Sex-specific activity of hesperidin, diosmin and genistein on human umbilical vein

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Abstract
Flavonoids intake may be associated to beneficial effects on the cardiovascular system. Aim of this study was to investigate biological activities and mechanism of action of two Citrus-derived flavonoids, hesperidin and diosmin, and the estrogenic isoflavone genistein on vascular tone of human umbilical vein (HU). Umbilical cords obtained at delivery from healthy women were divided into two groups according to the sex of the new-born. Experiments with flavonoids were carried out in seroton (5-HT)-precontracted HUV rings, before and after tissue pre-treatment with L-NAME (NOS inhibitor) or glybenclamide (K_ATP channel blocker). Hesperidin and genistein evoked larger NO-dependent vasorelaxation than diosmin. Endothelium-dependent responses to hesperidin and genistein, but not to diosmin, were gender-specific, being larger in female- as compared with male-derived HUV. Pre-exposure of tissues to L-NAME significantly impaired relaxant responses to hesperidin and genistein and completely blocked those to diosmin. Residual vasorelaxations elicited by hesperidin and genistein after L-NAME pre-treatment were blocked by incubation with glybenclamide.

In conclusion, while all flavonoids studied evoked NO-dependent vasorelaxation in HUV rings, the effects of hesperidin and genistein were gender-specific and greater than that of diosmin. Hesperidin, diosmin, and genistein might exert a beneficial effect on the vascular and endothelial function of HUV. This information may be important for foetal medicine and practice regarding the effects on foetal development of umbilical vascular functions.

Introduction
Natural polyphenols are the most common dietary antioxidants abundantly present in many fruits, vegetables and seeds; they exert a wide range of beneficial effects related to the prevention or the modulation of chronic diseases including gestational diabetes [1] and cardiovascular disease [2]. Observational studies have also shown an inverse correlation between the consumption of some classes of polyphenols and risk of stroke [3]; furthermore, preeclampsia occurred more frequently among women who rarely consumed daily serving of fruit and vegetables during pregnancy [4].

Polyphenols prevent endothelial dysfunction improving the release of NO [5,6] and protect cell constituents against oxidative damage [7]. Appropriate antioxidant capacity could prevent or attenuate the severity of those disorders induced by oxidative stress, such as preeclampsia [8] and intrauterine growth restriction [9]. Hesperidin is the predominant flavonoid found in many citrus species and it is presumed to have a role in the vascular protective effects of several citrus products and beverages in humans [10,11].

A recent clinical trial reported that 4-week consumption of orange juice (500 ml/day) as well as of control drink plus hesperidin resulted in a significantly decrease of diastolic blood pressure in healthy volunteers [12].

However, little is known about flavonoid-induced activity on human umbilical vein (HU) [12] and no conclusive results or clear evidence have been still obtained to support their use during gestation [13–15]. Many hypotheses have been considered for the lack of activity of flavonoids at vascular placental circulation; for instance, not all flavonoids have biological activity on endothelial cells [16] or cross the placental barrier. In addition, anatomical and physiological characteristics of HUV are significantly different from other venous vessels; it carries oxygenated blood from placenta to fetus and the vascular tone is not under nervous control [17]. Furthermore, acetylcholine as well as histamine mediate vasoconstriction instead of vasodilation as observed in almost all kinds of intact blood vessels [18].

Thus, the present study was designed to elucidate if Citrus-derived flavonoids hesperidin and diosmin had vasorelaxant activity on HUV, and in addition if their responses were sex-specific. Results were compared with those to genistein, an estrogen-like isoflavone contained in soya shoots but also widely distributed in leguminous plant foods such as beans, lentils and green peas.

Materials and methods
Human umbilical cords were collected by the local Hospital Macedonio Melloni (20129, Milan). Informed consent was provided according to the Declaration of Helsinki and the use was approved by the Ethics Committee at the University of Milan.

Ex-vivo experiments on isolated human umbilical vein rings
The umbilical cords were obtained from anonymous healthy pregnancy women (N= 90, 37–41 weeks) after delivery and were kept...

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at 5°C until the experiment, usually the same day of delivery. A rapid disruption of endothelium associated to increased response to 5-HT and absence of vasorelaxant responses were observed in preparations the day after delivery.

Umbilical cords were divided into male- and female-derived according to the sex of new-born. Medial segment of umbilical vein was carefully isolated from adherent connective tissue and cut into 4-5 mm long rings. From each vein, up to six vascular rings were prepared; care was taken not to touch the endothelial cell layer during the preparations of the HUA rings and thus it was assumed that the endothelium remained undamaged. The rings were mounted in 5 ml organ baths and perfused with Krebs solution (118 mM NaCl, 4.7 mM KCl, 1.2 mM KH₂PO₄, 1.1 mM MgSO₄, 2.5 mM CaCl₂, 25 mM NaHCO₃, 5.5 mM glucose, pH 7.4) at 37°C, continuously bubbled with 95% O₂ and 5% CO₂. Tension was recorded using a PowerLab/8SP (ADInstruments, UK) connected to isometric force transducers. Tissues were equilibrated for 1 h under a resting tension of 2.0 g. Preliminary experiments were performed to verify that 5-hydroxytryptamine (5-HT) at 1 µM (approximate EC₅₀ concentration) induced a stable and long-lasting contraction of the preparations. Upon recovery of a stable basal tone, concentration-response curves were obtained by cumulative addition of U46619 (thromboxane mimetic) (10⁻⁹ to 10⁻⁴ mol/l), histamine (10⁻⁸ to 10⁻⁴ mol/l), 5-HT (10⁻⁹ to 10⁻⁴ mol/l), adrenaline (10⁻⁶ to 10⁻⁴ mol/l), angiotensin-II (10⁻⁶ to 10⁻⁴ mol/l). Concentration-response curves to vasodilatory agonists: adenosine (10⁻⁴ to 10⁻³ mol/l), sodium nitroprussiate (10⁻⁴ to 10⁻⁶ mol/l), different flavonoids (10⁻⁸ to 3 x 10⁻⁴ mol/l), β-estradiol (10⁻⁶ to 3 x 10⁻⁴ mol/l), were obtained in endothelium-intact preparations of the HUA, after development of stable tone by 5-HT (EC₅₀). Each preparation was challenged with one agonist only, and time controls were carried out to ensure that responses were reproducible. Concentration-response curves to β-estradiol in the absence and presence of a competitive ERα- and ERβ-antagonist (ICI 182,780) were carried-out in male- and female-derived HUV. The effect of ER-antagonist on β-estradiol response was measured by calculating the pA₂ value, calculated by Schild regression analysis with the program GraphPad Prism 5.0.

Contractile responses to agonists were measured as increase in tone above baseline (mN/mg tissue), whereas relaxant responses were expressed as percentage of relaxation on 5-HT-precontracted tissues.

A separate set of experiments were carried out using preparations devoid of endothelium. The removal of endothelium was obtained by gentle scraping the lumen of umbilical vein ring with a needle (size: 1 mm), and its absence was assessed by an increased vasoconstriction to 5-HT and by the lack of vasodilation to hesperidin (1 µM).

**Drugs and chemicals**

Acetylcholine hydrochloride, adrenaline hydrochloride, adenosine, 17β-estradiol, glybenclamide, histamine, ICI 182,780, Nitro-L-arginine methyl ester hydrochloride (L-NAME), sodium nitroprusside (SNP), 5-hydroxytryptamine (5-HT), 9,11-Dideoxy-11α,9α-epoxymethanoprostaglandin F₂α (U46619), hesperidin, genistein, diosmin, were purchased from Sigma-Aldrich. All compounds were freshly dissolved in distilled H₂O, except flavonoids which were dissolved in DMSO.

**Statistical analysis**

All data were expressed as mean ± S.E.M. of eight experiments and represent unpaired data. Concentration-response curves were fitted and compared by analysis of variance (ANOVA) using GraphPad Prism 5.0. If P values were less than 0.05 the treatment affected the response over the tested range of concentration [19]. Maximal responses and pD₂ values (-log EC₅₀) for each agonist were compared by one-way ANOVA followed by Tukey’s post hoc test using GraphPad InStat.

**Results**

**Autacoids with contractile activity**

Many autacoids including histamine and neurotransmitters such as ACh caused concentration-dependent constriction in HUV (Figure 1) at resting tone. The stable thromboxane-A₂ agonist U46619 (10⁻⁹ to 10⁻⁶ mol/l) elicited larger concentration-dependent contractions than those evoked by histamine (10⁻⁴ to 10⁻⁴ mol/l), 5-HT (10⁻⁴ to 10⁻⁴ mol/l) and ATP (10⁻⁷ to 10⁻⁷ mol/l). However, U46619 caused long lasting vasoconstrictions of HUV preparations which did not return at resting tone even after several washes with fresh Krebs solution. The concentration-response curve to U46619 significantly differed from curves to exogenous serotonin (P = 0.01, n = 8), histamine (P = 0.005, n = 8) and ATP (P = 0.001, n = 8). Tissue sensitivity expressed as the pD₂ value between U46619 (6.94 ± 0.14; n = 8) and 5-HT (6.68 ± 0.11; n = 8) was comparable, but differed significantly from those of histamine (5.66 ± 0.04; n = 8) and ATP (4.08 ± 0.04; n = 8). No significant differences between vasoconstrictions in female- and male-derived HUV were observed. Adrenaline (10⁻⁵ to 10⁻⁴ mol/l) and angiotensin-II (10⁻⁶ to 10⁻⁴ mol/l) did not induce contractile responses at the concentrations tested, whereas acetylcholine induced vasoconstriction at resting tone.

**Vasorelaxant effects of exogenous sodium nitroprusside and adenosine**

The NO donor, sodium nitroprusside (SNP) added cumulatively (10⁻⁴ to 10⁻⁴ mol/l) to the 5-HT-constricted HUV rings produced endothelium-independent vasorelaxations that were not gender specific (Figure 2). Tissue sensitivity to SNP, expressed as the pD₂ value, was comparable in female- (6.54 ± 0.10) and in male-derived tissues (6.86 ± 0.28).

Adenosine (ADO, 1 nM-10 µM) induced concentration-dependent relaxations on 5-HT-precontracted rings of HUV (Figure 2); tissue sensitivity to ADO, expressed as the pD₂ was (6.54 ± 0.10). On the contrary, acetylcholine (ACh, 10⁻⁷ to 10⁻⁷ mol/l) was ineffective on 5-HT-precontracted preparations.

**The effect of ER-antagonist on β-estradiol response**

17β-estradiol (10⁻⁷ to 3x10⁻⁷ mol/l) caused comparable vasorelaxation in male- and female-derived 5-HT-precontracted rings of HUV (Figure 3). If preparations were pre-treated with the ER-antagonist ICI 182,780 (10⁻⁷ M for 30'), the concentration-response curves to 17β-estradiol were significantly shifted to the right in both male- and female-derived HUV. The estimated tissue sensitivity of ICI 182,780 to estrogen receptors, expressed as the pA₂ value was comparable in both groups being (6.87 ± 0.03) and (6.67 ± 0.09) in male- and female-derived HUV, respectively.

**Effect of hesperidin**

Figure 4 shows the concentration-response curves to hesperidin (10 nM-0.3 mM) in 5-HT-precontracted preparations. Female-derived HUV exhibited significantly greater vasorelaxations in response to hesperidin (P < 0.01) than male-derived tissues. Maximal responses were (54.0 ± 4%) and (35.0 ± 5%) in female- and male-derived tissues, respectively. Pre-treatment with the NO-synthase inhibitor, nitro-L-arginine methyl ester (L-NAME, 10 µM, 30'), significantly impaired hesperidin-mediated vasorelaxation (P < 0.01) both in male- and female-derived HUV.
female-derived rings. The NO component of the hesperidin-mediated vasodilation evaluated as percentage of inhibition induced by L-NAME was more evident in female- than in male-derived tissues. Human umbilical vein rings devoid of endothelium exhibited decreased vasorelaxations in response to hesperidin, which were comparable to those obtained in tissues pretreated with L-NAME. While pre-treated with both L-NAME (10 µM, 30’) and the ATP-sensitive K+ channel inhibitor glybenclamide (1 µM, 30’) rings showed a complete lack of response to hesperidin. Furthermore, pinacidil (1 µM) an ATP-sensitive K+ channel opener, induced vasodilation in 5-HT-precontracted rings (Figure 4).

Effect of diosmin

Concentration-dependent relaxations to diosmin (10 nM-0.3 mM) on 5-HT-precontracted HUV rings were not gender-specific and the two concentration-response curves obtained from male- and female-derived rings. The NO component of the hesperidin-mediated vasodilation evaluated as percentage of inhibition induced by L-NAME was more evident in female- than in male-derived tissues. Human umbilical vein rings devoid of endothelium exhibited decreased vasorelaxations in response to hesperidin, which were comparable to those obtained in tissues pretreated with L-NAME. While pre-treated with both L-NAME (10 µM, 30’) and the ATP-sensitive K+ channel inhibitor glybenclamide (1 µM, 30’) rings showed a complete lack of response to hesperidin. Furthermore, pinacidil (1 µM) an ATP-sensitive K+ channel opener, induced vasodilation in 5-HT-precontracted rings (Figure 4).

Effect of genistein

Genistein (1 nM-0.1 mM) elicited significantly (P < 0.01) increased concentration-dependent vasorelaxations in 5-HT-precontracted rings from female- and male-derived human umbilical vein (HUV). Maximal response was larger in female- (50.2 ± 4%) than in male-derived tissues (28.9 ± 7%) and differed significantly (P < 0.02). Pre-incubation with L-NAME (10 µM, 30’) in both female- and male-derived tissues impaired genistein-induced vasorelaxation, whereas pre-incubation with both L-NAME (10 µM, 30’) and glybenclamide (1 µM, 30’) completely blocked vasorelaxation (Figure 6).

Discussion and conclusion

In the present study we observed that hesperidin, and diosmin differed in their responses on HUV preparations; hesperidin elicited...
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[43x348]periments, unless occluded by symbol. Statistical significance for curves: *P

Figure 5. Cumulative concentration-response curves to diosmin in rings from female- and male-derived HUV. Vasorelaxations were expressed as percent inhibition of the 5-HT-induced (EC$_{50}$) contraction. Points show mean ± S.E.M. of 8 experiments, unless occluded by symbol. Statistical significance for curves: *P < 0.05 female-derived vs. female-derived + L-NAME and male-derived vs. male-derived + L-NAME.

Figure 6. Cumulative concentration-response curves to genistein in rings from female- and male-derived human umbilical vein (HUV). Vasorelaxations were expressed as percent inhibition of the 5-HT-induced (EC$_{50}$) contraction. Points show mean ± S.E.M. of 8 experiments, unless occluded by symbol. Statistical significance for curves: *P < 0.05 female-derived vs. female-derived + L-NAME and male-derived vs. male-derived + L-NAME.

2) hesperidin and genistein induced both endothelium-dependent (through eNOS-NO pathway) and endothelium-independent vasorelaxation. The increased vasodilation induced by hesperidin and genistein in female-derived HUV was not associated to an increased tissue sensitivity to NO, since relaxation to sodium nitroprussiate was not gender-specific. Our observations agreed well with those of Cattaneo et al. [24] who found a sex-specific difference in expression, activity, and function of eNOS in HUVEC.

A sex-specific diversity in response was not limited to umbilical vein; differences in reactivity to vasoconstrictor and vasodilator stimuli, and sex-specific eNOS expression and NO release have also been observed in other vascular beds [25,26].

On the contrary, diosmin evoked a weaker (25% at the higher concentration tested) and not gender-specific NO-induced vasorelaxation, which was completely blocked by L-NAME. Supraphysiological concentrations were also required to induce in vitro vasorelaxations to hesperidin and genistein. We assumed that flavonoids such as hesperidin and diosmin, in the form of glycosides, diffused through membranes of endothelial cells less rapidly and less efficiently than their aglycones; alternatively, although maximal care was taken to prepare HUV rings, we could not exclude that a deterioration in the venous blood flow (and thus in the vein endothelium) might occur during labour sometime.

In our experimental conditions, endothelium-independent vasorelaxations in response to hesperidin and genistein were completely blocked by glybenclamide; the presence of smooth muscle ATP-sensitive K$^+$ channel was also supported by the observation that pinacidil induced vasorelaxation in our preparations. ATP-sensitive K$^+$ channels are thought to play an important functional role in vascular responses of several vascular beds [27], including HUV [28].

Estrogen receptors (ER) beside their classical function of regulators of gene expression, have been associated to numerous non-genomic functions [29], including the regulatory function of eNOS activity [30]. In our experiments 17β-estradiol induced not gender-specific vasorelaxation that was antagonized by the ER-antagonist ICI 182,780. Thus, the increased eNOS activity in female-derived HUV in response to hesperidin and genistein was unlike to be accounted for an estrogenic effect; moreover, comparable levels of steroids, including 17β-estradiol, have been measured in fetal cord blood from male and female foetuses [31], and studies in twin pregnancies confirmed that maternal serum steroid levels were unrelated to the fetal sex [31]. Further studies would be necessary to elucidate this point; it is possible that the slight difference in chemical structure (hesperidin has saturated pyran ring while diosmin has unsaturated pyran ring) might explain the lower activity of diosmin.

Recent observations indicated that placenta was not a barrier for certain polyphenols, indeed, a transplacental transfer of flavonoids, estrogenic isoflavones, and their metabolites in humans [32] have been described.

Despite evidence for beneficial effects of flavonoids- and soy-rich diets in humans, potential adverse effects might occur in response to high flavonoid concentration in placental-fetal circulation [33], or in the case of a high consumption of phytoestrogens [34].

In conclusion, our data suggested that Citrus-derived hesperidin and diosmin may exert beneficial effects on placental circulation through different mechanisms. In particular, hesperidin has been found to be more active than diosmin, especially in female derived HUV.

In our experimental conditions, hesperidin and genistein elicited a significantly larger NO-dependent vasorelaxation than diosmin, comparable to that obtained by the isoflavone genistein.

Previous studies provided evidence that hesperidin induced rapid phosphorylation of eNOS, enhancing its activity and NO production in HUVEC [20]. Hesperidin was also highly involved, more than other Citrus-derived flavonoids, in the blood pressure lowering effects of orange products consumption [21] and a deficiency of hesperidin was associated to capillary leakiness [22]. Not all citrus-derived flavonoids were equally effective in inducing eNOS activity; for instance, naringenin (the aglycone form of naringin that remained after hydrolysis of phenolic glycoside) exhibited lower activity than hesperetin (the aglycone form of hesperidin) in HUVEC cells [23].

In our experimental conditions, hesperidin and genistein elicited a significantly wide endothelium-dependent vasodilation in female- than in male-derived HUV rings. The pre-incubation with L-NAME impaired vasodilation to hesperidin and genistein. Residual responses were comparable among male- and female-derived HUV, and in preparations devoid of endothelium. Taken together, these results suggested that

1) the NO component of the hesperidin- and genistein-mediating vasodilation was larger than that of diosmin, and was enhanced in female-derived HUV;
Therefore, a regular but moderate consumption (3-4 daily serving) of Citrus fruits, vegetables or orange juice might confer protection to both mother and foetus, reducing blood pressure, and the risk of some fetal vascular complications such as preeclampsia and intrauterine growth restriction.

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