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Original Article Rapid changes in plaque composition and morphology after intensive lipid lowering therapy: study with serial coronary CT angiography

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Abstract: Although intensive lipid lowering by statins can enhance plaque stability, few data exist regarding how early statins change plaque composition and morphology in clinical setting. Therefore, to examine early changes in plaque composition and morphology in clinical setting. Therefore, to examine early changes in plaque composition and morphology by intensive lipid lowering with statins, we evaluate coronary plaques from acute coronary syndrome (ACS) before and 3 weeks after lipid lowering by coronary CT angiography. We enrolled 110 patients with suspected ACS and underwent coronary CT. We defined plaque as unstable when CT number of plaque< 50HU and remodeling index (lesion diameter/reference diameter) >1.10. Rosuvastatin (5 mg/day) or atorvastatin (20 mg/ day) were introduced to reduce low density lipoprotein cholesterol (LDL-C). Then, CT was again performed by the same condition 3 weeks after lipid lowering therapy. Total 10 patients (8 men, mean age 72.0 years), in whom informed consent regarding serial CT examination was obtained, were analyzed. Among them, 4 patients who denied to have intensive lipid lowering were served as controls. In remaining 6 patients, LDL-C reduced from 129.5 \pm 26.9 mg/dl to 68.5 \pm 11.1 mg/dl after statin treatment. Under these conditions, CT number of the targeted plaque significantly increased from 16.0 \pm 15.9 to 50.8 \pm 35.0 HU (p<0.05) and remodeling index decreased from 1.22 \pm 0.11 to 1.11 \pm 0.06 (p<0.05), although these values substantially unchanged in controls. These results demonstrate that MDCT-determined plaque composition as well as volume could be changed within 3 weeks after intensive lipid lowering. This may explain acute effects of statins in treatment of acute coronary syndrome.

Keywords: Computed tomography, HMG-CoA reductase inhibitor, plaques, acute coronary syndrome

Introduction

Acute coronary syndromes (ACS) such as unstable angina and acute myocardial infarction is caused by the rupture of the atherosclerotic plaque and subsequent thrombosis [1]. Under these conditions, antiplatelet drugs, angiotensin converting enzyme inhibitor, β -blocker and HMG -CoA reductase inhibitors (statins) were shown to decrease the secondary cardiovascular event significantly [2-6]. Particularly statins strongly decrease low-density lipoprotein-cholesterol (LDL-C) and stabilize vulnerable plaque [6, 7]. Indeed, early use of the statin in ACS decreased the plaque volume determined by intravascular ultrasound 6 months after treatment [8]. In not only ACS but also in stable coronary syndrome,

we previously reported that statins reduced coronary plaque volume in Japanese patients [9]. However, few data exist regarding how early statins can change plaque composition and volume in clinical settings.

Recently, coronary plaque composition can be evaluated with not only invasive methods such as IVUS and optical coherence tomography but also with non-invasive one such as coronary CT angiography (CCTA) [10, 11]. Previous studies demonstrated CCTA exhibited good agreement with intravascular ultrasound in terms of evaluating plaque composition [12, 13]. In addition, repeated examination in short term could be done by CCTA except problems regarding use of contrast agent and radiation exposure. The aim

	Age	Sex	LDL(mg/dl)	HDL(mg/dl)	CT attenuation(HU)	Remodeling index
1	75	male	102	66	21.26	1.13
2	71	male	111	25	11.25	1.12
3	63	male	146	40	-8.43	1.36
4	76	male	113	39	17.00	1.36
5	71	male	174	43	14.25	1.20
6	79	female	131	57	40.83	1.13
72.0±5.6			129.5±26.9	45±14.5	16. 0±15.9	1.22±0.11

Table1. Patients' characteristics before statin treatment

of this study was to evaluate changes in plaque composition and morphology in early phase such as 3 weeks after intensive lipid lowering by CCTA.

Materials and methods

Participants and study design

We enrolled 110 patients who were suspected of ACS and were examined by 64-slice CCTA from Janurary 2009 to March 2011. Base line characteristics included age, sex, LDL-C, highdensity lipoprotein cholesterol (HDL-C) and LDL-C/HDL-C. When unstable plaque defined as described below was detected by CCTA, we administered statins such as rosuvastatin of 5mg/day or atorvastatin of 20 mg/day and re-evaluated the previously-determined coronary plaques 3 weeks later.

Image acquisition

The 64-slice CCTA scanning (Light Speed VCT, GE, USA) was performed with retrospective ECG gating. β-blocker such as propranolol of 20mg to 60mg was injected before scanning to decrease heart rate to 60 beats/min in average. Nitroglycerin of 0.3 mg was used for coronary artery dilatation. All image acquisitions were performed during a single breath-hold after inspiration. Scanning was performed with condition of effective tube current from 220 mA to 780 mA. effective tube voltage of 120 kV. 0.625 mm slice acquisition, and gantry rotation time of 350 ms. We performed contrastenhanced scanning using 0.8 ml/kg contrast medium (lopamidol 370, Bayer Healthcare, Berlin, Germany) during an inspiratory breathhold.

Image analysis

CCTA images were analyzed using commercially

available soft ware (Advantage Workstation Volume Share 4.3). For plaque detection, both cross-sectional view, stretched view and curved multiplanar reformation images were used for analysis. Under these conditions, we chose the coronary segment with $\geq 2 \text{ mm}$ in diameter. The coronary arterial remodeling was defined as a ratio of vessel diameter at the plaque site in comparison with that in the reference segment set proximal to the lesion in a normal-appearing vessel segment (reference segment). The remodeling index (lesion diameter/reference diameter) in cross-sectional view on CCTA was calculated and reported as positive remodeling when the diameter at the plaque site was at least 10% larger than the reference segment [13, 14]. We defined plaque attenuation as minimum CT density of the average of each 3 slices, and defined plaques as unstable when CT value ≦50HU [10]. Three weeks after statin therapy, same evaluation before statin therapy was repeatedly performed for follow up patients.

Data analyses

The quantitative variables are presented as mean \pm standard deviation and categorical data are shown as proportions. Mean values were compared using the paired t test. P-value < 0.05 indicated a significant difference.

Results

Among enrolled 110 patients, informed consent regarding the intensive statin therapy and repeated CCTA examination in 3 week was obtained in 6 patients (5 men, mean age 72.0 years, **Table 1**). The remaining 4 agreed with only repeated CT examination without intensive lipid lowering. There were 2 patients with diabetes mellitus, one patient with hypertension, 4 patients with hyperlipidemia and 2 patients with habit of smoking. There were 3 patients prescribing rosuvastatin and 3 patients prescribing

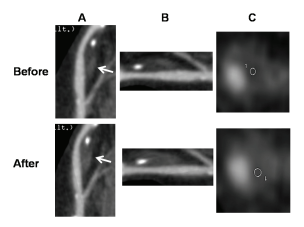


Figure 1. Coronary CT images of curved view (A), stretched view (B), and cross sectional view (C) before and after intensive lipid lowering therapy. Arrows indicate the target lesion and circles indicate the region of interests.

atorvastatin. Six of all had coronary plaques which were characterized by positive remodeling and low attenuation.

Representative case

A 63 year-old Japanese male had acute myocardial infarction and performed percutaneous coronary intervention for left anterior descending coronary artery. We performed CCTA after PCI and atorvastatin of 20 mg was administrated. Three weeks after statin therapy, LDL-C decreased from 146mg/dl to 71mg/dl, HDL-C increased from 33mg/dl to 40mg/dl, yielding reduced LDL-C/HDL-C from 3.86 to 2.15. Under these conditions, cross sectional view on MDCT showed the increased signal of target lesion from -8 HU to 39 HU with decreased remodeling index from 1.36 to 1.19 (**Figure 1**).

A В С 200 120 1 50 180 100 160 Remodeling index 1.30 CT attenuation (HU) 80 140 -DL-C (mg/dl) 120 60 100 1.10 40 80 20 60 0.90 40 0 20 p<0.05 p<0.05 p<0.05 0 70 -20 0 Before After Before After Before After

Lipid profiling and plaque analysis

In all 6 patients, LDL-C decreased significantly from 129.5 \pm 26.9 mg/dl to 68.5 \pm 11.1 mg/dl, or by 47% (p<0.05) (**Figure 2**). HDL-C substantially unchanged from 39.2 \pm 8.8 mg/dl to 45.0 \pm 14.5 mg/dl. Thus, LDL-C/HDL-C decreased significantly from 3.14 \pm 1.10 to 1.80 \pm 0.39, or by 42.6%.

CT number of the plaque increased significantly from 16.0 \pm 15.9 HU to 50.8 \pm 35.0 HU (p<0.05) and remodeling index decreased from 1.22 \pm 0.11 to 1.11 \pm 0.06 (p<0.05) (**Figure 2**). In controls, there were no significant changes in these values during 3 weeks. Although it is interesting to correlate changes in LDL-C levels and those in plaque composition and/or morphology, there were no correlations between changes in LDL-C and CT values in the present study.

Discussion

The present study demonstrates that CCTAdetermined plaque composition and morphology can be changed in only 3 weeks after starting intensive lipid lowering with statins. This may explain early effects of statin-induced intensive lipid lowering on prevention of cardiovascular events early after the occurrence of ACS.

Previous clinical trials showed that clinical benefit of statin therapy was observed within one month in patients with ACS [15-17]. However, few data exist demonstrating changes in plaque composition and morphology in early phase after statin therapy except findings observed by carotid ultrasound and virtual histology ultra-

> sound [15, 16]. The present study demonstrates that intensive LDL-C reduction by 47% in average can cause changes in CCTA-determined plaque composition and morphology within only 3 weeks.

Figure 2. Changes in low density lipoprotein-cholesterol (LDL-C) (A), in CT attenuation; (B) and in remodeling index (C) before and after intensive lipid lowering therapy Open circles indicate mean values of each data. In the experimental study, high cholesterolinduced atherosclerotic endothelial dysfunction could be diminished in 2 weeks after statin treatment [18]. In the present study, there were no correlations between changes in LDL-C levels and those in plaque morphology and compositions probably due to small number of patients, although previous invasive studies demonstrated correlation between them [8, 17]. This may be explained by the period of repeated CT examination is too short to obtain the mature changes in plaque morphology and composition.

In the evaluation of plaques, we used CCTA instead of intravascular ultrasound or optical coherence tomography both of which were already established as methods for plaque evaluation. It is reasonable to consider that CCTA was apparently less invasive than intravascular imaging techniques. In addition, recent studies showed the plaque exhibiting positive remodeling and low-attenuation on CCTA was consistent with vulnerable plaque on intravascular ultrasound [12, 19, 20]. Soeda et al evaluated coronary plaque with CCTA before and after statin therapy for 24 weeks [21], and found that CTdetermined plaque composition could be changed 24 weeks or 6 months after statin therapy.

Clinical implications and limitation

The most important clinical implication of the present study is how rapid plaque composition and/or morphology could change after intensive lipid lowering. Although experimental studies indicated that atherosclerotic plaque composition changes quickly after discontinuation of high cholesterol diet or stating statin therapy, this should take longer time in clinical setting. Indeed, previous imaging studies were performed by evaluating plaque at least 3 months for intervals. Particularly for ACS patients, we should give statins to establish intensive lipid lowering for 3 weeks, and plaque stabilization can be obtained.

There remain, however, several limitations in the present study, First, the study size was small mainly due to difficulty to obtain the informed consent for the present study in which repeated CT examinations with radiation exposure after intensive lipid lowering therapy were performed. However, it is evident that CCTAdetermined plaque composition and morphology could be changed rapidly after intensive lipid lowering in ACS patients, although a largescale randomized control study should be performed to confirm our result. Second, although spatial resolution of 64-slice CCTA has been improved compared to previous CT system, partial volume effect and CT attenuation of coronary lumen influence the CT attenuation of target plaque. We expect more advancement in CT technology to improve spatial resolution, which enable the evaluation of the plaque component precisely.

Conclusions

In the present study, we demonstrate that CCTAdetermined plaque composition as well as morphology could be immediately changed after starting intensive lipid lowering in ACS. We suggest that in ACS coronary narrowing as well as plaque composition and morphology should be evaluated by CCTA for possible indication of intensive lipid lowering therapy in these cohorts.

Declearation

None

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