Oligomerized, filamentous surface presentation of RANTES/CCL5 on vascular endothelial cells

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Vascular endothelial cells present luminal chemokines that arrest rolling leukocytes by activating integrins. It appears that several chemokines must form higher-order oligomers to elicit proper in vivo effects, as mutants restricted to forming dimers have lost the ability to recruit leukocytes to sites of inflammation. Here, we show for the first time that the chemokine RANTES/CCL5 binds to the surface of human endothelial cells in a regular filamentous pattern. Furthermore, the filaments bound to the surface in a heparan sulfate-dependent manner. By electron microscopy we observed labeling for RANTES on membrane projections as well as on the remaining plasma membrane. Mutant constructs of RANTES restricted either in binding to heparin, or in forming dimers or tetramers, appeared either in a granular, non-filamentous pattern or were not detectable on the cell surface. The RANTES filaments were also present after exposure to flow, suggesting that they can be present in vivo. Taken together with the lacking in vivo or in vitro effects of RANTES mutants, we suggest that the filamentous structures of RANTES may be of physiological importance in leukocyte recruitment.
coupled\(^7,18\). For example, RANTES and several other chemokines may oligomerize on heparin beads. Dimeric forms of many chemokines have also been reported to have higher affinity for GAGs than their monomeric variants and GAG binding may involve, or induce, oligomeric structures larger than dimers. For example, although MCP-1/CCL2 forms a dimer in solution, a heparin octasaccharide shifts the equilibrium toward MCP-1 tetramers\(^19\).

In the present study, we have analysed the distribution of endogenous and recombinant RANTES on vascular endothelial cells. We show that secreted RANTES bound at the cell surface, not in a diffuse or aggregated pattern, but surprisingly as specific, well ordered filaments that elongated over time. The filamentous expression of RANTES was not found when expressing mutants that are restricted to form dimers or tetramers, or when expressing a mutant with reduced GAG affinity. These data add further support to the merging view that formation of higher order oligomers of RANTES is crucial to elicit its fully biological function in activation of leukocytes.

Results

**RANTES is organized in filament-like structures on the endothelial cell surface.** In a screen for chemokines that sort to the regulated secretory pathway in endothelial cells, we observed by means of immunofluorescent staining and confocal microscopy that RANTES was expressed in patterns distinct from those of other chemokines\(^20,21\) (Øynebråten et al., unpublished data). When HUVECs were cultured in vitro and stimulated with TNF\(\alpha\) in combination with IFN\(\gamma\) before fixation and immunostaining, RANTES mainly localized in elongated, filamentous structures (Fig. 1A) and\(^26\). Five different antibodies towards RANTES were tested, and they all labeled elongated structures of RANTES. Analysis at different time points after exposing HUVECs to TNF\(\alpha\) and IFN\(\gamma\) revealed that RANTES was distributed in puncta and short elongated structures after 12 h. In the course of analysis these structures elongated from an average length of 2 \(\mu\)m at 24 h to 15 \(\mu\)m after 60 h of stimulation (Fig. 1A). Based on these observations, we suggest that short structures of RANTES can develop into long filaments in cultures of endothelial cells activated by pro-inflammatory stimuli. To elucidate whether the filaments were present on the cell surface, we stained live HUVECs kept on ice, observing that RANTES filaments are subject to surface presentation on endothelial cells (Fig. 1B).

Several types of membrane projections have been described for endothelial cells\(^8,22–24\), and indeed, RANTES as well as IL-8/CXCL8 have been detected on microvillous-like extensions on the luminal endothelial cell surface\(^8\). We therefore asked whether RANTES filaments are subject to surface presentation on endothelial cells (Fig. 1B).

Filament formation does not depend on TNF\(\alpha\) + IFN\(\gamma\)-stimulation. In agreement with a previous study\(^25\) we observed that RANTES was most strongly induced in HUVECs by simultaneous stimulation with TNF\(\alpha\) and IFN\(\gamma\). Because we did not observe filamentous organization of chemokines in resting or IL-1\(\beta\) stimulated-HUVECs\(^20,23\), we asked if...
observing at low and medium concentrations of heparin (0.9 and 10 ng/ml) TNFα in combination with 1 ng/ml IFNγ for 24 h and immunostained with an antibody towards MCP-1. The inset shows immunostaining of RANTES from the same experiment, in HUVECs stimulated with 10 ng/ml TNFα + 1 ng/ml IFNγ. (B) Left image; HUVECs were incubated with 1 µg/ml recombinant RANTES before fixation and immunostaining. Right image; HUVECs were not stimulated but electroporated with a DNA plasmid encoding RANTES before fixation and immunostaining of RANTES with a rabbit anti-RANTES antibody. Labelling with ulex (green) was included to visualize individual cells. Scale bars, 10 µm. All images were acquired by confocal microscopy.

Figure 2 | Filaments of RANTES form independently of TNFα and IFNγ-stimulation. (A) HUVECs were stimulated with 1 ng/ml IL-1β, or 10 ng/ml TNFα in combination with 1 ng/ml IFNγ for 24 h and immunostained with an antibody towards MCP-1. The inset shows immunostaining of RANTES from the same experiment, in HUVECs stimulated with 10 ng/ml TNFα + 1 ng/ml IFNγ. (B) Left image; HUVECs were incubated with 1 µg/ml recombinant RANTES before fixation and immunostaining. Right image; HUVECs were not stimulated but electroporated with a DNA plasmid encoding RANTES before fixation and immunostaining of RANTES with a rabbit anti-RANTES antibody. Labelling with ulex (green) was included to visualize individual cells. Scale bars, 10 µm. All images were acquired by confocal microscopy.

Extracellular molecules can polymerize RANTES. We next examined whether RANTES alone can polymerize and form filaments or whether other molecules are necessary (Fig. 3). First, we incubated RANTES (1 µg/ml) with cell growth medium (MCDB131) without serum, or with conditioned media harvest ed from unstimulated HUVECs. Heparin was included to visualize individual cells. Scale bars, 10 µm. All images were acquired by confocal microscopy.

Figure 3 | RANTES is dependent on other molecules to form organized structures. RANTES (1 µg/ml) was incubated either in cell growth medium without serum or in conditioned cell growth medium containing serum. The conditioned medium was harvested from unstimulated cultures of HUVECs. Heparin was added and its final concentration is indicated in each image. After 35 h, the samples were fixed and immunostained with a rabbit antibody toward RANTES. The images were acquired by widefield microscopy. Inserts show high magnification of squared areas. Scale bars, 10 µm.

1.7 µg/ml, respectively) no evidence of filament formation of RANTES (Fig. 3, left panel). However, following incubation with the highest concentration of heparin (3.4 µg/ml), RANTES organized in a structured pattern that nevertheless differed from that observed for RANTES in HUVEC cultures (Fig. 3, left panel, lower image). In the next series we added increasing concentrations of heparin to conditioned media, observing even in the absence of heparin that RANTES was distributed in an organized and distinct pattern (Fig. 3, right panel, upper image). When increasing the concentration of heparin (1.7 µg/ml) we observed that RANTES was organized in a pattern reminiscent of the RANTES filaments in HUVECs, and when reaching the highest level (3.4 µg/ml), RANTES was found in puncta (Fig. 3, right panel, lower image). Taken together, these experiments suggest that RANTES cannot alone organize into filaments and instead depends on helper or scaffold molecules. In addition, RANTES can organize into different patterns, likely depending on the type of molecules that are present.

Heparan sulfate is involved in cell surface immobilization of RANTES. It is well established that GAGs can immobilize chemokines on cell surfaces11,26,27. Because of this knowledge, and the finding that incubation of RANTES with heparin leads to formation of organized structures of RANTES (Fig. 3, left panel), we wanted to...
and overlapping with those of RANTES (Fig. 4B, upper panel), but overt co-localization was not observed. Another pattern appeared to be present at the cell border or between cells. At these sites, the antibody labeled long thread-like structures, and filaments of RANTES was observed along these structures (Fig. 4B, middle panel, and Fig. S2). Finally, because incubation of RANTES in conditioned media indicated that large scaffold molecules could organize RANTES (Fig. 3, right panel; upper image), we also labeled HUVEC cultures with biotinylated hyaluronan binding protein (bio-HABP). HABP shows high affinity for a decasaccharide unit of hyaluronan, which can form large polymers up to 20,000 kDa that organize into a wide variety of molecular architectures including fibrils and cable-like structures31. Except from some regions with large clusters of RANTES and HABP associated with HUVECs, we observed no co-localization between RANTES and HABP (Fig. 4B, lower panel). Between cells, HABP was found in large, round structures that were somewhat irregular and negative for RANTES (data not shown). Based on these experiments, we concluded that heparan sulfate molecules bind RANTES in HUVEC cultures, but the importance of other molecules for immobilization of RANTES cannot be excluded.

**Filamentous distribution of RANTES depends on the ability to form higher-order oligomers.** RANTES can self-associate and form higher-order oligomers in a concentration-dependent manner32,33. In contrast, two mutants of RANTES, E26A and E66A/E66S, show strongly reduced ability of such oligomer formation, and are instead restricted to form tetramers (E26A) and dimers (E66A/E66S)31,32,33. Furthermore, the 40s loop (“RKNR”) is suggested to be important for oligomerization as R44 may exert stabilization forces to the dimer interface, and R47 is shown to interact with the neighbouring molecule in RANTES structures16,34.

To elucidate whether the filamentous pattern of RANTES might be related to the properties of oligomer formation we electroporated HUVECs with DNA plasmids encoding E26A, “AANA”, or E66A. We also examined the mutant Y3A which shows in vivo properties similar to “AANA” but has unknown oligomerization status34. Microscopy after electroporation and immunostaining of the mutants E26A, “AANA”, and E66A revealed a pattern substantially different from that of wild type (wt) RANTES (Fig. 5A). The mutants mainly appeared in small granular structures as well as in the Golgi. “AANA” and E66A were also apparent throughout the cytoplasm, reminiscent of endoplasmic reticulum staining (Fig. 5A). Only 10–17% of the mutant-expressing HUVECs showed filamentous RANTES compared to 98% for the wt (Fig. 5C). The mutant Y3A distributed in a pattern similar to that of the other mutants suggesting that it had lost the ability to form higher-order oligomers (Fig. 5A).

To exclude that the dramatic reduction of filament formation was caused by low RANTES expression, we performed several control experiments. First, we compared the intensity of the signals for the immunolabeled mutants towards that of the wt by recording images at identical exposure times. By this comparison, we observed that the signal intensity of filament-forming wtRANTES varied and that it was present both at lower and higher levels than the signal of the mutants on a per cell basis. Even cells expressing wtRANTES at low levels (based on a very weak signal) showed a filamentous RANTES pattern similar to the wt by recording images at identical exposure times. By this comparison, we observed that the signal intensity of filament-forming wtRANTES varied and that it was present both at lower and higher levels than the signal of the mutants on a per cell basis. Even cells expressing wtRANTES at low levels (based on a very weak signal) showed a filamentous RANTES pattern similar to the wt by recording images at identical exposure times. By this comparison, we observed that the signal intensity of filament-forming wtRANTES varied and that it was present both at lower and higher levels than the signal of the mutants on a per cell basis. Even cells expressing wtRANTES at low levels (based on a very weak signal) showed a filamentous RANTES pattern similar to the wt by recording images at identical exposure times. By this comparison, we observed that the signal intensity of filament-forming wtRANTES varied and that it was present both at lower and higher levels than the signal of the mutants on a per cell basis. Even cells expressing wtRANTES at low levels (based on a very weak signal) showed a filamentous RANTES pattern similar to the wt by recording images at identical exposure times. By this comparison, we observed that the signal intensity of filament-forming wtRANTES varied and that it was present both at lower and higher levels than the signal of the mutants on a per cell basis. Even cells expressing wtRANTES at low levels (based on a very weak signal) showed a filamentous RANTES pattern similar to the wt by recording images at identical exposure times. By this comparison, we observed that the signal intensity of filament-forming wtRANTES varied and that it was present both at lower and higher levels than the signal of the mutants on a per cell basis. Even cells expressing wtRANTES at low levels (based on a very weak signal) showed a filamentous RANTES pattern similar to the wt by recording images at identical exposure times. By this comparison, we observed that the signal intensity of filament-forming wtRANTES varied and that it was present both at lower and higher levels than the signal of the mutants on a per cell basis. Even cells expressing wtRANTES at low levels (based on a very weak signal) showed a filamentous RANTES pattern similar to the wt by recording images at identical exposure times. By this comparison, we observed that the signal intensity of filament-forming wtRANTES varied and that it was present both at lower and higher levels than the signal of the mutants on a per cell basis. Even cells expressing wtRANTES at low levels (based on a very weak signal) showed a filamentous RANTES pattern similar to the wt by recording images at identical exposure times. By this comparison, we observed that the signal intensity of filament-forming wtRANTES varied and that it was present both at lower and higher levels than the signal of the mutants on a per cell basis. Even cells expressing wtRANTES at low levels (based on a very weak signal) showed a filamentous RANTES pattern similar to the wt by recording images at identical exposure times.
two-thirds and half the amount, respectively, of that measured for wt (Fig. 5D). Altogether, these data confirmed that the different patterns were not a result of various expression levels per cell or by the culture. Therefore, our findings clearly suggest that organization into filaments depends on the ability to form higher-order oligomers.

Next, to examine whether the mutant constructs of RANTES were present on the endothelial cell surface or restricted to intracellular compartments, we immunolabelled live HUVECs kept on ice and compared the pattern towards that of fixed and permeabilized cells. As shown in Fig. 5B (upper panel), the mutants Y3A and E26A distributed in a granular pattern throughout the HUVEC surface similar to that observed in permeabilized cells, suggesting that their binding to the surface was not disrupted. We also calculated the ratio between the number of cells expressing the mutant on the surface and the number of cells expressing the mutant after permeabilization (Fig. 5E). By this analysis we found that the ability of surface expression was not significantly affected for E26A or Y3A in comparison to the wt. In contrast, we found dramatically reduced surface expression for 44AANA47 and E66A.

RANTES does not co-localize with ICAM-1. Pre-existing tetraspanin-enriched microdomains (TEMs) containing adhesion receptors such as ICAM-1, VCAM-1, and CD44 are suggested to function as adhesive platforms on the endothelial cell surface35,36.
We wanted to examine whether RANTES localized in such platforms, using ICAM-1 as a marker. Consistent with previous findings, activation of HUVECs with TNFα or TNFα combined with IFNγ upregulated the ICAM-1 expression in all cells. Interestingly, the brightest cells had a speckled surface expression of ICAM-1 reminiscent of the pattern seen for RANTES. However, paired immunostaining for RANTES and ICAM-1 showed spots of overlapping signal but generally no co-localization (Fig. 6A). Previous studies have demonstrated an impressive redistribution of ICAM-1 during para- or transcellular leukocyte migration across the endothelium. ICAM-1 was enriched in vertical microvilli-like projections that embraced the leukocyte and drove redistribution of their integrins into linear tracks parallel to the direction of diapedesis. To elucidate whether RANTES distributed together with ICAM-1 into such a docking structure or transmigratory cup upon leukocyte addition, we reproduced the experiments by transfecting HUVECs to express RANTES, exposing them to TNFα, and adding peripheral blood mononuclear cells to such monolayers 20 min before fixation. While ICAM-1 was indeed observed in projections surrounding the leukocyte, the same projections were negative for RANTES (Fig. 6B). However, RANTES could be observed close to ICAM-1. RANTES was also observed at sites of docking or transmigration, but the strongest signal was typically observed in areas which were weak or negative for ICAM-1 (Fig. 6B, lower panel). Taken together, these experiments suggest that RANTES is not present on ICAM-1 positive mononuclear cells. RANTES-positive platelets could be observed in close proximity to

The RANTES filaments prevail flow forces. Because of the blood flow, molecules on the endothelial cell surface or at sites of vascular injury may be exposed to shear stress. To examine whether RANTES filaments can persist or will be disrupted by shear forces, we electroporated HUVECs with DNA encoding RANTES, cultured on cover slips before stimulation with TNFα + IFNγ for 30 h. RANTES was labeled with an anti-RANTES antibody (clone 21418), and the cover slips were mounted in a laminar flow chamber. The flow rate was adjusted to mimic vessel wall shear stress of 1 dyne/cm². The images were acquired by confocal microscopy before (left image, 0 min) and after 4 min with exposure to flow forces (right image, 4 min). HUVECs were treated as described in (A) except that they were not labeled with anti-RANTES antibody before exposure to flow. Human peripheral blood mononuclear cells were labeled with an anti-CD45 antibody, resuspended in medium and applied by a pump to the flow chamber (1 dyne/cm²). After 10 min, the cells were labeled with a rabbit anti-RANTES antibody. Images were acquired by sequential scanning confocal microscopy. Arrows and arrow heads indicate RANTES filaments and RANTES positive platelets, respectively. Original magnification in all panels, × 100. Scale bars, 10 µm.

**Figure 6** | RANTES shows distinct localization from that of ICAM-1. (A) Localization of RANTES compared to that of ICAM-1 after electroporation of HUVECs with plasmid DNA encoding wtRANTES and stimulation with TNFα before fixation, permeabilization and immunolabelling. Corner insets show high magnification of framed areas. Scale bar, 10 µm. (B) HUVECs were treated as indicated in (A) before incubation with MCP-1, followed by addition of peripheral blood mononuclear cells for 20 min followed by fixation, permeabilization, and immunolabelling. Corner insets show high magnification of framed areas, which are areas where one leukocyte has transmigrated. Arrows in the lower panel indicate two leukocytes that may have transmigrated. Original magnification in all panels, × 100. Scale bars, 10 µm. A rabbit anti-RANTES antibody was utilized in immunolabelling of RANTES. All samples were analyzed by use of sequential scanning confocal microscopy.

**Figure 7** | RANTES filaments are present after exposure to shear stress. (A) HUVECs were electroporated with DNA encoding RANTES, cultivated on cover slips before stimulation with TNFα + IFNγ for 30 h. RANTES was labeled with an anti-RANTES antibody (clone 21418), and the cover slips were mounted in a laminar flow chamber. The flow rate was adjusted to mimic vessel wall shear stress of 1 dyne/cm². The images were acquired by confocal microscopy before (left image, 0 min) and after 4 min with exposure to flow forces (right image, 4 min). (B) HUVECs were treated as described in (A) except that they were not labeled with anti-RANTES antibody before exposure to flow. Human peripheral blood mononuclear cells were labeled with an anti-CD45 antibody, resuspended in medium and applied by a pump to the flow chamber (1 dyne/cm²). After 10 min, the cells were labeled with a rabbit anti-RANTES antibody. Images were acquired by sequential scanning confocal microscopy. Arrows and arrow heads indicate RANTES filaments and RANTES positive platelets, respectively. Original magnification in all panels, × 100. Scale bars, 10 µm.
adhering mononuclear leukocytes, we could not observe RANTES-positive structures resembling a transmigratory cup (Fig. 7B). Taken together, the RANTES filaments with or without exposure to flow forces appeared to be similar, suggesting that they can be present in vivo.

**Discussion**

Several lines of evidence support a role for higher-order, oligomerized chemokines in leukocyte recruitment. First, wtRANTES but not disaggregated mutants, recruits leukocytes to the peritoneal cavity. Second, disaggregated mutants of RANTES fail to support leukocyte arrest to cultured endothelial cells under flow conditions, and some of these mutants are powerful anti-inflammatory agents. Taken together, these findings point to a possible role for oligomerized RANTES at the endothelial cell surface. Here we show for the first time morphological evidence that RANTES indeed binds to the endothelial cell surface in a regular, filamentous pattern. This depends on the oligomerization state of RANTES as all disaggregated mutants (identical or complementary to those tested under flow or in vivo) failed to appear as filaments on cultured endothelial cells.

Our observation that the mutant 44AANA did not form filaments (likely because it failed to bind the endothelial cell surface) fits well with its rapid appearance in peripheral blood after intraperitoneal injection, indicating that the mutant is not trapped in the tissue/extracellular matrix as efficiently as the wt and/or does not bind to the endothelial cell surface after abluminal to luminal transcytosis. This hypothesis would also be compatible with the independent observation that recombinant 44AANA-RANTES added to cultures of human microvascular endothelial cells does not bind to the cell surface. Taken together with the observation that the number of filaments was reduced after heparinase treatment, these data are consistent with the view that “RKNR” constitutes the principal GAG-binding site, and that binding to GAG is crucial for immobilization to the endothelial cell surface.

In our experiments, filament formation generated by the tetramer-restricted mutant E26A was reduced by more than 90% compared to the wt. Following injection into the mouse peritonitis model, E26A and wtRANTES appeared to be equally efficient in recruitment of cells, and it was concluded that the smallest leukocyte-recruiting units larger than dimers, the affinity for E66A could be lower than for the other constructs and may explain its reduced surface presentation.

Interestingly, Wang et al. proposed a model for how RANTES organizes into oligomers based on detailed structural analyses. The RANTES dimer is the building block, and long linear polymeric chains can form by contacts between residues of the second β-strand and residues at the C-terminal helix from one monomer of a dimer and similar residues in the neighbouring dimer. Consistent with the reduced ability of E26A and E66A to oligomerize and form filaments, both E26 and E66 appear to exert stabilization forces on the interaction between RANTES dimers. Wang et al. observed long oligomers at pH 7, and suggested that binding to sulfated GAGs would further promote length, as negatively charged GAG can neutralize electric repulsion forces between RANTES dimers. In our cell cultures, presence of RANTES filaments was at least partially dependent on heparan sulfate as there were fewer filaments following heparinase treatment. A crucial role for GAGs in promoting filament length, was supported by the finding that in the absence of cells, heparin was needed for filaments to appear.

During the initial events of leukocyte extravasation, membrane projections would probably increase the accessibility of the endothelium towards that of the leukocyte. Luminal endothelial membrane projections have been given names such as microvilli, microvilli, filopodia, protrusions, and nanotubes, and some of these are likely different structures with distinct functions. Similar to what was recently reported, we observed long membrane projections lateral to the endothelial cell surface as well as shorter membrane projections. Interestingly, Whitall et al. found that leukocytes interacted with both the long and short projections. By immunogold labeling of RANTES and electron microscopy, we observed a tendency of more labeling on the membrane projections than on the remaining plasma membrane. Unfortunately, the lower staining intensity obtained by immunogold labeling than by immunofluorescence, combined with the lost orientation of cells after scraping, did not allow us to draw firm conclusions on where the RANTES filaments localized at an ultrastructural level. Nevertheless, chemokines localized to membrane projections have been reported by others. For example, IL-8 immunoreactivity was detected at tenfold higher levels on vascular luminal projections than on the remaining plasma membrane. RANTES was also detected on such membrane projections.

Upon binding to leukocyte chemokine receptors, chemokines can trigger complex signaling transduction cascades leading to activation of integrins and, ultimately, to adhesion via binding to adhesion molecules such as ICAM-1. As this occurs rapidly and integrin activation is a reversible process, co-localization of RANTES and ICAM-1 could increase the possibility for the activated leukocyte to interact with ICAM-1. However, we were unable to observe overt co-localization both during the process of peripheral blood mononuclear cell extravasation and in the absence of leukocytes. By scanning electron microscopy we observed membrane projections resembling microvilli or small villi on the HUVEC surface. By immunogold labeling we observed membrane projections resembling microvilli or small villi on the HUVEC surface. (Oynebråten et al., unpublished data). This pattern was reminiscent of the ICAM-1 distribution observed after immunofluorescence, and our immunofluorescence data are very similar to what has been reported by others with ICAM-1 being present on microvilli or microvilli-like projections. Taken together, our data suggest that RANTES and ICAM-1 are present at different membrane sites.

We have stained for numerous chemokines after cytokine-stimulation or transfection of chemokine-encoding DNA into cultures of endothelial cells. In our study, the filaments were unique to RANTES, and due to high positive charge, it is proposed that RANTES is hindered from forming the more common organization of chemokine high order oligomers, i.e., globular complexes. However, it cannot be excluded that other chemokines than those we tested, for example MIP-1α/CCL3 and MIP-1β/CCL4, can form such elongated structures, although...
their charge and their binding to GAGs differ from that of RANTES'8,9. RANTES organized in different structures depending on the conditions in the well (presence of cells, heparin, conditioned media). Therefore, our data suggest that RANTES binding molecules are crucial for the organization of RANTES and its presentation to chemokine receptors. Given that filaments are formed in vivo, our data imply that presence of filaments may vary between types of endothelial cells, the tissue site, and inflammatory status.

In conclusion, our data together with biochemical analyses strongly suggest that RANTES filaments can form and be present at physiological conditions in vivo. Moreover, that filaments of RANTES can be of functional importance is supported by a study that shows reduced ability of the RANTES tetramer E26A to arrest monocytes on endothelial monolayers22. What might be the advantage of presenting RANTES in long filaments on the vascular surface? Clearly, the inherent ability to focus low numbers of molecules for presentation in a patch of concentrated chemokine to rolling leukocytes appears intuitively pleasing. Additionally, a closely related function of RANTES filaments could be to increase the accessibility of the ligand towards the receptor. In fact, Proudfoot et al. suggested that oligomerization of chemokines might be important for those whose GAG binding sites overlap with the receptor binding sites, as is the case for RANTES, MCP-1 and MIP-1αβ so that while some chemokine subunits bind to GAGs others can be exposed to the receptor1. The ability to organize into various forms from monomers to long filaments may also generate functionally distinct ligands11,12,22,30. Moreover, it is well documented that chemokines exert effects through dimers or oligomers of G-protein coupled receptors. Although the general view is that the receptor dimerizes shortly after synthesis in the endoplasmic reticulum43,44, a possible function of the RANTES filaments could be to facilitate ligand binding of two or more receptors. This could increase the number of integrins that are activated and the time for which a leukocyte presents activated integrins, and thereby promote the probability of leukocyte interaction with adhesion molecules and subsequent arrest to the endothelium.

**Methods**

**Antibodies and reagents.** Fetal bovine serum (FBS), gentamicin, fungizone, L-glutamine, MCDB 131, and Opti-MEM I were purchased from Life Technologies (Paisley, UK), and trypsin-EDTA was from Bio-Whittaker (Walkersville, MD). Recombinant human TNFa and IFNγ, recombinant human epidermal growth factor (EGF), recombinant human basic fibroblast growth factor (bFGF), recombinant human MCP-1, mouse and goat anti-human RANTES antibodies (MAB678, clone 21418, and B4E278, respectively) were purchased from R&D Systems (Abingdon, UK). A second mouse anti-human RANTES antibody was a kind gift from M. Sticherling (Klinikum der Christian-Albrechts-Universität zu Kiel, Germany), and mouse anti-human RANTES clone VL1 was from Biosource (Camarillo, CA). Rabbit anti-human RANTES (500–356) was from Peprotech (Rocky Hill, NJ), and mouse anti-heparan sulfate, clone 10E4, and hyaluronan binding protein was from Sekigakaku Corporation (Tokyo, Japan). The secondary rabbit anti-goat antibody used for electron microscopy was from DAKO (Glostrup, Denmark). The fluorescent (alexa 488 or 594) secondary anti-mouse or anti-rabbit antibodies were from Molecular Probes (Poortgebouw, The Netherlands), fluorescein ulex europaeus agglutinin I, and biotinylated horse anti-mouse IgG from Vector Laboratories (Burlingame, CA), streptavidin-Cy2 from Amersham Pharmacia Biotech (Piscatway, NJ), and streptavidin-Cy3 from Jackson (West Grove, PA). Protein A coupled to gold particles of different sizes was purchased from Bio Rag (Heilbronn, Germany). Heparin was from LeoPharma (Ballenrup, Denmark). All other reagents were from Sigma Chemical (St Louis, MO).

**Constructions.** RANTES was amplified from cdNA derived from TNFa/IFNγ-activated HUVECs. Primers (forward 5’-CTCTCCCAAGCCTACATGAGGATTCT-3’ and reverse 5’-GAAACTGATGCATCTCCTCCGTAAGGTTCTAGTACT-3’) were designed to introduce HindIII and XbaI restriction sites (underlined, respectively), to clone the RANTES-encoding DNA fragment into the pcDNA3.1(+) vector (Invitrogen, Carlsbad, CA). Based on this construct, alnane exchange of selected amino acids was performed using the QuickChangeTM Site-Directed Mutagenesis Kit (Stratagene, La Jolla, CA) according to the instructions by the manufacturer. The following primers were used to introduce the mutations, giving the sequence of the sense primers with mutated nucleotides written with small letters: Y3A: 5’-CTCTCCCAAGCCTACATGAGGATTCTAGTACT-3’ (vector). HUVECs were used for transfection by electroporation using 20 μg DNA according to the protocol 0394 from BTX (Holliston, MA). Following transfection, the cells were cultivated for approximately 24 h before fixation for examination by immunofluorescence microscopy or for 48 h before incubation with a mixture of heparinase I, II, and III (0.5 U/ml) for 2 h at 37°C and subsequently fixed as described above.

**Electroporation.** HUVECs were trypsinized, washed and resuspended in OptiMEM I containing 2.5% FBS, before transfection by electroporation using 20 μg DNA according to the protocol 0394 from BTX (Holliston, MA). Following electroporation, the cells were cultivated for approximately 24 h before fixation or staining of cell surface associated RANTES.

**Dose of heparan sulfate.** To examine whether RANTES was bound to the surface in a GAG dependent manner, monolayers of HUVECs were stimulated with 10 ng/ml TNFa combined with 1 ng/ml IFNγ for 48 h before incubation with a mixture of heparinase I, II, and III (0.5 U/ml) for 2 h at 37°C and subsequently fixed as described above. Sterile testicular hyaluronidase (1.0 mg/ml stock in DMEM; type IV; Sigma) or Streptomyces hyaluronidase [100 turbidity reducing units (TRU)/ml stock in DMEM; Seikagaku] was added directly to cells to yield a final concentration of 20, 50, or 100 μg/ml and 20 TRU/ml, respectively, and incubation continued for 3 h at 37°C in a 5% CO2-containing atmosphere.

**Adhesion of peripheral blood mononuclear cells.** HUVECs were electroporated with wtRANTES and cultivated on gelatine-coated chamberslides (Lab-Tek) for approximately 24 h before addition of 100 ng/ml TNFa (final concentration). After 12 h of stimulation, the cells were cultivated with 200 ng/ml recombinant human MCP-1 for 20 min. The cultures were then washed and peripheral blood mononuclear cells were added to the HUVECs 20 min before fixation. Mononuclear cells were isolated by centrifugation on Lymphoprep according to instructions by the manufacturer (Nycomed, Oslo, Norway).

**Immunostaining protocols and fluorescence microscopy.** HUVECs were cultivated on gelatine (1%/w/v) coated 10 × 10 mm glass coverslips or Lab-Tek chamber slides (Nunc, Roskilde, Denmark). For labelling of permeabilized HUVECs, cells were fixed in 4% paraformaldehyde for 10–15 min before washing in phosphate-buffered saline (PBS). For immunostaining, the fixed monolayers were permeabilized with 0.05% saponin before incubation with the antibodies, alternatively, 0.1% saponin was included in all solutions. In another set of experiments, recombinant RANTES was added to gelatine (1%/w/v) coated chamber slides in the absence of HUVECs for 35 h. RANTES was incubated in endothelial cell growth medium MCDB 131 without FBS or in conditioned MCDB 131 with FBS. Alternatively, heparin (LeoPharma, Ballerup, Denmark) was added in different concentrations before fixation. After secondary antibody labelling with primary antibodies and secondary reagents, the slides were mounted in mowiol combined with DABCO or polyvinyl alcohol. Labelling of cell surface-associated RANTES was performed on ice with cold (4°C) antibody solution. The primary biotinylated antibody was added for 45 min before fixation in paraformaldehyde followed by washing in PBS and sequential labelling with streptavidin-Cy3 or anti-rabbit Cy2. The immunostained cells were examined by an Axiphot 2 imaging Zeiss microscope using Plan-NEOFLUAR 40× and 100× oil objectives or a confocal laser scanning microscope (Leica TCS, Heidelberg, Germany) with A Plan apochromat 100×/1.4 oil objective equipped with an Ax (488 nm) and a He/Ne (543 and 633 nm) laser. Cells that expressed two or more RANTES filaments, were defined as filament-forming cells in the experiments shown in Fig. 5.

**Cryo-electron microscopy and immunogold labelling.** HUVECs were grown in 10 cm diameter culture dishes and cytokine-stimulated for 36 h before fixation in 0.1 M PBS containing 4% paraformaldehyde alone or a combination of 0.1% gluteraldehyde and 4% paraformaldehyde for 3 h at room temperature. After washing in 1 × PBS, cells were scraped-off and spun down. Cell pellets were embedded in 1 × PBS/12% gelatin and after infiltration with 2.3 M sucrose overnight at 4°C, cut into small blocks, mounted on pins and frozen in liquid nitrogen. Ultrathin cryosections of about 60–70 mm thickness were obtained by cutting at −80°C C with a Reichert Ultrotat ultramicrotome from Leica (Heidelberg, Germany). Cryosections were picked up in a 1 : 1 mixture of 2% methylcellulose and 2.3 M sucrose. Cryosections were then sequentially incubated with the goat anti-human RANTES antibody, the rabbit anti-gold antibody, and protein A-gold particles diluted in 1 × PBS/1% BSA for 30 min at room temperature with extensive washing between the incubations. (In an alternative immunolabeling protocol, clone 1D2/A12 was utilized.) Finally, the cryosections were contrasted with a 1 : 3 mixture of 3% uranyl-acetate and 2% methylcellulose before examination. When quantifying the distribution of RANTES, 30 pictures were utilized and gold particles associated with membrane projections versus the remaining plasma membrane were counted.
RANTES filaments at flow conditions. Glass coverslips were coated with gelatine (1% w/v). HUVECs were added and cultivated at standard conditions for 24 h before stimulation for approximately 40 h with TNFα (10 ng/ml) and IFNγ (1 ng/ml). Next, the cover slips were mounted in a flow chamber which was placed on a microscope stage for live imaging. The stage was enclosed by an incubator with temperature 37°C and CO2 adjusted to 6%. By use of a pump, medium with temperature 37°C was applied to the flow chamber corresponding to shear stress 1 dyn/cm². The flow rate was calculated by use of the formula T = 3μQ/2ba², where T = wall shear stress, μ = coefficient of viscosity (0.7 centipoise), Q = volumetric flow rate (cm³/s), a = half channel height (127 × 10⁻⁶ cm), and b = channel width (0.8 cm). Confocal images were acquired using an Olympus Fluoview1000 inverted microscope with a PlanApo × 60/1.42 objective (Olympus, Hamburg, Germany). ImageJ (NIH, Bethesda, MD, USA) and Adobe Photoshop (Adobe systems, San Jose, CA, USA) were used to process and prepare the images.

Statistical analyses. Statistical analyses were performed by use of GraphPad Prism version 6.04. Responses for each experimental group are presented as means with SEM. Differences between groups were analyzed by using one-way ANOVA with the Sidák method for multiple comparisons. p-values < 0.05 were considered significant.

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Author contributions
I.O., N.B., T.B., A.M.K. performed experiments; I.O. and N.B. prepared figures, all authors analyzed data; G.H. initiated the study; I.O., N.B., O.B. and G.H. designed the study, I.O. and G.H. wrote the manuscript.

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