ORIGINAL ARTICLE
PLASMA APELIN LEVEL IS DECREASED IN PATIENTS WITH CORONARY ARTERY DISEASE

Hiroaki Yokoyama1, Shin Saito1, Takumi Higuma1, Hiroyuki Hanada1, Tomohiro Osanai1, Kazuyuki Daitoku2, Ikuo Fukuda2 and Ken Okumura1

Abstract Adipose tissue secretes various bioactive molecules (adipokines), and apelin is one kind of adipokines. Recently, it was shown that plasma apelin level is decreased in patients with chronic heart failure, and apelin might play an important role in the pathogenesis of cardiovascular disease. However, plasma apelin level in coronary artery disease (CAD) or other heart disease such as valvular heart disease (VHD) has not been elucidated. We enrolled 31 patients with CAD and 14 patients with VHD who underwent elective cardiac surgery. We also examined plasma apelin level in 20 healthy subjects (Control). Blood samples were obtained before the surgery. Paired samples of visceral and subcutaneous adipose tissues were harvested during surgery. Plasma apelin level was lower in both CAD and VHD than in Control. When compared between CAD and VHD, it was lower in CAD than in VHD, and was not affected by treatment with HMG-CoA reductase inhibitors (statins) which was shown to increase adiponectin level. Left ventricular ejection fraction (LVEF) was lower in CAD than in VHD. There was no correlation between plasma apelin level and LVEF. Gene expression of apelin in visceral adipose tissue was higher than that in subcutaneous adipose tissue, but it was similar between two groups. These suggest that plasma apelin level was decreased in patients with cardiac diseases, especially in those with CAD. Its role in the pathophysiology of CAD remains to be elucidated.

Key words: apelin; coronary atherosclerosis; adipose tissue; chronic heart failure

冠動脈疾患における血中アペリン濃度の減少

横山 公章1） 齋藤 新1） 猛熊 拓未1） 花田 裕之1）
長内 智宏1） 大徳 和之2） 福田 幹夫2） 奥村 謙1）

抄録 脂肪細胞は様々な生理活性をもつアディポカインを分泌するが、アペリンはその一つである。血中アペリン濃度は心不全患者で低下することが報告され、心血管疾患で重要な役割を演じている可能性がある。しかし、冠動脈疾患（CAD）や弁膜症疾患（VHD）などの心疾患における血中アペリン濃度は解明されていない。我々は待機的術手を行ったCAD患者31例とVHD患者14例を対象に検討した。心疾患のない健常例20例においても検討した。術前術中の採血を行い、術中に内臓・皮下脂肪を採取した。血中アペリン濃度はCADとVHDで低下していた。アペリン濃度とCADやVHD間で比較すると、CADで有意に低下しており、HMG-CoA還元酵素阻害薬（スタチン）の影響を受けなかった。左室収縮能（LVEF）はCADで低下していた。血中アペリン濃度とLVEFの相関関係は認めなかった。内臓脂肪のアペリンmRNA発現は皮下脂肪より増加していたが、2群間に差を認めなかった。以上より、血中アペリン濃度はCADで低下するが、その病態生理学的意義については今後の検討を要する。

キーワード：アペリン；冠動脈の動脈硬化；脂肪細胞；慢性心不全

1) Department of Cardiology Hiroaki University Graduate School of Medicine, Hiroaki, Japan
2) Department of Thoracic and Cardiovascular Surgery, Hiroaki University Graduate School of Medicine, Hiroaki, Japan

Correspondence: T. Osanai
Received for publication, December 14, 2009
Accepted for publication, January 4, 2010

弘前大学 61:58—64. 2010
1. Introduction

It is now widely recognized that adipose tissue is not only a reservoir for energy storage but also an endocrine tissue. Adipose tissue secretes various bioactive molecules (adipokines) including adiponectin, tumor necrosis factor (TNF)-α, and interleukin (IL)-6. Apelin, one kind of adipokines, is a novel peptide identified as the endogenous ligand to the angiotensin receptor-like 1 (APJ) receptor, which resembles the angiotensin receptor, but does not bind to angiotensin II1. It is expressed in the human heart, lung, kidney, adrenal gland, adipose tissue, and large conduit vessels including coronary artery2. In contrast to angiotensin, a potent vasopressor and anti-diuretic hormone, apelin lowers blood pressure via a nitric oxide-dependent mechanism, produces diuresis by inhibition of arginine vasopressin activity and release, and has a positive inotropic effect3,8.

Recently, it was shown that plasma apelin level is decreased in patients with chronic heart failure (CHF)3,9, indicating that apelin might play an important role in the pathogenesis of cardiovascular disease. There are many studies suggesting the relation between plasma apelin and CHF, but there are only few studies suggesting the relation between plasma apelin and atherosclerosis. Hashimoto et al. reported that APJ might play an important role in the progression of oxidative stress-linked atherosclerosis in mice, and might be a therapeutic target for atherosclerosis10.

To our knowledge, plasma apelin level in coronary artery disease (CAD) or other heart disease such as valvular heart disease (VHD), and the gene expression of apelin in the visceral and subcutaneous adipose tissues have not been investigated. The aim of this study is to investigate the plasma apelin level in the patients with CAD and VHD, and the gene expression in the visceral and subcutaneous adipose tissues, and to clarify the relationship between atherosclerosis and apelin.

2. Methods

2.1. Patient’s profile

All patients gave written, informed consent before the study. The study protocol was approved by the Ethics Committee on Human Research at our institution (2008-151). This study enrolled 45 patients undergoing cardiac surgery at Hirosaki University Hospital. There were 31 patients with CAD (mean age 66 years, ranging from 50 to 81) who underwent elective coronary artery bypass graft surgery (CABG) (CAD group), and 14 patients with VHD (mean age 62 years, ranging 42 to 77) who underwent elective valve replacement or plasty (VHD group). The VHD patients were confirmed to have no CAD by coronary angiography. No patients had severe renal or hepatic dysfunction, or any history of neoplastic or autoimmune diseases. We further enrolled 20 healthy subjects (mean age 39 years, ranging from 27 to 61) who had no risk factors such as hypertension, diabetes mellitus and dyslipidemia, and were treated with no medications (Control subjects). All control subjects gave written, informed consent before the study.

Patients treated with antihypertensive agents or those with blood pressure > 140/90 mmHg were diagnosed as having hypertension. Diabetes mellitus was diagnosed according to the criteria of the World Health Organization11. Body mass index (BMI) was calculated and was used to stratify obesity degree. Insulin sensitivity was assessed by homeostasis model assessment (HOMA) index. Serum total cholesterol, low-density lipoprotein (LDL) cholesterol, high-density lipoprotein (HDL) cholesterol, triglyceride, and fasting blood sugar concentrations were measured using standard methods. The plasma level of B-type natriuretic peptide (BNP) was measured by immunoradiometric assay (Shionoria BNP RIA kit, Tokyo, Japan).
2.2. Blood sampling and measurement of plasma apelin

Blood samples were obtained from the peripheral vein before the induction of general anesthesia and the infusion of any perioperative metabolic substrates at surgery. After centrifugation at 3000 × g for 15 min at 4 °C, the supernatants were stored at -80 °C until use. Plasma apelin level was determined using a commercially available enzyme immunoassay (Phoenix Pharmaceuticals, CA, USA) according to the manufacturer’s instruction.

2.3. Adipose tissue sampling

Paired samples of the visceral and subcutaneous adipose tissues were harvested during surgery. In this study, adipose tissue beneath the sternum including the thymus tissue replaced by adipose tissue was used as the visceral adipose tissue. To measure the gene expression, the sample was immediately submerged in RNealater™ for storage to stabilize and protect cellular RNA. These samples were stored at -30 °C until use.

2.4. Measurement of gene expression of apelin in adipose tissues

Adipose tissue RNA was extracted using a RNeasy Lipid Tissue minikit (Qiagen, GmbH, Hiden, Germany), according to the manufacturer’s instruction. A two-step real time reverse transcription-polymerase chain reaction (RT-PCR) was carried out according to the instruction supplied with the TaqMan Gold RT-PCR kit (Applied Biosystems Foster City, CA, USA). All oligonucleotide primers and TaqMan probes for human apelin and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) were purchased from Applied Biosystems. Values were averaged from duplicate data and normalized with human GAPDH.

2.5. Statistical analysis

All data were expressed as mean ± SEM. The data were analyzed with unpaired Student’s t test between two groups. One way analysis of variance (ANOVA) was used for the comparison of 3 or more variables followed by Tukey-Kramer’s post hoc test. Probability values < 0.05 were considered to be statistically significant.

3. Results

3.1. Clinical characteristics of subjects

None of age, gender, BMI, and risk factors such as hypertension, diabetes mellitus, smoking, and medications including angiotensin converting enzyme inhibitor (ACE-I), angiotensin II type 1 receptor blocker (ARB), and diuretics differed between two groups (Table 1). Aspirin, β-blockers, and HMG-CoA reductase inhibitors (statins) were less administered to VHD patients compared with CAD patients. The incidence of dyslipidemia was

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Comparison of clinical profiles</th>
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<tbody>
<tr>
<td></td>
<td>CAD (n=31)</td>
</tr>
<tr>
<td>Age</td>
<td>66 ± 2</td>
</tr>
<tr>
<td>Male (n, %)</td>
<td>28 (90%)</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>24.4 ± 0.6</td>
</tr>
<tr>
<td>Risk factor (n, %)</td>
<td></td>
</tr>
<tr>
<td>hypertension</td>
<td>11 (87%)</td>
</tr>
<tr>
<td>diabetes mellitus</td>
<td>13 (42%)</td>
</tr>
<tr>
<td>dyslipidemia</td>
<td>29 (94%)**</td>
</tr>
<tr>
<td>smoking</td>
<td>16 (52%)</td>
</tr>
<tr>
<td>Medication (n, %)</td>
<td></td>
</tr>
<tr>
<td>aspirin</td>
<td>31 (100%)**</td>
</tr>
<tr>
<td>ACE-I</td>
<td>4 (13%)</td>
</tr>
<tr>
<td>ARB</td>
<td>20 (65%)</td>
</tr>
<tr>
<td>β-blocker</td>
<td>26 (84%)**</td>
</tr>
<tr>
<td>statin</td>
<td>19 (61%)**</td>
</tr>
<tr>
<td>diuretics</td>
<td>14 (45%)</td>
</tr>
<tr>
<td>LVEF (%)</td>
<td>48 ± 2*</td>
</tr>
</tbody>
</table>

CAD, patients with coronary artery disease; VHD, patients with valvular heart disease; ACE-I, angiotensin converting enzyme inhibitor; ARB, angiotensin II type 1 receptor blocker; BMI, body mass index; LVEF, left ventricular ejection fraction; statin, HMG-CoA reductase inhibitor

Variables are shown as mean ± S.E.M.

* P<0.05 vs VHD, ** P<0.01 vs VHD
higher in CAD patients than in VHD patients. Left ventricular ejection fraction (LVEF) assessed by left ventriculography (LVG) was lower in CAD than in VHD. There were no differences in the serum levels of total cholesterol, triglyceride, blood sugar, HbAlc, HOMA index, BNP and high sensitive C-reactive protein (hs-CRP) between two groups (Table 2). Serum levels of LDL-cholesterol and HDL-cholesterol in CAD patients were lower than those in VHD patients.

### 3.2. Level of plasma apelin

As shown in Figure 1, the plasma apelin level was significantly lower in CAD and VHD patients compared with Control subjects, respectively (p<0.01). Moreover, its level was significantly lower in CAD patients compared with VHD patients (p<0.05). There was no difference in plasma apelin level between CAD patients with and without β-blockers (β-blockers (+); 0.41 ± 0.06 vs (-); 0.45 ± 0.06 ng/ml). Statins also did not affect plasma apelin level in CAD patients (Statin (+); 0.41 ± 0.06 vs (-); 0.43 ± 0.11 ng/ml).

### 3.3. Gene expression of apelin in the visceral and subcutaneous adipose tissues

As shown in Figure 2, apelin gene expression was higher in visceral adipose tissue than in subcutaneous adipose tissue in VHD patients (P<0.05). The gene expression in visceral adipose tissue tended to be higher than that in subcutaneous tissue in CAD patients (P=0.1). There was no difference in apelin gene

**Figure 1** Comparison of the plasma apelin level among the patients with coronary artery disease (CAD) and valvular heart disease (VHD), and healthy subjects (Control). The plasma apelin level was lower in CAD and VHD patients than that in Control subjects respectively, and its level was lower in CAD patients than that in VHD. *P<0.05, **P<0.01.

**Figure 2** Gene expressions of apelin normalized by human glyceraldehydes-3-phosphate dehydrogenase (GAPDH) in the visceral and subcutaneous adipose tissues. Boxes show medians and interquartile ranges, with whiskers representing the 10th/90th percentiles. *P<0.05.
expression in visceral and subcutaneous adipose tissue between two groups, respectively.

3.4. Relationship between left ventricular function and plasma apelin level

There were no significant correlations between plasma apelin level and plasma BNP level (Figure 3) or LVEF (Figure 4).

4. Discussion

4.1. Plasma apelin level and coronary atherosclerosis

The major findings of this study were that the plasma apelin level was significantly lower in CAD and VHD patients than that in Control subjects respectively, and that its level was lower in CAD patients than that in VHD patients. Our result that BNP level was increased in CAD and VHD patients respectively supports the previous findings that plasma apelin level is decreased in patients with CHF$^{19}$. We also found that LVEF in CAD patients was lower than that in VHD patients. A recent study showed a positive correlation between LVEF and plasma apelin level in patients with CHF$^{3}$. It seems reasonable to presume that the difference of left ventricular function explains the finding that plasma apelin level in CAD patients was lower than that in VHD patients, although there was no significant correlation between apelin level and LVEF. The disparity of the relation between apelin level and LVEF may be dependent on the small number of subjects and the administration of medicines for CHF. Second, the decrease in plasma apelin level in CAD patients is likely to be dependent on coronary atherosclerosis. Recently, Li et al. showed that the plasma apelin level in patients with stable angina pectoris was lower than in controls$^{3}$. Since VHD patients were confirmed to have no CAD by coronary angiography, the difference in the degree of coronary atherosclerosis between CAD and VHD patients might be associated with the plasma apelin level. Our result that hs-CRP tended to be higher in CAD patients than in VHD patients also supports this idea. Overall, plasma apelin level appears to be a useful predictor of the progression of coronary atherosclerosis.

The difference of medication between two groups might affect the plasma apelin level. However, there was no difference in plasma apelin level between CAD patients with and without β-blockers. Statins also did not affect plasma apelin level in CAD patients. Recent study has shown that LDL-cholesterol lowering in healthy subjects with isolated dyslipidemia results in an increase in plasma apelin level$^{14}$. Therefore, the decrease in plasma apelin level is
independent of the difference in medication.

4.2. Gene expression of apelin in adipose tissue

We investigated apelin gene expression in visceral and subcutaneous adipose tissues, but no difference was found between two groups. Therefore, the decrease in plasma apelin level is independent of apelin gene expression in adipose tissue. Apelin was identified from bovine stomach extracts for the first time and was known existing in various tissues such as central nervous system, heart, lung, stomach, pancreas, adrenal gland and adipose tissue. Boucher et al. showed that the gene expression of apelin in the heart and the kidney is higher than that in adipose tissue. Moreover, recent study showed that apelin gene expression is 200-fold higher in atrial tissue than in ventricular tissue, suggesting that the heart may be the major source of circulating apelin in humans. There was no difference in apelin gene expression in visceral and subcutaneous adipose tissue between two groups, respectively. However, to our knowledge, this is the first report indicating that gene expression of apelin in visceral adipose tissue is higher than that in subcutaneous adipose tissue in humans.

4.3. Apelin as the therapeutic target of chronic heart failure

The plasma apelin level in CAD patients was decreased compared with that in VHD patients, and LVEF in CAD patients was also lower than that in VHD patients. In rats with CHF due to post-myocardial infarction, apelin infusion restores EF and increases cardiac output. Therefore, apelin might be useful for not only the predictor of coronary atherosclerosis but also the therapeutic target of CHF. Further studies are needed to establish the effectiveness and safety of apelin infusion for the therapy of CHF in humans.

4.4. Study limitations

We showed that plasma apelin level was decreased in patients with CAD, but the major source of circulating apelin still remains unclear. Since the heart is a possible candidate for the major source, further studies for apelin expression in the heart are needed.

5. Conclusions

Plasma apelin level in CAD patients was lower than that in VHD patients, and it may be associated with coronary atherosclerosis. Gene expression of apelin in visceral adipose tissue was higher than that in subcutaneous adipose tissue in humans. However, adipose tissue might not be the major source of plasma apelin.

References

3) Chong KS, Gardner RS, Morton JJ, Ashley EA, McDonagh TA. Plasma concentrations of the novel peptide apelin are decreased in patients with chronic heart failure, Eur J Heart Fail 2006;8:355-60.


