

Original articles

Increased *DAPK1* but decreased *CCL2* plasma levels of nucleic acids in patients with stable angina

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Abstract

Introduction: We hypothesized that patients with stable angina have increased plasma levels of mRNA from genes responsible for atherosclerotic plaque development and destabilisation, *i.e.* from death-associated protein kinase (*DAPK1*) and monocyte chemotactic protein-1 (*CCL2*).

Materials and methods: Nucleic acids were isolated from plasma of patients with stable angina and healthy subjects as controls. mRNAs were transcribed to cDNAs, quantified by real-time PCR and standardized to the amount of a reference gene. Reagents for PCR quantification are declared to be mRNA specific, but in our test conditions DNA was found to interfere in both assays.

Results: Patients had 5.1-times higher plasma level of *DAPK1* nucleic acids (mRNA and DNA) than controls ($P < 0.001$) and the highest levels were associated with the presence of diabetes. However, plasma levels of *CCL2* tended to be lower than in controls, and in statin-treated patients the decrement reached significance (-66.3%; $P = 0.041$).

Conclusion: The estimated levels are explicable in terms of current knowledge. Further studies with specific assays for mRNA PCR quantification are reasonable to access whether this approach offers non-invasive *in vivo* assessment and monitoring of gene expression profile in atherosclerotic vascular beds.

Key words: *DAPK1*; *CCL2*; cell-free nucleic acids; plasma; stable angina

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Introduction

Death-associated protein kinase (encoded by *DAPK1*) is a proapoptotic, calmodulin-regulated serine/threonine kinase that carries a C-terminal death domain. Apoptosis has been identified as a prominent feature of atherosclerosis (1) and *DAPK1* mRNA and *DAPK1* are increased in carotid atherosclerotic plaques (2). Monocyte chemotactic protein-1 (encoded by *CCL2*) is involved in the formation, progression, and destabilization of atherosclerotic plaques (3). A follow-up study in hypercholesterolemic rabbits showed that the increase of *CCL2* mRNA in arterial wall correlated with atherosclerotic plaque development (4).

Cell-free nucleic acids are nucleic acids (DNA, mRNA) present in plasma, serum and cell-free fractio-

ns of various other biological fluids (5). As to coronary artery disease, the prognostic value of nucleic acids in the plasma of patients with acute coronary syndrome has been established (6,7) and patients with coronary atherosclerosis have increased plasma cathepsin S mRNA levels (8). We proposed that it should be possible to measure *DAPK1* mRNA and *CCL2* mRNA levels, *i.e.* from genes responsible for plaque development and destabilization, in plasma of patients with coronary atherosclerosis, and we hypothesized that these levels are increased. If the answers were positive, it would mean that plasma mRNA estimation could offer non-invasive *in vivo* assessment and monitoring of gene expression profile in atherosclerotic vascular beds.

Materials and methods

Patients

Materials and methods (8) and optimisation of protocols (9,10) were in detail described previously. Briefly, patients were consecutively enrolled from those with stable angina or acute coronary syndrome undergoing routine coronary angiography. Stable angina and acute coronary syndrome were diagnosed by clinical assessment, EKG evaluation and laboratory findings. Inflammatory, thyroid, liver, endocrine, neoplastic and renal diseases (serum creatinine concentration values $\geq 120 \mu\text{mol/L}$) were excluded by detailed history and clinical examination. The study was approved by the national ethics committee. Informed consent was obtained from all participating subjects.

Methods

Peripheral blood was taken into EDTA tubes (Becton Dickinson, Plymouth, UK) after at least 12 hours of fasting and before coronary angiography. To reduce the bias resulting from instability of mRNA the same double-spin procedure was used in all subjects to obtain plasma nucleic acids. Sample processing took place within one hour of collection. Haemolysed samples were not used for analysis.

RNA was isolated from plasma using QIAamp MinElute Virus Spin Kit (Qiagen, Hilden, Germany) according to the manufacturer's recommendations, but with the following modifications: 1) the incubation time for proteolysis was 5 min; 2) RNA was eluted from the column with 55 μL RNase-free water and the collected eluate used for a second elution of RNA from the same column. Standard methods (A260/A280 on a NanoDrop or Agilent RNA 6000 Nano kit) are inappropriate for assessment of RNA yield and quality in the collected eluate because RNA carrier that is used in the process of plasma RNA isolation yields the signal, which is too dominant relative to the signal of RNA species of interest. As a consequence, RNA amounts based on starting volume rather than RNA quantity is used in all analytical procedures. Eluted mRNA was transcribed to cDNA with SuperScript VILO cDNA

Synthesis Kit (Invitrogen, Carlsbad, CA, USA). First, RNA eluate was incubated in a water bath (65 °C) for 5 min, then immediately placed on ice for another 5 min. The manufacturer's protocol was followed from then on.

The following TaqMan Gene Expression Assays (Applied Biosystems, Foster City, CA, USA) were used for cDNA quantification: Hs00234489_m1 for *DAPK1* (exon boundary 3-4, amplicon length 105), Hs00234140_m1 for *CCL2* (exon boundary 1-2, amplicon length 101) and Endogenous Control 4326316E for cyclophilin A (*PPIA*). cDNA was amplified by PCR according to the manufacturer's protocol (Roche Diagnostics, GmbH, Mannheim, Germany), using a 1.5 μL (*DAPK1* and *PPIA*) or 3.75 μL (*CCL2*) of the cDNA solution and the kit components in a reaction volume of 25 μL . Although the assay kits for cDNA quantification are declared to be mRNA specific, in our test conditions DNA was found to interfere in both assays. All reactions were run in triplicate and the average threshold cycle of triplicates (Ct) calculated. Between-run imprecision for the whole analytical procedure (from isolation to quantification), expressed as coefficient of variation of Ct, was less than 2%. To exclude the possibility of external nucleic acids contamination negative control (blank) was analysed in each PCR run. For clinical evaluation, the amount of each investigated transcript was standardized to the amount of a reference gene. *PPIA* was chosen as the most appropriate reference gene among 11 candidates available on TaqMan Human Endogenous Control Plate (Applied Biosystems), based on the smallest geometric Ct of individual candidates and results from geNorm, as in detail described previously (9).

Statistical analyses

Data are presented as counts; age is presented as median and range. Statistically significant difference between patients and controls for quantitative variables (age and plasma levels of nucleic acids) was tested using nonparametric Mann-Whitney test.

P values < 0.05 were considered statistically significant.

Statistical analyses were performed using SPSS 17.0 statistical software package (SPSS, Inc., Chicago, IL, USA).

Results

The clinical characteristics of the participants are summarized in Table 1. Healthy subjects were younger (median 55 vs. 61 years; $P = 0.029$). Age did not correlate with any observed parameter within either patient or control group (data not shown).

Although the assay kits for cDNA quantification are declared to be mRNA specific, in our test conditions DNA was found to interfere in *DAPK1* and *CCL2* assay, as confirmed by: 1) analysis of the size of the amplified PCR products on agarose gel electrophoresis; 2) running a PCR reaction with human DNA; and 3) running a PCR reaction with subjects' samples but omitting the reverse transcription (data not shown). The DNA interference was confirmed after finishing all the experiments. Interestingly, the new lot number of reagents purchased after informing the producer about the uncertainty, no longer showed DNA interference (data not shown). Because we used all cDNA samples, it was not possible to repeat the measurements with the new lot number of reagents. On the other hand, it was not possible to study in detail the relative contribution of DNA and cDNA to the signals obtained with the old lot number of reagents, because we used all of them.

The laboratory findings are summarized in Figure 1. *DAPK1* and *CCL2* plasma levels of nucleic acids (mRNA and DNA) were successfully determined in all patients and in 31 (out of 33) controls. Patients had, on average, 5.1-times higher plasma levels of *DAPK1* nucleic acids than controls (median and interquartile range 33.82 (11.40-77.44) vs. 6.63 (2.87-11.47); $P < 0.001$). Patients with diabetes had 5.5-times higher levels than patients without diabetes (68.23 (34.12-99.18) vs. 12.38 (7.09-35.48); $P = 0.006$), and the latter still had 86.7 % higher levels than the controls (12.38 (7.09-35.48) vs. 6.63 (2.87-11.47); $P = 0.014$). On a contrary, plasma levels of *CCL2* nucleic acids tended to be lower in patients

TABLE 1. Clinical characteristics of subjects.

Parameter	Stable angina patients	Healthy subjects
Gender (males; N/total)	16/29	21/33
Age (years)	61 (46-83)*	55 (38-81)*
Family history of CAD (N/total)	7/29	0/33
Smoking status (current or former; N/total)	8/29	0/33
Diabetes (N/total)	12/29	0/33
Coronary arteriography (number of diseased vessels)		/
0	2	
1	16	
2	4	
3	7	
Statin therapy (N/total)	24/29	0/33
Aspirin therapy (N/total)	25/29	0/33
ACE inhibitor therapy (N/total)	17/29	0/33

* median (min-max); CAD - coronary artery disease.

ts than in controls (12.55 (6.77-38.61) vs. 36.25 (5.90-74.54); $P = 0.062$) and, when only statin-treated patients were compared to the controls, the decrease reached the significance (-66.3%; 12.21 (6.41-26.60) vs. 36.25 (5.90-74.54); $P = 0.041$). It is noteworthy that 22 statin-treated patients (out of 24) were receiving aspirin and 16 patients ACE inhibitors.

Levels of nucleic acids in patients' plasma did not correlate with the number of diseased coronary arteries or with any other observed parameter. In the controls, plasma levels of nucleic acids correlated neither with plasma lipids nor with any other observed parameter. Importantly, plasma levels of nucleic acids did not correlate with leukocyte or platelet counts.

Discussion

Our study shows the existence of cell-free nucleic acids from *DAPK1* and *CCL2*, i.e. from genes responsible for atherosclerotic plaque development and

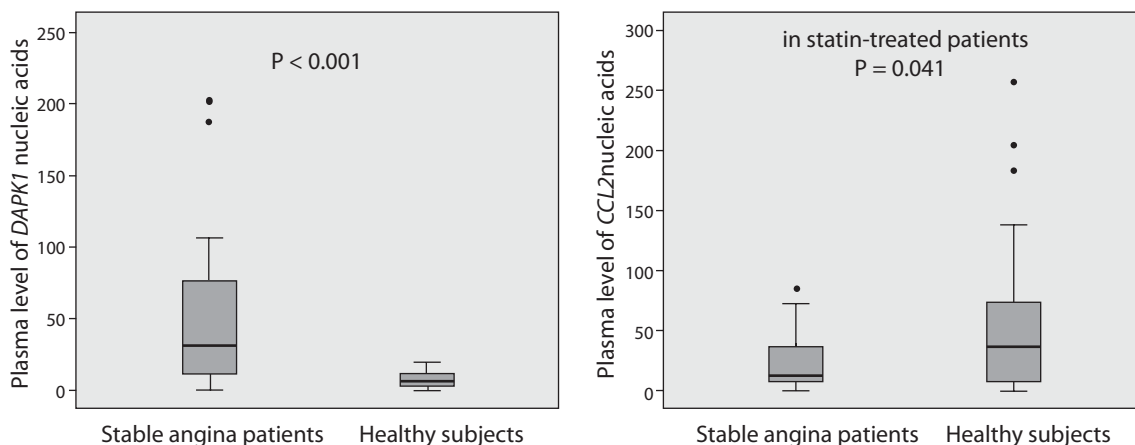


FIGURE 1. Plasma *DAPK1* and *CCL2* nucleic acids levels in patients with stable angina (N = 29) and in healthy subjects (N = 33). Plasma levels were standardized to the amount of reference gene (*PPIA* mRNA). Box plots demonstrate interquartile range (25th and 75th percentiles), and line inside the box denotes the median bar. The whiskers demonstrate the largest and the smallest values inside the 1.5 times interquartile range. Outliers (•) are the values between the 1.5 and the 3 times interquartile range. Extreme values are not presented.

destabilization, in plasma of patients with coronary atherosclerosis. Plasma nucleic acids analysis is an emerging field for non-invasive *in vivo* assessment and monitoring of pathologic processes in tissues of their origin. It represents an alternative or a supplement to protein analysis. Nowadays, quantitative and qualitative analysis of cell-free nucleic acids is considered to be clinically useful in non-invasive prenatal diagnosis, and in diagnosis and monitoring of numerous cancers (5), but novel applications are emerging also in other critical care settings, such as acute coronary syndrome (6–8,11). These nucleic acids can be measured precisely in only 200 µL of plasma of patients with stable angina as well as of healthy subjects (8,11). We believe that with the use of higher plasma volumes for nucleic acids isolation (commercial kits are available) and with further improvement of specificity of assays for PCR quantification there will be possible to reliably measure plasma mRNA from *DAPK1* and *CCL2*, thus offering a new technique for non-invasive *in vivo* assessment and monitoring of gene expression profile in vascular beds.

Although conclusions are not as expected and DNA interference introduces uncertainty to the data, the results of our study are interesting and explicable in the light of current knowledge. High

plasma levels of *DAPK1* nucleic acids in patients with stable angina are consistent with the current evidence that apoptosis is a prominent feature of atherosclerotic plaque, especially in regions with a high density of macrophages (1,12). Apoptosis is an important source of nucleic acids (mRNA and DNA) in plasma (5). Furthermore, *DAPK1* is a well known positive mediator of apoptosis in a wide array of apoptotic systems (13) and *DABK1* was found to be highly expressed in carotid atherosclerotic plaques (2). Finally, prolonged hyperglycaemia and/or impaired insulin signalling were also shown to induce apoptosis (14). In this respect it should be noted that our study design does not exclude other sources of *DAPK1* nucleic acids. *DAPK1* nucleic acids containing apoptotic bodies in non-vascular tissues may escape the elimination by reticuloendothelial system in those tissues and thus protrude into the blood. This could be an additional reason for extremely high plasma *DAPK1* nucleic acids levels in our patients with diabetes. Unexpectedly, stable angina patients had lower plasma levels of *CCL2* nucleic acids than controls. Here it is noteworthy that if DNA had had a major contribution to the signals obtained, it would have been difficult to explain the divergent results for plasma nucleic acids levels from *CCL2* and *DAPK1*. Lower plasma nucleic acids levels from *CCL2* in patients with

stable angina could be explained as a consequence of statin pleiotropy (15). Twenty-four patients (out of the 29) were receiving statin therapy. Statin-treated patients had lower plasma concentrations of CCL2 than those without statin therapy (16), and statin treatment significantly diminished neointimal and medial CCL2 in a rabbit model of atherosclerosis (17). Finally, statin treatment reduced CCL2 mRNA levels in peripheral blood monocytes *in vivo* (18).

To conclude, cell-free nucleic acids from genes responsible for atherosclerotic plaque development and destabilization are possible to quantify in plasma of patients with stable angina and of healthy

subjects and the estimated levels are explicable in terms of current knowledge. Further studies with specific assays for PCR quantification are reasonable.

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Potential conflict of interest

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Povišena razina DAPK1 te snižena razina CCL2 nukleinskih kiselina u plazmi kod bolesnika sa stabilnom anginom pektoris

Sažetak

Uvod: Postavili smo hipotezu da bolesnici oboljeli od stabilne angine pektoris imaju povišenu razinu mRNA gena odgovornih za stvaranje i destabilizaciju aterosklerotskog plaka, odnosno iz proteinske kinaze povezane sa smrću stanice (engl. *death-associated protein kinase, DAPK1*) i monocitnog kemotaktskog proteina-1 (engl. *monocyte chemotactic protein-1, CCL2*) u plazmi.

Materijali i metode: Iz plazme bolesnika oboljelih od stabilne angine pektoris te zdravih ispitanika izolirali smo nukleinske kiseline. Učinjena je transkripcija molekula mRNA u cDNA, te su kvantificirane lančanom reakcijom polimerazom u stvarnom vremenu (engl. *real-time PCR*) i standardizirane na količinu referentnog gena. Reagensi za kvantifikaciju metodom PCR deklarirani su kao specifični za mRNA, no u našim uvjetima ispitivanja ustanovili smo da DNA interferira u oba testa.

Rezultati: Razina *DAPK1* nukleinskih kiselina (mRNA i DNA) u plazmi bila je 5,1 puta viša kod bolesnika nego kod kontrolnih ispitanika ($P < 0,001$), a najviše su razine bile povezane sa šećernom bolesti. Razina *CCL2* u plazmi nije bila statistički značajno niža kod skupine bolesnika nego kod kontrolnih ispitanika, dok je kod bolesnika liječenih statinima sniženje razine postalo statistički značajno (- 66,3%; $P = 0,041$).

Zaključak: Dobivene se vrijednosti mogu objasniti trenutnim spoznajama. Potrebna su daljnja istraživanja specifičnim testovima za kvantifikaciju mRNA metodom PCR kako bi se moglo procijeniti pruža li taj pristup neinvazivno *in vivo* određivanje i monitoring razine ekspresije gena u aterosklerotskim lezijama.

Ključne riječi: CCL2; DAPK1; slobodne nukleinske kiseline; plazma; stabilna angina pektoris