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The natural history of human atherosclerosis : a histopathological approach

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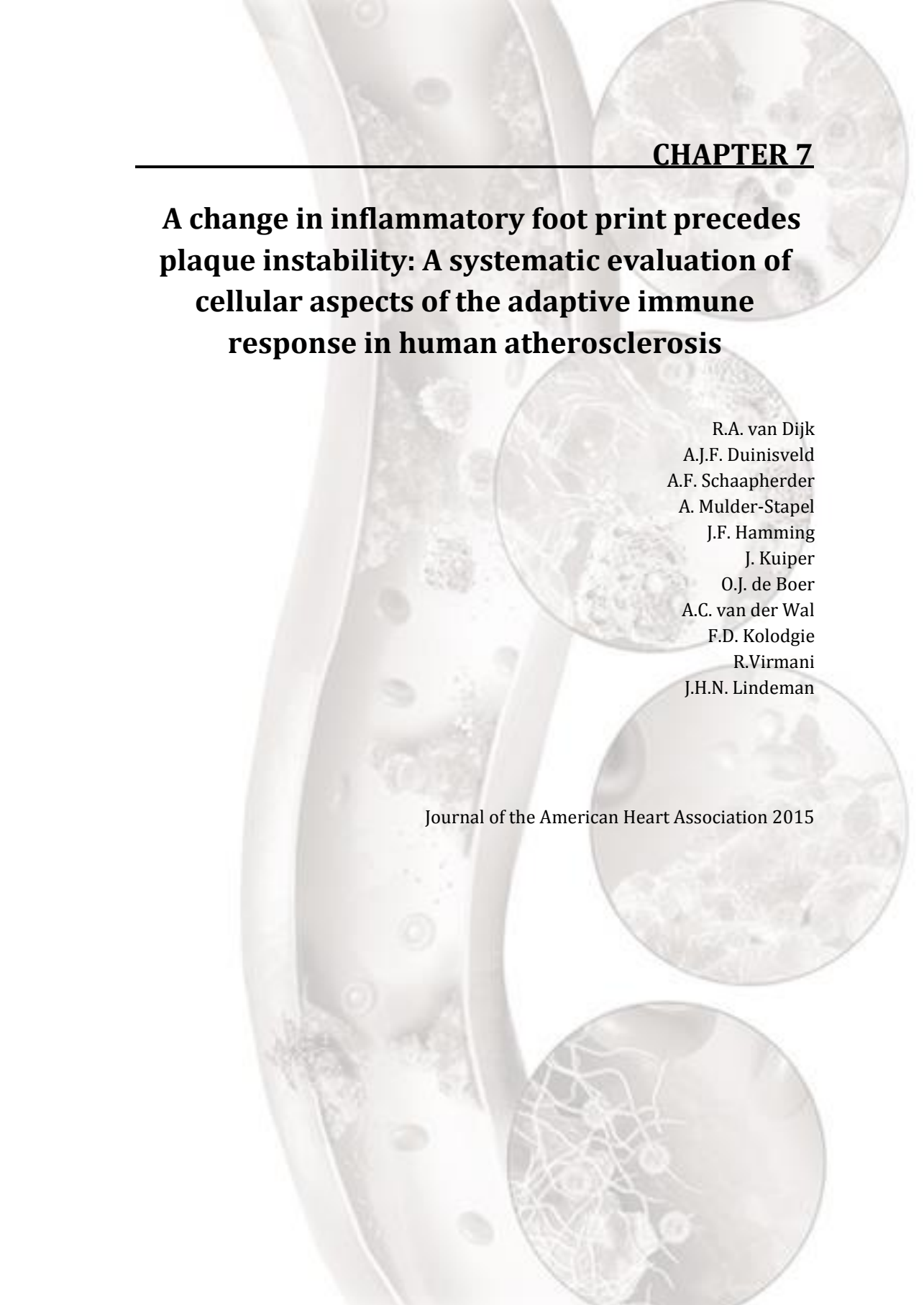


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CHAPTER 7

A change in inflammatory foot print precedes plaque instability: A systematic evaluation of cellular aspects of the adaptive immune response in human atherosclerosis

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ABSTRACT

Background: Experimental studies characterize adaptive immune response as a critical factor in the progression and complications of atherosclerosis. Yet, it is unclear whether these observations translate to the human situation. This study systematically evaluates cellular components of the adaptive immune response in a biobank of human aortas covering the full spectrum of atherosclerotic disease.

Material and methods: A systematic analysis was performed on 114 well-characterized peri-renal aortic specimens with immunostaining for T-cell subsets (CD3/4/8/45RA/45RO/FoxP3) and the Th1/non-Th1/Th17 ratio (CD4+T-bet+/CD4+T-bet-/CD4+/IL-17+ double staining). CD20 and CD138 were used to identify B-cells and plasma cells, while B-cell maturation was evaluated by AID/CD21 staining and expression of lymphoid homeostatic CXCL13.

Results: Scattered CD4 and CD8 cells with a T memory subtype were found in normal aorta and early, non-progressive lesions. The total number of T-cells increases in progressive atherosclerotic lesions (~1:5 CD4/CD8 T-cell ratio). A further increase in medial and adventitial T-cells is found upon progression to vulnerable lesions. This critical stage is further hallmarked by de-novo formation of adventitial lymphoid-like structures containing B-cells and plasma cells, a process accompanied by transient expression of CXCL13. A dramatic reduction of T-cell subsets, disappearance of lymphoid structures, and loss of CXCL13 expression characterize post-ruptured lesions. FoxP3 and Th17 T-cells were minimally present throughout the atherosclerotic process.

Conclusion: Transient CXCL13 expression, restricted presence of B-cells in human atherosclerosis, along with formation of non-functional extranodal lymphoid structures in the phase preceding plaque rupture, indicates a “critical” change in the inflammatory footprint before and during plaque destabilization

INTRODUCTION

Atherosclerosis is a highly dynamic^{1,2} metabolic disease with a strong inflammatory component^{3,4,5}. Compelling evidence, largely based on data derived from murine models of atherosclerotic disease, implies extensive involvement of the adaptive immune response in the initiation, progression and complications of atherosclerotic disease⁶. As a consequence, the adaptive immune response is considered a potential target for medical interventions.

An open question is how these observations translate to the human situation. There are fundamental immunological and inflammatory differences between mice and men^{7,8,9}. Moreover, murine models of accelerated atherosclerotic disease require an immunologic background with exaggerated Th1-cell responsiveness in order for atherosclerosis to develop^{8,9}. Hence, observations from mouse models may be skewed towards Th1 response. Extrapolation of the murine data is further complicated by absence of vulnerable lesion formation in the current models of atherosclerotic disease. As a result, data on a possible involvement of the adaptive immune response in the advanced stages of atherosclerotic disease is missing.

Available human data on the other hand largely relies on surgical specimens. Yet, it is important to note that this material generally represents the final stages of the disease process^{10,11}. As such, data from these human studies is not representative for the earlier and intermediate phases of atherosclerotic disease; hence knowledge on the nature of the adaptive immune response in the human atherosclerotic process is limited.

Given the above considerations, we regarded an evaluation of the adaptive immune response within the process of atherosclerotic lesion formation, progression and stabilization relevant. To that end, we performed comprehensive and systematic histological assessment of cellular aspects of the adaptive immune response in tissue samples from a biobank of arterial tissue that covers the full spectrum of human atherosclerotic disease. Results from this explorative study confirm extensive and dynamic presence of cellular components of the adaptive immunity in the human atherosclerotic process, and reveal profound changes in the inflammatory foot print immediately prior to and during the process of plaque destabilization.

MATERIAL AND METHODS

Patients and tissue sampling

Tissue sections were selected from a tissue bank of aortic wall patches that were obtained during liver, kidney and pancreas transplantation with grafts derived from cadaveric donors. Details of this bank have been described previously by van

Dijk *et al* ¹². All patches were from grafts that were eligible for transplantation (*i.e.* all donors met the criteria set by The Eurotransplant Foundation). Due to national regulations, only transplantation relevant data for donation is available, as such background information on the specimens is limited. Sample collection and handling was performed in accordance with the guidelines of the Medical and Ethical Committee in Leiden, Netherlands and the code of conduct of the Dutch Federation of Biomedical Scientific Societies (<http://www.federa.org/?s=1&m=82&p=0&v=4#827>).

Each tissue block in the bank was Movat and H&E stained and classified according to the modified American Heart Association (AHA) classification as proposed by Virmani *et al* by two independent observers with no knowledge of the characteristics of the aortic patch ^{12,13}. The tissue block showing the most advanced plaque was used for further studies. In order to obtain balanced and representative study groups, we selected the first 100 aortic wall samples from the tissue bank. Due to the fact the vulnerable lesions (*i.e.* thin cap fibroatheroma and plaque ruptures) were under-represented we randomly selected several aortic wall samples classified as vulnerable lesions from the remaining 250 patches in order to obtain approximately 10-12 samples from each atherosclerotic stage. A total of 114 cases were selected for further examination. Demographic data and the causes of death are summarized in Table I.

Characterization of the lesions and histological definitions

Aortic samples not showing any signs of intimal thickening and intimal inflammation were classified as normal. The other samples were classified based on the most advanced lesion type present in the section. Lesions were defined as adaptive intimal thickening (AIT), intimal xanthoma (IX), pathological intimal thickening (PIT), early fibroatheroma (EFA), late fibroatheroma (LFA), thin cap fibroatheroma (TCFA), acute plaque rupture (PR), healed plaque rupture (HR) and fibrotic calcified plaque (FCP). For detailed descriptions concerning a complete overview on plaque processing, morphological analysis and additional information concerning the studied population are provided in reference 12.

Immunohistochemistry

All specimens used for immunohistochemistry were washed in phosphate-buffered saline (PBS), formalin fixed and decalcified (Kristenssens solution) and paraffin embedded using standard procedures. The details of the antibodies used for immunohistochemistry are listed in Table II. Conjugated biotinylated horse anti-mouse (1:400 dilution; Vector laboratories, Amsterdam, The Netherlands) or Envision® + System – HorseRadish Peroxidase (HRP) labeled polymer anti mouse/anti rabbit (prediluted; Dako, Heverlee, Belgium) functioned as secondary antibodies. Sections were developed with Nova Red® or DAB and counterstained

with Mayer's haematoxylin or methylgreen. Positive and negative controls were always included and performed by adding or omitting the primary antibody on 3-Aminopropyltriethoxysilane (APES)-slides containing human tonsils.

The Foxp3/CD3 and IL-17/CD3 double staining was performed to identify Regulatory T cells and Th17cells, respectively (Table I). Conjugated polymer anti mouse/ anti rabbit (Immunologic®) functioned as secondary antibodies. After the goat-derived anti-IL17 incubation, a rabbit anti-goat secondary antibody (Southern Biotech, Birmingham, AL, USA) served as a bridge reagent for the next step with alkaline phosphatase (AP) anti-rabbit polymer (Immunologic®, Duiven, The Netherlands). All immunohistochemical stainings were performed as previously described¹⁴. Despite many attempts and various techniques, we did not succeed to use GATA-3 as the T-helper 2 marker. We therefor had to turn to double staining for CD4/Tbet to identify the T-helper lineage (Lineage: Th1= CD4⁺/Tbet⁺ and non-Th1= CD4⁺/Tbet⁻) having regard to the provision that non-Th1 may also include Th0, Th17 and Th22 populations.

Table 1. Demographic data of the 114 studied aortic samples

	Male		Female	
N	64		50	
Mean age (years) [SD]	49.4	[14.3]	43,8	[17.3]
Mean length (cm) [SD]	179.9	[11.9]	166.6	[12.1]
Mean weight (kg) [SD]	80.1	[16.3]	64.9	[15.3]
Mean BMI (kg/m2) [SD]	24.8	[3.1]	22.9	[4.4]
Patients with known history of nicotine abuse [percentage]	21	[36.8%]	17	[39.5%]
Patients with known history of hypertension* [percentage]	13	[22.8%]	10	[23.3%]
Patients with known diabetes [percentage]	0	[0.0%]	1	[2.3%]
Cause of Death [n, percentage]				
Severe head trauma	11	[17.1%]	7	[14.0%]
Cerebral vascular accident (CVA)	8	[12.5%]	10	[20.0%]
Subarachnoid bleeding (SAB)	15	[23.4%]	14	[28.0%]
Cardiac arrest	7	[10.9%]	0	[0.0%]
Trauma	1	[1.5%]	1	[2.0%]
Other	6	[9.3%]	3	[6.0%]
Unknown	16	[25.0%]	15	[30%]
Medication [n, percentage]				
Anti-hypertensives	11	[19.3%]	6	[14.0%]
Statins	1	[1.7%]	1	[2.3%]
Anti-coagulants	1	[1.7%]	2	[4.7%]
Other	5	[8.8%]	8	[18.6%]
None	31	[54.4%]	22	[51.2%]
Unknown	11	[19.3%]	9	[20.9%]

*Known antihypertensive medication / systolic blood pressure >140mmHg and diastolic >90mm Hg in the period preceding death. Abbreviations: SD = Standard Deviation.

Table 2. Antibodies used in the present study

Antibody, clone	Host isotype; subclass	Specificity	Pretreatment		Dilution	Reference/source
CD3, polyclonal	Rabbit	Pan T cells	10x Tris/EDTA	pH 9,2	1:200	DAKO
CD4, 4B12	Mouse, IgG1- κ	T helper-cells	10x Tris/EDTA	pH 9,2	1:200	DAKO
CD8, C8/144B	Mouse, IgG1- κ	Cytotoxic T-cells	10x Tris/EDTA	pH 9,2	1:200	DAKO
CD45RA, HI100	Mouse, IgG2b- κ	Naive T cells	Citrate	pH 6,0	1:2000	BioLegend
CD45RO, UCHL1	Mouse, IgG2a- κ	Memory T cells	Citrate	pH 6,0	1:1000	BioLegend
CD20, L26	Mouse	Pan B cells	Citrate	pH 6,0	1:1000	DAKO
CD21, IF8(4)	Mouse, IgG1- κ	Follicular Dendritic Cells, mature B cells	Citrate	pH 6,0	1:400	DAKO
CD138, MI15	Mouse	Plasma cells	10x Tris/EDTA	pH 9,2	1:1000	DAKO
CCR7, ab191575	Rabbit, polyclonal	Activated T-lymphocytes	Citrate	pH 9,2	1:200	Abcam
Tbet, polyclonal	Rabbit, sc-21003	T helper1-cells	10x Tris/EDTA	pH 9,2	1:800	Santa Cruz
IL-17, polyclonal	Goat	Th-17 cells	-	pH 9.0	1:50	R&D
FoxP3, clone 236A/E7	Mouse	Regulatory T cells	-	pH 9.0	1:50	Abcam
CXCL13, cat no AF801	Goat	Lymphorganogenic chemokine	Citrate	pH 6,0	1:100	R&D
anti-AID, EK2 5G9	Rat	Activation-induced cytidine deaminase	Citrate	pH 6,0	1:16.000	Cell Signalling

Table 3. Histological classification of aortic tissue according to the modified AHA classification proposed by Virmani *et al*

Morphological description	Abbreviation	Male			Female			Total
		N	Mean age	[SD]	N	Mean age	[SD]	N
Normal aorta	N	7	23.8	[18.9]	5	10.0	[3.7]	12
Non-progressive intimal lesions								
Adaptive intimal thickening	AIT	5	40.5	[10.0]	5	30.4	[9.0]	10
Intimal xanthoma	IX	7	37.3	[10.0]	5	34.2	[14.8]	12
Progressive atherosclerotic lesions								
Pathological intimal thickening	PIT	5	46.0	[8.9]	7	55.4	[11.5]	12
Early fibroatheroma	EFA	7	50.9	[5.9]	5	46.0	[3.5]	12
Late fibroatheroma	LFA	5	55.0	[3.6]	7	51.1	[11.6]	12
Vulnerable atherosclerotic lesions								
Thin cap fibroatheroma	TCFA	7	57.2	[5.2]	4	60.3	[5.7]	11
Plaque rupture	PR	7	56.9	[2.6]	3	45.0	[13.1]	10
Stabilizing lesions								
Healing rupture	HR	8	58.1	[11.1]	4	57.0	[0]	12
Fibrotic calcified plaque	FCP	6	62.7	[6.8]	5	53.0	[11.4]	11
		64	49.4	[14.3]	50	43.8	[17.3]	114

*Abbreviations: SD = Standard Deviation.

Assessment of immunolocalization of T cells (and T cell subset), B cells and plasma cells

The tissue block showing the most advanced plaque was used for further studies. When the aortic sample contained more than one lesion, the most advanced lesion on the slide was used. The lesion was defined as the area between the endothelium and the first elastic lamina over a distance of 1mm. In the presence of a necrotic core the distance was expanded with another 500µm on both sides of the necrotic core to include the adjacent shoulder region¹². The regions of interest (ROI) were the intima (and in the presence of an early or late necrotic core the intima was divided in a cap and both shoulder regions), the media and the adventitia (figure 1). Within the ROIs (*i.e.* intima, media and adventitia), three representative adjacent images were made at a 200x magnification. A total of 9 images per atherosclerotic lesion were separately analyzed from the lesion. Immunostaining intensity was quantitatively assessed with an image processing program (Image J; plug-in Cell counter). Representative examples of the immunological stainings are provided in figure 2.

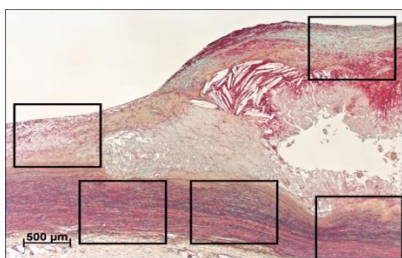


Figure 1. Example of the regions of interest as used for the assessment of immunolocalization of T cells (and T cell subset), B cells and plasma cells in a healed rupture. The regions of interest are drawn in the fibrous cap, in one of the shoulders and three times in the media in this particular example. For every lesion (normal or ruptured) a total of nine images were made (3 in the intima, 3 in the media and 3 in the adventitia) and the immunostaining was quantitatively assessed with an image processing program (Image J; plug-in Cellcounter).

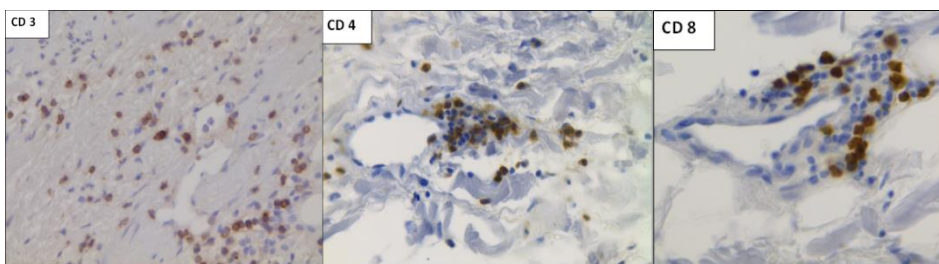


Figure 2. High resolution examples of the immunohistochemical staining for CD3, CD4 and CD8 cells. The high resolution images are examples of the quality of the immunohistochemical staining for CD3, CD4 and CD8. All images were taken at a 400x magnification at the medial adventitial border. Within all the images one can see the clear positively stained Tcells mainly located near the *vasa vasorum*. All sections were developed with DAB and counterstained with Mayer's haematoxylin.

Statistical analysis

Data in figures are presented as mean $\pm$ SEM. Spearman's correlation was used to demonstrate the relationship between the amount of positive cells in the intima, media and adventitia within one phase of the atherosclerotic disease and compared with the previous and/or next phase in atherosclerosis (SPSS 20.0; Chicago, IL). The study population consisted of 114 individuals. However during the staining process some tissue samples were lost. Therefore the total number of samples for each staining may be less than 114. The Wilcoxon-Mann-Witney test was used to analyze the non-normally distributed data of the mean total number of cells within each atherosclerotic phase and the Kruskal Wallis test was used to control for a type 1 error. A value of $p < 0.05$ was considered statistically significant.

RESULTS

Study population

Characteristics of the studied population are provided in Table I. The male/female ratio was evenly distributed (57% male), as was the mean age for each sex (~49-years). There was a strong correlation between donor age and atherosclerosis progression¹² (Table 3). The mean donor age for the aortic wall samples classified as normal was 23-years, whereas the mean age for aortas with fibrotic calcified plaques, representing the final stage of the disease, was 62-years. Nearly 40% of patients had a known history of smoking and 2 patients received statin therapy. Twenty-three patients had a known history of hypertension; 17 of these patients received anti-hypertensive medication.

Normal aorta and non-progressive atherosclerotic lesions

The intima and media layers of the normal (non-atherosclerotic) aortic wall are devoid of CD3⁺ T cells. Scattered T cells found at the medial-adventitial border are mainly CD45RO⁺ (memory) T cells. The CD4⁺ (Thelper) to CD8⁺ (cytotoxic) T cell ratio is approximately 1:2 (figure 3A). Th₁ cells and non-Th1 cells were also expressed in a 1:2 ratio (Figure 3B).

Naïve T cells (CD45RA⁺), Regulatory T cells (CD3⁺/FoxP3⁺), Th17 cells (CD3⁺/IL17A⁺), B cells (CD20⁺) and plasma cells (CD138⁺) are all absent in aorta's classified as normal.

Isolated T cells in the thickened intima, and progressive numbers of T cells in the medial/adventitial border characterize so-called non-progressive lesions (*i.e.* adaptive intimal thickening and intimal xanthomas). Memory T cells remain the dominant phenotype, and the subset ratios remains unchanged (CD4⁺/8⁺; Th1/TH2). Regulatory T cells, Th17 cells, naïve T cells, B cells and plasma cells are absent in these early lesions (figure 4).

Progressive atherosclerotic lesions

A significant increase is seen in intimal CD3⁺ T cells ($p < 0.0001$) in progressive lesions (early fibroatheroma and late fibroatheroma) with beginning infiltration of the medial wall as well. The predominance of cytotoxic T cells persists (CD4⁺: CD8⁺ T cell ratio 1:5) in the early fibroatheroma, and T helper cells are dominated by the Th₁ lineage at a 1:3 ratio. Transition from an early to late fibroatheroma stage is accompanied by the accumulation of T cells in the intima shoulder regions of the lesion, and the CD4⁺/CD8⁺ T cell ratio shifts from 1:5 in early fibroatheroma to ~1.5:1 in late fibroatheroma.

Memory T cells remain the dominant phenotype in advanced lesions, yet scattered naïve T cells (CD45RA⁺) start to appear within the medial-adventitial border in lesions classified as EFA. A further increase in naïve T cells and T memory cells is seen in late fibroatheroma with a notable accumulation of T cells alongside infiltrating *vasa vasorum* in the outer layers of the media underlying the necrotic core. In general, B cells and plasma cells are absent, although present in a minority of individuals (Figure 9). Regulatory T cells and Th17 cells are not identified in the progressive lesions.

The vulnerable lesions

Thin cap fibroatheromas and ruptures are hallmarked by a peak in T cell infiltration in the adventitia underneath to the culprit lesion. The thin cap remains devoid of T cells and the number of T helper cells decreases resulting in an approximate 1:1 CD4/CD8 cell ratio. Non-Th1 cells remain the dominant T-helper lineage in the vulnerable phase, due to doubling of the number of non-Th1 cells (Figure 6). Dispersed memory T cells and naïve T cells are abundantly present in the media and adventitia near the *vasa vasorum*. Absence of CD45RO/CCR7 double positive cells indicates that all CD45RO⁺ T cells within the lesion and infiltrates are effector memory T cell and the only few central memory T cells are identified in the aortic wall located within the *vasa vasorum* of the adventitia (figure 10).

A unique finding in the vulnerable lesions is the appearance of B-cells and occasional plasma cells in tertiary follicle-like structures. Staining for CXCL13, a homeostatic, lymphorganogenic chemokine that is critical for organization follicle-like structures, shows CXCL13 expression exclusively in vulnerable lesions, (*viz.* no CXCL13 staining was found in the other stages of the disease) and reveals a particular staining pattern of a dendritic network with extension from the follicles (Figure 9). Additional staining for mature B cells (CD2) and Activation-induced cytidine deaminase (AID) was mainly negative; indicating the lack of B cell maturation. Scattered FoxP3⁺ T cells are found within the intima and underlying adventitia in 2 of 9 aortic sections of thin cap fibroatheroma (figure 6). On the other hand, Th17 cells are absent in vulnerable lesions.

Stabilized lesions

Healing plaque ruptures exhibit a dramatic decrease in T cells ($P<0.001$), naïve T cells, and disappearance of B cells and plasma cells. The decline in T cells is more outspoken for the CD4+ cells than for the CD8+ cells for all vascular layers. Within the TCFA, non-Th1 cells dominate over Th1 with a five-fold-increase and in stable lesions the Th1/non-Th1 cell ratio returns to 1:2 as seen in normal aorta's (figure 3B). The decline in memory T cells predominantly reflects a reduction localized to the intima ($P<0.01$). The overall number of memory T cells in the media and adventitia remains stable. Native T cell numbers are highly variable. The follicles-like structures vanish and CXCL13 disappeared from the arterial wall. No regulatory T cells or Th17 cells are detected in the stabilized lesions.

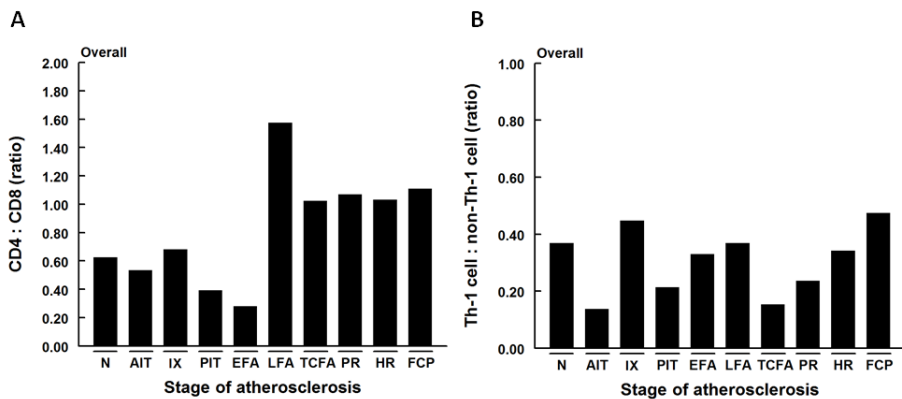


Figure 3. The CD4/CD8 and Th1/non-Th1 ratio per stage of atherosclerosis. **A.** The number of CD8 cells dominate the overall T cell count in normal aorta's and in the early stages of atherosclerosis up until the early fibroatheroma. In late fibroatheroma, vulnerable atherosclerosis (*i.e.* thin cap fibroatheroma and plaque ruptures) and post-ruptured plaques the total number of CD4 and CD8 cells are practically in balance. **B.** Non-Th1 cells dominate the Thelper lineage not only in normal aorta's but also during the entire development of atherosclerotic lesions. Within thin cap fibroatheroma the non-Th1 cells dominate the Th1 cells with a factor 5 and as the lesion stabilizes the Th1/non-Th1 cell ratio returns to 0.4-0.5 as seen in normal aorta's. The solid bars represent the mean number of CD4 (Figure A) or Th1 (Figure B) cells per stage of atherosclerosis divided by the mean number of CD8 (Figure A) or non-Th1 (Figure B) cells per stage of atherosclerosis.

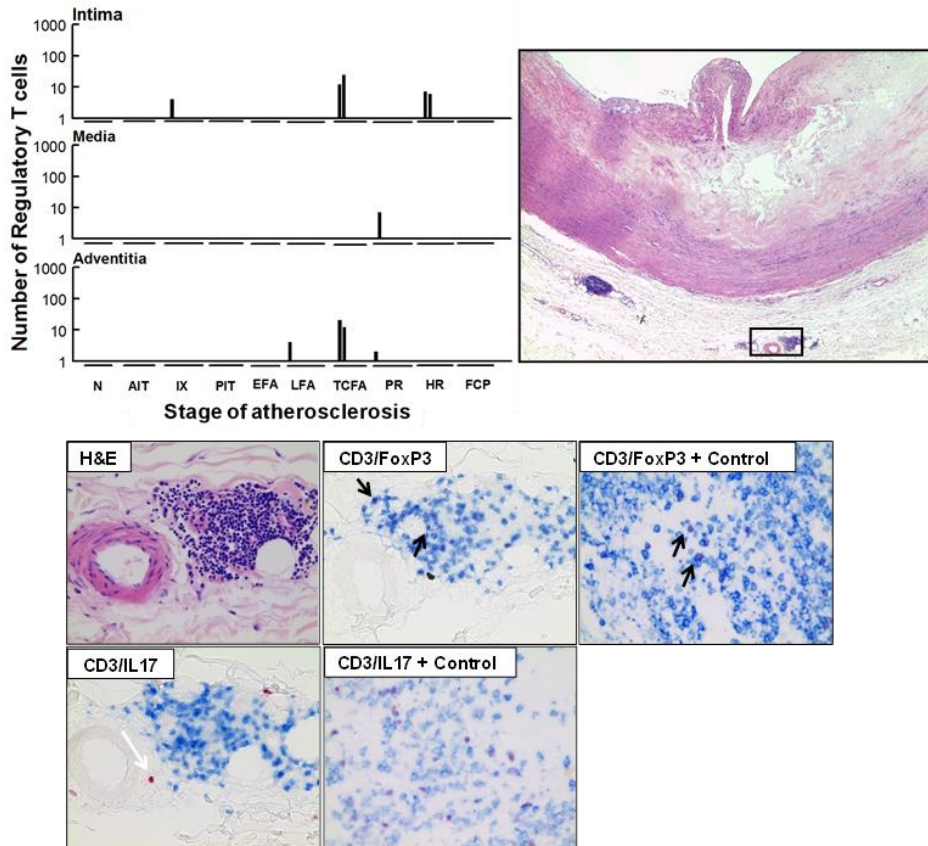


Figure 4. Number of regulatory T cells within the various layers of the aortic wall per stage of atherosclerosis. Only one aortic sample with an intimal xanthoma stained positive for FoxP3+ T cells (4 scattered cells). Within the aortic samples classified as a thin cap fibroatheroma two lesions showed 12 and 24 FoxP3+ T cells in the intima with respectively 20 and 12 FoxP3+ T cells in the adjacent adventitia. A few scattered FoxP3+ T cells were seen in the intima of two healed ruptured plaques. The additional figures show an H&E stained adventitial follicle in a thin cap fibroatheroma with the consecutive sections stained for regulatory T cells (CD3+/FoxP3+) and Th17 cells (CD3+/IL17+). There are a limited number of regulatory T cells (black arrow) in the vulnerable phase and Th17 cells were not identified. A few scattered CD3-/IL17+ cells (white arrow) were seen in the adventitia that, as previous studies showed, can be addressed to macrophages¹⁵. A human tonsil (not associated with the aortic biobank) functioned as a positive control for the CD3/IL17 staining and showed numerous Th17 cells. Ferangi blue was used to develop CD3 and alkaline phosphatase (AP) for FoxP3 and IL17. The solid bars represent the number of FoxP3+ T cells within the various layers of the aorta per stage of atherosclerosis. Total number of cases: 95 (N (9), AIT (9), IX (11), PIT (11), EFA (10), LFA (9), TCFA (9), PR (10), HR (8) and FCP (11)). For abbreviations and a detailed description concerning the classification see Material & Methods section. All images were taken at a 400x magnification.

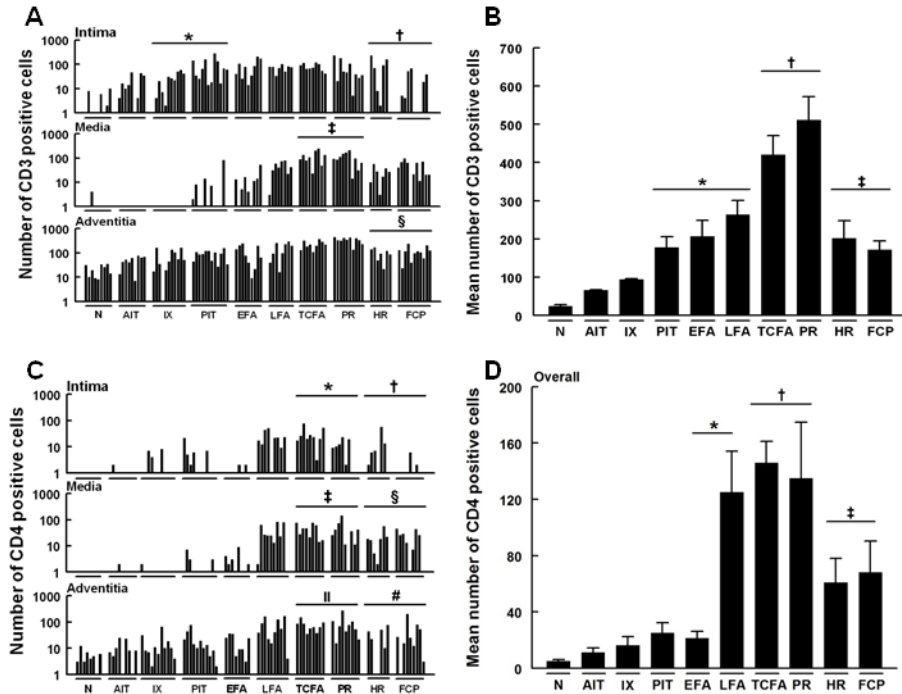


Figure 5. Number of CD3+, CD4+ and CD8+ T cells within the various layers of the aortic wall. **A.** Intimal CD3+ T cells are present in all stages of atherosclerosis. There is a significant influx of CD3+ T cells in intimal xanthomas and pathological intimal thickening when compared to normal aortic wall samples (* $P < 0.009$). The number stabilizes as the lesion becomes vulnerable (*viz.* TCFA and PR). A significant decrease ($\dagger P < 0.014$) is seen in the stabilizing lesions (*viz.* HR and FCP) compared to vulnerable lesions (*viz.* TCFA and PR). The media remains clear of CD3+ T cells until pathological intimal thickening and shows a remarkable increase in CD3+ expressing T cells ($\ddagger P < 0.0001$) in vulnerable lesions. There are an increasing number of CD3+ T cells in the adventitia with advancing atherosclerosis. Stabilized plaques are hallmarked by a significant decrease in adventitial CD3+ T cells ($\S P < 0.0001$) when compared to the vulnerable lesions. **B.** The total number of CD3+ T cells in all vascular layers increases with advancing atherosclerosis. Progressive atherosclerotic lesions (*viz.* PIT, EFA and LFA) show significantly more CD3+ T cells compared to non-progressive lesions (AIT and IX; * $P < 0.0001$). A further increase is seen in vulnerable lesions ($\dagger P < 0.0001$) whereas lesion stabilization is hallmarked by a dramatic decrease ($\ddagger P < 0.0001$). Total number of cases in figures A and B: 95 (N (9), AIT (9), IX (11), PIT (12), EFA (9), LFA (8), TCFA (9), PR (10), HR (7) and FCP (11)). **C.** The intima remains devoid of CD4+ T cells up until late fibroatheroma. There is a significant increase of T-helper cells in vulnerable lesions compared to PIT, EFA and LFA followed by a significant decrease in stabilizing lesions (* $P = 0.008$; $\dagger P < 0.004$). Similar patterns are seen within the media and adventitia ($\ddagger P < 0.0001$; $\S P < 0.003$; $\parallel P < 0.002$ and $\# P < 0.008$). **D.** T-helper cells significantly increase in progressive and vulnerable lesions in comparison to PIT, EFA and LFA followed by a decrease in stabilized lesions (* $P < 0.021$; $\dagger P < 0.0001$ and $\ddagger P < 0.001$). Total number of cases in figures C and D: 91 (N (8), AIT (9), IX (11), PIT (11), EFA (8), LFA (10), TCFA (9), PR (9), HR (7) and FCP (9)).

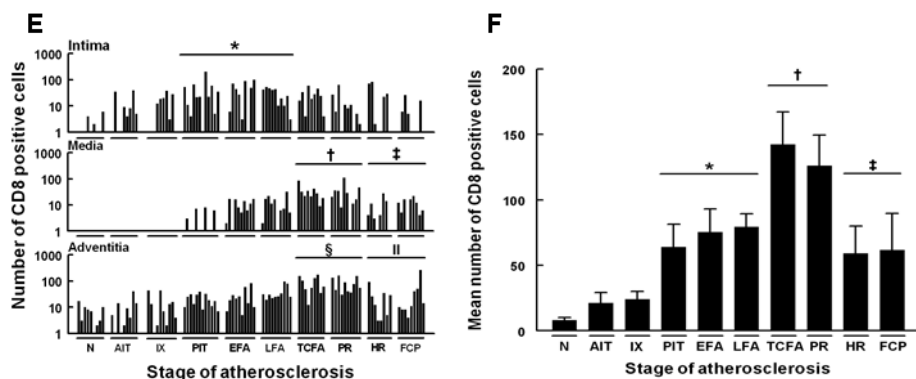


Figure 5 Continued. E. Cytotoxic T cells are more abundantly present in the intima in the progressive stages of atherosclerosis (* $P<0.001$) when compared to normal, AIT and IX. The media and adventitia show an increasing number of cytotoxic T cells within progressive atherosclerotic lesions (PIT, EFA en LFA) compared to the prior phases followed by significant rise within the vulnerable plaques and a significant decrease as the lesions stabilize († $P<0.0001$; ‡ $P<0.0001$; § $P<0.0001$ and ¶ $P<0.0001$). **F.** The total number of cytotoxic T cells follows a similar pattern as the T-helper lineage in figure B; a significant increase within progressive and vulnerable lesions followed by a decrease in stabilized lesions (* $P<0.002$; † $P<0.0001$ and ‡ $P<0.0001$). Total number of cases in figures E and F: 96 (N (9), AIT (9), IX (10), PIT (12), EFA (10), LFA (10), TCFA (9), PR (10), HR (8) and FCP (9)). The vertical axis of figures A, C and E is presented as a log-scale. Each solid bar in figure A, C and E represents the number of positively stained T cells within the intima, media and adventitia of one aortic plaque. The solid bars in figures B, D and F represent the mean total number of cells within the entire aortic wall per stage of atherosclerosis $\pm$ SEM. Abbreviations: N: normal, AIT: adaptive intimal thickening, IX: Intimal xanthoma, PIT: Pathological intimal thickening, EFA: early fibroatheroma, LFA: late fibroatheroma, TCFA: thin cap fibroatheroma, PR: plaque rupture, HR: healed rupture and FCP: fibrotic calcified plaque. For a detailed description concerning the classification see Material & Methods section.

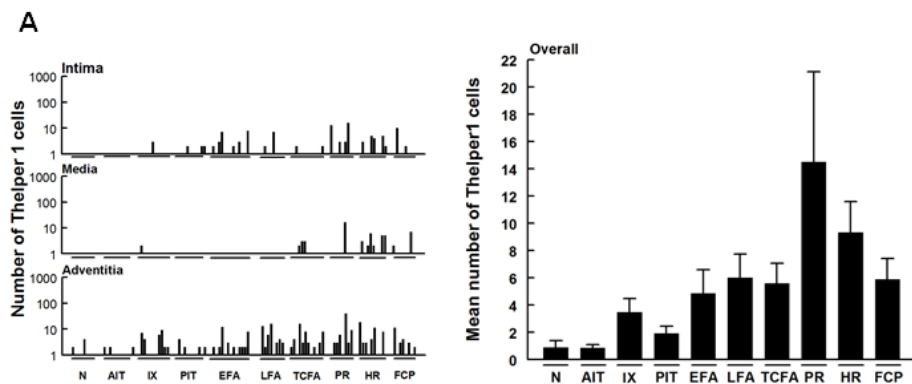


Figure 6. Number of Thelper1 and non-Thelper1 cells within the various layers of the aortic wall. **A.** There is a gradual increase in Thelper1 cells in the intima and adventitia with advancing atherosclerotic lesions. An increase is seen in the amount of Thelper1 cells within ruptured plaques; note the fact that these cells are mainly located in the adventitia. Total number of cases in figure A: 97 (N (9), AIT (9), IX (11), PIT (11), EFA (12), LFA (8), TCFA (12), PR (8), HR (9) and FCP (8)).

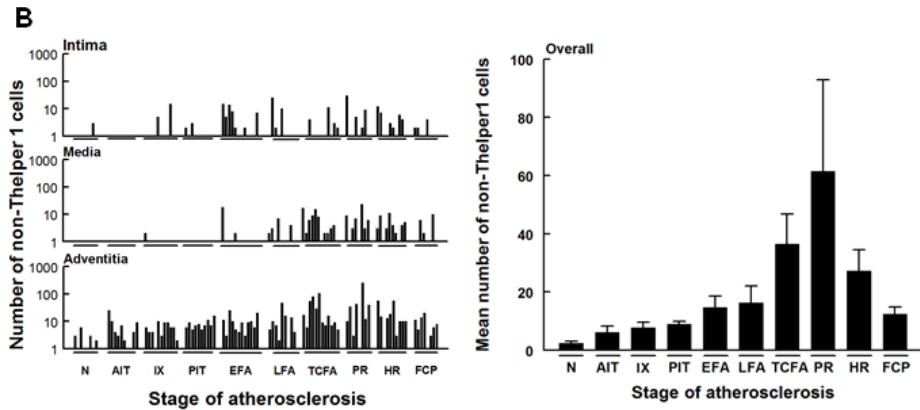
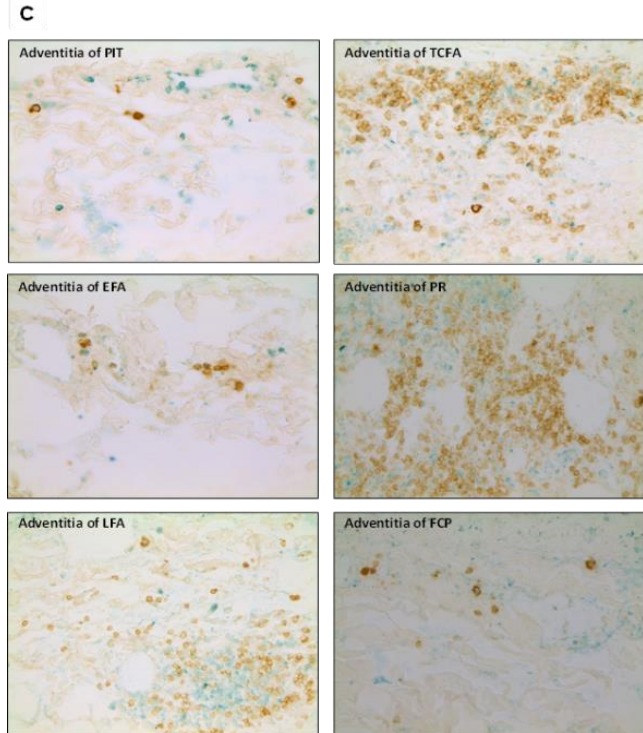


Figure 6. Continued. **B.** There are far more non-Thelper1 cells within the aortic wall compared to the number of Thelper1 cells. However, the same gradual increase is seen in number as the atherosclerotic lesions progress towards the vulnerable thin cap fibroatheroma. Again, an increase is seen within ruptured plaques. Total number of cases in figures B: 98 (N (8), AIT (10), IX (11), PIT (10), EFA (12), LFA (9), TCFA (12), PR (8), HR (10) and FCP (8)).

C. Illustrative images of the adventitia adjacent to the intimal plaque in the various stages of the atherosclerotic process with CD4/Tbet double-staining corresponding with the presented graphs A and B. Thelper1 cells are identified by staining positive for CD4 (brown; diaminobenzidine chromogen) and Tbet (methylgreen). Non-Thelper1 cells are CD4 positive and Tbet negative. Note that there are numerous cells that only stain Tbet positive. The vertical axis of the figures with the intima, media and adventitia separated is presented as a log-scale and the solid bars represent the number of Thelper1 or non-Thelper1 cells within the various layers of one aortic plaque. The solid bars in the adjacent figures represent the mean total number of Th1 cells or non-Th1 cells within the entire aortic wall $\pm$ SEM.



For abbreviations and a detailed description concerning the classification see Material & Methods section. All images were taken at a 400x magnification.

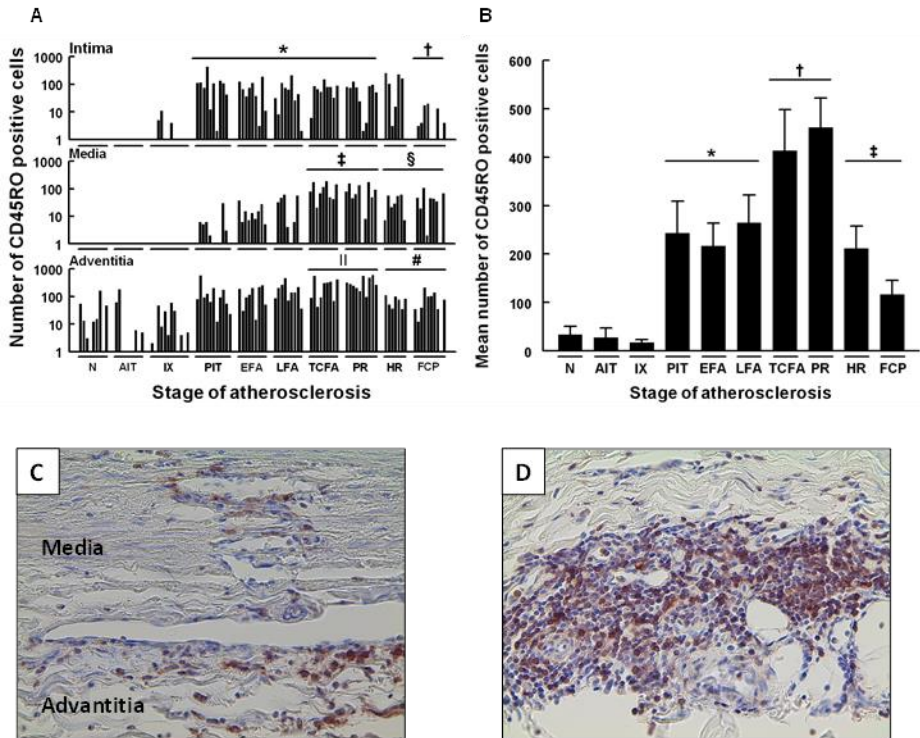


Figure 7. Number of memory T cells within the various layers of the aortic wall. **A.** Progressive atherosclerotic lesions and vulnerable plaques represent a high amount of CD45RO+ T cells in the intima (* $P < 0.011$). Only fibrotic calcified lesions show a significant decrease compared to vulnerable lesions ($\dagger P < 0.001$). In comparison with normal, AIT and IX the media and adventitia show an increasing number of memory T cells within progressive atherosclerotic lesions followed by significant rise within the vulnerable plaques and a significant decrease as the lesions stabilize ($\dagger P < 0.0001$; $\ddagger P < 0.0001$; $\S P < 0.0001$ and $\eta P < 0.0001$). **B.** Memory T cells significantly increase in number within progressive and vulnerable lesions followed by a decrease in stabilized lesions (* $P < 0.0001$; $\dagger P < 0.0001$ and $\ddagger P < 0.0001$). **C. D.** Are illustrative images of the CD45RO staining of respectively the adventitia of a LFA and the adventitia of a TCFA corresponding with the presented graphs. All sections were developed with DAB and counterstained with Mayer's haematoxylin. The vertical axis of figure A is presented as a log-scale. The solid bars in figure A represent the number of CD45RO+ T cells within the various layers of one aortic plaque. The solid bars in figures B represent the mean number of CD45RO+ T cells within the entire aortic wall of one aortic plaque $\pm$ SEM. Total number of cases in figures A and B: 95 (N (9), AIT (9), IX (12), PIT (11), EFA (9), LFA (9), TCFA (9), PR (10), HR (7) and FCP (10)). For abbreviations and a detailed description concerning the classification see Material & Methods section. All images were taken at a 400x magnification.

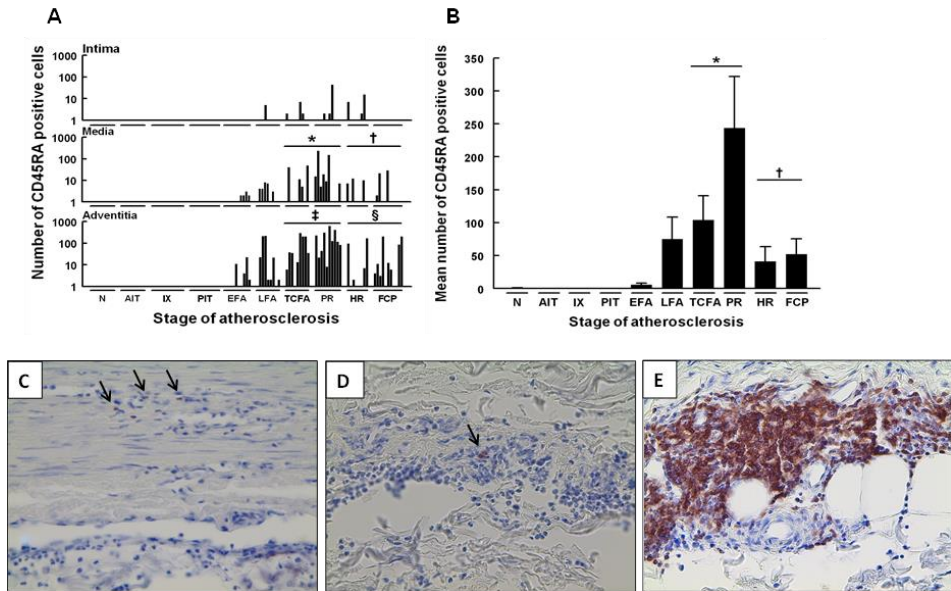


Figure 8. Number of naïve T cells within the various layers of the aortic wall. **A.** Limited numbers of CD45RA+ T cells are detected in the intima of vulnerable lesions. However, the media and adventitia contain a significant increase in naïve T cells within the vulnerable lesions compared to the progressive lesions (EFA and LFA) followed by a decrease in stabilized lesions (* $P < 0.001$; † $P < 0.006$; ‡ $P < 0.0001$ and § $P < 0.0001$). **B.** Naïve T cells increase significantly in number in vulnerable plaques (* $P < 0.0001$) and decrease dramatically in number when the lesions stabilize († $P < 0.0001$). The vertical axis of figure A is presented as a log-scale. The solid bars in figure A represent the number of CD45RA+ T cells within the various layers of one aortic plaque. Notice the significant increase of adventitial naïve T cells in the vulnerable lesions (**D** and **E**) compared to the progressive lesions (**C**) and the naïve T cells alongside the *vasa vasorum* (black arrows). All sections were developed with DAB and counterstained with Mayer's haematoxylin. The solid bars in figures B represent the mean number of CD45RA+ T cells within the entire aortic wall of one aortic plaque $\pm$ SEM. Total number of cases in figures A and B: 97 (N (9), AIT (9), IX (11), PIT (11), EFA (10), LFA (9), TCFA (9), PR (10), HR (8) and FCP (11)). For abbreviations and a detailed description concerning the classification see Material & Methods section. All images were taken at a 400x magnification.

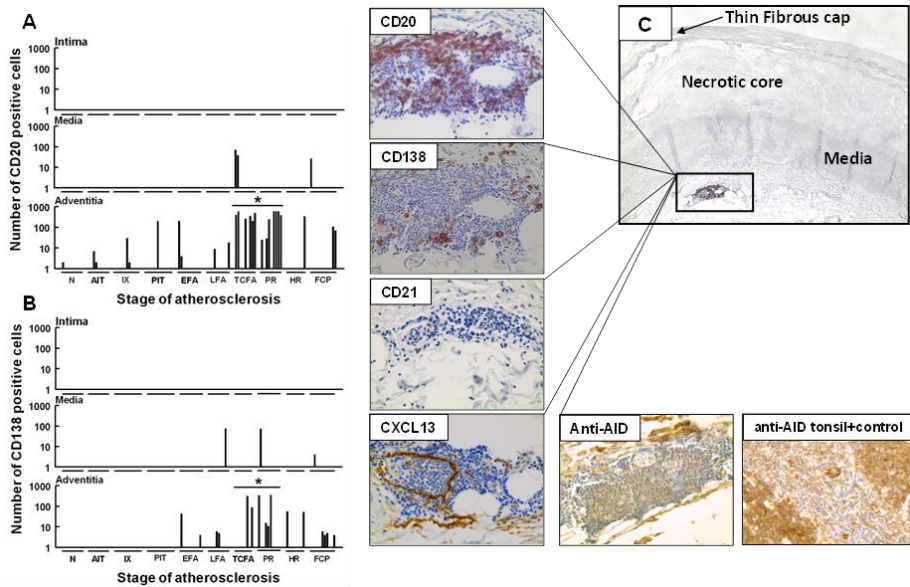


Figure 9. Number of CD20+ B cells and CD138+ plasma cells in the various layers of the aortic wall. The intima and media of the aortic wall remain largely deprived of CD20+ B cells and CD138+ plasma cells (resp. figure A and B), although incidental B cells are found in the adventitia. There is a significant increase in the amount of B cells and plasma cells in the adventitia in close proximity to the intimal plaque in the vulnerable lesions compared to the prior phase (* $P < 0.0001$ and * $P < 0.003$). Total number of cases: 91 (N (9), AIT (9), IX (11), PIT (11), EFA (10), LFA (9), TCFA (9), PR (10), HR (8) and FCP (11)). Figure C shows an adventitial follicle in a thin cap fibroatheroma consecutively stained for CD20, CD21 and CXCL13. Notice the abundance of (CD21 negative) B cells clustered around the CXCL13 positive radiating pattern of tube-like structures. Anti-AID stained negative within the adventitial infiltrates. Due to the disturbing amount of background staining, a high resolution image of a positive control for the anti-AID staining in a human tonsil is provided to illustrate the mature B-cell. All images were taken at a 400x magnification. The vertical axis is presented as a log-scale. The solid bars represent the number of CD20+ B cells or CD138+ plasma cells within the various layers of one aortic plaque. For abbreviations and a detailed description concerning the classification see Material & Methods section.

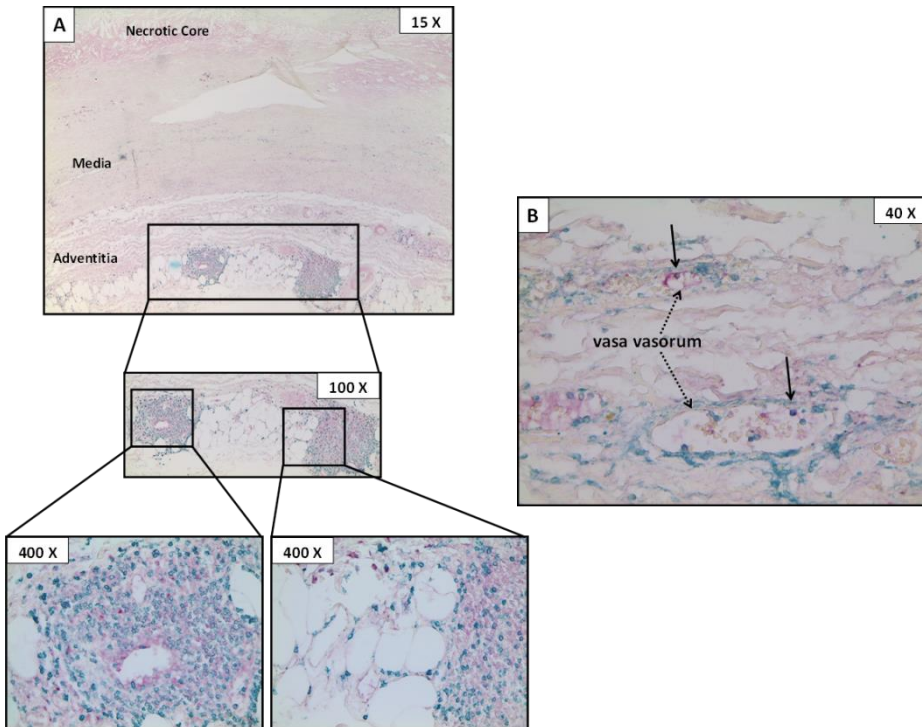


Figure 10. Thin cap fibroatheroma stained for CD45R0/CCR7¹⁶. Immunohistochemical double staining for CCR7/CD45R0 was done to differentiate between effector memory T cells (CCR7-) and central memory T cells (CCR7+). **A.** The adventitia of the aortic wall containing a thin cap fibroatheroma consists of numerous T cells in follicle-like structures. All the CD45R0+ T cells (chromogen Ferangi blue) within the lesion and infiltrates are effector memory T cell (CCR7-; otherwise warp red (klinipath, Duiven, Netherlands)). Only a few cells stain positive for CCR7 but negative for CD45R0. **B.** The only few central memory T cells (CD45R0+/CCR7+) identified in the aortic wall are located within the *vasa vasorum* of the adventitia. The image was taken at a 400x magnification.

DISCUSSION

A wealth of preclinical evidence supports a critical role of the adaptive immunity in the initiation and progression of atherosclerosis^{17,18}. This study, based on histological observations in human aorta sections, confirms extensive and progressive involvement of cellular components of the adaptive immune response in atherosclerosis¹⁹. Findings point to profound changes in the nature of the response in the lead up to plaque destabilization and show extinguishing of inflammatory processes upon plaque stabilization. This study reveals fundamental differences between the human atherosclerotic disease *and* mouse models of the disease; particularly with respect to a very limited presence of regulatory T cells, absence of Th17 cells throughout the atherosclerotic process, and lack of B-cells in the early-, intermediate-, and final stages of the process.

Insight in the atherosclerotic process highly depends on observations from murine models of the disease²⁰. Indeed, genetically modified mouse models have been critical for understanding the atherosclerotic process. Yet, by virtue of the metabolic adaptations in the lipoprotein metabolism, necessary to induce atherosclerotic lesion formation, the process in these animals is essentially lipid driven; a situation that may not fully mimic the human situation^{21,22}. Translation of rodent findings is further obscured by critical dependence on genetic backgrounds with Th1 dominated immune responses in order for atherosclerosis to develop; by the fundamental and intrinsic differences in inflammatory and immune responses between mice and men and by failure of the experimental lesions to progress to culprit lesions (vulnerable plaque) formation^{23,24,25}. Consequently, information provided by these models may be biased, and is incomplete at least with respect to vulnerable lesions. As result the preclinical observations may not directly translate to the human situation^{8,9}.

Data on human atherosclerosis is also limited, a situation largely reflecting the fact that most observations are made on material obtained during surgical procedures (e.g. endarterectomy). This material typically represents the final stage(s) of the disease and, in the case of an endarterectomy material, will not provide information on the outer media and the adventitia, both major interphases in vessel wall inflammation. With this in mind, we set up a biobank of aortic wall samples from organ grafts designated for transplantation. Material from this bank almost covers the full life span (5-80 years) and shows a nearly equal sex distribution. The relatively healthy pre-mortal status of the donors is reflected by minimal use of statins and antihypertensive drugs. Classification was done for all individual tissue sections in the bank, *viz.* each individual tissue block was Movat and haematoxylin stained, and histologically staged using an established adapted version of the AHA classification system^{12,13}. Modifications in the adapted classification system highlight specific critical morphological events in the final stages of the disease process. This allows for a more precise interpretation of processes occurring during plaque destabilization and subsequent healing. An earlier systematic evaluation of material in the biobank showed that the bank covers the full spectrum of atherosclerotic disease¹². Exact morphologic descriptions and examples of the different lesions have been published and discussed previously¹².

Immunohistochemical staining for CD3, CD4 and CD8 shows progressive T-cell accumulation during the atherosclerotic process. The earlier phases are dominated by diffuse cytotoxic T cell infiltration but progressive quantities of T helper cells are found during progression of the disease resulting in an increase in the CD4⁺/CD8⁺ T cell ratio during disease progression. These observations are in line with an earlier report on renal artery atherosclerosis, and with observations from

other progressive inflammatory disorders^{26,27}. Due to inherent limitations of paraffin embedded tissue, we were unable to test whether these shifts reflect an (auto)immune phenomenon as has been proposed in the context of advanced atherosclerotic disease^{28,29}.

CD4/T-Bet double staining was used to identify Th1 cells. T-bet is the lineage defining transcription factor for Th1 cells³⁰. In the absence of Th17 and with minimal regulatory T cells throughout the atherosclerotic spectrum we defined the CD4+/T-bet negative cells as non-Th1 cells. Quantification of the Th1/non-Th1 ratio on basis of the CD4/T-Bet double staining did not confirm Th1 dominance in human atherosclerotic process. This may indicate that the presumed Th1 dominance in atherosclerosis reflects an artificial phenomenon related to the requirement of Th1 dominated backgrounds in mouse models in order for atherosclerosis to develop. Yet, alternative explanations are that identification based on T-Bet positivity underestimates the number of Th1 cells or, vice versa, that identification based on cytokine profiles overestimates the number of Th1 cells³¹.

Findings for the CD45RO (T-memory cells) and CD45RA (naïve T-cells) largely follow observations for other solid organs (predominance of T-memory cells) and extend those made in rodent models of atherosclerotic disease showing increasing T-cell infiltration during disease progression, with a clear maximum during plaque destabilization followed by a rapid decline during plaque stabilization (plaque healing)³². A remarkable and novel observation is the change in the inflammatory foot print that accompanies plaque destabilization, and that fully resolves during plaque stabilization. This change appears to precede plaque destabilization, is confined to the area adjacent to the culprit lesion. On the cellular level this change is characterized by a sharp increase in the number of T-helper cells, emergence of naïve T-cells, and the appearance of B-cells and occasional plasma cells in tertiary follicle-like structures. B-cells in the follicles are largely Activation-induced cytidine deaminase (AID) and CD21 negative indicating gross absence of B-cell maturation^{33,34}. These observations suggest that signals promoting B-cell homing and follicle assembly accompany (and may even precede) plaque destabilization, but that essential signals critical for maturation of B-cells are missing.

The chemokine CXCL13 is a pivotal homeostatic signal for follicle formation^{35,36}. As of the appearance of follicle-like structures we performed immunostaining for CXCL-13 and observed CXCL-13 expression exclusively in association with the follicle-like structures in the vulnerable lesions, *viz.* CXCL-13 was not detected in the other stages of the disease. CXCL13 distribution shows a remarkable radiating pattern of tube-like structures that may reflect expression in dendritic cells³⁷.

In the light of the remarkable association between CXCL13 and plaque instability, we tested an association between plasma CXCL13 levels and acute cardiovascular

events (acute myocardial infarction) in plasma samples of the Mission study³⁷. CXCL13 was largely undetectable and no association was found between CXCL-13 and cardiovascular events (results not shown). Absence of such an observation may reflect the highly localized character of CXCL13 expression in the vulnerable lesion.

B-cells are considered critical players in the atherosclerotic process, albeit their role (protective, detrimental) is still under debate^{38,39}. Our findings of a very restricted presence of B-cells in human atherosclerosis follow observations from Frostegård *et al*⁴⁰. Hence, these findings and gross absence of signs of B cell maturation in the infiltrating B cells, exclude an autocrine or paracrine role of B-cells in the human atherosclerotic process. Yet, humoral factors released by B-cells in para vascular lymph nodes or more distant locations may well be involved in the disease process.

Regulatory T cells were incidentally found in a subset of the vulnerable lesions, whereas Th17 cells were fully absent in the aortic sections. This latter observation follows results of a comprehensive analysis of cytokines in atherosclerotic wall samples that failed to detect IL-17 in all samples tested (detection threshold of the assay <1 pg/L)⁴¹. Although these observations do not exclude a distant role for these cell populations, they seem to contradict observations from murine models implying these cells as local key-players in the atherosclerotic process.

Limitations: this study was performed on aortic sections of deceased individuals, as such seemingly the continuous data in this study is composed of incidental findings from a large series of patients and the data may therefore not necessarily reflect longitudinal data. Another limitation of the study is the fact the all findings are based on IHC using paraffin-embedded tissue sections. Although IHC has the advantage of showing the spatial relationships, multiple stainings are elaborative, and only allow for a very limited marker sets. Consequently, findings in this study should be considered at a global level. *I.e.* the apparent mismatch between the total number of CD3 count and the sum of CD4 and CD8 may reflect abundance of other CD3 expressing populations such as NK and NK-T cells but also reflect technical limitations of IHC with different epitope availabilities and antibody efficacies. Yet, it is likely that the latter phenomena would equally apply to all tissue sections, and it is thus unlikely that this would influence the conclusions of the study.

In conclusion, this study shows clear changes in the cellular components of adaptive immune system in anticipation of and during plaque destabilization. Observations that suggest that changes in the inflammatory foot print precede and accompany plaque instability. It is tempting to speculate that delineation of this chain of events may provide clues for early culprit lesion detection and plaque stabilization.

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