Effects of Milk Casein Derived Tripeptides on Endothelial Enzymes In Vitro; a Study with Synthetic Tripeptides

Abstract

In the fermentation of milk by certain lactic acid bacteria, casein is degraded into bioactive tripeptides shown to lower blood pressure in experimental animal models and in mildly hypertensive humans. This effect is suggested to result mainly in inhibition of angiotensin converting enzyme 1 (ACE-1). Due to the complexity of renin-angiotensin system (RAS), several other enzymes than ACE-1 can participate in the production of vasoactive components. Therefore, in the present study we investigated effects of tripeptides isoleucine-proline-proline (IPP), valine-proline-proline (VPP) and leucine-proline-proline (LPP) on some endothelial enzymes that are important in RAS or otherwise have a role in the endothelial function. The enzymes investigated were renin, chymase, neutral endopeptidase (NEP), prolyl oligopeptidase (POP), cathepsin G, endothelin converting enzyme 1 (ECE-1), and cyclooxygenase 1 and 2 (COX-1 and COX-2). The tripeptides inhibited prolyl oligopeptidase (POP) dose-dependently. IPP was the most potent inhibitor (IC\textsubscript{50} 486 ± 95 \textmu M). Contrary, cathepsin G was activated by IPP, VPP and LPP as well as the amino acids proline and isoleucine. The other investigated enzymes were not affected. Inhibition of POP and activation of cathepsin G do not explain the blood pressure lowering effects of the tripeptides. Thus the inhibition of ACE-1 remains the most plausible mechanism of the antihypertensive effects of the tripeptides.

Introduction

Blood pressure and vascular functions are regulated by numerous circulating and local factors. The main regulatory system is renin-angiotensin system (RAS) (\textcircled{\textup{\textcircled{1}}}, Fig. 1), but also other endothelial enzymes produce highly active compounds such as prostacyclin and nitric oxide. The most biologically active substance in RAS is angiotensin II (Ang II), which is broken down from angiotensin I (Ang I) mainly by angiotensin converting enzyme 1 (ACE-1) but also by chymase and cathepsin G. ACE-1 also degrades bradykinin, a vasodilatory peptide into inactive fragments (\textcircled{\textup{\textcircled{2}}}). RAS contains a number of other enzymes besides ACE-1 which have an important role in blood pressure regulation. Some of these enzymes have a direct effect on Ang II but they also participate in the conversion of substances such as angiotensin-(1–7) (Ang-(1–7)) with potent vasodilatory and cardioprotective effects. [1–3] In addition to classical AT-1 receptors, new focus has also been set to vasodilatory AT-type 2 and Ang-(1–7)-Mas-receptors [3,4]. Ang-(1–7) also increases the physiological impact of bradykinin. [3–5]

When milk is fermented e.g. with L. helveticus or S. cerevisiae, casein is degraded into bioactive tripeptides isoleucine-proline-proline (IPP), leucine-proline-proline (LPP) and valine-proline-proline (VPP). These tripeptides have been found to lower blood pressure and improve vascular function in hypertensive rats [6–9] and in humans [10–12]. Their blood pressure-lowering effect is believed to result mainly from the inhibition of ACE-1 activity [6,13], but some of their favorable vascular functions are different from those of captopril, a standard ACE inhibitor [8]. This could be related e.g. to inhibition of arginase [13] which increases local substrate concentration for nitric oxide production. The effects of the casein derived tripeptides cannot be explained solely by ACE-1 inhibition due to their manifold pharmacological effects. Therefore we investigated their effects on other blood pressure and vascular function regulating enzymes in vitro.
Materials and methods

Tripeptides IPP, VPP and LPP, substrates dansyl-D-Ala-Gly-4-Nitro-Phe-Gly-0H (DAGNP), Z-Gly-Pro-AMC and N-Suc-Ala-Ala-Pro-Phe-pNA were purchased from Bachem (Weil am Rein, Germany). NEP was purchased from Innovative Research (Novi, MI, USA). POP (Recombinant Human), ECE-1 (Recombinant Human) and MCA-Arg-Pro-Pro-Gly-Phe-Ser-Ala-Phe-Lys-(DNP)OH were purchased from R&D Systems (Minneapolis, MO, USA). Cathepsin G (Human neutrophil) was purchased from Merck Chemicals (Nottingham, UK).

Renin assay

Renin (Human recombinant) activity was measured using a commercial fluorometric kit (Sensolyte® 520 Renin Assay Kit, AnaSpec, San Jose, CA, USA). The activity of renin was determined using 5-FAM/QXL™520 fluorescence resonance energy transfer (FRET) peptide as a substrate which renin degrades into 2 separate fragments. The fluorescent can be monitored at ex/em = 490/520 nm (Perkin Elmer Life Science, WALLAC VICTOR² 1420, Turku; Finland). Incubation time was 60 min at 37 °C including 10 measurements at room temperature. The substrate (10 μM) was Tris (25 mM)-NaCl (250 mM)- dithiotreitol (DTT, 2.5 mM) which was the assay buffer. The enzyme (0.005 μg, 50 μM and 1 μM–3.3 mM) were solved and diluted in the assay buffer. The enzyme concentrations with all tripeptide concentrations (1 μM–3.3 mM). The tripeptides were solved and diluted in the assay buffer.

Chymase assay

Chymase (Human recombinant) activity was measured using commercial kit (Chymase activity kit, Sigma-Aldrich, Jerusalem, Israel). The method is based on the ability of chymase to degrade N-Succinyl-Ala-Ala-Pro-Phe-p-nitroanilide into a colorimetric product which can be monitored at wavelength 405 nm (Multiscan RC, Labsystems, Helsinki, Finland). Incubation time was 50 min at 37 °C including 9 measurements with all tripeptide concentrations (1 μM–1 mM). The tripeptides were solved and diluted in the assay buffer.

Neutral endopeptidase (NEP) assay

NEP activity was measured using the method by Florentin et al. [14] with small modifications. Assay substrate was DAGNP which was diluted first in methanol and thereafter in assay buffer Tris-HCl (50 mM, pH 7.4), the final concentration of the substrate in the reaction was 50 μM. Methanol concentration in the reaction was 0.2%. The method was based on NEPs ability to degrade DAGNP of which fluorescence was monitored at ex/em = 342/580 nm (Perkin Elmer Life Science, WALLAC VICTOR² 1420). Incubation time was 120 min at 37 °C including 7 measurements with all tripeptide concentrations (1 μM–1 mM). The tripeptides and the enzyme were solved and diluted in the assay buffer.

Prolyl oligopeptidase (POP) assay

POP activity was measured using the method design by Stanzio et al. [15] and Atack et al. [16]. Assay substrate was Z-Gly-Pro-AMC from which POP can cleave AMC group, a fluorescent substance which can be measured in ex/em = 355/460 nm (Perkin Elmer Life Science, WALLAC VICTOR² 1420). The assay buffer was Tris (25 mM)-NaCl (250 mM)- dithiotreitol (DTT, 2.5 mM) buffer (pH 7.5). The enzyme, the tripeptides and the substrate were diluted in the assay buffer. The final concentrations of the enzyme, the substrate and the tripeptides in reaction were 0.005 μg, 50 μM and 1 μM–3.3 mM, respectively, and the incubation time was 60 min, including 10 measurements at room temperature.

Cathepsin G assay

Cathepsin G activity was measured using the method by Barret [17] with a small modifications. The method is based on the ability of cathepsin G to degrade the substrate N-Suc-Ala-Ala-Pro-Phe-pNA into a colorimetric product that can be measured spectrophotometrically at 410 nm (Multiscan RC, Labsystems, Helsinki, Finland). The substrate was diluted in dimethyl sulfoxide (DMSO). The final concentration of the substrate and DMSO in the reaction was 200 μM and 25%, respectively. The enzyme (0.1–0.2 mU/reaction) was diluted in ice cold distilled water. The tripeptides were solved and diluted in HEPES-NaOH buffer (100 mM, pH 7.5) which was the assay buffer. The incubation time was 70 min at 37 °C including 11 measurements with all tripeptide concentrations (0.1 μM–3.3 mM).

Endothelin converting enzyme I (ECE-1) assay

ECE-1 activity was measured using Activity Assay Protocol by R&D systems. The assay procedure is based on the ability of ECE-1 to degrade MCA-Arg-Pro-Pro-Gly-Phe-Ser-Ala-Phe-Lys-(DNP)OH and the development of the fluorescent products can be measured in ex/em = 320/405 nm (Perkin Elmer Life Science, WALLAC VICTOR² 1420). The assay buffer was 0.1 mM 2-(N-morpholino)ethanesulfonic acid (MES) – 0.1 mM NaCl –buffer (pH 6). The substrate (10 μM), the enzyme (0.005 μg) and the tripeptides (1 μM–3.3 mM) were solved and diluted in the assay buffer. The incubation time was 60 min at room temperature including 10 measurements with all tripeptide concentrations.
Cyclooxygenase 1 (COX-1) and cyclooxygenase (COX-2) assays
The activities of COX-1 and COX-2 were measured using commercial fluorescence-based assay (COX Fluorescent Inhibitor Screening Assay Kit, Cayman Chemical Company, Ann Arbor, MI, USA). The method includes 2 reactions: COX-1 and COX-2 form prostaglandin G₂ (PGG₂) from arachidonic acid in the first reaction and in the second reaction PGG₂ reacts with 10-acetyl-3,7-dihydroxyphenoxazine (ADPH) forming fluorescent product resorufin which can be measured at ex/em=530/590 nm. The tripeptides (1 μM-3.3 mM) were solved and diluted in the assay buffer. The incubation time was 60 min at room temperature including 10 measurements with all tripeptide concentrations.

Statistical analysis
The results are presented as mean ± SEM. Statistical analyses were performed using GraphPad Prism software (version 4.02). One way analysis of variances (ANOVA) followed by Dunnett's multiple comparison tests were used to compare area under a curve (AUC) results. AUC was calculated from the dose-response curves. The activities of the enzymes in different groups were presented as percentage from the control (group without the tripeptide) activity. Difference was considered significant if p-value < 0.05.

Results
IPP, VPP and LPP inhibited POP dose-dependently at high micromolar concentrations (Fig. 2). IC₅₀-values for IPP, VPP and LPP were 486 ± 95, 761 ± 146 and 875 ± 192 μM (mean ± SEM), respectively. The amino acids proline and isoleucine showed no inhibitory effects on POP (data not shown).

Interestingly, all the investigated tripeptides increased the activity of cathepsin G (Fig. 3). Also amino acids proline, isoleucine and arginine elevated cathepsin G activity (Fig. 4). Activation was up to 50% over the control values with all the investigated tripeptides and amino acids. Proline activated cathepsin G at lower concentrations (0.1–10 μM) than IPP, VPP and LPP and amino acids arginine (all 1 μM–3.3 mM) and isoleucine (1–100 μM).

The tripeptides IPP and VPP showed no inhibition or activation of the other investigated enzymes renin, chymase, NEP, ECE-1, COX-1 and COX-2 (n=3–11) at the concentrations between 1 μM–3.3 mM (Data not shown).
In the present study, IPP, VPP and LPP inhibited POP dose-inactive fragments (otensin converting enzyme 2 (ACE-2) can create Ang-(1–7). ACE-Ang-(1–7) from Ang I and Ang II and NEP from Ang I. Also angi- be carried out by chymase and cathepsin G. Cathepsin G is also Ang I which is further converted to Ang II by ACE-1. This can also (COX-1 and COX-2) and endothelin converting enzyme 1 (ECE- 1). Renin originating from kidney cleaves angiotensinogen to Ang I which is further converted to Ang II by ACE-1. This can also be carried out by chymase and cathepsin G. Cathepsin G is also able to split Ang II directly from angiotensinogen. POP forms Ang-(1–7) from Ang I and Ang II and NEP from Ang I. Also angiotensin converting enzyme 2 (ACE-2) can create Ang-(1–7). ACE- 1, NEP and POP participate in the degradation of bradykinin into inactive fragments (Fig. 1) [1, 2, 18, 19]. In the present study, IPP, VPP and LPP inhibited POP dose-dependently. This, however, does not explain their blood pressure lowering effects because POP converts a vasodilatory product Ang[1–7] from Ang I and Ang II. Ang(1–7) is able to stimulate recently describe Mas-receptors (Fig. 1). Furthermore, relatively high concentrations of the tripeptides were needed for inhibition of POP compared to ACE-1 inhibition [6, 13]. IC_{50}-values were higher than found in plasma after a single dose to the rat [20]. However, IPP seems to accumulate in the tissues and thus in vivo give slight pharmacological effect.

Bradykinin is also a substrate for POP (Fig. 1), in addition to Ang I and Ang II at least in in vitro conditions [19, 22, 23]. It may be possible that Ang-(1–7), produced by POP, balances its bradykinin degradation effects in the regulation of blood pressure and vascular function, and therefore POP does not markedly affect blood pressure. However, POP-like activity has been shown to be increased during ACE-1 inhibition in rats [23]. Similarly, also Ang-(1–7) levels are increased during ACE-1 inhibition in rats and in humans [24]. Incubation of mesenteric arteries with IPP improved Ang-(1–7) – and bradykinin-induced mesenteric artery relaxation [5]. This may be due to POP/NEP/Ang-(1–7)-Mas-receptor axis and due to ACE-1 inhibition caused by IPP. It is also possible that POP and other Ang-(1–7) forming enzymes, such as NEP and ACE-2, play a physiological role during ACE-1 inhibition or in special pathological situations. Despite numerous studies [18, 19, 21, 22, 24, 25], the exact physi- ological role of POP is unclear. POP cleaves short naturally occurring proline containing peptides on the carboxy side of proline residues. For that reason it may have a role in modulating neuronal peptides and hormones containing proline residues such as substance P and arginine vasopressin. It has been identified as a potential target in cognitive function and in some neurodegen- erative disease and POP inhibitors have been utilized for drug development mostly for neurological disorders and as memory enhancers [24, 25].

Interestingly, the three tripeptides and all investigated amino acids elevated cathepsin G activity. The results disagree with a previous study which measured the effects of tripeptides on cathepsin G [13]. In that study, tripeptides had no effect on cathepsin G but the incubation time was only 10 min while in the present study it was 70 min. It may be possible that the cathepsin G activating effect was not to seen in the short incubation. This is supported by our own in vitro vascular tests [8]. In addition, the effects of amino acids were not tested in our previous study [13]. Concerning in vivo condition amino acids isoleu- cine and proline stimulated cathepsin G already at such low concentrations which could mean physiological relevance of the findings.

Activation of cathepsin G cannot explain the decrease of blood pressure and the improvement of vascular function because cathepsin G converts Ang II from angiotensinogen and Ang I (Fig. 1). These findings support the assumption that main mechanism for the antihypertensive effect and vascular function improvement by the tripeptides is inhibition of ACE-1. Physiological and pathophysiological compounds can activate cathepsin G. Brecher et al. [26] found that acetaldelyde activated cathepsin G at high millimolar concentrations by ca. 50% similarly to the peptides and amino acids in the present study. One limitation of the present study is that it was carried out using pure peptides and enzymes in vitro, thus not in physiological conditions. However, these findings show that bioactive, antihypertensive milk-derived peptides are not only ACE-1 inhibitors but have also other effects on the components of RAS. Furthermore, the present study indicates the complexity of the...
RAS with its numerous enzymes of which inhibition or activation can change the balance between vasoconstrictive and vasodilating components to unexpected directions. Ex vivo and in vivo experiments are warranted to clarify possible physiological relevance of the findings.

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Conflict of Interest

Riitta Korpela was as an employee and Heikki Vapaatalo was consulting for Valio Ltd. at the time when the present study measurements were done.

References