic) stimuli.

### Methods

#### Ethical approval

The medical ethical committee of Maastricht University Medical Center + approved the study. Each subject gave verbal and written informed consent prior to participation in the study. All procedures conformed to the standards of the Declaration of Helsinki.

#### Subjects

Sixteen young adults (18–28 years; 8 M, 8 F) and sixteen elderly (68–78 years, 8 M, 8 F) were included (characteristics in Table 1). All subjects were healthy, non-obese, non-smokers, and not taking medications. All young females were tested during the early follicular phase of their menstrual cycle. Subjects refrained from caffeinated or alcoholic beverages in the morning prior to the test, but were allowed to eat a small breakfast.

#### Protocol

Because of increased sympathetic vascular tone in the morning (Panza et al. 1991) experiments were conducted in the afternoon. After subjects were seated skin sites were cleaned with alcohol and air-dried. The studied skin sites were the dorsal side of the hand between the 2nd and 3rd fingers.

### Table 1 Selected subject characteristics

| Subject characteristics | Y (n = 16) | E (n = 16) | P value |
|-------------------------|-----------|-----------|---------|
| Age (year)              | 23.19 ± 0.73 | 71.33 ± 0.61* | P < 0.01 |
| Gender (Male/female)    | 8/8       | 8/8       |         |
| Height (m)              | 1.74 ± 0.02 | 1.67 ± 0.02* | P < 0.04 |
| Mass (kg)               | 64.36 ± 2.32 | 65.77 ± 2.29 | N.S. |
| BMI (kg/m²)             | 21.25 ± 0.73 | 23.42 ± 0.53* | P < 0.01 |
| Systolic BP (mmHg)      | 119.00 ± 2.69 | 129.47 ± 3.63* | P < 0.03 |
| Diastolic BP (mmHg)     | 66.38 ± 1.17 | 69.07 ± 2.28 | N.S. |
| Mean BP (mmHg)          | 83.92 ± 1.24 | 89.20 ± 2.12* | P < 0.04 |
| Mean skin temp. (°C)    | 33.74 ± 0.04 | 32.9 ± 0.06* | P < 0.01 |
| Whole body fat (%)      | 22.48 ± 2.00 | 27.40 ± 1.81 | N.S. |
| Leisure activity level  | 3.31 ± 0.12 | 3.55 ± 0.11 | N.S. |

Values are presented as mean ± SEM

* Y young adults, E elderly, BP blood pressure

* P < 0.05 versus young adult
metacarpal, the ventral hand at the base of the thumb, and the ventral forearm halfway the wrist and inner side of the elbow. To reduce the influence of age differences in systemic sympathetic adrenergic tone (Wilson et al. 2004) bretylium tosylate (10 mM, Bio-Connect, the Netherlands), diluted in glycol propylene, was delivered via iontophoresis (PerIont, Sweden). Iontophoresis is a non-invasive method of delivering a drug through the skin using a small electric charge. Bretylium (BR) is a positively charged drug, which is propelled through the cutaneous barrier by the anode of the iontophoretic chamber. Glycol propylene was chosen as a carrier as it is non-conducting by itself; hence it does not interfere with the drug delivery (Kellogg et al. 1989). Bretylium is known to selectively block local endogenous release of noradrenergic neurotransmitters (Kellogg et al. 1989). Because of inherent differences in skin permeability between glabrous and non-glabrous skin, iontophoretic characteristics were modified between sites to deliver a dose that caused a similar physiologic effect. Aging per se may affect the cutaneous barrier properties (Carmeli et al. 2003). Thus, the iontophoresis dose scheme was set at a level to be maximally effective for both groups. From literature (Johnson et al. 1995; Kellogg et al. 1989; Pergola et al. 1993; Yamazaki and Sone 2006) and a pilot study it was concluded that for effective bretylium treatment, different duration and current strength of iontophoresis was needed for different sites, as shown in Table 2. Control measurements (BR−) of perfusion response to local cooling were performed ipsilaterally to the locations treated with bretylium (BR+). After iontophoresis of bretylium, sites were given 1 h for hyperemia to subside and sympathetic blocking to take effect. Skin temperature was measured in 1 min intervals using iButtons (type DS1921H; Maxim/Dallas Semiconductor Corp., USA) (van Marken Lichtenbelt et al. 2006) at the 14 positions of the adapted Mitchel/Wyndham equation for mean skin temperature (Choi et al. 1997). Next, subjects were dressed in a water-perfused suit (TUBEsuit, DTI) to control whole body skin temperature (Blanketrol II, Cincinnati Sub-zero). The effectiveness of the bretylium treatment was tested via whole body cooling by setting the temperature of the water perfused suit to 26°C. Whole body cooling elicits reflex vasoconstriction through release of noradrenalin and cotransmitters from sympathetic nerve terminals; effective blockade of presynaptic nerve terminals abolishes the response, while at non-treated sites perfusion drops (Kellogg et al. 1989). During the rest of the experiment, subjects remained in thermal comfort by perfusing the suit with water at 33°C (Parsons 2003). Room temperature was kept at 24°C. Laser Doppler flowmetry (Perimed PFS000, Sweden) was used for measurement of skin perfusion. Custom-made Peltier elements in the casing of the probes allowed for local temperature control of skin sites at 24°C and 33°C. Measurements were divided into three phases (see Fig. 1). After a 30 min baseline period at 33°C, skin sites were cooled to 24°C (5°C/min) for 15 min, until perfusion reached a nadir. Next noradrenalin, diluted in glycol propylene (Bio-Connect, the Netherlands), was delivered transcutaneously via iontophoresis to simulate maximal response to noradrenergic stimulation (details in Table 2, example recording in Fig. 2).

Table 2  Current and duration of iontophoresis (0.5 cm²) and concentration of drugs at treated skin sites

| Iontophoresis properties at treated skin sites | Current (mA) | t (min) | Concentration (mM) |
|-----------------------------------------------|-------------|--------|-------------------|
| Bretylium                                      |             |        |                   |
| Dorsal hand                                    | 0.24        | 15     | 10                |
| Ventral hand                                   | 0.30        | 25     | 10                |
| Ventral forearm                                | 0.24        | 10     | 10                |
| Noradrenalin                                   |             |        |                   |
| Dorsal hand                                    | 0.24        | 1      | 2.5               |
| Ventral hand                                   | 0.30        | 5      | 5.0               |
| Ventral forearm                                | 0.24        | 1      | 2.5               |

Fig. 1 Timeline of measure protocol repeated at each skin site. Blood pressure was measured at start of baseline and at end of noradrenalin phase.

Fig. 2 Individual recording of skin perfusion. Local skin temperature was 33°C during baseline, 24°C during cooling and administration of noradrenalin.
At control sites (BR−) noradrenalin was delivered to a subset of subjects only (young: n = 11 and elderly: n = 13).

A study by Thompson (Thompson et al. 2005a) has shown that local cooling of skin and local administration of NA does not alter mean arterial pressure (MAP). Therefore, and to minimize the influence of the blood pressure (BP) measurement itself to the perfusion, BP measurement was restricted to before baseline and after administration of noradrenalin via digital blood pressure monitoring (CRESTA, Taiwan). Cutaneous vascular conductance (CVC) was calculated as perfusion divided by mean arterial pressure (MAP = 1/3 systolic BP + 2/3 diastolic BP). Whole body fat percentage was measured through Dual-X-ray Absorptiometry (DXA, HOLOGIC, USA). Daily leisure activity of subjects was measured through a Baecke (Baecke et al. 1982) questionnaire.

Data analysis

Perfusion data was sampled at 8 Hz. Baseline values of CVC were obtained by averaging 3 min of perfusion data at baseline. Vasoconstriction measures were obtained by averaging 3 min during local cooling or after administration of noradrenalin. Data for local cooling alone were normalized to baseline; total vasoconstriction response (local cooling + NA) was normalized to baseline. SPSS16.0 for Mac was used for statistics. Subject characteristics were tested for differences via Student-t testing. A MANOVA was used to test for differences in %ΔCVC among age and gender groups. Data were presented as mean ± SEM unless stated otherwise. Statistical significance level was set at P < 0.05.

Results

Elderly subjects differed significantly from young adult subjects in BMI, baseline mean arterial pressure (MAP) and mean skin temperature (P < 0.05) (Table 1). Since no significant gender differences were evident, data of both males and females were pooled. Daily leisure activity did not differ significantly between both age groups. Because BMI, baseline MAP, and mean skin temperature showed no significant relation to vasoconstriction measures and leisure activity was equal over age groups, we assumed that observed effects were mainly related to age. Due to technical malfunction during iontophoresis, testing on n = 3 young adults and n = 1 elderly was aborted.

To test the effectiveness of the bretylium treatment, whole body rapid cooling was applied. This resulted in a decrease of perfusion units (PU) at control sites, but not at bretylium-treated sites for both age groups: (young control −73 ± 4%; young bretylium −8 ± 6%; elderly control −65 ± 4%; elderly bretylium −5% ± 3%).

At bretylium-treated sites baseline CVC values did not differ significantly between age groups. Control sites also showed no age effect except for the ventral hand (young adults 0.66 ± 0.12 PU/mmHg vs. elderly 0.31 ± 0.04 PU/mmHg; P < 0.02).

Non-noradrenalin treated skin sites

Bretylium-treated sites at the ventral forearm and dorsal side of the hand (non-glabrous skin) showed a significantly smaller percentage change in CVC due to local cooling than non-treated sites (see Table 3). At the glabrous skin site (ventral hand) vasoconstriction to local cooling at BR+ sites was not significantly different than vasoconstriction at BR− sites, indicating that local mechanisms dominated. At bretylium-treated sites no significant age effect to local cooling was found. At control sites (BR−), only the ventral forearm showed reduced vasoconstriction in elderly (P < 0.05).

Noradrenalin-treated skin sites

After iontophoresis of a supra-physiological dose of noradrenalin perfusion decreased significantly at all measurement sites (Table 2). The change in CVC was significantly smaller in elderly as compared with young adults at the bretylium-treated sites of the ventral forearm (P < 0.01), indicating a reduced response to noradrenalin in elderly. However, at the ventral and dorsal sides of the hand no significant age effect was found. The effect of noradrenalin at control (BR−) compared with bretylium-treated (BR+) sites was not significantly different at the hand sites, but lower at the ventral forearm (P < 0.05).

Discussion

This study explored age effects in vasoconstriction during local cooling at three different sites of the skin. For the first time, the dorsal and ventral sides of the hand were measured, in addition to the ventral forearm that is traditionally studied. The primary findings of this study are

1. There was no significant age effect on vasoconstrictor response to local cooling at both glabrous and non-glabrous skin of the hand.
2. In contrast to the results from the ventral forearm skin, elderly did not show a blunted response to noradrenalin during local cooling at hand skin sites. This indicates that at hand skin the noradrenergic mechanism of vasoconstriction is maintained with age.
Ventricular forearm

Studies by Thompson et al. (Thompson et al. 2005a; Thompson-Torgerson et al. 2007b) have shown that at the ventral forearm vasoconstrictor response to local cooling is not different between young adults and elderly. Yet the mechanisms that mediate vasoconstriction are different. Elderly have a decreased noradrenergic vasoconstriction response, which is compensated by increased vasoconstriction through local mechanisms. In the present study, however, we observed a blunted vasoconstriction in elderly to local cooling at the control skin site (P < 0.05). On the other hand, our observation that the neural (noradrenergic) response is reduced in elderly was in line with earlier studies that also indicate reduced sensitivity to noradrenaline at the ventral forearm (Thompson et al. 2005a, b). This was strengthened by the result that the vasoconstriction response of noradrenaline at bretylium-treated skin (BR+ NA+), was significantly smaller in the elderly.

At ventral forearm skin that had sympathetic blockade (BR+), we observed no age effect in vasoconstriction to local cooling. This indicates that the elderly did not show increased vasoconstriction through local mechanisms.

Dorsal hand

The dorsal hand (non-glabrous skin) showed no age effect of vasoconstriction to local cooling at both bretylium-treated and control sites. Even the application of exogenous noradrenaline did not result in an age effect. Both in young adults and elderly, the bretylium-treated skin sites showed a significantly smaller vasoconstrictor effect to local cooling alone than the control site. This indicates that at the dorsal hand, similar to the ventral forearm, vasoconstriction during early phase local cooling was obtained through noradrenergic and non-adrenergic pathways. In combination with the absence of an age effect during local cooling alone at the control site, this indicates that aging per se does not alter noradrenergic sensitivity. This was strengthened by the results that no age effect is found after administration of noradrenaline (BR+ NA+).

Ventral hand

The ventral hand, like the dorsal hand, showed no age effect of vasoconstriction either to local cooling or to vasoconstriction after administration of noradrenaline. Glabrous skin consists of capillaries and arterio-venous anastomoses (AVA). Non-glabrous skin only has capillaries. These AVAs, which are solely innervated by sympathetic adrenergic nerves, are capable of shunting blood directly from the arteriole to the venous bed, thereby bypassing the capillaries and drastically reducing cutaneous blood flow (Kellogg 2006). Suprisingly, despite greater sympathetic innervation at glabrous skin sites (Defrin et al. 2009) no significant difference in vasoconstriction was observed between the BR+ and BR− site during local cooling. This indicates that at the ventral hand, vasoconstriction by local mechanisms was able to compensate for lack of vasoconstriction by noradrenergic mechanisms. An interesting parallel may be drawn with another study by Wilson et al. (2005), who found that glabrous skin with sympathetic blockade was still able to autoregulate flow during blood pressure variations, whereas non-glabrous skin lost its capability of auto-regulation.

From a functional perspective of thermoregulation, the ventral side of the hand, as compared with the dorsal side and the ventral forearm, is most adapted as a heat exchanger through the abundance of AVAs (DeGroot and Kenney 2007; Grahn et al. 2009; Van Someren 2007) and...
high sensitivity to temperature changes (Defrin et al. 2009). Various studies underscore the importance of glabrous skin areas (Gilbert et al. 1999; Grahn and Craig Heller 2004; Krauchi and Wirz-Justice 1994). Therefore, maintaining the functionality as heat exchanger of these skin areas is more important than, for instance, at the less exposed forearm. Analogous to the beneficial effects of physical exercise on vascular reactivity (Lenasi and Strucl 2004), naturally occurring exposure to cold of the hands might keep the skin of elderly “fit” to respond to daily changes in environmental temperatures.

Limitations

Other studies on local regulation of skin perfusion cited in this article in general used 34°C for baseline whole body mean skin temperature and local skin temperature. Although we used bretylium to avoid effects of systemic sympathetic tone, this may still have influenced results on non-treated sites. Furthermore, relatively fit and healthy elderly were selected. Despite age in years was indicative of senescence, the biological age could have been much younger. Therefore, subject selection based on biological age would make sense. Yet, the difficulty remains how to define the biological age. Since the majority of elderly in western societies are taking drugs that affect blood pressure (69.6% of aged over 65 years in the Netherlands (CBS 2009)), it may be interesting to study whether these elderly, whose biological age might better reflect their age in decades, do show a greater effect of age in vascular responses. Furthermore, the delivery of drugs through iontophoresis might induce a vasodilator response, by stimulating c-fibers, which has to be overcome before actual vasoconstriction is measured (Drummond and Lipnicki 1999). In this study, no age effect was found for the small iontophoretic vasodilation at any of the measured locations. Furthermore, despite differences in protocols, vasoconstriction values found in our study matched that of Thompson et al. (Thompson et al. 2005a) with respect to vasoconstrictor response to noradrenalin imposed on local cooling.

Summary

In summary, there was no significant age effect on vasoconstrictor response to local cooling at both glabrous and non-glabrous skin of the hand. This is in line with the results from earlier studies on the ventral forearm. Contrary to earlier studies, elderly did not show a blunted response to noradrenalin during local cooling. This indicates that at hand skin sites the noradrenergic (neural) mechanism of vasoconstriction is maintained with age.

Conclusion

In contrast to the results from literature covering ventral forearm studies, our study on hand skin sites shows that upon cooling both local (non-neural) and noradrenergic (neural) mechanisms of vasoconstriction are maintained in the elderly.

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Conflict of interest statement None.

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