Origin of neointimal endothelium and α-actin–positive smooth muscle cells in transplant arteriosclerosis

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The development of transplant arteriosclerosis (TA) is today’s most important problem in clinical organ transplantation. Histologically, TA is characterized by perivascular inflammation and progressive intimal thickening. Current thought on this process of vascular remodeling assumes that neointimal vascular smooth muscle (VSM) cells and endothelium in TA are graft-derived, holding that medial VSM cells proliferate and migrate into the subendothelial space in response to signals from inflammatory cells and damaged graft endothelium. Using MHC class I haplotype-specific immunohistochemical staining and single-cell PCR analyses, we show that the neointimal α-actin–positive VSM cells in rat aortic or cardiac allografts are of recipient and not of donor origin. In aortic but not in cardiac allografts, recipient-derived endothelial cells (ECs) replaced donor endothelium. Cyclosporine treatment prevents neointima formation and preserves the vascular media in aortic allografts. Recipient-derived ECs do not replace graft endothelium after cyclosporine treatment. We propose that, although it progresses beyond the needs of functional repair, TA reflects the activity of a normal healing process that restores vascular wall function following allograft-induced immunological injury.


Introduction

The development of chronic transplant dysfunction (CTD) has become a limiting factor for long-term graft survival and is today’s most important problem in clinical organ transplantation after the first perioperative year. To date, CTD cannot be prevented by current immunosuppressive protocols (1, 2). Depending on the type of organ transplanted (liver, kidney, heart, lung), the incidence of CTD 3 years after engraftment varies from 4 to >50% (3–5). Irrespective of the organ grafted, graft vessels eventually develop so-called transplant arteriosclerosis (TA), which is, however, most prominent in cardiac allografts (6, 7). This vascular remodeling process consists of progressive concentric intimal thickening (intimal hyperplasia or neointima formation) coinciding with ongoing perivascular inflammation. TA has generally been accepted as the main cause of progressive deterioration in graft function. Pathogenesis of TA seems to be multifactorial, but precise mechanisms involved in the development of this remodeling process still remain obscure. Risk factors appear to include cold ischemia time and reperfusion injury, MHC disparity between donor and recipient, number of rejection episodes, and infection (especially by cytomegalovirus) (2, 6).

Current thought on the process of TA holds that in response to signals from inflammatory cells (perivascularitis) and activated or damaged graft endothelium donor-derived medial vascular smooth muscle (VSM) cells of affected arteries start to proliferate and migrate from the media into the subendothelial space just beneath the endothelial cell (EC) layer (8–10). According to this concept, neointimal VSM cells replaced donor endothelium. Cyclosporine treatment prevents neointima formation and preserves the vascular media in aortic allografts. Recipient-derived ECs do not replace graft endothelium after cyclosporine treatment. We propose that, although it progresses beyond the needs of functional repair, TA reflects the activity of a normal healing process that restores vascular wall function following allograft-induced immunological injury.
 novo formation of a new vessel wall (25, 26). In time, a neointima and neomedia were found to develop inside the lumen of the graft. Since the graft itself only consists of synthetic material, the cellular components forming the neomedia and the neointima, in this model, must be recipient-derived. Based on these observations we hypothesized that also after regular allogeneic organ transplantation a similar process of de novo vessel wall formation (vascular remodeling) will occur in the arteries of the graft, constituting cells originating from the recipient (27).

To test this hypothesis, we used two experimental transplant models. First, we used the allogeneic aorta transplant model in rats, which is a well-described and reproducible model to study TA (28). Second, we used a cardiac transplant model in which MHC-incompatible cardiac allografts are transplanted following intrathyric immune modulation and short-term immunosuppression (29). Cardiac allograft survival is significantly prolonged, but development of TA in the coronary artery (CA) tree cannot be prevented, resembling clinical outcome after cardiac transplantation (30). Using both transplant models we analyzed the origin of neointimal ECs and VSM cells, using MHC class I haplotype-specific immunohistochemistry and two independent PCR-based analyses (31, 32). Recipient- and donor-specific PCR analyses were performed on 70-µm-thick tissue samples, microdissected neointimal cell clusters, and single-VSM-cell nuclei (aorta allografts), and on microdissected whole CAs and single-VSM-cell nuclei (cardiac allografts). Similar analyses were performed on aortic allografts that were γ-irradiated prior to transplantation, and grafts from recipients that were treated with cyclosporine (CsA).

Our results indicate that after allogeneic aorta transplantation neointimal ECs and VSM cells are of recipient and not of donor origin. CsA treatment prevented neointima formation and preserved the vascular media. After CsA treatment the ECs were still of donor origin and had not been replaced by recipient-derived cells. Although the neointimal VSM cells in cardiac allografts are of recipient origin, recipient-derived cells do not replace donor ECs.

Thus, in contrast to current thought, VSM cells constituting the hyperplastic neointima in aortic as well as cardiac allografts are of recipient and not of donor origin. EC replacement is not a general phenomenon after allogeneic organ transplantation and might be related to the severity of the initial damage of graft endothelium. We propose that TA is part of a normal healing process trying to restore function of the damaged vessel wall. Recipient-derived ECs and VSM cells are involved in this repair process, which, however, does not seem to be controlled adequately and eventually may lead to total occlusion of the vessels.

Methods

Animals. Specific pathogen–free Lewis (Lew, RT-1<sup>l</sup>), Brown Norway (BN, RT-1<sup>n</sup>), Pibold Viral Glasco (PVG) (RT-1<sup>g</sup>), and Albino Oxford (AO) (RT-1<sup>u</sup>) rats were obtained from Harlan (Zeist, The Netherlands). The rats were 7–10 weeks of age and were used for aorta and cardiac transplantation. All transplantations were performed from female donor to male recipient rats. Animals were maintained under clean conventional conditions and fed standard rat chow and acidified water ad libitum. All animals received care in compliance with NIH principles of laboratory animal care and the Dutch Law on Experimental Animal Care.

Aorta transplantation. Aortic allografts (10–12 mm) were transplanted orthotopically as described previously (28). Total ischemic time was consistently less than 30 minutes. Transplantations were performed in the following MHC-incompatible strain combinations: PVG to AO, BN to Lew, and Lew to BN. Isograft controls consisted of aortic grafts transplanted from AO to AO and BN to BN rats. Grafts were removed at autopsy 1 month (BN to Lew and Lew to BN) or 3 months (PVG to AO and isograft controls) after transplantation and processed for histological evaluation or microdissection.

Cardiac transplantation. Cardiac allografting was performed from female PVG to male AO recipient rats. To prevent acute rejection, recipient rats were intrathyrically immune-modulated as described elsewhere (29). Immediately after cardiac transplantation (day 0), 1 ml rabbit anti-rat lymphocyte serum (Accurate Chemical & Scientific Corp., Westbury, New York, USA) was injected intraperitoneally. CsA (Sandoz Pharma AG, Basel, Switzerland) was injected intramuscularly on days 1, 2, and 3 after transplantation (15 mg/kg). Cardiac allografts thus transplanted show significantly prolonged graft survival but eventually develop TA indicative of CTD (30).

Treatment groups. Aortic allografting was performed from female PVG donor to male AO recipient rats. One group was left untreated (n = 4), whereas in the other group aortic allografts were ex vivo 50 Gy γ-irradiated (137Cs, IBL 673; CIS Bio International Gif-Sur-Yvette, France) prior to transplantation in order to prevent proliferation and migration of donor-derived medial VSM cells (n = 3). Three AO isograft controls were performed (n = 3). To study the effect of CsA treatment on the development of TA, another set of aorta transplantations was performed in the female Lew to male BN combination. Recipient rats received either no treatment (n = 5) or CsA (Sandoz Pharma AG) at 5 mg/kg/d (n = 5) or 10 mg/kg/d (n = 4) for 28 days. These concentrations were shown to reduce intimal hyperplasia and preserve medial nuclear density after allogeneic aorta transplantation in rats (33). CsA was administered as a liquid in olive oil by intramuscular injections. BN to Lew allografts (n = 3) and BN isografts (n = 5) served as control groups.

To analyze TA in cardiac allografts, three female-derived PVG grafts were transplanted into male AO recipient rats followed by intrathyric immune modulation as described above (n = 3). Grafts were harvested 530 days (19 months) after transplantation and processed for histological analysis and microdissection.
CsA blood levels. Twenty-four-hour trough whole blood CsA levels were determined 2 and 4 weeks after aortic transplantation. Blood was analyzed using the Emit 2000 cyclosporine-specific assay (Dade Behring, Marburg, Germany).

Immunohistochemical analyses. At the time of sacrifice, the explanted aortic and cardiac grafts were either fixed in Bouin’s fixative and embedded in paraffin or immediately frozen in liquid nitrogen and stored at -80°C for cryostat sections. For evaluation of vascular changes, paraffin cross sections (7 µm) from the center of the grafts were prepared, and double staining was performed for elastin (Lawson solution; Klinipath, Duiven, The Netherlands) and α-actin to detect neointimal VSM cells in arteriosclerotic lesions. Briefly, α-actin–positive intimal cells were detected by an indirect immunoperoxidase technique using mAb asm-1 directed against VSM cell α-actin isoform (34, 35) (Roche Molecular Biochemicals, Almere, The Netherlands). Incubation with the primary asm-1 antibody was followed by incubation with a second-step horseradish peroxidase–conjugated rabbit anti-mouse antibody (DAKO A/S, Glostrup, Denmark). Subsequently, chromogen 3-amin-9-ethyl carbazole was applied, followed by counterstaining with Mayer’s hematoxylin. Sections were coverslipped with Kaiser’s glycerol gelatin (Merck, Amsterdam, The Netherlands).

To detect MHC class I expression on ECs and graft-infiltrating inflammatory cells the following mAb’s were used: OX18 (pan–class I) (36), OX27 (PVG [RT-1] and BN [RT-1] class I) (37), and U9F4 (AO [RT-1] class I) (38). To detect ECs and infiltrating macrophages, mAb’s RECA-1 (39) and ED1 (40) were used, respectively. MHC class I, EC, and macrophage staining was performed on 7-µm cryostat sections using the DAKO EnVision staining kit (DAKO A/S). Sections were counterstained with Mayer’s hematoxylin and coverslipped with Depex mounting medium (BDH Laboratory Supplies, Poole, United Kingdom).

Quantitation of TA. After histological staining of tissue sections for elastin and α-actin, presence of TA was quantified using a computerized morphometric analysis system (QWin Software; Leica Microsystems B.V., Rijswijk, The Netherlands). Total surface area of arteriosclerotic lesions (neointima) was calculated and expressed as µm².

Microdissection of neointimal tissue. Microdissection of neointimal tissue was performed on α-actin–stained 7-µm cryostat sections (taken from the center of the graft) from aortic as well as cardiac allografts. α-Actin staining ensured positive identification of neointimal VSM cells in aortic allografts and affected CAs in cardiac allografts. After counterstaining with Mayer’s hematoxylin, tissue sections were placed in PBS and stored overnight at 4°C. Subsequently, microdissection was performed using a micromanipulator (Leica Microsystems B.V.) from neointimal lesions containing large numbers of α-actin–positive VSM cells. Cell clusters consisting of 20–30 cells were microdissected from neointimal lesions in aortic allografts as well as in CAs. To prevent potential contamination of the cell clusters with infiltrating mononuclear cells, single nuclei from α-actin–positive neointimal cells in aortic allografts and CAs were microdissected as well. For single cell picking, nuclei were randomly selected among α-actin–positive cells having an easily visible nucleus surrounded by α-actin–positively stained cytoplasm. Microdissected cell clusters and isolated single nuclei were directly transferred into PCR reaction tubes containing 20 µl Milli-Q water (Millipore B.V., Etten-Leur, The Netherlands, USA) and stored at -20°C until PCR analysis.

PCR analyses. The first PCR that was used to analyze the origin of neointimal cell types after aortic and cardiac allografting was based on polymorphic regions present in the Tap2 gene encoding a subunit of a peptide transporter involved in loading of MHC class I molecules with peptides (31). Because the Tap2 gene is located within the MHC, all MHC haplotypes are associated with a specific Tap2 allele. To determine the Tap2 genotype of isolated DNA, a PCR procedure developed by Joly et al. was used based on differences in restriction sites found in the sequences of the Tap2 cDNAs (31). Oligonucleotides used were as follows: 360 5′-ccgggctc-tagatgcagactgttagaagggc-3′ and 361 5′-ccgggcaacctctgggagt-3′ (sequences in lower case do not correspond to Tap2 sequences but were introduced for cloning of the various alleles by PCR). Since microdissected cell clusters and single nuclei did not contain sufficient DNA to give a positive signal (not shown), the Tap2 PCR was performed on 70-µm-thick tissue samples obtained from aortic allografts. In addition, microdissected CAs showing severe TA from one graft were pooled (10–20 CAs) to get sufficient template DNA to perform the Tap2 PCR. After amplification, 10 µl of the PCR reaction mixture was analyzed on a GelStar (BMA, Rockland, Maine, USA) 2% agarose gel. Sizes of the PCR products of the different rat strains were: Lewis, 380 bp; BN, 360 bp; PVG, 340 bp; and AO, 360 bp. Since all transplantsations were performed in female (donor) to male (recipient) combinations, we could also use sex markers as indicators for cellular origin. Therefore, a nested PCR procedure was developed that specifically enabled identification of a Y chromosome–associated MHC sequence (European Molecular Biology Laboratory [EMBL]/GenBank data libraries accession no. X80155) (41) at the single-cell level and thereby the (male) origin of cells (described in detail elsewhere; ref. 32). The following Y chromosome–specific oligonucleotides were used: first round 5′-caggtctgtaaatctgagggtaatgatggaggtgagatagta-3′ (forward) and 5′-ccccatgtttggctgactatagagta-3′ (reverse), and second round 5′-gtcttgattttcagttcgtgagttta-3′ (forward) and 5′-aatgttgactgactgccgatc-3′ (reverse). The first and second round of PCR amplification resulted in 549-bp and 128-bp DNA fragments, respectively. After amplification, 10 µl of the 2nd round reaction mixture was analyzed on a GelStar (BMA) 2% agarose gel. All amplifications were carried out using an Amplitron II...
**Thermolyne thermocycler** (Barnstead/Thermolyne, Dubuque, Iowa, USA).

**Restriction enzyme analysis.** Since Tap2 PCR fragments derived from Lew DNA bear a HindIII restriction site and BN-derived DNA fragments do not, further discrimination between Lew- and BN-derived DNA could be made by HindIII restriction analysis. After Tap2 PCR analysis, 5 µl PCR product was added to 5 µl restriction mix containing 5 U HindIII (Life Technologies B.V., Breda, The Netherlands). After incubation for 1 hour at 37°C, DNA samples were analyzed by 2% agarose gel electrophoresis. Lew-derived DNA was cut into two fragments (276 and 104 bp), whereas BN-derived DNA remained uncut.

Since PVG- and AO-derived DNA both lack the HindIII restriction site in their Tap2 sequence, discrimination could only be made by difference in size of the PCR products (340 and 360 bp respectively). Therefore, those PCR products were electrophoresed on a 12% polyacrylamide gel using a Mini-PROTEAN II Bio-Rad system (Bio-Rad Laboratories B.V., Veenendaal, The Netherlands). After electrophoresis the gel was soaked in ethidium bromide–containing TBE buffer to visualize PCR products.

**Statistical analysis.** To analyze differences in surface area of arteriosclerotic lesions for statistical significance, a Mann-Whitney U test was performed using SPSS 7.5.2 for Windows (SPSS, Chicago, Illinois, USA). Differences were considered to be statistically significant when P values were less than 0.05.

**Results**

**Effect of irradiation and CsA treatment on TA.** Transplantation of fully MHC-mismatched aortic allografts in the PVG to AO rat strain combination resulted in the development of severe TA. Whereas AO isografts were unaffected or only slightly affected, allografts showed significant (P < 0.05) neointima formation 3 months after transplantation, indicating that TA in this model is mainly caused by an alloantigen-driven process (Figure 1a). We then asked whether aortic allografts, which contain medial VSM cells that have lost migratory and proliferative capacities, still lead to development of neointimal lesions after transplantation. To answer this question, PVG aortic allografts were ex vivo 50 Gy γ-irradiated prior to engraftment into AO recipients. As shown in Figure 1a γ-irradiated allografts developed neointimal lesions to the same extent as nonirradiated allografts, indicating that graft medial VSM cells probably do not contribute to the process of TA.

Subsequently, the influence of immunosuppressive therapy on the development of TA after allogeneic aorta transplantation was analyzed. Aortic grafts were transplanted in the Lew to BN strain combination. From previous studies it became obvious that severe TA develops within 4 weeks after transplantation in this strain combination (Figure 1b) (42). Recipient rats received either no treatment or CsA at 5 mg/kg/d or 10 mg/kg/d for 28 days. After 14 and 28 days, 24-hour trough whole blood CsA levels were determined.
in three animals from either group. No differences were observed between the 14- and 28-day time points, and therefore data from both time points were pooled. Animals treated with 5 mg/kg/d and 10 mg/kg/d CsA had mean trough levels of 1016 ± 268 ng/ml and 2383 ± 327 ng/ml, respectively. The nontreated group had trough levels below detection level (<25 ng/ml). Treatment with 5 mg/kg/d and 10 mg/kg/d both resulted in complete inhibition of neointima formation 28 days after transplantation (P < 0.01 and P < 0.05, respectively) (Figure 1b). After CsA treatment, no differences were observed compared with the isograft (BN to BN) controls.

Representative micrographs of vascular wall morphology are shown in Figures 1, c–f. The nontreated group displayed severe neointima formation with presence of large numbers of α-actin–positive VSM cells (Figure 1c). The media had become acellular, and medial α-actin–positive VSM cells had disappeared. In addition, graft-infiltrating ED1+ macrophages were present in both the adventitia and the neointima (Figure 1d).

After CsA treatment, however, morphology appeared to be normal (no neointima formation) in both treatment groups and was similar to nontransplanted aortic tissue. Grafts from CsA-treated recipients were characterized by a high nuclear density in the media, which appeared to be α-actin–positive VSM cells (Figure 1, e and f).

**Origin of ECs in nontreated and CsA-treated aorta recipients.** To analyze the origin (donor vs. recipient) of ECs after allogeneic aorta transplantation, MHC class I haplotype-specific immunohistochemistry was performed (Figure 2). PVG allografts transplanted in AO recipient rats showed severe neointima formation 3 months after transplantation (Figure 1a). Presence of ECs was confirmed by positive staining using mAb RECA-1 (not shown). Serial tissue sections were stained using mAb’s OX18, OX27, and U9F4. Neointimal ECs express MHC class I antigens as indicated by intense staining with mAb OX18 (pan–class I), whereas neointimal VSM cells do not express MHC class I antigens sufficiently to allow detection by this staining (Figure 2a). Note the absence of nuclei in the media and the swollen appear-

**Figure 2**
Neointimal ECs in aortic allografts are of recipient origin. Immunohistochemical staining using MHC class I haplotype-specific mAb’s was performed to determine the origin (donor vs. recipient) of neointimal ECs after allogeneic aorta transplantation. PVG allografts transplanted in AO recipients showed severe TA. (a) MHC class I expression on ECs (OX18-positive). Note the absence of VSM cells in the media (m). Donor endothelium had disappeared (OX27-negative) (b) and had been replaced by recipient-derived ECs (U9F4-positive) (c). Lew allografts transplanted into BN recipient rats also showed severe TA. (d) MHC class I expression on ECs (OX18-positive). (e) Donor endothelium had been replaced by recipient-derived ECs (OX27-positive). After CsA treatment, no neointima formation was observed 1 month after transplantation. (f) ECs covering the internal elastic lamina expressed MHC class I antigens (OX18-positive). However, ECs were OX27 (recipient class I)-negative (g), indicating that after CsA treatment EC replacement does not occur. Arrowheads indicate positively stained ECs. Original magnification: ×400.
ance of the ECs. Staining with mAb OX27 (donor type) was negative (Figure 2b), whereas mAb U9F4 (recipient type) clearly showed positive staining of ECs (Figure 2c), indicating that donor ECs had been replaced by recipient-derived ECs. Similar results were obtained after haplotype-specific MHC class I staining of PVG allografts that were 50 Gy γ-irradiated prior to transplantation into AO recipient rats (not shown).

To further analyze whether EC replacement had also taken place after CsA treatment, serial tissue sections from aortic grafts transplanted in the Lew to BN strain combination were stained using mAb’s OX18 (pan–class I) and OX27 (recipient type). Again, presence of ECs was confirmed by positive staining using mAb RECA-1 (not shown). Neointimal ECs from untreated animals showed abundant MHC class I expression as indicated by positive OX18 staining (Figure 2d). Also, in this strain combination, donor endothelium had been replaced by recipient-derived ECs. Similar results were obtained after haplotype-specific MHC class I staining of PVG allografts that were 50 Gy γ-irradiated prior to transplantation into AO recipient rats (not shown).

VSM cell α-actin staining (Figure 3f) confirmed presence of severe neointima formation and luminal narrowing in the CAs in PVG allografts explanted 530 days after transplantation. RECA-1 staining showed presence of ECs (Figure 3, b and g). Note the swollen and activated appearance of the neointimal ECs (Figure 3, h). MHC class I expression on ECs was confirmed by positive staining using OX18 (Figure 3, c and h). Neointima-infiltrating leukocytes were virtually absent in the arteriosclerotic lesions. However, perivascular mononuclear cell infiltrates (MHC class I–positive) were present in the PVG allografts. Staining with recipient-specific mAb U9F4 showed positive staining of the mononuclear cell infiltrates (which, as expected, are recipient-derived) (Figure 3i). However, the neointimal ECs in the PVG allografts are negative, suggesting that, in contrast to aortic allografts, recipient ECs have not replaced donor endothelium in CAs with severe TA. This was confirmed by positive staining of neointimal ECs with the donor-specific mAb OX27.
Similar findings were obtained in CAs in PVG allografts showing only mild TA as well as CAs showing no TA at all (not shown).

**Tap2 PCR analysis of aortic and cardiac tissue.** Since neointimal VSM cells do not express MHC class I antigens sufficiently to allow MHC class I-specific immunohistochemistry to study the origin of neointimal VSM cells in aortic and cardiac allografts, another (PCR-based) approach was followed to define the origin of α-actin-positive neointimal cells.

The first PCR that was used to analyze the origin of neointimal cell types after aortic and cardiac allografting was based on polymorphic regions in the Tap2 gene. Because of the relatively low sensitivity of the Tap2 PCR, analysis was performed on 70-μm-thick tissue samples of aortic allografts transplanted in the BN to Lew, Lew to BN, and Lew to BN + CsA (5 and 10 mg/kg/d CsA) combinations. Three individual grafts from either group were included, and of each graft four 70-μm-thick tissue samples were analyzed. Nontransplanted BN and Lew aortic tissue served as controls for specificity of this Tap2 PCR. As shown in Figure 4a, control BN and Lew DNA resulted in DNA fragments of 360 bp and 380 bp, respectively. Restriction analysis with HindIII showed cleavage of the Lew-derived 380-bp fragments into 276-bp and 104-bp fragments, whereas the BN-derived 360-bp fragments remained uncut, thereby proving the specificity of this Tap2 PCR. Subsequently, this PCR was applied to tissue samples obtained from grafts transplanted from BN donor to Lew recipient rats. Tap2 PCR followed by HindIII restriction showed cleavage of the DNA fragments into two fragments, indicating Lew (recipient) origin of the majority of the DNA present in these samples. This was confirmed by analysis of tissue obtained from grafts transplanted into the reciprocally combined (Lew to BN). After HindIII restriction the DNA fragments remained uncut, indicating BN (recipient) origin of the DNA present in these samples. Finally, the same analysis was performed on tissue from Lew grafts transplanted into BN recipient rats followed by CsA treatment (5 mg/kg/d) for 28 days. HindIII restriction showed cleavage of the DNA fragments into two fragments, indicating Lew (donor) origin of the majority of the DNA after CsA treatment (Figure 4a). Similar results were obtained from analysis of aortic allografts after treatment with 10 mg/kg/d CsA (not shown). These results suggest that both neointimal ECs and VSM cells are recipient-derived in aortic allografts.

To study whether the same holds true for neointimal cells in cardiac allografts, Tap2 PCR analysis was performed on microdissected CAs showing severe TA. To obtain sufficient DNA, microdissected CAs from one graft were pooled prior to PCR analysis. Since PVG- and AO-derived DNA both lack the HindIII restriction site in their Tap2 sequence, discrimination could only be achieved using HY PCR analysis of microdissected neointimal cell clusters from allografts transplanted in the female PVG to male AO strain combination. Grafts were 50 Gy-γ-irradiated prior to transplantation or left untreated. Representative results of first- and second-round PCR products (549 and 128 bp) of two irradiated and three untreated grafts are shown. All samples show the male-specific DNA fragments after first- and second-round PCR.

**Figure 4**

PCR analysis of 70-μm-thick tissue samples, microdissected cell clusters, and microdissected single nuclei of α-actin-positive neointimal VSM cells after allogeneic aorta transplantation. (a) Tap2 PCR analysis of 70-μm-thick tissue samples from aortic allografts transplanted in the BN to Lew, Lew to BN, and Lew to BN + CsA combinations. Lew-derived DNA is cut after HindIII restriction, whereas BN-derived DNA remains uncut. Restriction analysis indicates recipient origin in untreated allografts. After CsA treatment, restriction analysis indicated donor origin. (b) HY PCR analysis of microdissected neointimal cell clusters from allografts transplanted in the female PVG to male AO strain combination. Grafts were 50 Gy-γ-irradiated prior to transplantation or left untreated. Representative results of first- and second-round PCR products (549 and 128 bp) of two irradiated and three untreated grafts are shown. All samples show the male-specific DNA fragments after first- and second-round PCR. (c) Micrograph (double staining for α-actin and elastin) on an aortic allograft after isolation of single nuclei using a micromanipulator. Arrowheads indicate remaining gaps after isolation of nuclei of three separate α-actin-positive neointimal cells. Note absence of VSM cells in the media. a, adventitia; m, media; ni, neointima. Asterisk shows an α-actin-positive VSM cell. Original magnification: ×400. (d) HY PCR performed on eight single nuclei microdissected from two aortic allografts transplanted from female PVG to male AO recipient rats, showing the male-specific (recipient-specific) 128-bp DNA fragment after nested PCR. bp, 100-bp ladder; H2O, water control (i.e., no template DNA present in the PCR reaction mixture).
PCR-based analysis was developed as described in detail elsewhere (32). This highly sensitive and specific nested PCR procedure enabled the identification at the single-cell level of a 128-bp Y chromosome–associated DNA sequence and thereby of the (male) origin of cells. Since aortic and cardiac allografting was performed not only between MHC-incompatible rat strains but in a female (donor) to male (recipient) combination as well, we could use this HY marker as an indicator for male cellular origin. First, this PCR was applied to microdissected cell clusters derived from neointimal lesions in aortic allografts with or without 50 Gy γ-irradiation transplanted from female PVG donor to male recipient rats. Three individual grafts of either group were studied, and of each graft five microdissected cell clusters were analyzed. HY PCR performed on microdissected cell clusters resulted in appearance of Y chromosome–specific DNA fragments in the nonirradiated as well as the 50 Gy γ-irradiated aortic allografts (Figure 4b). First-round PCR already resulted in detectable amounts of specific DNA fragments (549 bp). Additional nested PCR confirmed this observation by the amplification of a 128-bp DNA fragment. Subsequently, this PCR was applied to microdissected single CAs obtained from three cardiac allografts. Of either graft, four single CAs (with severe TA) were microdissected and analyzed individually for the presence of male-derived (recipient-derived) cells. All samples analyzed showed the male-specific DNA fragments after the first round, as well as after nested PCR. Representative results of the nested PCR performed on three individual CAs microdissected from three different grafts, as well as specificity controls (male- and female-derived DNA), are shown in Figure 5b. Results obtained with the HY-specific as well as with the Tap2 PCR thus indicate the presence of recipient-derived cells, both in microdissected neointimal cell clusters obtained from aortic allografts and in microdissected whole CAs showing severe TA. However, although neointimal tissue predominantly consists of α-actin–positive VSM cells (Figures 1c and 3f), neointimal lesions can also contain an appreciable number of recipient-derived infiltrating leukocytes, especially macrophages (43). The neointimal lesions present in aortic allografts in this study indeed contained considerable numbers of ED1+ macrophages (Figure 1d). Neointimal lesions in CAs appeared to contain few if any infiltrating inflammatory cells (Figure 3); however, one cannot exclude the possibility that such inflammatory cells were present. Since the HY PCR is sufficiently sensitive to detect male-derived cells on the single-cell level, false-positive results could be obtained by neointima-contaminating recipient-derived inflammatory cells.

HY PCR analysis of microdissected single VSM cell nuclei. To avoid contamination of the samples with neointima-infiltrating inflammatory cells, HY PCR analysis was performed on microdissected single nuclei from α-actin–positive neointimal cells from aortic and car-
Mendelian inheritance patterns, we could test the efficacy of the transplantation procedure and subsequent PCR analysis. In total, 13 isolated nuclei from two grafts were analyzed, of which 12 (92.3%) showed the Y chromosome–specific (male-specific) 128-bp DNA fragment (not shown). Subsequently, the same procedure was performed on nuclei isolated from female PVG grafts transplanted to male AO recipients. Three grafts were analyzed, and from each graft eight nuclei were isolated. PCR analysis revealed that 21/24 (87.5%) of the samples contained DNA and, of these, 19/21 (90.5%) displayed the male-specific 128-bp DNA fragment. These results indicate that at least 90% of α-actin–positive VSM cells in transplant arteriosclerotic lesions after allogeneic aorta transplantation are of recipient origin.

Coronary arteries. Subsequently, this PCR was applied to single nuclei microdissected from CAs showing severe TA. Again, three grafts were analyzed and from each graft eight nuclei were isolated. After allogeneic cardiac transplantation, neointimal α-actin–positive VSM cells are recipient-derived. Figures 4d and 5c show representative results of the HY PCR applied to microdissected single nuclei from aortic and cardiac allografts, respectively.

The nucleotide sequence of the 128-bp PCR products was determined to confirm the Y-chromosomal nature of the amplified material. The obtained sequences were compared with the sequence data available from EMBL/GenBank data libraries, accession no. X80155 (clone 9.1ES8). The sequence data of the PCR products were identical to the sequence data available from the database, indicating that the 128-bp DNA fragment belongs to the Y chromosome (not shown).

Discussion
CTD is the primary cause of allograft loss after the first perioperative year and is today’s most important problem in clinical organ transplantation. The hallmark of CTD is development of TA; i.e., progressive intimal thickening (neointima formation) consisting of α-actin–positive VSM cells intermingled with recipient-derived mononuclear cells (T cells and macrophages) (2). Current thought on the process of TA holds that the Y chromosome–specific immunohistochemistry and PCR analyses of DNA sequences allowing discrimination between donor and recipient origin were performed.

First we analyzed development of TA after transplantation of aortic grafts that were ex vivo 50 Gy γ-irradiated (to inhibit proliferation and migration of medial VSM cells) prior to transplantation. It has been shown that γ-irradiation of cultured VSM cells in vitro inhibits proliferation of these cells after stimulation (45). Three months after transplantation, irradiated grafts showed severe TA, similar to nonirradiated grafts, suggesting recipient origin of neointimal cells. Aziz et al. performed similar experiments using liquid nitrogen to destroy medial VSM cells in aortic allografts prior to transplantation. These frozen allografts showed TA to the same extent as unmodified grafts, also suggesting recipient origin of neointimal cells (22).

Subsequently, aorta recipients were treated with 5 mg/kg or 10 mg/kg CsA for 28 days to study the effect of immunosuppressive treatment on the development of TA. CsA treatment completely blocked neointima formation and preserved the vascular media, confirming that development of TA is primarily alloimmune-mediated. Neointima formation is dependent on, and proportional to, the extent of medial VSM cell damage (46). By suppressing the alloimmune response by CsA treatment, no medial VSM cell (and possibly also no EC) damage occurs; thereby development of TA is prevented. However, Stoltenberg et al. showed that administration of the same amounts of CsA, resulting in similar 24-hour CsA trough levels, resulted in significant inhibition, but no prevention of neointima formation 3 and 6 months after aorta transplantation in the AxCgg35Irish (ACI) to Lew strain combination (33). Differences between the rat strains used and the follow-up period might explain the different results.

To confirm the origin (donor vs. recipient) of neointimal endothelial and VSM cells in rat aortic allografts displaying extensive TA, we first performed recipient MHC class I specific immunohistochemistry. Results indicate that ECs indeed are of recipient origin in nontreated and irradiated aortic allografts. CsA treatment prevented neointima formation, and MHC class I staining showed that ECs layering the internal elastic lamina are still of donor type and are not being replaced by recipient-derived ECs. In contrast to the aortic allografts, neointimal ECs are still of donor type in cardiac allografts showing severe TA 530 days after engraftment. Similar results were found by Hasegawa et al. (47), who showed that after MHC-mismatched cardiac allografting in mice ECs in CAs with neointimal lesions are still of donor origin at 12 weeks after transplantation by using MHC class II–specific immunohistochemistry.

The origin of ECs (donor vs. recipient) after organ allografting has been debated since Woodruff (48) and Medawar (49) suggested that replacement of donor ECs with recipient ECs (graft adaptation) might be an
and microdissected CAs with severe TA, we found that between donor and recipient-derived cells. For MHC class II antigens on neointimal VSM cells (47), detection. A similar lack of expression has been described for sufficient amounts of MHC class I antigens for neointimal and medial haplotype-specific immunochemistry did not allow dis-

experimental transplant models (21–23). MHC class I indicating recipient origin of neointimal VSM cells in patients using fluorescence in situ hybridization on tissue sections (23). This is supported by findings of Hruban et al., who showed donor origin of neointimal VSM cells in two patients using fluorescence in situ hybridization on tissue sections (20). However, data have also been reported indicating recipient origin of neointimal VSM cells in experimental transplant models (21–23). MHC class I haplotype-specific immunochemistry did not allow dis-
tinction between donor and recipient origin of neointimal VSM cells in our transplant models, since both neointimal and medial α-actin–positive VSM cells do not express sufficient amounts of MHC class I antigens for detection. A similar lack of expression has been described for MHC class II antigens on neointimal VSM cells (47). Therefore we addressed the question of whether neointimal VSM cells are truly graft-derived using two unrelated PCR-based analyses (Tap2 and HY) on aortic and cardiac allografts, enabling discrimination between donor and recipient-derived cells.

Using both types of PCR analysis on aortic allografts and microdissected CAs with severe TA, we found that at least a significant part of the cells in these tissues displays recipient-type DNA. Similar data were obtained for irradiated aortic allografts. However, neointimal lesions have been reported to contain an appreciable number of recipient-derived infiltrating leukocytes (43). Also, the aortic allografts analyzed in this study contained neointima-infiltrating mononuclear cells, consisting predominantly of ED1+ macrophages and some T cells. Such contaminating recipient-derived leukocytes can account for the recipient-specific signal in our analysis of aortic allografts and microdissected CAs, especially for the male-specific analysis, since the HY PCR is much more sensitive than the Tap2 PCR. Similar problems may also have influenced the results of Plissonnier et al. (21) and more recently Brazelton et al. (23) on the origin of VSM cells in vascular allografts. These groups used allospecific immunofluorescent staining of cell suspensions (21) and tissue sections (23) of aortic and femoral arterial allografts. Although their conclusion, that neointimal VSM cells are of recipient origin, in itself is probably correct, their type of analysis cannot distinguish between the infiltrating (recipient-derived) leukocytes and neointimal VSM cells.

Since our male-specific nested HY PCR allows detection of male-derived cells at the single-cell level (32), we analyzed microdissected single nuclei of neointimal VSM cells. Nuclei were isolated from neointimal lesions of both aortic allografts and CAs with severe TA by positive selection (α-actin staining), thereby eliminating the risk of contaminating our samples with infiltrating recipient-derived mononuclear cells. HY PCR analysis of single nuclei of α-actin–positive neointimal cells showed that at least 90% of these cells in neointimal lesions of both aortic allografts and CAs from cardiac allografts are of recipient origin. Although Hasegawa et al. claim that VSM cells in neointimal lesions of CAs after cardiac transplantation in mice are probably still of donor origin at 12 weeks after transplantation, their type of analysis, MHC class II specific immunohistochemistry, does not allow determination of origin of neointimal VSM cells because of lack of MHC class II antigen expression on these neointimal VSM cells (47). Since our data show recipient origin of neointimal VSM cells (aortic and cardiac allografts) and endothelium (aortic allografts), the question arises: what is the anatomical origin of these neointimal cells? Kouchi et al. demonstrated bloodstream origin of endothelial and α-actin–positive smooth muscle cells present on Dacron grafts transplanted in dogs (51). Recently, recirculating bone marrow–derived CD34+ EC progenitor cells (EPCs) have been identified in the peripheral blood of humans (52, 53). The frequency of circulating EPCs increases in response to regional ischemia in both mice and rabbits. This increased frequency of circulating EPCs contributes to neovascularization of ischemic regions (54), indicating that EPCs are mobilized from the bone marrow into the circulation and subsequently “home” to sites needing neovascularization. Using bone
marrow chimeric dogs, Shi et al. demonstrated donor (bone marrow) origin of ECs attached on implanted Dacron grafts, also indicating existence of circulating bone marrow–derived ECs (55). In another study, Bhat-tacharyya et al. recently showed enhanced re-endothe-
lization of Dacron grafts after seeding with bone mar-
orow–derived CD34+ cells prior to implantation (56).
Taken together, these results suggest an important role
for bone marrow–derived CD34+ progenitor cells as a
source for ECs.

The anatomical source of the neointimal VSM cells
is still unknown. One possibility might be ingrowth of
medial VSM cells from the recipient side of anastomosis,
as suggested by Aziz et al. (22). VSM cell destruction
by freezing the recipient aortic ends at both sides near
the anastomosis resulted in less pronounced intimal
thickening compared with untreated aortic allografts.
However, the frozen recipient aortic ends remained
devoid of α-actin–positive VSM cells during the obser-
vation period of 60 days. If the neointimal VSM cells
originate from the recipient media, one should expect
migration of α-actin–positive VSM cells along the
frozen aortic ends toward the mid-graft region. This
was, however, not clear from the study by Aziz et al (22).
A second possibility is transdifferentiation from ECs
into α-actin–positive VSM cells. Studying quail
embryos, de Ruiter et al. showed transdifferentiation of
embryonic ECs into α-actin–expressing mesenchymal
cells in vivo and in vitro (57).

The third possibility for the anatomical origin of
neointimal VSM cells is the existence of VSM progeni-
tor cells that differentiate into α-actin–positive VSM
cells. Circulating VSM progenitor cells, however, have
not been identified so far. Bucala et al. reported the exis-
tence of a non–bone marrow–derived circulating cell
population with fibroblast properties that specifically
enters sites of tissue injury (58). In addition, Kouchi
et al. described the presence of α-actin–positive smooth
muscle cells on Dacron grafts, which apparently origi-
nated from the bloodstream (51). Using whole vessel
wall organ cultures, Slomp et al. showed migration of
adventitial myofibroblasts to the luminal surface, where
they transdifferentiated into endothelial-like cells as
well as into a VSM cell phenotype in intimal thickening
(59). These results suggest the existence of a myofi-
broblast cell lineage that can serve as an endothelial and
smooth muscle stem cell population. In our aortic
transplant model, sequential immunohistological
analysis of developing TA at an early stage after trans-
plantation indeed suggests a blood-borne influx of
regenerating VSM cells (not shown).

In conclusion, in contrast to current thought about
the process of TA, the α-actin–positive VSM cells in
the hyperplastic neointima in aortic and cardiac allografts
are of recipient and not of donor origin. Neointimal
ECs in aortic allografts are replaced by recipient-
derived endothelium, whereas in cardiac allografts
neointimal ECs are not replaced and are still of donor
origin. So, independent of EC replacement, donor VSM
cells in the graft are replaced by VSM cells of recipient
origin in these two transplant models.

We propose that the development of TA is an
attempt to restore vascular function upon immuno-
logical injury (alloreactivity) and is essentially part of
a normal healing process. Basically, as a result of the
immunological injury, medial VSM cells disappear,
leaving the elastin network as a scaffold allowing
restoration of the vessel wall by recipient-derived cells.
However, in TA this remodeling does not seem to stop,
eventually leading to total occlusion of the vessels
involved. According to this concept, at least two win-
dows in time seem to exist that would allow therapeu-
tic intervention to prevent the development of severe
TA. The first is early after transplantation, when pri-
mary vascular graft damage through either ischemia
or alloreactivity should be preventable. Perhaps cur-
rent immunosuppressive therapies should even be
intensified or adapted to suppress nonclinical rejec-
tion episodes causing vascular damage eventually lead-
ing to TA. The second window for intervention would
be later, when TA starts to develop. Our data suggest
that TA is the result of a repair process proceeding
beyond the needs of functional repair. Future inter-
vention strategies at this stage should aim at regaining
proper control of this dysregulated process.

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