

Serial optical coherence tomography imaging of ACS-causing culprit plaques

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KEYWORDS

- acute coronary syndrome
- optical coherence tomography
- plaque erosion
- plaque rupture

Abstract

Aims: The aim of this study was to understand better the mechanisms of repair of plaque complications causing acute coronary syndrome.

Methods and results: We used OCT in the acute phase and at follow-up (one to seven months) to investigate the plaque healing in 10 culprit plaques: five ruptured fibrous cap (RFC) and five intact fibrous cap (IFC) which were not treated with stent deployment and caused ST-segment elevation myocardial infarction (n=8) and non-STEMI (n=2). At follow-up OCT, the margins of the evacuated cavity in RFC plaques showed a smooth surface and a morphology similar to that of baseline images, while IFC plaques showed a smoothed intimal border and a double layering indicating organising thrombus incorporated in the superficial layers of the plaque.

Conclusions: In the months following a successfully dissolved acute thrombosis, OCT revealed that the cavity of RFC plaques persists and is bordered by a smooth “neointima”, while IFC plaques showed features suggesting partial incorporation of the deepest layers of thrombus in the plaque.

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Introduction

The natural history of culprit plaques, both with a ruptured fibrous cap (RFC) and with an intact fibrous cap (IFC), causing an acute coronary syndrome (ACS) has not been systematically investigated *in vivo*¹. In 10 patients with ACS, we used serial optical coherence tomography (OCT) to explore the repair and healing morphology of non-critically narrowed culprit plaques, both with RFC and IFC, which were treated with thrombus aspiration or systemic thrombolysis alone without angioplasty or stent implantation.

Methods

This is a hypothesis-generating prospective study which was performed using 10 patients with ACS (Table 1). Twenty-five patients with ACS were scrutinised over four months. Ten fulfilled the following criteria of enrolment: onset of symptoms <12 hours, achievement of “optimal reperfusion” with a Thrombolysis In Myocardial Infarction (TIMI) 3 flow with non-critical residual angiographic narrowing, and a small plaque/thrombus prolapse on OCT², after thrombus aspiration or thrombolysis. Seven patients received thrombus aspiration, two were treated with IV abciximab and one with systemic thrombolysis (Table 2). Baseline OCT study of culprit lesions

was obtained after 5.2±2.6 days, whilst the follow-up OCT study was repeated at a time point between one and seven months. Patient characteristics and procedural details are described in Table 1 and Table 2.

All procedures were performed according to the current guidelines. Patients received aspirin, thienopyridine, and unfractionated heparin (5,000-10,000 IU) (Table 2). After discharge, patients underwent coronary angiogram and OCT assessments as per local practice. The repeat angiogram with OCT was done at two, four, six and seven months in four, one, four and one patient, respectively. Written informed consent was obtained before follow-up angiographic OCT procedures that were deemed clinically useful.

Coronary angiograms were analysed off-line using quantitative coronary angiography (QCA)³.

The OCT culprit lesion morphology was described according to previously reported criteria^{2,4,5}.

Statistical analysis

Continuous variables are reported as mean (standard deviation) and were compared with the Student's t-test, while categorical variables are reported as n (%) and were compared using Fisher's exact test. Statistical significance was set at the two-tailed 0.05 level.

Table 1. Patient characteristics.

ID	Gender	Age	Diabetes	Dyslipidaemia	Hypertension	Smoker	Family history	Prior MI	Prior PCI	LVEF (%)	Clin. syndr.	Culprit lesion	n. diseased vessels
1	M	45	NO	YES	NO	NO	NO	YES	YES	55	STEMI	LAD	2
2	M	51	NO	NO	NO	NO	NO	YES	NO	67	STEMI	LAD	1
3	M	37	NO	NO	NO	YES	YES	NO	NO	48	STEMI	LAD	1
4	M	24	NO	YES	NO	YES	NO	NO	NO	50	STEMI	LAD	1
5	M	43	YES	NO	NO	YES	NO	NO	NO	62	NSTEMI	RCA	1
6	M	71	NO	NO	NO	YES	NO	NO	NO	45	NSTEMI	LAD	1
7	M	35	NO	YES	NO	YES	NO	NO	NO	50	STEMI	LAD	1
8	M	42	NO	YES	NO	NO	YES	NO	NO	50	STEMI	RCA	1
9	M	66	NO	YES	YES	YES	NO	NO	NO	70	STEMI	LAD	2
10	M	21	NO	NO	NO	YES	NO	NO	NO	40	STEMI	LAD	1

Clin. syndr: clinical syndrome; LAD: left anterior descending; LVEF: left ventricular ejection fraction; NSTEMI: non-ST-elevation myocardial infarction; RCA: right coronary artery; STEMI: ST-elevation myocardial infarction

Table 2. Procedural details.

ID	Systemic thrombolysis	Thrombus asp. device	IIb/IIIa inhib.	ADP antag.	Basal DS (%)	Post-interv. % DS	Basal TIMI flow	Post-interv. TIMI flow	Tot. isch. time (hours)	Clinical syndrome
1	NO	YES	YES	PRASUGR	100	46	0	III	2	STEMI
2	NO	YES	YES	CLOPID	100	40	0	III	1	STEMI
3	YES	NO	NO	CLOPID	53	20	II	III	3	STEMI
4	NO	YES	YES	PRASUGR	90	33	I	III	1.5	STEMI
5	NO	YES	YES	PRASUGR	78	41	I	III	12	NSTEMI
6	NO	NO	YES	PRASUGR	53	25	II	III	2.5	NSTEMI
7	NO	NO	YES	PRASUGR	45	30	II	III	1.5	STEMI
8	NO	YES	NO	PRASUGR	92	22	I	III	1.5	STEMI
9	NO	YES	YES	PRASUGR	59	23	II	III	2.5	STEMI
10	NO	YES	YES	PRASUGR	95	43	I	III	13	STEMI

ADP antag: ADP antagonist; CLOPID: clopidogrel; DS: diameter stenosis; IIb/IIIa inhib: IIb/IIIa inhibitors; PRASUGR: prasugrel; Tot. isch. time: total ischaemic time

Results

Eight patients had an ST-elevation myocardial infarction (STEMI), and the remaining two patients suffered a non-ST-elevation myocardial infarction (NSTEMI). Of the eight patients with STEMI, thrombus aspiration was performed in six, systemic thrombolysis in one, and the remaining patient was treated only with IV abciximab. Of the two patients with NSTEMI, one was treated with thrombus aspiration and one with IV abciximab (Table 1, Table 2).

Pre- and post-intervention plaque morphology

The mean total ischaemic time was 2.9 ± 3.5 hours; baseline angiography is described in Table 2. The pre-intervention MLD and RVD were 0.30 ± 0.42 mm and 3.2 ± 0.45 mm, respectively.

Thrombus aspiration (n=7) or medical therapy (n=3) led to a mild residual diameter stenosis with final TIMI 3 flow in all cases.

Baseline OCT assessment of culprit lesions demonstrated RFC in five patients and IFC in the remaining five patients, with OCT aspects of thrombosis being revealed in all cases.

Follow-up plaque morphology and clinical outcome

Angiography confirmed the presence of a TIMI 3 flow in all cases without significant luminal compromise. The morphology on OCT of the five culprit lesions with RFC did not change during follow-up (Table 3) with all lesions showing clear communication between the plaque cavity and the lumen (Figure 1), both early (one to two months after the baseline assessment) and late (four and six months). Fibrous cap thickness, minimal lumen area and mean lumen area remained unchanged during follow-up (+12.7%, p=ns; -1.56% p=ns and -0.10% p=ns, respectively). The quantitative measurements of the longitudinal and circumferential extension of the cavity also remained unchanged (Table 4). New tissue apposition (neointima-like) was seen to develop at follow-up.

The morphology of the IFC changed over time. The irregular inner border of the thrombus in the acute phase evolved to a homogeneous and smooth profile in all OCT cross-sections of the four cases that were serially imaged after at least 42 days (Table 3). On the other hand, the images of the patient who was followed up at 28 days were similar to the baseline morphology (Figure 2). At follow-up, the organised thrombus showed a rim of circumferential transition exhibiting a low backscatter (neointima) in all of the cross-sections (Figure 2). In the five patients with IFC, a non-significant reduction in minimal and mean lumen area was observed (-8.5 and -9.9 percent, respectively).

During a mean clinical follow-up of 26.9 ± 10.4 months (range one to four years), all 10 patients have remained asymptomatic and have not developed further acute cardiovascular events.

Discussion

This hypothesis-generating serial OCT study addresses the natural history of the evolution of plaque morphology of non-critically narrowed culprit lesions both with RFC and IFC in patients

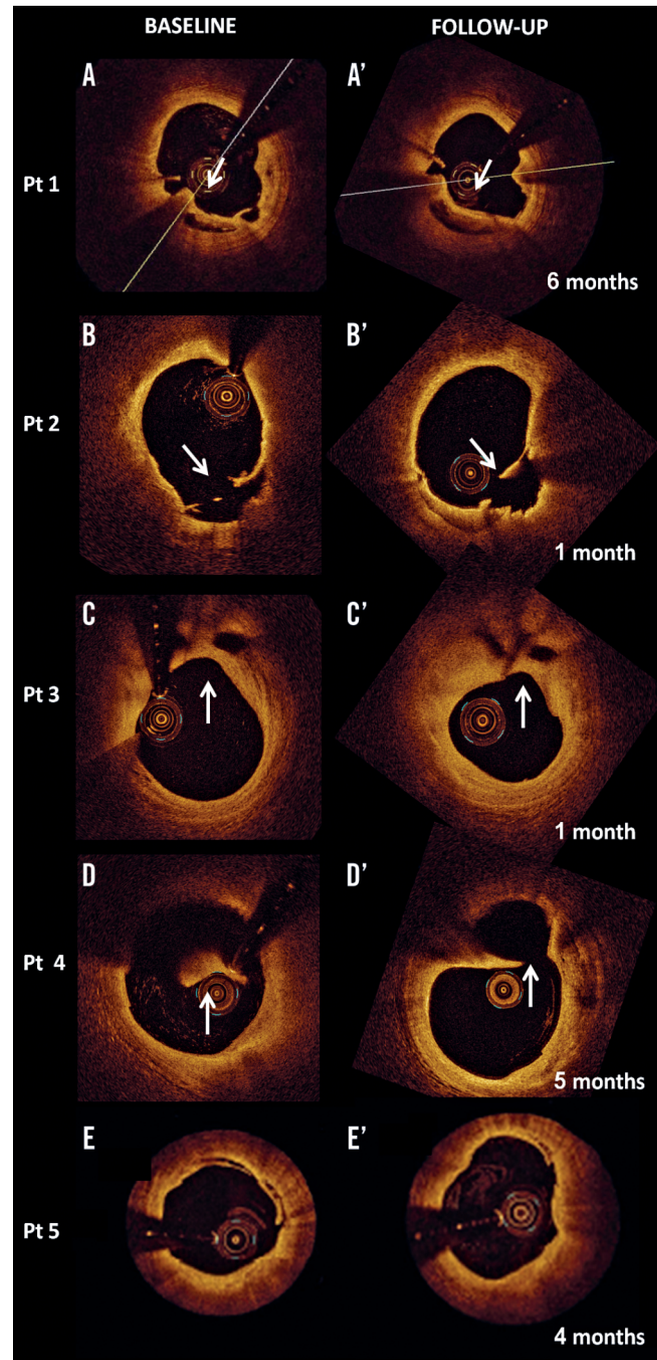


Figure 1. OCT assessment of culprit lesions with ruptured fibrous cap (RFC) at baseline and inherent changes over time. A-A' and B-B'. Examples of culprit lesions with RFC (arrows in the left panel) exhibiting a clear communication between the inner (necrotic) core of the plaque and the lumen. Plaque morphology is unchanged at follow-up from the acute event with plaque ulceration still visible (arrow in the right panels). C-C'. The presence of RFC can be seen at follow-up. In this case the lumen gets smaller probably due to a process of organisation of the thrombus remnants. D-D'. Plaque ulceration imaged just proximal to a lipid pool. The sign of plaque rupture can be seen at follow-up. Please note that the presence of thrombus on the fibrous cap hampers the clear visualisation of the cavity in the acute phase. E-E'. A superficial plaque dissection can be seen in the acute phase (left panel) and at follow-up (right panel).

Table 3. OCT qualitative analysis.

ID	Mechanism of plaque instability	Acute/subacute assessment			Follow-up assessment		
		OCT time point after AMI (days)	Thrombus detection	Culprit lesion with RFC	OCT time point after AMI (days)	Organised thrombus	Culprit lesion with RFC
1	IFC	1	YES	NO	28	YES	NO
2	IFC	3	YES	NO	187	YES	NO
3	IFC	8	YES	NO	160	YES	NO
4	IFC	2	YES	NO	183	YES	NO
5	IFC	4	YES	NO	42	YES	NO
6	RFC	8	YES	YES	205	YES	YES
7	RFC	5	YES	YES	41	NO	YES
8	RFC	6	YES	YES	33	YES	YES
9	RFC	7	YES	YES	177	YES	YES
10	RFC	8	YES	YES	116	YES	YES

IFC: intact fibrous cap; RFC: ruptured fibrous cap

Table 4. Quantitative OCT analysis.

ID		Acute phase				Follow-up				Difference			
		MLA mm ²	Mean LA mm ²	FC thick μ m	LP Arc ^o	MLA mm ²	Mean LA mm ²	FC thick μ m	LP Arc ^o	MLA mm ²	Mean LA mm ²	FC thick μ m	LP Arc ^o
1	IFC	6.33	9.95	160	210	4.6	7.74	160	210	1.73	2.21	0	0
2	IFC	7.16	8.94	0	0	5.92	8.75	0	0	1.24	0.19	0	0
3	IFC	2.36	3.75	120	208	1.9	3.27	120	190	0.46	0.48	0	18
4	IFC	5.62	7.52	140	92	5.92	7.17	135	75	-0.30	0.35	5	17
5	IFC	2.77	6.09	0	0	3.85	5.72	0	0	-1.08	0.37	0	0
	Mean	4.85	7.25	84.00	102.00	4.44	6.53	83.00	95.00	0.41	0.72	1.00	7.00
	STD	2.16	2.44	77.97	104.65	1.67	2.13	77.10	100.87	0.49	0.31	0.87	3.78
6	RFC	5.62	7.73	40	245	5.92	7.42	45	250	-0.30	0.31	-5.00	-5.00
7	RFC	6.86	9.12	40	306	6.82	9.69	61	258	0.04	-0.57	-21.00	48.00
8	RFC	4.85	6.7	70	200	3.47	5.5	75	200	1.38	1.20	-5.00	0.00
9	RFC	7.01	10.25	88	220	7.46	13.13	90	220	-0.45	-2.88	-2.00	0.00
10	RFC	4.49	6.59	69	180	4.71	4.61	75	170	-0.22	1.98	-6.00	10.00
	Mean	5.77	8.08	61.40	230.20	5.68	8.07	69.20	219.60	0.09	0.01	-7.80	10.60
	STD	1.14	1.58	20.95	48.74	1.61	3.44	16.98	36.20	-0.46	-1.85	3.97	12.53

LA: lumen area; LP: lipid pool; FC: fibrous cap; IFC: intact fibrous cap; MLA: minimal lumen area; RFC: ruptured fibrous cap; STD: standard deviation

presenting with ACS. OCT images showed that the plaque cavity which is the marker of culprit lesions with RFC may remain open several months after an acute coronary event. The border of the cavity may appear smoother as compared to the acute phase, suggesting a superficial healing process with the growth of a neointimal layer. OCT images in IFC plaques, wherein the cavity is typically absent, show the presence of a two- (or more) layer structure with low backscatter inner rim(s) of circumferential transition from the surface to original plaque. This suggests that the layer of residual thrombotic material is probably incorporated in the plaque and shows a neointimal-like evolution. The plaque demonstrates a slight increase of the intraluminal burden. At a short to mid interval from the acute event, plaques which caused local occlusive thrombosis (thrombogenic plaques) show the OCT morphological features of stable plaques.

Most knowledge on coronary lesion development, progression and complications is based on histopathology studies, which provide a single time point profile of the coronary pathology⁶. Intravascular imaging offers the unique opportunity of interrogating coronary plaques during serial studies, thus potentially addressing research on plaque changes and evolution *in vivo*⁴. Studies aimed at assessing plaque instability in the coronary tree of patients with ACS^{7,8} have identified multiple complex lesions but did not prove the simultaneous destabilisation of more than one plaque. Disruption of multiple plaques reflects a diffuse or multifocal coronary instability, a well-established concept which rests on both post-mortem data⁶ and biomarker studies demonstrating widespread inflammation⁹. What is still unknown is the time lapse between the rupture of a plaque and its healing and whether healed plaques can be considered as clinically stable and non-thrombogenic. The studies which

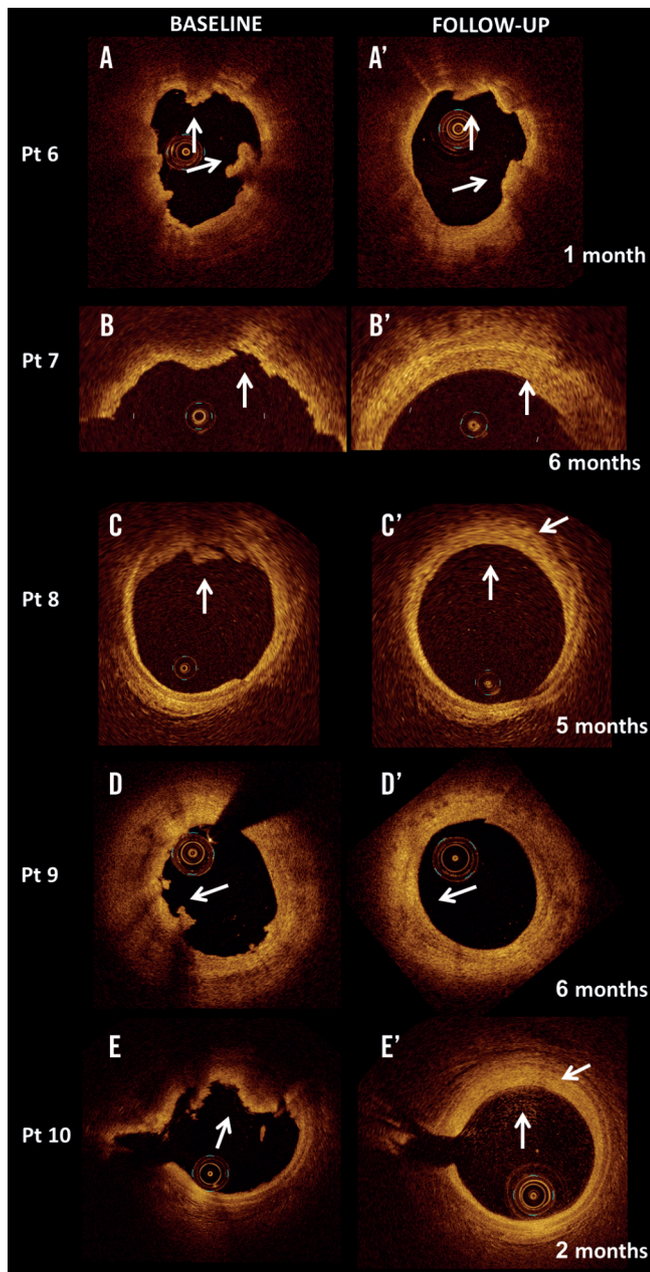


Figure 2. OCT assessment of culprit lesions with intact fibrous cap (IFC) at baseline and inherent changes over time. A-A'. Patient with STEMI who had an early 28-day follow-up assessment. In the acute phase OCT shows multiple thrombus appositions (arrows in the left panel), which are still seen at follow-up (arrows in the right panel). The remaining panels refer to the four culprit lesions with IFC (B, C, D, E), which were studied in the follow-up time period (B', C', D', E') of between 42 and 187 days. Irregular contours due to thrombus apposition are visible in the acute phase (arrows in the left panel) and become smooth during follow-up. Of note, signs of organised thrombus can be seen at follow-up (arrows in the right panel) with a two-layer appearance and a rim of circumferential transition exhibiting a low backscatter.

have identified multiple ruptured plaques have consistently implied that the plaque ulceration is a marker of acute plaque instability¹⁰. However, our preliminary OCT data suggest that the unhealed

ulcers may not necessarily imply clinical instability, especially in the presence of effective antiplatelet therapy. Our findings are supported by a recent observation, showing that the aspect of quiescent plaque rupture remains stable during long-term follow-up¹¹, and that finding multiple ruptured plaques may not directly translate into multilocus coronary instability.

Finally, the present study suggests that patients with non-critically narrowed culprit plaques could be treated simply with thrombus removal to reduce the risk of no-reflow related to balloon angioplasty and stent deployment.

Limitations

The limitations of this study include the low number of patients, the different intervals of time which elapsed between baseline OCT evaluation and follow-up as well as the non-identical treatment options in the 10 patients, in whom, however, dissolution of the thrombus was equally and successfully achieved.

The penetration capability of OCT limits the assessment of the deeper portion of culprit plaques, particularly in the presence of superficial components which can attenuate the signal^{2,4}.

Conclusions

The preliminary data of this hypothesis-generating study indicate that, in ACS patients with mildly narrowed culprit plaques with RFC, the cavity may persist for months and may gradually develop neointima repair. In the culprit plaques with IFC, the residual thrombus is probably incorporated into the plaque, generating the double layer seen in OCT images.

Impact on daily practice

This serial OCT study was carried out in patients with ACS, treated with thrombus removal without the adoption of technical solutions that affect the mechanisms of plaque repair, such as balloon dilatation and stenting. Plaque cavities in ruptured fibrous cap lesions tend to remain in contact with the lumen, even several months after an acute coronary event, while, in the presence of intact fibrous cap, residual thrombotic material tends to organise and be incorporated in the plaque. Importantly, in both cases, plaque size does not increase significantly with respect to baseline. Therefore, in daily practice, our results can support a strategy of thrombus removal only, without stent implantation to treat culprit lesions, regardless of the mechanism of local thrombosis. Furthermore, plaques with signs of ulceration and without local thrombosis on OCT should be considered quiescent plaques that do not require any specific treatment.

Conflict of interest statement

The authors have no conflicts of interest to declare.

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