T-cadherin is present on endothelial microparticles and is elevated in plasma in early atherosclerosis

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Aims
The presence of endothelial cell (EC)-derived surface molecules in the circulation is among hallmarks of endothelial activation and damage in vivo. Previous investigations suggest that upregulation of T-cadherin (T-cad) on the surface of ECs may be a characteristic marker of EC activation and stress. We investigated whether T-cad might also be shed from ECs and in amounts reflecting the extent of activation or damage.

Methods and results
Immunoblotting showed the presence of T-cad protein in the culture medium from normal proliferating ECs and higher levels in the medium from stressed/apoptotic ECs. Release of T-cad into the circulation occurs in vivo and in association with endothelial dysfunction. Sandwich ELISA revealed negligible T-cad protein in the plasma of healthy volunteers (0.90 ± 0.90 ng/mL, n = 30), and increased levels in the plasma from patients with non-significant atherosclerosis (9.23 ± 2.61 ng/mL, n = 63) and patients with chronic coronary artery disease (6.93 ± 1.31 ng/mL, n = 162). In both patient groups there was a significant (P = 0.043) dependency of T-cad and degree of endothelial dysfunction as measured by reactive hyperaemia peripheral tonometry. Flow cytometry analysis showed that the major fraction of T-cad was released into the EC culture medium and the plasma as a surface component of EC-derived annexin V- and CD144/CD31-positive microparticles (MPs). Gain-of-function and loss-of-function studies demonstrate that MP-bound T-cad induced Akt phosphorylation and activated angiogenic behaviour in target ECs via homophilic-based interactions.

Conclusion
Our findings reveal a novel mechanism of T-cad-dependent signalling in the vascular endothelium. We identify T-cad as an endothelial MP antigen in vivo and demonstrate that its level in plasma is increased in early atherosclerosis and correlates with endothelial dysfunction.

Keywords
Atherosclerosis • Endothelial dysfunction • Plasma biomarkers • Endothelial microparticles • T-cadherin

Introduction
Atherosclerosis is a chronic inflammatory disease of large arteries that is initiated by activation and injury of the vascular endothelium upon exposure to multiple cardiovascular risk factors. There is mounting interest in detecting and influencing both the development of atherosclerosis at its non-significant stages and the shift from ‘indolent disease’ to acute disease. Studies have demonstrated that endothelial activation and dysfunction detected by measuring changes in endothelium-dependent vasodilatation or by analysing plasma levels of soluble inflammatory markers (e.g. high-sensitivity C-reactive protein, CD40L) can be a prognostic tool predicting future coronary events.¹–⁴

Among the hallmarks of endothelial activation and damage is the presence of endothelial cell (EC)-derived surface molecules in the circulation. Plasma levels of soluble forms of endothelial membrane proteins (e.g. VCAM-1, E-selectin and P-selectin, LOX-1 receptor) have variously been shown to reflect the severity and the stage of
the disease. EC membrane proteins (e.g. E-selectin, ICAM-1, VE-cadherin) are also present in the circulation on microparticles (MPs), defined as vesicles <1 μm in size shed from the plasma membrane by exocytotic budding in response to cell activation, injury, or apoptosis. Cellular origin (ECs, leucocytes, lymphocytes, erythrocytes, and platelets), amount, and protein composition of MPs in the circulation have been suggested to be useful clinical markers for a great variety of diseases with vascular involvement and hypercoagulability. In addition to their biological actions in inflammation, immune responses and coagulation MPs are capable of directly stimulating intracellular signalling and eliciting cellular responses, such as proliferation, survival, adhesion, chemotaxis, and intercellular communication.

Accumulating evidence supports a role for cell surface T-cadherin (T-cad) survival responses of vascular cells to activation and injury of the vasculature. T-cad expression is increased in vivo in atherosclerotic lesions from human aorta, in experimental restenotic lesions of the carotid artery, and in proliferating ECs and smooth muscle cells in vitro. Upregulation of T-cad in ECs also occurs during oxidative and endoplasmic reticulum stress and functions to diminish stress-induced apoptosis. Overexpression and ligation of T-cad on the EC surface activate signalling pathways (PI3-kinase, Akt, GSK3β) important for cell survival, proliferation, angiogenesis, and motaxis.

The foregoing studies utilized artificial ligands (i.e. antibodies or recombinant protein) to induce T-cad ligation. The present study examines (i) whether T-cad might also be shed from ECs under conditions of stress and in what form, (ii) whether the shed T-cad might be a functionally relevant and naturally occurring ligand for cell surface-expressed T-cad, and (iii) whether T-cad might be present in the circulation in association with endothelial activation/damage.

**Methods**

**In vitro studies**

Established protocols were used for culture of EC, viral transduction, Western blotting, MP isolation, flow cytometry, electron microscopy, sandwich ELISA, and angiogenesis in vitro (see Supplementary material online).

**Clinical study**

The study complies with the Declaration of Helsinki and was approved by the regional ethical committee. Informed consent was obtained from all subjects. They include 225 consecutive patients who underwent coronary angiography at the Luzerner Kantons spitale and a group of 30 volunteers younger than 35 years and without any cardiovascular risk factors. Intravascular ultrasound (IVUS) and quantification of IVUS in the patients was carried out using Volcano equipment. Based on important differences in clinical and angiographic presentation, patients were divided into two groups of those with chronic coronary artery disease (CAD) (n = 163) or with non-significant atherosclerosis (n = 63). Definition of chronic CAD was angiographically and IVUS-proven relevant coronary stenosis (lesions greater than 50%) of at least one vessel and with either angina and/or ST depression during exercise. Definition of non-significant atherosclerosis was diffuse luminal irregularities without relevant lesions (lesions less than 50%) as determined angiographically and by IVUS. The indication for exam in these patients was symptoms of angina or dyspnoea or in the course of investigation of arrhythmias or palpitations. In the group of patients with chronic CAD, mean grade of stenosis was 66.7 ± 26.1%, mean plaque burden was 48.6 ± 10.5%, and mean necrotic core volume (% of plaque volume) was 17.2 ± 8.7%. In the group of patients with non-significant atherosclerosis, mean grade of stenosis was 6.1 ± 15.2%, mean plaque burden was 31.6 ± 12.4%, and mean necrotic core volume (% of plaque volume) was 9.5 ± 8.1%; all P < 0.001 in comparison with the group of patients with chronic CAD.

Measurements of reactive hyperaemia-peripheral arterial tonometry (RH-PAT) and of T-cad in plasma were performed following protocols detailed in Supplementary material online.

**Statistical analysis**

For in vitro studies, differences were variously determined with one-way or two-way repeated-measures analysis of variance (ANOVA) with Tukey’s or Bonferroni’s multiple comparison test, respectively, using Prism 5.0 software (GraphPad Software, San Diego, CA, USA), or three-way repeated-measures ANOVA with Bonferroni’s multiple comparison test using STATA Version 9.2 (STATA Corporation, College Station, TX, USA). Statistical analysis of data from patients and healthy volunteers was performed using STATA Version 9.2. Continuous variables were compared using Student’s t-test assuming normal distributions or the Wilcoxon rank sum test for variables with non-normal distributions. Dichotomous variables were compared using the x²-test or Fisher’s exact test when cell counts were <5. In patients, Spearman’s correlation using continuous variables and logistic regression models using dichotomous variables were used to detect associations between T-cad plasma levels and the degree of ED as assessed by RH-PAT. For the purpose of logistic regression analysis, RH-PAT index was dichotomized at a value of 2.0 and T-cad values were dichotomized at a detection limit of 1 ng/mL. A significance level of 0.05 was assumed for all tests.

**Results**

**T-cadherin can be detected in culture supernatants from endothelial cells**

To determine whether T-cad protein is released from EC, conditioned media were harvested from monolayer cultures of parietal human umbilical vein endothelial cells (HUVECs) and HUVEC transduced with native T-cad protein or T-cad-c-myc-tagged protein following a 24 h culture period. Immunoblotting demonstrated the presence of T-cad protein in media (Figure 1A). Different agents inducing endothelial stress/apoptosis (TNF-α, hydrogen peroxide, and thapsigargin) increased the content of T-cad in the medium (Figure 1B). Release of T-cad into the medium was not prevented by inclusion of various compounds that can inhibit the catalytic cleavage of membrane
proteins from the cell surface, including protein kinase inhibitors—
bisindoylmaleimide, staurosporine, and phorbol myristate
acetate; protease inhibitors—phenylmethylsulphonyl fluoride,
Sigma proteolytic cocktail, and aprotinin; divalent cation scavenger
ethylenediamine tetraacetic acid; lipid-raft disrupting agent
filipin; phospholipase D metalloproteinase inhibitor 1,10-
phenanthroline; phospholipase C inhibitor U73122 (Figure 1C).

However, ultracentrifugation of conditioned media resulted in
the loss of T-cad from supernatants with simultaneous accumu-
lation of the protein in the pellets (Figure 1A), suggesting that the
major fraction of T-cad was released in association with MPs
rather than as soluble or cleaved protein.

**T-cadherin is present on the surface of microparticles released from
endothelial cells**

To determine whether endothelial MPs harbour T-cad, MPs were
isolated from TNF-α-treated HUVEC culture supernatants and
categorized by electron microscopy (Figure 2A), immunoblotting
(Figure 2B), and flow cytometry (Figure 2C–F). Isolated MPs had
diameters of less than 1 μm (Figure 2A). The number of collected
MPs was 190 ± 35/10³ cells (mean ± SD of five isolations)
(Figure 2C). T-cad was present on MPs collected from culture
supernatants of parental and T-cad-overexpressing HUVEC
(Figure 2B). Flow cytometry analysis showed that MPs from par-
ental HUVEC stained positively for annexin V, a characteristic
feature of these apoptotic vesicles exposing phosphatidylserine
on their outer membrane surface, and annexin binding was
decreased in the absence of Ca²⁺, confirming the specificity of
the staining (Figure 2D). MPs isolated from HUVEC culture super-
natants were, as expected, double-positive for CD31 and annexin
V (53% of all MPs) (Figure 2E). With immunolabelling for T-cad and
annexin V, 67% of all the MP population were double-positive, con-
trasting with minimal double-positivity in control samples labelled
with either annexin V in the absence of Ca²⁺ (8%) or isotype
control IgG (14%) (Figure 2F).
Endothelial microparticles expressing T-cadherin on their surface can induce T-cadherin-dependent signalling and angiogenic behaviour in endothelial cells

In vitro homophilic ligation of cell surface T-cad by recombinant T-cad protein or agonistic antibody induces intracellular signalling and the angiogenic phenotype in ECs. How such ligation might occur in vivo is unclear, since in intact EC monolayers T-cad, like other GPI-anchored molecules, is present on the apical surface of cells and not at sites of intercellular contacts.

MPs are recognized as diffusible vectors for the transfer of biological information from one cell to another (homotypic or heterotypic) within proximal or remote tissues. To test the postulate that T-cad-carrying MPs represent ligands inducing T-cad-dependent signalling and behaviour in target ECs, we performed gain-of-function experiments. MPs collected from T-cad-overexpressing ECs (MP-T) and control vector-transduced ECs (MP-E) were analysed for their ability to induce Akt phosphorylation in HUVEC, human aortic endothelial cells (HAECs), and human microvascular endothelial cell line (HMEC-1). Comparable results were obtained for all EC types. Both MP-E and MP-T induced a quick Akt phosphorylation response in parental EC, which was observed within 5 min after addition of MPs to cell cultures, reached its peak within 10–15 min (Figure 3A), and was normalized within 45 min (not shown). The response to MP-T during the first 10 min was significantly more pronounced than that induced by MP-E (Figure 3A).

To exclude that signalling might be induced by traces of foetal calf serum or cytokines in MP preparations, we treated HUVEC with equal volumes of the final phosphate-buffered saline (PBS) supernatants collected during the preparation of

Figure 2 T-cad is present on microparticles (MPs) released from apoptotic ECs. (A) Electron microscopic characterization of MPs isolated from TNF-α-treated (4 μmol/L, 24 h) HUVEC. (B) Immunoblotting for T-cad in MPs isolated from TNF-α-treated parental (wt) and T-cad-overexpressing HUVEC (T-cad+). (C–F) Flow cytometry of MPs from TNF-α-treated HUVEC. (C) MPs were defined on a dot-plot histogram (region R2) and counted using an internal standard of calibrating TruCOUNT beads (R1). (D) Staining with annexin V–FITC conjugate with the confirmation of specific annexin V binding by staining in the absence of Ca²⁺. (E) MPs are double-positive for annexin V and CD31. Isotype control IgG-PE conjugate was used as negative control. (F) MPs are double-positive for annexin V and T-cad. Negative controls: labelling with isotype control IgG and annexin V in the absence of Ca²⁺. Experiments were performed on at least three separate occasions.
MP-E and MP-T (sup-E and sup-T, respectively). No Akt phosphorylation was observed in response to either sup-E or sup-T (Figure 3B).

To validate T-cad-specific effects of MP-T on Akt phosphorylation, we performed gain-of-function and loss-of-function experiments using T-cad-overexpressing and T-cad-silenced HMEC-1.

Firstly, we examined the responses of T-cad-silenced (shT) and control shRNA-transduced (shC) HMEC-1 to MP-T and MP-E derived from T-cad-overexpressing and control-transduced HMEC-1, respectively. The Akt phosphorylation response of shC-HMEC-1 to MP-T was greater than that to MP-E (Figure 3C). In contrast, for shT-HMEC-1, the Akt phosphorylation responses to MP-E and MP-T were comparable (Figure 3C). Secondly, we examined the response of parental HMEC-1 to MPs collected from shT-HMEC-1 or shC-HMEC-1. MPs from T-cad-silenced HMEC-1 induced a markedly weaker and delayed response than...
MP from shC-HMEC-1 (Figure 3D). These data support that activation of intracellular signalling in target EC by MP-T depended upon T-cad homophilic interactions.

Next, we investigated whether effects of T-cad-carrying MP on intracellular signalling might translate into relevant biological effects. Angiogenic behaviour of EC was selected as the functional read-out since in vitro and in vivo studies have identified T-cad as a proangiogenic molecule. Moreover, an important action of MP in the vascular system is their ability to induce angiogenesis. MP-E and MP-T were added to HMEC-1 monolayers and their effects on angiogenic behaviour were monitored by time-lapse videomicroscopy. In parental HMEC-1, both types of MP induced a proangiogenic phenotype and formation of net structures, but for MP-T the total length of tubular structures was greater (Figure 4A), and onset of network formation occurred earlier (Figure 4B). The angiogenic response of T-cad-silenced HMEC-1 (shT) to MP-T was significantly lower than that of control HMEC-1 (shC) (Figure 4C). These data demonstrate that MP-harboured T-cad can elicit a biologically relevant and T-cad-dependent (i.e. angiogenic) response, and further confirm homophilic interaction of MP-T with T-cad molecules expressed on the surface of target cells.

Figure 4 Endothelial MPs expressing T-cad on their surface facilitate angiogenesis in vitro. MP-E or MP-T were added (10 μg/mL) to parental HMEC-1 monolayers (A, B), T-cad-silenced (shT), and control-transduced (shC) HMEC-1 (C). Formation of tubular structures was monitored by time-lapse videomicroscopy over 36 h. Morphometric analysis of the length of tubular structures (A, C) was performed on either videomicroscopy still-frames or after fixation/staining of cells. Images depict fixed/stained cells (A, 36 h) and still-frames after 0, 24, and 36 h (B) or 24 h (C). Scale bars = 0.1 μm (A, B) or 0.2 μm (C). Data in histograms (mean ± SEM, n = 3) present morphometric analysis at 24 h. One-way (A) and two-way (C) ANOVA; **P < 0.01.
T-cadherin is present in human plasma at increased levels during early stages of atherosclerosis and correlates with endothelial dysfunction

We next addressed whether T-cad might be present in the plasma of human donors and whether plasma levels of T-cad reflect in vivo ED and/or injury. The clinical characteristics of the study populations are shown in Table 1; there was a population of young healthy volunteers, a population of middle-aged patients with intermediate atherosclerotic risk and non-significant atherosclerosis, and a slightly older population with highly prevalent cardiovascular risk factors and documented chronic CAD.

ELISA of plasma samples demonstrated the presence of T-cad in 13.3% of healthy volunteers, in 42.9% of patients with non-significant atherosclerosis, and in 34.6% of patients with chronic CAD (Figure 5A). Levels of T-cad in both patient groups were significantly higher than healthy volunteers (Figure 5A and 8), even with exclusion from the analysis of patients with outlying high T-cad plasma levels (Figure 5B). Levels of T-cad between the patient groups did not differ significantly (Figure 5B).

In patients with chronic CAD or non-significant atherosclerosis, Spearman’s correlation using continuous variables revealed a significant (P = 0.043) dependency of T-cad and degree of ED as measured by RH-PAT (Figure 5C). Patients with elevated T-cad had a lower RH-PAT index (r <- 0.187), further suggesting a link between ED and T-cad release into the circulation. To further evaluate the relationship between T-cad and RH-PAT, we divided the RH-PAT scores into three categories: low (≤ 4), medium (5-7), and high (> 7), and performed a chi-square test of independence on the counts. The Fisher’s exact test showed a P-value of 0.046.

In multivariate logistic regression adjusted for age, gender, cardiovascular risk factors (smoking, diabetes, hypertension, hyperlipidaemia, and family history), and statin use, only a lower RH-PAT index was significantly associated with elevated T-cad (OR 2.2 for elevated T-cad in patients with lower vs. higher RH-PAT; Table 2).

Table 1: Clinical characteristics of healthy volunteers and participating patients

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<td><strong>Age (years)</strong></td>
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<td><strong>Gender</strong></td>
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Microparticles isolated from plasma express T-cadherin on their surface

To analyse whether T-cad is released into the circulation on the surface of endothelial MP, we isolated and characterized MP from the plasma of human donors. Analysis was performed on several plasma samples (n = 7) identified as positive for T-cad protein in ELISA. Figure 6 shows representative stainings of samples from patients with non-significant atherosclerosis. ELISA of plasma samples demonstrated the presence of T-cad in 13.3% of healthy volunteers, in 42.9% of patients with chronic CAD (Figure 5A). Levels of T-cad in both patient groups were significantly higher than healthy volunteers (Figure 5A and 8), even with exclusion from the analysis of patients with outlying high T-cad plasma levels (Figure 5B). Levels of T-cad between the patient groups did not differ significantly (Figure 5B).

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negative control labelling is shown in lower panels), supporting the EC origin of T-cad harbouring MPs in plasma.

**Discussion**

We have previously demonstrated that T-cad protein expression on EC is upregulated under conditions leading to EC activation and damage. Here, we provide evidence that T-cad can be shed from EC and in amounts reflecting the extent of activation or damage. Firstly, *in vitro* studies demonstrated that T-cad protein is present on MP released from stressed/apoptotic EC, and that these MP can induce T-cad-dependent signalling and a proangiogenic functional response in target cells. Secondly, release of T-cad from EC occurs *in vivo* in association with ED.

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**Figure 5** T-cad in the plasma of human donors is increased at early stages of atherosclerosis: T-cad levels in the plasma of healthy subjects (*n* = 30), patients with non-significant atherosclerosis (*n* = 63), patients with chronic CAD (*n* = 162) was measured by sandwich ELISA. (A) Error bars in the scatter plot are mean ± SEM. (B) T-cad levels are given as mean ± SEM. *P*-values indicate significance of difference between subject groups as determined with two-sample Wilcoxon rank-sum (Mann–Whitney) test; comparison with healthy volunteers, comparison between the patient groups either without or with exclusion of patients with outlying high plasma T-cad levels. (C) The correlation between plasma T-cad levels and the degree of ED in the patients (*n* = 225) as measured by RH-PAT was estimated with the Spearman’s method using continuous variables (*r* = −0.19, *P* < 0.05) and is depicted using linear regression with 95% confidence bands (interrupted lines).
negligible T-cad protein was detected in the plasma of healthy human donors, but its level was increased in the plasma from patients with non-significant atherosclerosis and patients with chronic CAD.

Cell surface molecules can be released into the circulation by several mechanisms. Proteolytic cleavage of the extracellular domains of transmembrane polypeptides by matrix metalloproteases, serine proteases, and disintegrins, known as ectodomain shedding, has been demonstrated for growth factor receptors, proteoglycans, and adhesion molecules. Lipid-anchored molecules, such as GPI-proteins or gangliosides can be cleaved from the plasma membrane by phospholipases. In addition to being a mechanism of regulating surface protein expression, enzymatic cleavage is involved in the control of various cell functions since soluble protein fragments can retain their biological activity and act as ligands inducing signalling events in neighbouring cells.

Shedding of endothelial adhesion molecules modulates adhesive properties of luminal ECs and recruitment of inflammatory cells to the vessel wall, thus modulating inflammatory responses. Cleavage of extracellular domains of N- and E-cadherin modulates cell adhesive and migratory behaviour. Our in vitro data obtained using inhibitors of shedding-inducing enzymes and ultracentrifugation argues for T-cad release from EC not as soluble protein, but rather in association with some kind of cellular particles. Flow cytometry confirmed the presence of T-cad on the surface of endothelial apoptotic MP (or ectosomes), both in conditioned culture medium and in the plasma of human donors in vivo. Notwithstanding, proteolysis or lipolysis as alternative mechanisms for shedding surface T-cad cannot be completely excluded, since trace amounts of T-cad were occasionally detected in medium clarified by ultracentrifugation.

In addition to their characteristic loss of plasma membrane asymmetry and exposure of phosphatidylserine (annexin V positivity) on the outer leaflet, MP are positive for T-cad, permitting us to conclude that a significant pool of microvesicles carrying T-cad represent MPs.

Table 2

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<th>P-value</th>
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<td>Age (OR per year increase)</td>
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<td>Gender (OR for female vs. male)</td>
<td>1.61 (0.84–3.10)</td>
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<tr>
<td>Smoker (OR for smoker vs. non-smoker)</td>
<td>1.33 (0.73–2.44)</td>
<td>0.356</td>
</tr>
<tr>
<td>Diabetes (OR for diabetes vs. no diabetes)</td>
<td>0.92 (0.43–1.99)</td>
<td>0.840</td>
</tr>
<tr>
<td>Hypertension (OR for hypertension vs. no hypertension)</td>
<td>0.95 (0.53–1.72)</td>
<td>0.873</td>
</tr>
<tr>
<td>Hyperlipidaemia (OR for hyperlipidaemia vs. no hyperlipidaemia)</td>
<td>0.70 (0.37–1.33)</td>
<td>0.282</td>
</tr>
<tr>
<td>Family history (OR for family history vs. no family history)</td>
<td>1.04 (0.57–1.88)</td>
<td>0.907</td>
</tr>
<tr>
<td>Statin use (OR for statin vs. no statin)</td>
<td>1.00 (0.49–2.05)</td>
<td>0.998</td>
</tr>
<tr>
<td>RH-PAT index (OR for low vs. high)</td>
<td>2.19 (1.16–4.16)</td>
<td>0.016</td>
</tr>
</tbody>
</table>

The precise mechanisms leading to increased elevation of MP-bound T-cad in the plasma of patients with non-significant atherosclerosis and patients with chronic CAD are not clearly defined. T-cad is upregulated on ECs early upon activation and stress in vitro, is elevated on ECs during atherosclerosis, and can undergo redistribution within the plasma membrane of activated vascular cells. Therefore, local conditions of EC activation owing to endothelial shear stress together with the presence of proatherogenic inflammatory, thrombotic, apoptotic, or oxidative substances, and a possible concomitant upregulation of T-cad gene expression or even redistribution of protein within the plasma membrane are factors that might underlie increased membrane vesiculation and/or increased levels of T-cad in blebbing plasma membrane domains.

There is considerable interest in plasma biomarkers that provide information about the functional status of the cardiovascular system at early stages of atherogenesis before acute clinical manifestations take place. Since recent IVUS studies demonstrate high incidence of coronary atherosclerotic lesions even in asymptomatic teenagers and young adults, of special importance would be markers that help to detect not only intimal thickening but also the shift from silent to acute disease that manifests itself in functional abnormalities. Promising candidates include markers of acute or chronic inflammatory processes occurring in the coronary vessel wall during atherogenesis, such as soluble forms of endothelial adhesion molecules, or C-reactive protein for which even a small elevation was shown to be a powerful independent predictor of future vascular events in apparently healthy asymptomatic individuals. Accumulating evidence supports a prognostic role of ED in CAD, however, only von Willebrand factor and E-selectin are viewed as biochemical markers specifically reflecting ED. Since our patient subgrouping was based on angiographic and IVUS parameters, an elevation in plasma T-cad cannot be unequivocally interpreted as a criterion for distinguishing between indolent and functionally relevant disease. Nevertheless, ED represents one of the earliest functional disturbances that precede acute coronary events. We found both the presence of T-cad plasma in patients with non-significant atherosclerosis and an

Table 2 Multivariate logistic regression
association between T-cad levels and degree of ED. Therefore, it is plausible to consider T-cad as a marker of early-stage atherosclerosis, which is clinically silent but characterized by disturbances in endothelial function.

Interestingly, although ELISA clearly demonstrated elevation of T-cad in plasma in patients with atherosclerosis when compared with healthy subjects, T-cad levels between patients with chronic CAD and those with non-significant atherosclerosis (albeit slightly higher) were not significantly different. This might be explained by the assumption that T-cad levels reflect not so much the actual vessel occlusion but rather the state of endothelial activation and dysfunction. Our previous in vitro studies on oxidative stress and endoplasmic reticulum stress show that T-cad expression on EC is upregulated within 2–3 h of treatment with pro-inflammatory and pro-apoptotic agents and thereafter declines when manifestations of acute cell damage and death become obvious. Here, T-cad upregulation represents a protective mechanism that can shift the balance in cellular stress response to the prosurvival signalling branches, which is only later followed by alarm and apoptotic phases. In vivo, this phenomenon might

![Figure 6](image-url)
translate into an attempt of vascular cells to limit tissue dysfunction and damage at the onset of atherosclerosis.

Many studies have demonstrated that MP shedding is a hallmark of endothelial activation and injury. Circulating MP levels correlate with the degree of ED, impaired vasodilatation in patients with chronic and end-stage renal failure, and acute coronary syndromes, and are increased in patients with hypercholesterolaemia, hypertension, and diabetes.\textsuperscript{5–7} Two recent papers present evidence for potential usefulness of endothelial MP levels in the identification of patients at immediate risk of acute cardiovascular events. Assessment of endothelial dysfunction by measurement of endothelial MP independently predicted cardiovascular events in patients with coronary heart disease suggesting this readout as an important component of a multiple biomarker strategy of improving risk stratification.\textsuperscript{31} Bernard et al. showed an association between CD144$^+$-MP and the presence of coronary non-calcified vulnerable plaques indicating that elevation of endothelial MP may predict acute thrombotic events in type 2 diabetic patients.\textsuperscript{32}

Identification of T-cad as an endothelial MP antigen in vivo has potential value with respect to the detection of endothelial activation/injury. However, it must be recognized that considerable technical difficulties are associated with standardized isolation, measurement, and characterization of MP, and utilization of MP as a reproducibly quantifiable parameter is far from the mainstream clinical analysis. Although cell-free T-cad is mostly MP-bound, we successfully applied ELISA-based methodology to measure T-cad in human plasma. This methodology offers a more reliable and simple technical opportunity to detect ED characterized by the elevation of ‘biomarker’ endothelial MP in plasma.

A limitation of the study is that healthy subjects in the control group were significantly younger than patients in both groups with atherosclerosis. However, it seems unlikely that age is a potential confounder that results in elevated plasma T-cad in our patient groups. In our multivariate logistic regression analysis, age does not show up as a factor (OR 1.0). This is important since the other variables are age-prone factors. On the other hand, a lower RH-PAT index (ED) shows up with an OR 2.2 ($P < 0.016$), which underscores that the level of T-cad in plasma is a factor that relates to endothelial activation/dysfunction, and, irrespective of age, may reflect a prevailing state of vascular activation and injury.

Relevance for an association between T-cad in plasma and ED is underscored by the recent genome-wide association studies aimed at identification of genetic variants contributing to cardiovascular disease. The Framingham Heart Study\textsuperscript{33} and Org et al.\textsuperscript{34} demonstrated correlations between single nucleotide polymorphisms within intron 11 and promoter regions of CDH13 gene and long-term average diastolic and systolic blood pressure phenotypes and arterial stiffness, indicating a role for T-cad in progression of hypertension. The impact of T-cad within the complicated network of processes contributing to cardiovascular disorders is seemingly broad. T-cad has been identified as a receptor for adipose tissue-derived adiponectin,\textsuperscript{35} a hormone that profoundly affects progression of atherosclerosis and diabetes by modulating insulin sensitivity, lipoprotein metabolism, and EC function. T-cad also acts as a receptor for low-density lipoproteins,\textsuperscript{36} suggesting a role in the regulation of EC function in hypercholesterolaemia.

Our study has also revealed a novel mechanism of T-cad-dependent signalling in the vasculature. Like other members of the cadherin superfamily, T-cad is capable of homophilic–ligation interactions mediating intercellular adhesion and inducing intracellular signalling events. Downstream signalling pathways seem to be activated by both T-cad ligation and upregulation.\textsuperscript{13,14,16,18} Ligation of T-cad on the cell surface is equivalent in vivo to trans-interactions between T-cad molecules on neighbouring cells and can be mimicked in vitro by the addition of recombinant T-cad protein or agonistic antibody to T-cad-expressing cells. In the case of T-cad overexpression, downstream signalling might be induced either by cis-clustering of T-cad molecules within the plasma membrane or by increased engagement of T-cad-signalling adaptor molecules. In the intact endothelial monolayer, T-cad is expressed only on the apical surface and not in the intercellular contacts.\textsuperscript{19} Such a cellular localization pattern in vivo (i.e. facing the vessel lumen) restricts opportunities for T-cad ligation-induced signalling. By demonstrating that T-cad-harboured MP can induce Akt phosphorylation and the proangiogenic phenotype in target EC, we have identified MP as a novel physiological ligand capable of initiating T-cad-dependent signal transduction in the endothelium. These T-cad-mediated homophilic-based interactions may occur both in a paracrine or in an autocrine manner, i.e. induce signalling in distant ECs or ECs from which MPs were released.

Depending on the origin and protein composition, the outcome of MP binding to endothelial and blood cells might be adverse (e.g. promotion of thrombotic events) or beneficial (e.g. tissue regeneration).\textsuperscript{7,8} In vivo, the interaction of plasma-delivered T-cad-harbouring MP to the luminal endothelium, accompanied by an upregulation of surface T-cad expression on activated/injured EC, might function to regulate pathophysiological processes involving endothelial activation and stress, and therefore influence vessel remodelling.

Supplementary material
Supplementary material is available at European Heart Journal online.

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References


