Vascular Smooth Muscle Cell Senescence Promotes Atherosclerosis and Features of Plaque Vulnerability

Running title: Wang et al.; VSMC senescence in atherosclerosis

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Abstract

**Background**—Although vascular smooth muscle cell (VSMC) proliferation is implicated in atherogenesis, VSMCs in advanced plaques and cultured from plaques show evidence of VSMC senescence and DNA damage. In particular plaque VSMCs show shortening of telomeres, which can directly induce senescence. Senescence can have multiple effects on plaque development and morphology; however, the consequences of VSMC senescence or the mechanisms underlying VSMC senescence in atherosclerosis are mostly unknown.

**Methods and Results**—We examined expression of proteins that protect telomeres in VSMCs derived from human plaques and normal vessels. Plaque VSMCs showed reduced expression and telomere binding of Telomeric repeat-binding factor-2 (TRF2), associated with increased DNA damage. TRF2 expression was regulated by p53-dependent degradation of TRF2 protein. To examine the functional consequences of loss of TRF2, we expressed TRF2 or a TRF2 functional mutant (T188A) as either gain or loss of function studies *in vitro* and in ApoE−/− mice. TRF2 overexpression bypassed senescence, reduced DNA damage, and accelerated DNA repair, whereas TRF2<sup>188A</sup> showed opposite effects. Transgenic mice expressing VSMC-specific TRF2<sup>T188A</sup> showed increased atherosclerosis and necrotic core formation *in vivo*, whereas VSMC-specific TRF2 increased relative fibrous cap and decreased necrotic core areas. TRF2 protected against atherosclerosis independent of secretion of senescence-associated cytokines.

**Conclusions**—We conclude that plaque VSMC senescence in atherosclerosis is associated with loss of TRF2. VSMC senescence promotes both atherosclerosis and features of plaque vulnerability, identifying prevention of senescence as a potential target for intervention.

**Key words:** atherosclerosis; vascular disease; smooth muscle; senescence
Proliferation of vascular smooth muscle cells (VSMCs) is a central axiom of most models of atherosclerosis, promoting atherogenesis as a ‘response to injury’\(^1\) or inflammation\(^2\). However, most heart attacks are caused by rupture of a ‘vulnerable’ plaque with a thin VSMC-poor fibrous cap overlying a relatively large necrotic core\(^3,4\). Plaque repair requires VSMC proliferation, and is thus beneficial at this stage. However, VSMCs from advanced human plaques show poor proliferation and premature senescence in culture\(^5\) and \textit{in vivo}\(^6\); furthermore, fibrous cap VSMCs show extensive DNA damage, marked telomere shortening, and markers of senescence\(^7\). Although these findings suggest that VSMC senescence may be important in atherogenesis, its mechanisms and direct consequences are unproven.

Replicative cell senescence is mediated in part by telomeres, which shorten during replication and ultimately trigger a DNA damage response (DDR) and growth arrest. Telomeres comprise tandem DNA repeats that are maintained in a compact T-loop structure by the shelterin complex of telomere-associated proteins, including TRF1, TRF2, POT1, TIN2, RAP1 and TPP1. Shelterin proteins restrict access to telomerase and exonucleases/ligases, thus avoiding inappropriate telomere elongation and shortening respectively, and prevent exposure of chromosome ends that are recognised as double-stranded DNA breaks (DSBs). Although each of the shelterin proteins is important for telomere maintenance, Telomeric repeat-binding factor-2 (TRF2) has a particularly critical role. TRF2 regulates replicative senescence in part by reducing telomere length at senescence\(^8,9\), and can also stop the ataxia telangiectasia kinase (ATM) from initiating a DDR from functional telomeres\(^10\). Loss of TRF2 induces multiple features of senescence, including irreversible growth arrest, expression of senescence-associated \(\beta\)-Galactosidase (SA\(\beta\)G), and telomere dysfunction with chromosomal fusions\(^11-13\). TRF2 can also regulate cell longevity in a telomere-independent manner by direct association with multiple
DDR proteins, including ATM, Nijmegen Breakage Syndrome-1 (NBS1), and Checkpoint kinase-2 (Chk2)\textsuperscript{14-16}. ATM phosphorylates TRF2 after DNA damage on T188 in the dimerization domain\textsuperscript{17}; defective phosphorylation (e.g. thr > ala, T188A) prevents fast DNA repair following DSBs that is only part rescued by the phospho-mimetic glutamic acid (T188E)\textsuperscript{18}. Despite differences in mouse and human telomeres, TRF2 inhibition can induce senescence and apoptosis in mouse tissues in vivo\textsuperscript{19}.

We examined the expression of proteins that protect telomeres in human VSMCs derived from atherosclerotic plaques or normal arteries. We show that plaque VSMCs show reduced expression and telomere binding of TRF2, associated with increased markers of DNA damage. Gain and loss of function studies of TRF2 in ApoE\textsuperscript{-/-} mice show that VSMC senescence regulates both plaque development and morphology. We conclude that plaque VSMC senescence in atherosclerosis is regulated by TRF2, and promotes both atherosclerosis and features of plaque vulnerability.

**Methods**

**Human atherosclerotic plaque and normal vessels**

Plaques and normal aorta were obtained from patients undergoing carotid endarterectomy or coronary artery bypass/valve replacement respectively, under informed consent using protocols approved by the Cambridge or Huntingdon Research Ethical Committee. Age and sex-matched human normal aorta and carotid plaque VSMCs were cultured from tissue explants and studied at passages 2–5; VSMC cultures from individual patients were not pooled. Cells were cultured in Dulbecco's modified Eagle's medium (DMEM) containing 20% fetal calf serum (FCS) supplemented with 100U/ml penicillin, 100\(\mu\)g/ml streptomycin and 2mM L-glutamine.
Protein Extraction, Western blotting

Protein extraction from cultured cells and Western blotting were performed as previously described\(^{20}\) and Supplemental Methods.

ChIP

TRF2 association with telomeres was examined by ChIP as described previously\(^{21}\) and Supplemental Methods.

Immunocytochemistry

Immunocytochemistry for nuclear foci was performed as described previously\(^{20}\) and Supplemental Methods.

Immunohistochemistry

Immunohistochemistry of human and mouse arteries was performed as described previously\(^{22}\).

Plasmid Constructs

cDNAs of human TRF2, TRF2\(^{AB}\) (lacking N-terminal basic domain), TRF2\(^{AM}\) (lacking C-terminal Myb domain) and TRF2\(^{ABAM}\) (lacking both basic and Myb domains) in retrovirus pWZL-hygro-nMyc vectors were purchased from Addgene, Ma, USA (originally supplied by Dr. Titia de Lange, NY, USA). cDNAs from human TRF2\(^{T188A}\) (constitutively un-phosphorylated at T188) and TRF2\(^{T188E}\) (constitutively phosphorylated) in pTET vector were kind gifts from Dr. David Gilley (In, USA) and were subsequently subcloned into the pWZL-hygro-nMyc vector.

Generation of Stable Cell Lines

All cDNAs cloned into retrovirus pWZL-hygro-nMyc vectors (TRF2, TRF2\(^{AB}\), TRF2\(^{AM}\), TRF2\(^{ABAM}\), TRF2\(^{T188A}\), TRF2\(^{T188E}\)) or the empty vector were transfected into Bosc23 ecotropic packaging cells using HiPerfect transfection agent (Invitrogen). Replication-deficient virus was used to infect mouse ApoE\(^{-/-}\) VSMCs. Infected cells were selected with 250\(\mu\)g/ml hygromycin.
(Roche) and TRF2 expression confirmed by western blot.

**Cell Culture**

Mouse cell lines were cultured as described in Supplemental Methods.

**Time Lapse Videomicroscopy**

Cells were cultured overnight in 6-well plates at 3x10⁴ cells/well. Cells were incubated in 40 M HEPES (pH 7.2) for 1h prior to recording using OpenLab software, taking 3 image frames/well using a time-lapse BX51 microscope (Olympus), air-cooled CCD camera (CoolSnap) and imaging and analysis software (Soft Imaging Systems). Over 100 cells per cell line were analysed and cell proliferation and apoptosis assayed. Non-division was defined as no observed divisions over 48 hours (~4 cycles).

**SAβG**

Detection of SAβG was as previously described⁷.

**Comet Assay**

10⁵ cells were washed in PBS, re-suspended in 150μl 1% low melting point agarose (LMPA)/PBS and left to cool on Gelbond plastic films (Lonza) at 4°C for 10 min. After immersing in lysis buffer (2.5M NaCl; 100mM EDTA; 10mM Trizma base; pH 10) overnight at 4°C, gel-films were transferred into electrophoresis buffer (0.3M NaOH; 1mM EDTA; pH>13) for 30 min prior to electrophoresis at 24V, 300mA for 30 min at 4°C. Gel-films were washed 3 times in neutralization buffer (0.4M Tris, pH 7.5), once in distilled H₂O and incubated in ice-cold ethanol for 10 min. Films were dried overnight, rehydrated in distilled H₂O for 10 min and stained with 2μg/ml EtBr (Sigma) in the dark at room temperature for 2h. Multiple images per slide were analysed using a Zeiss microscope and comet tail length/moment measured using Comet Assay IV software (Perception Instruments).
Experimental Animals

All animal procedures followed UK Home Office licensing and were approved by the local animal ethical committee. To produce constructs for transgenesis, HA tags were inserted at the N-terminus of TRF2 and TRF2^{T188A} cDNA by enzymatic reaction. Tagged cDNAs were subcloned between the minimal SM22α promoter\(^2\) and polyA sequence in a pBS vector. Transgenic mice expressing SM22α-TRF2 or -TRF2^{T188A} were generated by pro-nuclear injection. Founders were bred to ensure transgene transmission and expression, then backcrossed five times with ApoE\(^{-/-}\) mice prior to experiments. Male and female control ApoE\(^{-/-}\), SM22α-TRF2/ApoE\(^{-/-}\) or SM22α-TRF2^{T188A}/ApoE\(^{-/-}\) mice were fed a high fat diet (HFD- 21% total fat, 0.2% cholesterol, 0% sodium cholate) from 6w-22w, and blood and tissues taken at sacrifice.

Telomere length and TEL-FISH

Telomere length and telomere fluorescence in situ hybridisation was undertaken on VSMCs derived from control and transgenic mice as described in Supplemental Methods.

Plaque morphometry

Atherosclerotic plaque morphometry was determined as described previously\(^2\) and in Supplemental methods.

Lipid measurements, blood counts, blood pressure

Blood was taken from experimental animals every 4 weeks, and serum lipids analysed for total cholesterol, triglycerides, LDL and HDL using a Dade-Behring Dimension autoanalyzer and LDL calculated using the Friedwald formula. Blood counts were analysed on a Coulter counter, and blood pressure determined by tail cuff.

Cytokine and chemokine detection using FlowCytomix assay

FlowCytomix assay (eBioscience) was used to determine chemokine (GM-CSF, MCP-1, MCP-3,
MIP-1α, MIP-1β, Rantes, CXCL1/KC) and cytokine (IL-6) concentrations in conditioned media from cell lines or mouse serum. Samples were analysed according to the manufacturer’s recommendations.

**Statistical Analysis**

Data that were normally distributed (e.g. cultured cell foci and mouse plaque sizes) were compared using one-way ANOVA (GraphPad Prism, Ca, USA) with no adjustments for multiple comparisons. Data that were not normally distributed were examined using Mann Whitney U tests. Cumulative data (e.g. cumulative cell proliferation) were examined by comparison of cell divisions at the end of the experiment using Mann Whitney U tests. Data are expressed as Means±SEM or Means±SD. p<0.05 was considered statistically significant.

**Results**

**TRF2 is reduced in human atherosclerotic plaque VSMCs**

As human atherosclerotic plaque VSMCs show shortened telomeres and premature senescence7, we first examined expression of shelterin complex proteins in VSMCs cultured from normal human aorta or atherosclerotic plaques. Most proteins showed similar expression, but TRF2 was reduced in multiple plaque VSMC isolates (Figure 1A). Relative expression of each protein in plaque vs. aortic VSMCs was: TRF1 (0.88±0.2, p=0.43), TRF2 (0.76±0.06, p=0.011), POT1 (0.68±0.3, p=0.06), TIN2 (1.06±0.1, p=0.69), RAP1 (1.03±0.1, p=0.82), TPP1 (0.88±0.13, p=0.48)(mean±SD, n=3). In addition, ChIP showed that plaque VSMCs had markedly reduced TRF2 bound to telomeres by (Figure 1B), indicating that telomeres in plaque VSMCs are unprotected by TRF2.

TRF2 is regulated at both mRNA and protein levels. There was no difference in TRF2
mRNA expression between aortic (n=3) or plaque (n=3) VSMCs (Figure 1C) by qPCR, despite the marked differences in TRF2 protein expression, suggesting post transcriptional regulation of TRF2 protein. In some cells the stability of TRF2 protein is regulated via ubiquitin-mediated proteasomal degradation, itself regulated by a p53-induced E3 ubiquitin ligase siah-1a. Consistent with previous studies, p53 was increased in whole lysates of human plaques compared with normal vessels, and also increased in SMA-positive cells in plaques compared with normal aorta (Supplemental Figure 1). p53 itself is also regulated by proteasomal degradation, suggesting that this mechanism may be active. Indeed, although p53 protein expression was undetectable in plaque and normal human VSMCs by Western blot (not shown), expression of siah-1a was increased 1.9-fold in plaque vs. aortic VSMCs, associated with 3.5-fold increased expression of the senescence marker p16 (n=3) (Figure 1D). To demonstrate whether p53 regulated TRF2 expression in VSMCs, we examined TRF2 levels in control or p53−/− VSMCs after treatment with the proteasomal inhibitor MG132. Proteasomal inhibition decreased TRF2 expression in control VSMCs (Figure 1E), associated with increased p53 expression. TRF2 was increased in p53−/− null VSMCs, consistent with a role for p53 in regulating TRF2 protein expression, but there was no significant effect of MG132 (Figure 1E).

We next examined whether reduced TRF2 expression in plaque VSMCs was associated with DNA damage. Cultured plaque VSMCs had reduced TRF2 nuclear foci, associated with increased foci of the DNA damage markers phosphorylated ATM (P-ATM) and γ-H2AX (Figure 2A-B). Despite reduced TRF2 foci, plaque VSMCs showed increased co-localization of TRF2 foci with P-ATM and γ-H2AX, indicating DNA damage at telomeres, and co-localization of γ-H2AX with P-ATM (Figure 2A-B). To confirm that TRF2 is also reduced in vivo, and its association with DNA damage markers, we examined cells in normal aorta or advanced plaque
(AHA Grade IV+). Cells expressing γ-SMA in human plaques in vivo also showed reduced TRF2 and increased P-ATM and γ-H2AX expression (Figure 2C-D). These findings suggest that TRF2 expression and binding to telomeres is reduced in plaque VSMCs, associated with increased DNA damage and DDR activation at telomeres.

**TRF2 bypasses senescence whereas TRF2188A promotes VSMC senescence**

Reduced TRF2 in plaque VSMCs, and the known role of TRF2 in protecting against senescence, suggest that TRF2 may be a critical regulator of VSMC senescence. If correct, manipulation of TRF2 might be used to increase or decrease senescence in VSMCs in culture and in vivo to study the consequences of VSMC senescence on plaque development. TRF2 knockout is embryonally lethal in mice. However, TRF2 contains different functional domains, including a C-terminal DNA-binding Myb domain that recognises telomeric tandem repeat sequences at chromosome ends, a flexible hinge domain involved in protein–protein interactions, a TRFH domain required for homodimerization, and a divergent N-terminal domain rich in basic residues (basic domain). Deletion of these regions both alone and together can promote senescence in transformed cells or those from newborns, both mouse and human, and thus act as a useful model for loss of TRF2 function. However, the effect of loss of TRF2 function varies widely between cell types (e.g. 28), and the effect on VSMCs is not known, particularly VSMCs from atherosclerosis models such as ApoE−/− mice.

To find the most effective way of increasing or reducing TRF2 function in VSMCs, we stably expressed full-length human TRF2 or truncations lacking the basic domain (ΔB), the Myb domain (ΔM), or both (ΔB/ΔM), or TRF2T188E/A point mutations in mouse Apolipoprotein E−/− VSMCs (Supplemental Figure 2A-B). Vector-expressing VSMCs senesced at ~passage 14; TRF2, TRF2ΔB, TRF2ΔM or TRF2ΔBΔM all promoted cell proliferation (Supplemental Figure
2C-D) and cells bypassed senescence until >passage 60. In contrast, TRF2\textsuperscript{T188A} or TRF2\textsuperscript{T188E} reduced cell proliferation and cells underwent premature senescence by passage 4, associated with increased SAβG expression (Supplemental Figure 2E), indicating that dynamic changes in TRF2 phosphorylation are particularly required for function in VSMCs. None of the TRF2 truncations/mutations significantly increased apoptosis (Supplemental Figure 2D).

**TRF2 promotes DNA repair, whereas TRF2188A delays repair**

We next examined DNA repair after t-BHP, an oxidant that promotes premature senescence in VSMCs in part through induction of DSBs and telomere damage\textsuperscript{7}. As TRF2, TRF2\textsuperscript{AB}, TRF2\textsuperscript{AM} and TRF2\textsuperscript{ABAM} showed similar effects on cell proliferation and senescence, subsequent studies focused on TRF2 and TRF2\textsuperscript{T188A/E}. TRF2 was not detectable in control VSMCs, but t-BHP induced transient TRF2 phosphorylation in cells expressing ectopic TRF2, but not TRF2\textsuperscript{T188A} or TRF2\textsuperscript{T188E}, again consistent with the known role of this residue in TRF2 phosphorylation (Figure 3A). t-BHP increased γ-H2AX expression in control, TRF2\textsuperscript{T188A} and TRF2\textsuperscript{T188E} VSMCs, but γ-H2AX expression was markedly reduced in TRF2 VSMCs. γ-H2AX returned to baseline by 16h in control and TRF2\textsuperscript{T188E} cells, but TRF2\textsuperscript{T188A} VSMCs maintained γ-H2AX expression to 16h, indicating long-term unrepaired damage (Figure 3A). Thus, TRF2 inhibits DNA damage (as seen by the lack of γ-H2AX expression at baseline and at any stage after DNA damage) and TRF2\textsuperscript{T188A} increases damage / inhibits repair (as seen by persistent γ-H2AX at 16h). TRF2 overexpression also reduced both γ-H2AX and P-ATM foci 1h after t-BHP (Figure 3B-D) and γ-H2AX foci in recovery (Figure 3E); in contrast, TRF2\textsuperscript{T188A} increased γ-H2AX foci 1h after t-BHP (Figure 3C), which persisted to >6h in recovery (Figure 3E). TRF2\textsuperscript{T188E} increased early γ-H2AX induction after t-BHP, but DNA damage normalized by 6h (Figure 3E).

DSBs can be quantified using an alkaline comet assay, with the kinetics of comet tail
shortening reflecting efficiency of DSB repair. t-BHP-induced DSBs were repaired by 6h in control VSMCs, but TRF2 accelerated repair with recovery by 2h; TRF-2^{T188E} slowed initial DNA repair but was complete by 6h, whilst TRF2^{T188A} VSMCs showed markedly delayed repair to >6h (Figure 3F, Supplemental Figure 3). Thus, TRF2 overexpression inhibits replicative arrest in VSMCs, accelerates DSB repair after redox stress, and reduces DNA damage.

TRF2^{T188A/E} induce premature replicative arrest, delay DNA repair and increase DNA damage; these effects are most marked with TRF2^{T188A}, and indicate that TRF2 and TRF2^{T188A} can be used to examine the effects of prevention or induction of VSMC senescence respectively.

**Generation and characterisation of VSMC-specific TRF2 or TRF2^{188A} transgenic mice**

To determine the role of VSMC senescence in atherogenesis and plaque stability, we generated transgenic mice that expressed TRF2 or TRF2^{T188A} from the minimal SM22a promoter (Supplemental Figure 4). The minimal SM22 promoter is expressed only in arterial VSMCs in adult mice, and not in visceral SMCs, cardiomyocytes or skeletal myocytes. 3-bp mutations were also introduced into the promoter CArG (CC(A/T)6GG-) boxes to prevent promoter down-regulation when VSMCs undergo phenotypic modulation. Recent studies have shown that bone marrow-derived myeloid cells can express ‘VSMC’ markers in culture and atherosclerosis, and vessel wall-derived VSMCs can also express ‘macrophage’ markers. However, we have shown that this promoter is expressed in <1% of myeloid cells that express ‘VSMC’ markers in atherosclerotic plaques in mice, indicating that expression of transgenes from this promoter are specific for vessel-wall derived VSMCs.

Two founders from each line were studied, which each behaved similarly. SM22α-TRF2 and SM22α-TRF2^{T188A} mice were born with the expected frequencies, the transgene was transmitted between generations, and expressed only in arteries (Figure 4A). Although primary
mouse VSMCs proliferate slower than mouse VSMC lines, VSMCs derived from SM22α-TRF2 mice showed the same features of increased proliferation (Figure 4B), reduced non-dividing % (Figure 4C), and accelerated DSB repair (Figure 4D), particularly seen with normalization of Comet lengths by 2h (Figure 4D). Similarly, SM22α-TRF2T188A VSMCs showed decreased proliferation, increased non-dividing %, and defective initial DSB repair, indicating that transgene expression in these mice recapitulates expression of TRF2 and TRF2T188A in ApoE−/− VSMCs (Supplemental Figure 2). Importantly, apoptosis was similar in all cell types (Figure 4C), such that functional consequences of transgene expression in vivo were not due to increased/decrease VSMC apoptosis. SM22α-TRF2 and SM22α-TRF2T188A mice showed no gross vascular or other abnormality, and γ-H2AX expression was similar in aortas of all mouse groups, indicating that this overall DNA damage was not affected; in contrast, the senescence marker p16 was markedly increased in SM22α-TRF2T188A mice (Figure 4E), confirming that TRF2T188A can induce markers of senescence in VSMCs in normal arteries in the absence of DNA damage.

The effect of TRF2 and TRF2T188A on telomeres, telomere length and chromosome structure in VSMCs from transgenic mice was examined using qPCR for telomere length and telomere fluorescence in situ hybridisation. Although TRF2 reduced telomere length, this was not associated with telomere deletions, and there was a trend towards reduced telomere fusions, consistent with previous studies indicating a protective role for TRF2 overexpression8,32. In contrast, TRF2T188A increased the incidence of telomere deletions, consistent with loss of telomere function (Supplemental Figure 5).

Effects of TRF2 or TRF2T188A on atherosclerosis

To examine the effect of VSMC senescence on atherosclerosis, SM22α-TRF2 and SM22α-
TRF2<sub>T188A</sub> mice were backcrossed with C57Bl6/J ApoE<sup>−/−</sup> mice, and male and female ApoE<sup>−/−</sup>, SM22α-TRF2/ApoE<sup>−/−</sup>, or SM22α-TRF2<sub>T188A</sub>/ApoE<sup>−/−</sup> littermates were fat fed from 6w-22w. Body weight, blood pressure, serum lipids, total leukocyte counts, and differential leukocyte % showed no differences between groups (Supplemental Figure 6). However, both total atherosclerotic plaque and necrotic core areas were significantly increased in aortic root lesions of SM22α-TRF2<sub>T188A</sub> mice (Figures 5 and 6, Table 1). Plaque area was also increased in brachiocephalic arteries of SM22α-TRF2<sub>T188A</sub> mice (0.60±0.08 mm<sup>2</sup>) and reduced in SM22α-TRF2/ApoE<sup>−/−</sup> mice (0.10±0.09 mm<sup>2</sup>) compared to controls (0.33±0.1 mm<sup>2</sup>, Mean±SEM p=0.047) (Supplemental Figure 7). Similar to human plaques, TRF2 was reduced in VSMCs in plaques of ApoE<sup>−/−</sup> mice vs. the undiseased media (Supplemental Figure 8).

In humans, rupture-prone plaques exhibit a thin fibrous cap and a relatively large necrotic core, we therefore undertook detailed analysis of plaque architecture in these mice. SM22α-TRF2 mice had significantly increased fibrous cap/plaque and cap/core ratios, and reduced core/plaque ratios compared with control ApoE<sup>−/−</sup> mice (Figure 6A-E, Table 1), indicating a relative increase in fibrous cap and decrease in core areas. Overall % SMA-positive, mac3-positive, and apoptosis frequencies were similar between groups (Table 1), although fibrous caps of SM22α-TRF2 mice showed more extensive SMA expression (Figure 5).

**Increased atherosclerosis in SM22α-TRF2<sub>T188A</sub> mice is not due to a SASP**

Premature senescence may promote atherogenesis by reducing the VSMC content of fibrous caps, a major factor in promoting plaque rupture and subsequent growth in humans<sup>33</sup>. However, senescence can also induce a senescence-associated secretome (SASP), a pattern of pro-inflammatory cytokines released when cells (including VSMCs) undergo replicative or stress-induced premature senescence<sup>34</sup> that might promote atherosclerosis by increasing monocyte
invasion. However, both mac3-positive content of plaques (Table 1) and serum levels of SASP cytokines were similar (MCP-3, MCP-1, Mip-1β, RANTES) or undetectable (Mip-1α, IL-1, IL-6) in all mouse groups at 22w (Supplemental Figure 9).

To determine whether mouse VSMCs expressing TRF2 or TRF2T188A showed differences in SASP cytokine secretion, we cultured VSMCs expressing these proteins and determined cytokine secretion after 2 days. Conditioned media of ApoE−/− VSMCs expressing TRF2 or TRF2T188A showed similar cytokine secretion to control cells (Supplemental Figure 10). The increased atherosclerosis in SM22α-TRF2T188A mice with a similar SASP profile in vivo and in culture thus suggests that VSMCs exert a direct protective effect on the vessel wall.

Discussion

VSMC proliferation is a fundamental component of both historical and current models of atherogenesis. For example, the ‘response to injury’ hypothesis viewed aberrant VSMC proliferation as a major primary event in plaque development. In contrast, more recent models suggest that VSMC proliferation is predominantly reparative, for example as a response to inflammation35,36. While increased VSMC proliferation is seen in atherogenesis, VSMCs in advanced human plaques are characterised by multiple markers of senescence, including telomere loss, expression of the cyclin-dependent kinase inhibitors p21 and p16, and expression of SAβC7. Although this indicates that VSMC senescence occurs in advanced plaques, it is not known whether senescence contributes to plaque development or the morphology of advanced lesions, or its underlying mechanisms.

We demonstrate that human atherosclerotic plaque VSMCs show reduced TRF2 expression in vitro and markedly reduced binding of TRF2 to telomeres, associated with
increased markers of DNA damage. Other shelterin proteins showed no consistently altered expression, suggesting that TRF2 is a critical regulator of VSMC senescence in atherosclerosis, and that plaque VSMCs are very sensitive to telomere damage. TRF2 was also reduced and associated markers of DNA damage increased in α-SMA-positive cells in human plaques in vivo. TRF2 protein was regulated by proteasomal degradation that depended upon p53. TRF2 reduced DNA damage, accelerated DSB repair, and inhibited replicative arrest in VSMCs, indicating that VSMC senescence may be due to both replicative exhaustion and stress-induced premature senescence (SIPS). Using novel gain and loss of function VSMC-specific expression of TRF2 or TRF2T188A, we show that VSMC senescence promotes atherosclerosis and features of plaque vulnerability, including decreased relative fibrous cap and increased necrotic core areas, without a significant SASP.

We show that TRF2 is unique of the shelterin proteins in being downregulated in plaque VSMCs. TRF2 protein is regulated by ubiquitin-mediated degradation, in part by the p53-induced E3 ubiquitin ligase siah1.25 VSMCs in plaques are characterised by multiple forms of DNA damage, including double and single strand breaks, oxidative lesions and mtDNA damage, and DNA damage markers and activation of multiple DNA repair pathways are maintained in cultured plaque VSMCs.7,20,38 Plaque VSMCs in culture and cells expressing α-SMA in human plaques showed down-regulated TRF2 associated with markers of both DNA damage (γ-H2AX) and DNA repair (p-ATM). In culture, plaque VSMCs showed reduced TRF2 and increased siah-1a, associated with increased p16. Although overall TRF2 expression was reduced after proteasomal inhibition, this was also associated with marked upregulation of p53, and the effect of proteasomal inhibition on TRF2 expression was dependent upon p53. This suggests a specific mechanism for TRF2 down-regulation in atherosclerosis by which DNA
damage-induced activation of p53 and subsequently siah-1a results in TRF2 degradation in plaque VSMCs.

Although studies on human plaque VSMCs do not indicate if reduced TRF2 is a cause or consequence of senescence, we show that overexpression of TRF2 from a retrovirus promoter or from transgenic mice using the SM22α minimal promoter blocks cell senescence in culture. The latter is not a strong promoter, indicating that endogenous levels of TRF2 may be sufficient to prevent senescence. This hypothesis is strengthened by our studies using TRF2^{188A} which induces senescence in VSMCs that express endogenous levels of wild type TRF2. The use of both gain and loss of function studies in vitro that show opposite effects strongly imply that TRF2 is a critical regulator of VSMC senescence, and that the observed loss of TRF2 is a major factor in VSMC senescence in atherosclerosis.

To determine the effects of manipulating TRF2 in VSMCs in vivo on atherogenesis and plaque morphology directly we generated two novel transgenic mouse models that express VSMC-specific TRF2 or TRF2^{188A} as gain or loss of function respectively. TRF2 and TRF2^{188A} were expressed at similar levels, a critical control to be able to compare their effects. TRF2 promoted proliferation, inhibited senescence, and accelerated DNA repair in VSMCs derived from SM22α-TRF2 mice, whereas TRF2^{188A} did the opposite. Furthermore, in normal aortas TRF2^{188A} increased p16 expression with no change in γ-H2AX, indicating that TRF2^{188A} can promote VSMC senescence in the absence of widespread DNA damage. TRF2^{188A} promoted plaque development associated with increased plaque and necrotic core areas, whereas TRF2 expression increased the fibrous cap area relative to both plaque and core areas, and reduced relative core size, with no change in apoptosis in either mouse group.

Recent genetic lineage tracing studies indicate that conventional markers of macrophages
can also be expressed by cells derived from the vessel wall\textsuperscript{31}, and markers of VSMCs can also be expressed by myeloid cells that differentiate in culture or migrate into the plaque\textsuperscript{30}. This potentially might mean that myeloid cells activate ‘VSMC’ promoters when they migrate into atherosclerotic plaques, confounding our interpretation that the effects we observe are due to the effects of loss of TRF2 on vessel-wall derived VSMCs. However, we and others have shown that bone marrow-derived cells expressing SMC markers are either infrequent\textsuperscript{30} or non-existent\textsuperscript{39} in plaques in ApoE mice, and could not be demonstrated in fibrous caps of advanced plaques at 22w\textsuperscript{30}. We have also transplanted bone marrow-derived cells into ApoE\textsuperscript{-/-} mice with transgenes expressed from the minimal SM22\alpha promoter, which showed that <1\% of cells in advanced plaques expressed the transgene, and again there was no expression in fibrous caps\textsuperscript{30}. Thus, the opposite effects of TRF2 and TRF2\textsuperscript{T188A} expression on plaque morphology argue strongly that changes in TRF2 function in these transgenic mice are mediated through senescence of vessel wall-derived VSMCs.

Premature senescence may promote atherogenesis by reducing the protective fibrous cap (as demonstrated here), or via a senescence-associated secretome (SASP), that might promote atherosclerosis by increasing monocyte invasion, or both. However, it has been difficult to demonstrate a functionally important SASP in mouse cells, and this often requires culture in physiological oxygen concentrations and extreme measures such irradiation\textsuperscript{40}. In addition, senescence induced by p16 (which is increased in SM22\alpha-TRF2\textsuperscript{T188A} VSMCs and human plaque VSMCs) is often not associated with a SASP\textsuperscript{41}. We find no differences in ‘classical’ SASP cytokines in serum of transgenic mice compared with controls, or in cultured VSMCs expressing TRF2 of TRF2\textsuperscript{T188A}. While it is possible that a local SASP may influence cellular composition and morphology of plaques \textit{in vivo}, other anti-atherogenic effects of VSMCs should also be
considered, including acting as a barrier to invasion of monocytes, or prevention of plaque rupture that leads to plaque growth. In particular, our findings illustrate the profound protective properties of normal VSMC function on the vessel wall, and a novel role of VSMC senescence in necrotic core formation. Our results may also explain why accelerated atherosclerosis and myocardial infarction are features of progeroid syndromes that show loss of VSMCs, DNA damage, and premature cellular senescence, and why genes that regulate senescence such as the INK4/ARF locus might be associated with heart attacks in GWAS studies.

Our data support a model of VSMC senescence in atherosclerosis involving telomere dysfunction, DNA damage, TRF2 and p53 (Figure 7). In atherogenesis, mitogens from platelets, invading myeloid cells, and the local vessel wall promote VSMC replication. Inflammation and reactive oxygen species (for example) promote DNA damage, including in telomeres of replicating cells. DNA damage also activates p53, resulting in siah1-mediated degradation of TRF2, which causes exposure of telomere ends. The combination of replication, loss of TRF2 protection, and direct DNA damage induce telomere dysfunction. Telomere dysfunction results in activation of a DNA damage response that further activates p53, and induction of p16. The p53 target p21 and p16 induce senescence, in part mediated through hypophosphorylation of pRB.

In summary, we show that TRF2 is a major regulator of VSMC senescence, and is downregulated in VSMCs from human atherosclerotic plaques. VSMC senescence promotes atherogenesis and necrotic core formation, while reducing cap formation. Protection against VSMC senescence may represent a novel mechanism to reduce both plaque formation and features of vulnerable plaques.
Acknowledgments: We thank members of the Molecular Cytogenetics Unit, Welcome Trust Sanger Institute, UK) for technical assistance with TEL-FISH.

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Conflict of Interest Disclosures: None.

References:


Table 1. Morphometric and immunohistochemical quantification of aortic root atherosclerotic plaques in Control ApoE\textsuperscript{-/-}, SM22\textalpha-TRF2/ApoE\textsuperscript{-/-}, and SM22\textalpha-TRF2\textsuperscript{T188A}/ApoE\textsuperscript{-/-} mice at 22w after fat feeding from 6-22w. Data are means±SD. *p<0.05 vs. control ApoE\textsuperscript{-/-} mice.

<table>
<thead>
<tr>
<th></th>
<th>Control (ApoE\textsuperscript{-/-}) (n=18)</th>
<th>SM22\textalpha-TRF2/ApoE\textsuperscript{-/-} (n=11)</th>
<th>SM22\textalpha-TRF2\textsuperscript{T188A}/ApoE\textsuperscript{-/-} (n=11)</th>
<th>p Value (ANOVA)</th>
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<tr>
<td>Plaque area ((\mu\text{m}^2))</td>
<td>386764 ± 159285</td>
<td>381339 ± 141513</td>
<td>531202 ± 196287*</td>
<td><strong>0.044</strong></td>
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<tr>
<td>Necrotic core area ((\mu\text{m}^2))</td>
<td>266685 ± 140666</td>
<td>217660 ± 110186</td>
<td>377240 ± 177253*</td>
<td><strong>0.046</strong></td>
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<td>Cap area ((\mu\text{m}^2))</td>
<td>123466 ± 43778</td>
<td>163679 ± 60618</td>
<td>159371 ± 43281</td>
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<tr>
<td>Cap/Plaque ratio</td>
<td>0.33 ± 0.099</td>
<td>0.45 ± 0.103*</td>
<td>0.33 ± 0.107</td>
<td><strong>0.013</strong></td>
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<tr>
<td>Cap/Core ratio</td>
<td>0.50 ± 0.211</td>
<td>0.87 ± 0.302*</td>
<td>0.53 ± 0.276</td>
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<tr>
<td>Core/Plaque ratio</td>
<td>0.67 ± 0.099</td>
<td>0.55 ± 0.107*</td>
<td>0.67 ± 0.107</td>
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<td>SMA (% area)</td>
<td>14.17 ± 6.60</td>
<td>14.26 ± 6.27</td>
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<td>Mac3 (% area)</td>
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<td>TUNEL (% cells)</td>
<td>0.061 ± 0.11</td>
<td>0.209 ± 0.26</td>
<td>0.129 ± 0.195</td>
<td>0.16</td>
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</table>

Figure Legends:

Figure 1. Human Plaque VSMCs show reduced TRF2 expression. (A) Western blot of shelterin proteins from normal human aortic (n=3) or plaque VSMCs (n=3). (B) Immunoprecipitation of TRF2 with telomere sequences in aorta (Ao) or plaque (Pl) VSMCs (n=3). (C) qPCR for TRF2 in normal aortic (n=3) or plaque VSMCs. Data are Means±SEM, n=3. (D) Western blot of plaque and normal VSMCS for siah1a and p16 (n=3). (E) Western blot of control or p53\textsuperscript{-/-} null mouse VSMCs after treatment with 10\(\mu\text{M}\) MG132 for 5 hours (n=2).

Figure 2. Human Plaque VSMCs show reduced TRF2 and increased DNA damage. (A) Nuclear foci of TRF2, \(\gamma\)-H2AX or P-ATM in aortic or plaque VSMCs. (B) Quantification of single or
combined foci (n=20). (C–D) Immunohistochemistry for α-sm-actin (SMA–blue) and either TRF2, γ-H2AX or P-ATM (brown) in normal aorta or plaque. Double-labeled panels are from areas indicated in H+E in aortic media and fibrous cap. Scale bars=50μm. Insets are high power views of outlined areas. Data are means±SEM, n=5.

**Figure 3.** TRF2 inhibits and TRF2T188A increases DNA damage. (A) Western blot for TRF2 and γ-H2AX for ApoE−/− VSMCs expressing vector alone (Cont), TRF2, TRF2T188A, or TRF2T188E, after treatment with 100μM t-BHP for 1 hour and recovery (Rec.) for either 2 (upper panels) or 16 hours (lower panels). (B) Immunocytochemistry of VSMCs from (A) 1h after t-BHP showing nuclear foci of TRF2/γ-H2AX, or TRF2/P-ATM. (C,D) Quantification of nuclear foci in (B) 1h after t-BHP. (E,F) γ-H2AX foci (E) or comet tail moment (F) in VSMCs 1h after t-BHP (time 0) and 1, 2 or 6 hours recovery. Data are means±SEM, n=3. *p<0.05, **p<0.01 vs. controls.

**Figure 4.** VSMCs from TRF2 and TRF2T188A transgenic mice show increased or decreased proliferation and DNA repair respectively. (A) qPCR for SM22α-TRF2/TRF2T188A transgene mRNA in two SM22α-TRF2 or SM22α-TRF2T188A founders (aorta), or gut. (B–D) Basal cell proliferation (B) or % cell populations (C), or Comet tail moment after 1h of 100μM t-BHP and 5h recovery (D) in VSMCs from control or transgenic mice. (E) Western blot for TRF2, γ-H2AX and p16 in aortas of control ApoE−/− (Control), SM22α-TRF2/ApoE−/− (TRF2) or SM22α-TRF2T188A/ApoE−/− (TRF2T188A) at 22w. Data are means±SEM n=3. *p<0.05 vs. controls.

**Figure 5.** Effects of TRF2 on plaque development and morphology. Histochemistry and immunohistochemistry of control, SM22α-TRF2 or SM22α-TRF2T188A mouse aortic roots for
Masson’s Trichrome, α-SMA and Mac3 at 22w after fat feeding from 6-22w. High power views represent areas outlined on corresponding low power views above. Scale bar=500 μm (low power), and 150μm (high power).

**Figure 6.** Morphometry of plaques in experimental mice. (A-E) Quantification of aortic root plaque, core and cap areas and ratios in Control ApoE<sup>−/−</sup>, SM22α-TRF2/ApoE<sup>−/−</sup>, and SM22α-TRF2<sup>T188A</sup>/ApoE<sup>−/−</sup> mice at 22w after fat feeding from 6-22w. Data are means±SEM. n=11-18.

**Figure 7.** Model of VSMC senescence in atherosclerosis. A full description is given in the text.
Figure 1
Figure 2

Aorta Plaque

TRF2 γ-H2AX

TRF2 P-ATM

γ-H2AX P-ATM

B

Foci number / nucleus

TRF2 P-ATM γ-H2AX TRF2/ P-ATM TRF2/ γ-H2AX P-ATM/ γ-H2AX

p=0.0001 p=0.032 p=0.001 p=0.012 p=0.007 p=0.031

C

H&E

Aorta Plaque

SMA TRF2

SMA P-ATM

SMA γ-H2AX

D

% Positive VSMCs

TRF2 P-ATM γ-H2AX

p=0.021 p=0.008 p=0.003
Figure 4
SUPPLEMENTAL MATERIAL

Supplemental Methods

Protein Extraction, Western blotting

Protein extraction from cultured cells and Western blotting were performed as previously described\(^1\). Tissue was dissected into 25mg pieces with 3 additional freeze-thaw cycles in lysis buffer in liquid nitrogen prior to incubation on ice for 12 minutes. Proteins were then extracted as per cultured cells. Primary antibodies were used as follows: mouse-TRF1 (#3529;1:200; Cell Signalling Technology), rabbit-POT1 (ab21382), mouse-TIN2 (ab13791), mouse-RAP1 (ab14404) (all 1:5000; Abcam), mouse-TPP1 (B01; Abnova), rabbit-\(\gamma\)-H2AX-ser139 (#2577), rabbit-TRF2 and mouse-TRF2 (D1Y5D)(#2645 and #13136), rabbit-\(\gamma\)-Tubulin (#2144), mouse-p53 (#2524; 1C12) (all 1:1000; Cell Signalling Technology) and rabbit-p16-INK4A (10883-1-AP, 1:5000; Proteintech). Goat-siah1 (1:1000; ab2237; Abcam), HRP-conjugated secondary antibodies were sheep anti-mouse (1:2000; GE Healthcare) and donkey anti-rabbit (1:5000; GE Healthcare) in 1% BSA/TBST.

ChIP

TRF2 association with telomeres was examined by ChIP. In brief, 1-5x10^6 cells were fixed in 1% formaldehyde for 10 min at room temperature. The reaction was quenched with 125mM glycine. Cross-linked cells were scraped, centrifuged and incubated in lysis buffer (1% SDS, 10mM EDTA pH 8.0, 50mM Tris pH 8.1 and protease inhibitors (Roche)) on ice for 10 min. The lysate was then sonicated (Diagenode Bioruptor® Standard) to shear chromatin and 50µg of chromatin used for immunoprecipitation. Samples were incubated overnight at 4°C with either anti-TRF2 (#2645, 1:100, Cell Signalling Technologies) or equivalent isotype control (Santa Cruz Biotechnology). Following immunoprecipitation, 40µL of protein A dynabeads (Life Technologies) were added to samples and rotated for 1h. Beads/chromatin complexes were washed with low salt buffer (2X), high salt buffer (2X), LiCl buffer (1X) and TE buffer (2X) and immunocomplexes eluted at 65°C with 10% SDS and 1 mM NaHCO3 buffer. After addition of 40mM NaCl and formaldehyde, cross-linking was reversed at 65°C
overnight. DNA was recovered using a mini-elute DNA purification kit (Qiagen). Telomere specific real-time PCR was performed as previously described².

**Immunocytochemistry**

Immunocytochemistry for nuclear foci was performed as described previously¹. Primary antibodies were rabbit anti-γ-H2AX<sup>ser139</sup>, P-ATM, and TRF2 (#9718, #4526 and #2645, all 1:100, Cell Signaling Technologies). Secondary antibodies were Alexa Fluro488 goat anti-mouse IgG and Alexa Fluro694 Goat anti-rabbit IgG (1:500; Invitrogen). γ-H2AX<sup>ser139</sup> foci intensity was assayed using ImageJ (NIH)

**Immunohistochemistry**

Immunohistochemistry of human and mouse arteries was performed as described previously³.

**Cell Culture**

Mouse cell lines were cultured in DMEM supplemented with 10U/ml Penicillin (Sigma), 10µg/ml Streptomycin (Sigma), 5µg/ml L-Glutamine (Sigma), 10% fetal calf serum (Sigma) ± hygromycin. Cells were treated with t-butyl hydroperoxide (t-BHP, 100µM) for 1 hour at 80% confluency prior to analysis at selected time points.

**Measurement of Telomere Length (Q-PCR)**

The length of mouse telomeres was measured by Q-PCR on a Rotor-Gene (Corbett Research) PCR platform. The single-copy control gene (36B4) was measured using forward and reverse primers 5’ ACT GGT CTA GGA CCC GAG AAG 3’ and 5’ TCA ATG GTG CCT CTG GAG ATT 3’, respectively. The primers used to measure average telomere length were 5’ CGG TTT GTT TGG GTT TGG GTT TGG GTT TGG GTT 3’ and 5’ GGC TTG CCT TAC CCT TAC CCT TAC CCT TAC TAC 3’. The master mix reaction for
36B4 contained 10µl SYBR Green JumpStart Taq ReadyMix (Sigma), 20ng purified DNA in 5µl, 300nM forward primer, 500nM reverse primer and PCR-grade H₂O (Invitrogen) in a 20µl final volume. The reaction mix for telomere length was similar to 36B4, except using 300nM of both forward and reverse primers. qPCR reaction conditions were 95°C for 15 minutes, followed by 35 cycles of elongation steps starting at 95°C for 15 seconds, 54°C for 60 seconds and 72°C for 30 seconds, and finished at 72°C for 10 minutes. ApoE⁻/⁻ mouse DNA, serially diluted from 60ng to 1.875ng, was used to generate a reaction standard curve. Each DNA sample was run in triplicate and two negative controls (no template DNA; no primers) were run in duplicate in parallel. The input amounts of telomere and 36B4 DNA were calculated based on the standard curve generated after averaging the threshold cycle number (Ct) values for each sample. The average telomere length ratio (ATLR) was then calculated by dividing the amount of telomeric DNA to the amount of 36B4 product.

**Chromosome harvest and and telomere FISH**

To prepare metaphase chromosomes, mouse VSMCs were treated with 0.03µg/mL of Colcemid (Gibco, Life Technologies) for 16h, followed a buffered hypotonic solution (0.4% KCl in 10 mM HEPES, pH7.4) for 8 min at 37°C. The cells were centrifuged and resuspended in 4:1(v/v) methanol:glacial acetic acid and kept at -20°C until use. Metaphase preparations were placed onto pre-cleaned microscope slides. Telomere FISH with a Cy3-TelC PNA probe was performed following the manufacturer’s protocol (PNA Bio inc.). Digital images were captured on a Leica DM5000 B microscope equipped with an ORCA-03G CCD camera (Hamamatsu) controlled by SmartCapture software (Digital Scientific UK).

**Analysis of Atherosclerosis**

Atherosclerosis extent and composition was analyzed as described previously² Briefly, sections from the aortic root and brachiocephalic arteries were stained with H&E, α-SMA,
Mac3, and TUNEL. Plaque extent and composition was measured using CellD analysis software. The fibrous cap and necrotic core was defined using Masson’s Trichrome.
Supplemental Figure 1

(A) Western blot for p53 and SMA of whole lysates from normal human aorta (Ao) or carotid plaque (Pl). (B) Quantification of Western blots from aortic or plaque lysates. n=4.

(C) Immunohistochemistry for p53 or SMA in normal human aorta or carotid plaque. Scale bars = 150 µm. Insets show high power views of areas outlined.

(D) Quantification of % of VSMCs that express p53 in aortic or plaque sections. n=4.

Supplemental Figures
Supplemental Figure 2

(A) Schematic of TRF2 constructs. Basic, dimerization and myb domains, and nuclear localisation signal (NLS) are marked. (B) Western blot for TRF2 in ApoE−/− VSMC lines, including control (uninfected) and vector-infected VSMCs. (C) Time-lapse videomicroscopic analysis of % cell proliferation over 24 hours in VSMC lines. (D) % cells undergoing cell division, apoptosis or senescence in VSMC lines. Data are means±SD. (E) SAβG expression in VSMC lines at passage 14 (vector) or passage 4 (TRF2, T188A/E). (Data are means, n=3). *p<0.05, **p<0.01 vs. vector lines.
Supplemental Figure 3
Comet assay of VSMCs expressing vector alone, TRF2, TRF2\textsuperscript{T188A} or TRF2\textsuperscript{T188E} at baseline (Time 0), 1 hr of BHP treatment (BHP) and either 2 or 6 hours after removal of BHP (recovery). Scale bar=25\textmu m.
Supplemental Figure 4
Map of transgenic targeting constructs
Supplemental Figure 5

(A) Telomere length in mouse ApoE−/− VSMCs (Control), SM22α-TNF2/ApoE−/− VSMCs (TRF2), or SM22α-TNF2T188A/ApoE−/− VSMCs (TRF2T188A). (B) Telomere FISH of VSMCs derived from ApoE− (Control), SM22α-TNF2/ApoE− (TRF2) and SM22α-TNF2T188A/ApoE− (TRF2T188A) mice. The Telomeric Cy3-labelled synthetic PNA-(CCCTAA)3 probe is shown in green and chromosomal counterstain (DAPI) in red. High magnification examples of end association and telomere deletions are shown on the right. White arrows indicate terminal deletion, blue arrows indicate end association. (C) Quantification of % chromosomes showing end association or terminal deletion. Scale bar=10μM. Data are means±SD. (n=400-1000 chromosomes/genotype).
Supplemental Figure 6

(A-F) Body weight (A), blood pressure (B), serum high density lipoprotein (HDL), low density lipoprotein (LDL), Triglycerides (TRI) or total cholesterol (TOT) (C), and leukocyte counts (D-F), in ApoE−/− (Control), SM22α-TRF2/ApoE−/− (TRF2) and SM22α-TRF2T188A/ApoE−/− (TRF2T188A) mice after fat feeding from 6-22w. Data are means±SEM, n=8-12.
Supplemental Figure 7
Brachiocephalic plaques from Control ApoE<sup>−/−</sup>, SM22α-TRF2/ApoE<sup>−/−</sup> and SM22α−TRF2<sup>T188A</sup>/ApoE<sup>−/−</sup> mice after fat feeding from 6-22w.

Supplemental Figure 8
(A) Immunohistochemistry for TRF2 (brown) and SMA (blue) in ApoE<sup>−/−</sup> mouse plaques or undiseased media. High power view of fibrous cap or medial VSMCs are shown on the right. Scale bar=150 µm. (B) Quantification of %TRF-2-positive VSMCs in ApoE<sup>−/−</sup> mouse plaques or undiseased media. Data are means±SD (n=8).
Supplemental Figure 9
Serum cytokines from experimental mice at 22w for MCP-3, MCP-1, MIP1β and RANTES after fat feeding from 6-22w. Data are means±SEM, n=8-15.
Supplemental Figure 10
Cytokines secreted from Control ApoE<sup>−/−</sup>, TRF2/ApoE<sup>−/−</sup> and TRF2<sup>188A</sup>/ApoE<sup>−/−</sup> VSMCs. Data are means±SEM. n=8-10.
Supplemental References


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