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1 Title: Involvement of an IgE/Mast cell/B cell amplification loop in

2 abdominal aortic aneurysm progression

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1 Abstract

2

IgE type immunoglobulins and their specific effector cells, mast cells (MCs), are associated with abdominal aortic aneurysm (AAA) progression. In parallel, immunoglobulin-producing B cells, organised in tertiary lymphoid organs (TLOs) within the aortic wall, have also been linked to aneurysmal progression. We aimed at investigating the potential role and mechanism linking local MCs, TLO B cells, and IgE production in aneurysmal progression.

8 Through histological assays conducted on human surgical samples from AAA patients, we 9 uncovered that activated MCs were enriched at sites of unhealed haematomas, due to 10 subclinical aortic wall fissuring, in close proximity to adventitial IgE+ TLO B cells. 11 Remarkably, *in vitro* the IgEs deriving from these samples enhanced MC production of IL-4, a 12 cytokine which favors IgE class-switching and production by B cells. Finally, the role of MCs 13 in aneurysmal progression was further analysed *in vivo* in ApoE^{-/-} mice subjected to angiotensin 14 II infusion aneurysm model, through MC-specific depletion after the establishment of 15 dissecting aneurysms. MC-specific depletion improved intramural haematoma healing and 16 reduced aneurysmal progression.

Our data suggest that MC located close to aortic wall fissures are activated by adventitial TLO B cell-produced IgEs and participate to their own activation by providing support for further IgE synthesis through IL-4 production. By preventing prompt repair of aortic subclinical fissures, such a runaway MC activation loop could precipitate aneurysmal progression, suggesting that MC-targeting treatments may represent an interesting adjunctive therapy for reducing AAA progression.

Keywords: Abdominal aortic aneurysm, mast cells, IgE antibodies, tertiary lymphoid organs,
B lymphocytes, aortic dissection.

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1 Abbreviations

- 2 AAA = abdominal aortic aneurysm; AngII = Angiotensin II; ApoE = apolipoprotein E; DT =
- 3 diphtheria toxin; MC = mast cell; NAA = non-aneurysmal aortas; RMB = red mast cells and
- 4 basophil; TLO = tertiary lymphoid organ.

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1 Introduction

Arteries are subjected to recurrent mechanical insults which rise from the luminal side of vessels (Caligiuri et al., 2019). Vessels have intrinsic capacities to ensure prompt healing of local injuries and are assisted in this task by resident and recruited inflammatory cells (Dutertre et al., 2014). However, the persistence of vascular inflammation can eventually amplify the arterial damage and lead to severe and life-threatening conditions including coronary artery disease, strokes or abdominal aortic aneurysms (AAAs) (Libby et al., 2018). Promoting arterial healing by targeting inflammation thus constitutes a major challenge in modern medicine.

9 In this study, we focused on AAA progression. Monitoring AAA enlargement is problematic 10 due to the discontinuous, so-called 'staccato' growth where month-lasting no-growth/healing 11 periods can be succeeded by a sudden enlargement and again a no-growth/healing period 12 (Kurvers et al., 2004). So far, no pharmacological treatments have been identified to reduce or 13 stop AAA expansion and subsequent arterial rupture, which causes up to 200,000 deaths 14 worldwide each year (Sakalihasan et al., 2018). The only therapeutic option when the aneurysm 15 diameter exceeds a certain value (55mm in men, 50mm in women) is aorta surgery. Therefore, 16 deciphering the molecular pathways involved in AAA healing and/or progression is essential 17 to set up pharmacological alternatives.

18 Chronic immune responses involve adaptive and innate immunity and their relationship. In 19 particular, we and others have observed that chronic immune stimulation in AAAs leads to local 20 tertiary lymphoid organ (TLO) development within the adventitia (Clement et al., 2015; Guedj 21 et al., 2014; Mohanta et al., 2014; Nicoletti et al., 2013). TLOs contain germinal centre B cells 22 corresponding to B cells undergoing differentiation into plasma cells (Clement et al., 2015). 23 We have reported that TLO development is associated with increased levels of antibodies in 24 the adventitia of human AAAs, in particular IgEs (Clement et al., 2015), which are also

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involved in chronic inflammation. Notably, increase in IgE blood concentration correlates with
 the progression of AAAs and other arterial diseases (Guo et al., 2016; Tuleta et al., 2017). These
 observations raised the possibility that IgEs are produced locally within the arterial wall.

In parallel, triggered by the binding of IgE at their surface, activated mast cells (MCs) release 4 5 potent proteases, cytokines and vasoactive molecules, such as leukotriene and histamine, 6 directly or indirectly favouring the detersion of the extracellular matrix and a massive loss of the contingent of medial smooth muscle cells, that could eventually lead to aneurysmal 7 8 expansion (Sun et al., 2007; Sun et al., 2009; Swedenborg et al., 2011). Interestingly, MC 9 number increases in diseased arteries (Kaartinen et al., 1994; Tsuruda et al., 2008). 10 Furthermore, MCs can promote B cell effector functions (Merluzzi et al., 2010). As a 11 consequence, MCs have been surmised to be detrimental in AAAs (Furubayashi et al., 2007; 12 Sun et al., 2007; Sun et al., 2009). Studies aiming at inhibiting MC activation with pemirolast, 13 a histamine H1 antagonist, were inconclusive regarding their effect on AAA reduction (Sillesen 14 et al., 2015). However, pemirolast cannot uncouple IgE-mediated effector processes, which are 15 by far the most potent and specific trigger of MC activation. Hence, to date, the role of MCs in aneurysmal progression remains elusive. 16

Here, we aimed at investigating the potential mechanism linking MC activity to IgEs and TLO
B cell in aneurysmal progression. To do so, we performed histological and *in vitro* assays using
human surgical samples from AAA patients. Furthermore, the putative pathogenic role of MCs
in aneurysmal growth was evaluated *in vivo*, by inducing MC depletion in Red Mast cell and
Basophil (RMB) - ApoE^{-/-} mice subjected to angiotensin II infusion during the phase of
aneurysm progression.

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1 **Results**

IgE producing-TLOs are associated with aneurysmal wall unhealed fissures in patients

4 Tissue infiltrated IgE were enriched in aneurysmal samples and soluble IgE plasma levels were elevated in AAA patients compared to control non-aneurysmal aortas (NAAs, Figure 1-figure 5 6 supplement 1). In parallel, TLOs were found in a majority of AAA samples (20 out of 25, 7 Figure 1A) whereas they were absent in all NAA, in agreement with previous findings (Clement et al., 2015; Guedj et al., 2014; Mohanta et al., 2014; Nicoletti et al., 2013). In particular, 8 9 adventitial TLOs were consistently found in samples from patients with larger aneurysms 10 (Figure 1B), suggesting their higher frequency in these patients. In these adventitial TLOs, the 11 proportion of germinal centre B cells was dramatically increased (Figure 1-figure supplement 12 2), suggesting that adventitial TLO B cells in large aneurysms are engaged into an Ab-13 producing program from which diverse Ig isotypes could emerge. Interestingly, we observed a strong staining for IgEs within the TLO "light zone" (where the centrocytes, i.e. B cells 14 15 differentiating in immunoglobulins-producing cells, are located (Pipi et al., 2018); Figure 1A) 16 in 45% (n=9) of the samples. Altogether these results suggest that germinal centre B cells within 17 adventitial TLOs situated within large AAAs may actively produce IgE immunoglobulins.

Intriguingly, combined histological analysis of Carstair's, Perl's + DAB and Orcein staining revealed the presence of recent or past wall fissuring with haematoma formation in the media in 92% AAA samples (Figure 1C, Figure 1–figure supplement 3, and Table 1). Indeed, 84% of total AAA tissues (n=21) presented an accumulation of red blood cells (Figure 1C), reflecting the recent occurrence of intramural haematomas. Of note, the diameter of the aneurysmal samples was significantly larger than 3 cm, implying a greater tension and mechanical stress, than the one of the non-aneurysmal aortas (diameter < 3 cm) (Sakalihasan et al., 2018) and an

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1 attentive evaluation of pre-surgery tomography angiograms consistently revealed the presence 2 of at least one detectable aneurysmal wall macroscopic fissure, with radiologic contrast penetrating via blood disruptions from the aortic lumen to the aortic wall through the 3 4 intraluminal thrombus, in 78% of patients with intramural haematomas (Figure 1D and Figure 5 1-figure supplement 3D). In some samples, we were able to observe the entry site of blood 6 from the lumen on AAA histological sections, suggesting that the fissures leading to intramural 7 haematomas were provoked by tears (micro-fissures) initiated from the aorta lumen (Figure 1-8 figure supplement 4A-B and 4D-E). Unhealed intramural haematomas could also be evidenced 9 as large areas of ferrous iron deposits next to modest red blood cell accumulation in the media 10 (Figure 1-figure supplement 3B) in 8% (n=2) of samples. Importantly, IgE+ TLOs were 11 frequently localized near these intramural haematomas (Figure 2A-C), suggesting that the maturation of B cells towards IgE-producing cells within adventitial TLOs and the occurrence 12 13 of micro-fissures are linked.

14

MCs are enriched at sites of aneurysmal wall fissures and IgE-producing TLOs in human AAAs

17 In parallel, we observed that the adventitia of AAA patients was significantly enriched in MCs compared to the adventitia of NAA organ donors (Figure 2E-F and Figure 2-figure supplement 18 19 1), in agreement with previous observations (Tsuruda et al., 2008). Moreover, MCs were often 20 located next to IgE+ TLOs (Figure 2B-C), and extracellular tryptase+ granules reflecting recent 21 MC activation could be observed around these MCs by immunostaining on AAA sections (Figure 2G-H). Indeed, as compared to NAA samples, adventitial MCs in AAA samples were 22 23 degranulating, as documented by the expression of the degranulation/MC activation marker CD107a analysed by flow cytometry (Figure 2E-F). Strikingly, MCs enriched in AAA tissues 24

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were consistently found at sites of haematomas on AAA sections (Figure 2D and Figure 1–
 figure supplement 4C and 4F) where their granule enzymes could favour arterial wall rupture.
 Altogether, these results suggest that MCs and IgE+ TLOs could concur and be linked together
 with the occurrence of arterial wall injuries and aneurysmal progression.

5

6 Soluble molecules from human AAA adventitia induce MC degranulation and IgE-

7 dependent production of IL-4

8 Given that MCs displayed an activated phenotype only in the adventitia of AAA patients 9 (Figure 2), we assessed the putative presence of MC activation triggers in the conditioned 10 medium prepared from the adventitia of AAA as compared to control NAA tissues. We 11 observed that conditioned medium from AAA adventitias induced more surface expression of the degranulating marker CD63 and more IL-4 mRNA transcription than conditioned medium 12 13 from NAA adventitia in cultured ROSA human MCs (Figure 3A-B). This is important because 14 the cytokine IL-4 is required to carry out an IgE antibody class switch recombination and hence 15 allow the production of IgE by B cells (Saunders et al., 2019). Therefore, these results point at 16 an unforeseen function of MCs in AAAs, whereby MCs could orientate the local adaptive 17 immune response generated in adventitial TLOs toward IgE production. In turn, the binding of 18 adventitial IgE on their high affinity receptor on MCs could be responsible for triggering MC 19 activation. To directly assess this hypothesis, we pre-incubated the conditioned medium with anti-IgE DARPin[®] protein bi53 79 (5 µM) before adding it to MCs (Figure 3C-D). DARPin[®] 20 21 protein bi53 79 is a soluble molecule that specifically binds to the IgE-Fc portion thereby preventing IgE binding to FccRI (Eggel et al., 2014). Whereas treatment of conditioned medium 22 23 with bi53 79 did not prevent MC degranulation (Figure 3C), it decreased IL-4 mRNA production compared to untreated conditioned medium from AAA tissues (Figure 3D). IgEs 24

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contained in the conditioned medium from AAA adventitias therefore emerge as instrumental
 in promoting MC IL-4 production.

To assess if IL-4 production, MCs, and IgEs are also linked *in vivo*, we measured IgE, tryptase, and IL-4 quantities in the conditioned medium from AAA adventitias. We found a statistically significant positive correlation between the concentration of IgEs and IL-4 (p<0.001), as well as between the concentration of IL-4 and tryptase (p<0.05, Figure 3E-F). No such correlations were observed in the plasma of paired patients (Figure 3–figure supplement 1). These observations support the existence of a local amplification loop in the adventitia of AAAs involving IL-4 producing MCs and IgE-producing B cells.

10

11 MCs aggravates aneurysmal progression in mice

12 Previous studies have identified MCs as directly involved in provoking arterial damage 13 (Furubayashi et al., 2007; Sun et al., 2007; Sun et al., 2009). Consistently, the presence of 14 medial/adventitial MCs with an activated phenotype was associated to micro-fissures in our 15 human AAA samples, suggesting that MCs could play a role in the aneurysmal remodelling of the aortic wall, upon the occurrence of tissue fissuring. Therefore, we asked whether specific 16 depletion of MCs could prevent the aneurysmal remodelling in ApoE^{-/-} mice subjected to 17 18 chronic infusion of Angiotensin II (AngII), a well-known mouse model of aortic dissection 19 eventually followed by aneurysmal progression (Saraff et al., 2003). In this setting, 20 pseudoaneurysm of the abdominal aorta starts within 10 days upon the formation of large 21 intramural haematomas due to the occurrence of multiple medial tears originating from the 22 lumen, at the origin of the main side branches (Trachet et al., 2017). In order to conditionally 23 induce specific MC depletion by DT injection after the occurrence of the aortic fissuring, we 24 crossed ApoE^{-/-} mice with RMB mice (ApoE-RMB mice) (Dahdah et al., 2014). ApoE-RMB

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mice were subjected to infusion with AngII for 28 days. We injected DT (or PBS as a control)
14 days after the beginning of AngII infusion. Thus, MCs were depleted after the occurrence
of the dissection in this model (Saraff et al., 2003), allowing to evaluate the effect of their
depletion on the remodelling of the dissected aortas and their potential effect on aneurysmal
progression 14 days later (Figure 4A).

6 Survival was similar in the two groups (Table 2). As anticipated, the proportion of mice 7 presenting an intramural haematoma at day 28 was equivalent in DT-treated mice (44%, n= 8/18) and in the control group (47%, n=7/17). DT injection almost completely depleted MCs 8 9 from the aortic tissue (Figure 4B-C) and the peritoneal cavity (Figure 4–figure supplement 1). 10 MC depletion reduced the expansion of the pseudoaneurysm (Figure 4D-F), comprising the 11 size of the intramural haematoma and of the perivascular adventitial cuff (Figure 4-figure 12 supplement 2A-C). Furthermore, on the cross-sections of the dissected aorta segments stained 13 with picrosirius red we observed that the aortic wall of the pseudoaneurysm of DT-treated mice

presented an increased density of collagen compared to control mice (Figure 4E and 4G), especially in the adventitial fibrous cuff (Figure 4–figure supplement 2D). Our data therefore show that MC depletion improves the adventitia fibrotic remodelling which is critical for providing an adequate stiffness and strength subsequently constraining the expansion of the haematoma and the reduction of the aneurysmal progression (McGloughlin, 2011).

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1 **Discussion**

This study reveals the existence of a potential amplification loop involving IgE+ TLO B cells
and MCs at site of tissue fissuring, which could constitute an inflection point in AAA evolution,
precipitating aneurysmal progression (Figure 5).

5 To our knowledge, this is the first time IgE+ germinal centre B cells have been reported in 6 TLOs. Indeed, while they appear early in primary immune responses, coinciding with the peak 7 of IL-4 production, IgE+ cells are unable to populate the long-lived B cell compartment, which 8 explains why IgE+ germinal centres are rarely seen in lymphoid organs (He et al., 2013). Our 9 data show that the observed IgE+ TLOs develop close to aortic wall fissures. Strikingly, the 10 proximity of MCs and IgE+ TLOs at site of tissue injury in the wall of progressive aneurysms 11 supports a possible role for these highly reactive inflammatory cells in the progression of the 12 disease and in the maintenance of the pathologic loop, through their production of IL-4 in 13 response to locally produced IgEs, which in turn sustains the IgE+ germinal centre cells in local 14 TLOs. The correlation between IgE and IL-4 concentrations on one side, and IL-4 and tryptase 15 concentrations on the other side strongly supports the putative interrelated activities of MCs 16 and B cells. Furthermore, the ROSA human MC line is activated by purified IgEs only if these 17 are cross-linked by an anti-IgE antibody (Saleh et al., 2014). Importantly, we did not induce 18 such cross-linking with the conditioned medium. The fact that MCs were nonetheless activated 19 in such experimental conditions implies that the IgEs contained in the conditioned medium 20 were likely cross-linked by their cognate antigens. On the contrary, the fact that blood IgEs 21 failed to activate MCs (data not shown) indicates that they were not circulating as immune 22 complexes and hence strongly suggests that the antigen recognised by IgEs might be 23 specifically present in the aneurysmal aorta. Locally produced IgEs may also directly contribute 24 to AAA progression by promoting the senescence of smooth muscle cells (Guo et al., 2019).

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Whereas MC IL-4 production was dependent on IgEs present in the conditioned medium from aneurysmal aorta, degranulation was not. This indicates that besides IgEs, other molecules from the adventitia stimulate MCs. Indeed, other classes of immunoglobulins, which are also enriched in aneurysmal aortas (Clement et al., 2015), cytokines, DAMP-containing molecules and inflammatory mediators released by the injured vascular stroma could also trigger MC degranulation. Further studies focusing on the relationship between IgEs and MC activation could lead to novel therapeutic strategies to reduce chronic inflammatory disorders.

Our pre-clinical in vivo data directly support a role for MCs in the fate of aneurysmal 8 9 development of arterial wall subjected to fissuring and formation of an intramural haematoma. 10 Importantly, for reaching this conclusion, we used the ApoE-RMB mouse model in which the 11 depletion of MCs, conditioned by the use of DT, started only after the initial trigger (aortic 12 dissection). Instead, the rodent models previously used to address this question were 13 constitutively defective in MCs (Sun et al., 2007; Swedenborg et al., 2011; Tsuruda et al., 2008), implying that the deficiency of MCs preceded, and possibly biased, the experimental 14 intervention itself. Furthermore, constitutive MC-deficient models such as Kit^{W/Wv} and Kit^{W-sh/W-} 15 sh mice display other hematopoietic abnormalities, notably neutrophilia, thrombocytosis, and 16 17 macrophage defects (Dahdah et al., 2014; Nigrovic et al., 2008) which could be responsible for 18 some of the effects attributed to MCs. In addition, as resident MCs are present in healthy aortas, 19 constitutive MC depletion could impact aorta homeostasis as well as the 20 formation/progression/rupture of AAAs in these models. By triggering MC depletion after the occurrence of dissections in our ApoE^{-/-} RMB mouse model, we were able to specifically tackle 21 22 the role played by MCs during the healing processes associated with the progression of AAAs 23 in this model. We found that AAAs in MC-depleted animals displayed improved arterial remodelling with an increased collagen content of the aneurysmal wall, supporting an active 24 25 role for MCs in collagen degradation and expansion of AAA (Abdul-Hussien et al., 2007;

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Dobrin et al., 1984). Mechanistically, MC chymase and tryptase could increase collagen 1 2 degradation by converting collagenases from their inactive to their active form (Swedenborg et 3 al., 2011). MCs could also inhibit the production of collagen by smooth muscle cells (Wang et 4 al., 2001). Hence, MCs could actively contribute to weaken the arterial wall at sites of micro-5 fissures and favour the iterative expansion of aneurysms, under the biomechanic stress of 6 intraluminal thrombus (Davis et al., 2019; Wilson et al., 2013). Furthermore, the unhealed 7 micro-fissures could allow the accumulation of plasma proteins into the arterial wall, which is 8 associated to AAA growth (Behr Andersen et al., 2018).

9 Importantly, the experimental setting we used here is similar to the clinical settings of a recently 10 undertaken clinical trial investigating whether pemirolast, a histamine H1 antagonist acting as 11 a MC stabilizer in allergic conditions, could retard the growth of medium-sized AAAs (Sillesen 12 et al., 2015). This trial in which the major end-point was the change in aortic diameter as 13 assessed by ultra-sound imaging, failed to report a significant improvement in the patients 14 receiving the MC inhibitor. Although this study was elegantly designed and performed, it did 15 not however bring a definitive conclusion regarding MC involvement in AAA pathophysiology 16 because i/ the pemirolast treatment failed to decrease the plasma tryptase levels, indicating a 17 critical dosing issue, ii/ the study interval might have been too short, and iii/ contrary to our 18 study, the effect that MC inhibition might have exerted on tissue composition could not be 19 studied in this clinical study. Hence, in light of our results, we believe it would be interesting 20 to re-evaluate the effect of perimolast (and/or other MC-inhibiting drugs) on AAA growth and 21 rupture rate by targeting patients presenting elevated levels of IgEs and focusing on adventitial 22 remodelling. Inhibiting other key steps of the amplification loop, for instance by using anti-IgE 23 antibodies, could be another promising therapeutic strategy for such patients.

Altogether, our data suggest that locally produced IgEs complexed with lesion-specific antigens
 activate MCs and trigger IL-4 production, which subsequently promote TLO B cell IgE class

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- 1 switching. The runaway of this self-sustained local loop could drive repetitive MC
- 2 degranulation and thus constitute an inflection point in AAA evolution, preventing arterial
- 3 healing and precipitating AAA rupture (Figure 5).

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1 Materials and Methods

2 Mice

3 Apolipoprotein E-deficient (ApoE^{-/-}) mice (RRID:IMSR JAX:002052) were crossed to RMB (B6.Ms4a2^{tm1Mal}) mice (Dahdah et al., 2014). RMB mice carry an additive transgene containing 4 5 the promoter of the high affinity IgE receptor β subunit controlling the human diphtheria toxin 6 (DT) receptor and the tomato red fluorescent protein (Dahdah et al., 2014). The heterozygous offspring were then intercrossed to generate homozygous ApoE^{-/-} RMB mice. Mice were 7 8 maintained on a C57Bl/6J background and fed a regular show diet. Induction of AAAs, 9 histology and flow cytometry were performed as described below. All investigations on mice 10 conformed to the Directive 2010/63/EU of the European Parliament, and review and approval 11 of the study was obtained from the Comité d'Ethique Paris Nord #121 (APAFIS #12027).

12 Human samples

13 AAA tissues and blood were obtained from patients undergoing surgery and enrolled in the RESAA protocol (REflet Sanguin de l'évolutivité des Anévrysmes de l'aorte abdominale) 14 15 (Caligiuri et al., 2006). As expected (Sakalihasan et al., 2018), AAA patients were 16 predominantly men, and presented risk factors such as age, hypertension, hyperlipidaemia and 17 smoking (Table 3). All patients gave written informed consent, and the protocol was approved 18 by the Comité Consultatif de Protection des Personnes dans la Recherche Biomédicale 19 (CCPPRB, Paris-Cochin, approval no. 2095). Our study complies with the Declaration of 20 Helsinki. Control aortas were sampled from dead organ donors with the authorisation of the 21 French Biomedicine Agency (PFS 09-007, Table 4). Depending on their sizes, samples were 22 cut in several pieces that were used to prepare conditioned medium, and/or digested for flow 23 cytometry analysis, and/or fixed in 3.7% paraformaldehyde (PFA) for histological studies, and

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stored in the Inserm human CV biobank (BB-0033-00029), included in the European network
 BBMRI-ERIC.

3 Histology and immunofluorescence on human samples

4 PFA-fixed aortic tissues were paraffin-embedded. Four um-thick sections were deparaffinised 5 in toluene and rehydrated in ethanol. Sections were subjected to Carstair's stain, orcein stain or 6 Perl's + diaminobenzidine (DAB) stain. Alternatively, sections were incubated with retrieval 7 reagent (R&D Systems), then immunostained using antibodies against IgE (goat polyclonal, 8 Vector Laboratories), MC tryptase (rabbit monoclonal, clone EPR8476, abcam; or mouse 9 monoclonal, clone AA1, abcam), CD20 (mouse monoclonal, clone L26, Dako; or goat 10 polyclonal, Thermofisher), CD117 (rabbit polyclonal, Dako), or glycophorin A (rabbit 11 monoclonal, clone EPR8200, abcam). Immunostaining with isotypes were conducted to verify 12 the specificity of the primary antibodies. Overnight incubation at 4°C with primary antibodies 13 were followed by incubation with fluorophore-coupled anti-species antibodies (Jackson Immunoresearch) for one hour at room temperature. References for antibodies are shown in 14 15 Supplementary File 1. Nuclei were then stained with Hoechst 53542, and slides were mounted 16 with fluorescent mounting medium (ProLong Gold, ThermoFisher). Slides were kept in the 17 dark at 4°C. Blind analysis was performed for the presence of IgE+ TLOs, MCs and 18 haematomas.

19 Preparation of conditioned medium and immunodetection of soluble molecules

For the preparation of conditioned medium, the adventitia of AAA and control aortas were separated from the media and cut into small pieces (5 mm³). The samples were then incubated for 24 hours at 37°C in a standardised volume (6 mL/g of tissue) of RPMI 1640 medium (Gibco) supplemented with antibiotics and antimycotics. The conditioned medium was then centrifuged and the supernatant aliquoted and frozen at -80°C until use.

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The IL-4 concentration in conditioned medium from adventitial tissues and serum was analysed by CBA (BD biosciences). Bead fluorescence was recorded on a LSRII flow cytometer (BD). Tryptase was quantified by ELISA (USCN). Immunoglobulins were analysed using a bio-plex pro human isotyping assay (Bio-Rad) according to manufacturer instructions. Beads were analysed on a Bioplex-200 analyser (Bio-Rad). IL-4 and Tryptase were analysed on a different set of patients.

7 ROSA mast cell line

The ROSAKIT WT human cell line (RRID:CVCL 5G49) was cultured in the presence of 8 9 recombinant human SCF (80 ng/mL, R&D Systems), as described previously (Saleh et al., 10 2014). Cells were stimulated in the presence of conditioned medium from AAA and control 11 adventitia, or PMA (10ng/mL, Sigma-Aldrich) and ionomycin (1µM, Sigma-Aldrich) 12 (hereafter P+I condition) or IgE (2µg/mL, BeckmanCoulter) for 15 minutes followed by anti-13 IgE antibody (5µg/mL). After 1 hour of stimulation, the expression of CD63 in MCs was 14 analysed by flow cytometry. Alternatively, after 4 hours of stimulation, cells were collected 15 and dry pellets were frozen for gene expression analysis. In some experiments, 5µM DARPin[®] 16 protein bi53 79 (Eggel et al., 2014) was added to the conditioned medium. Experiment was 17 repeated twice with similar results.

18 Gene expression analysis

Total RNAs were extracted from ROSA MCs using the PureLink RNA mini kit (Invitrogen),
and mRNA reverse transcription was performed using iScript reverse transcriptase (Bio-Rad).
Real time quantitative PCR (RT-qPCR) was performed on a CFX 100 thermocycler (Bio-Rad)
using the following primers: forward IL-4: GCAGTTCTACAGCCACCAT; reverse IL-4:
ACTCTGGTTGGCTTCCTTCA . One nanogram of cDNA from each sample was mixed with
forward and reverse primers (300 nM) and SYBR Green master mix (Bio-Rad). The

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1 amplification program was as follows: A first step of initial denaturation at 95°C for 3 minutes, 2 then 40 cycles of 3 steps: denaturation at 95°C 15 seconds, annealing at 57°C for 15 seconds, and a final extension at 72°C for 30 seconds. The data were analysed using the $2^{-\Delta\Delta Ct}$ formulas: 3 the Ct values of IL-4 were normalised to the average Ct values of RPS18 (forward RPS18: 4 5 GCGGCGGAAAATAGCCTTTG; reverse RPS18: GATCACACGTTCCACCTCATC) and 6 ACTB (forward ACTB: TCCCTGGAGAAGAGCTACG; ACTB: reverse 7 TTTCGTGGATGCCACAGGAC) and non-treated MCs were used as reference.

8 Ang-II abdominal aortic pseudoaneurysms in mice

9 Thirty-eight 28-week-old ApoE-RMB males were used for the experiments, as males are more 10 susceptible to develop AAAs after Angiotensin II (AngII) infusion (Manning et al., 2002). 11 AngII (#A9525, 1 mg/kg/day, Sigma-Aldrich, St Louis, Missouri) was continuously infused 12 into the experimental mice via osmotic mini-pumps (Model 2004, Alzet, Charles River 13 Laboratories) that were surgically implanted subcutaneously in the interscapular region under 14 anaesthesia induced by intraperitoneal injection of ketamine (100 mg/kg) and xylazine (20 15 mg/kg). Surgery was followed by a buprenorphine intraperitoneal injection (0.1 mg/kg) for 16 analgesia. Three mice (8%) died within the first 10 days. Fourteen days after inducing AAAs 17 through AngII infusion (Saraff et al., 2003), mice received two intraperitoneal injection of DT 18 $(1\mu g/\text{ injection, n=18})$ at a one-day interval (or PBS in the control group, n=17) in order to 19 deplete MCs. Randomization was achieved by having mice from each experimental group in 20 each cage. Mice were sacrificed at day 28 by intracardiac exsanguination under overdose of 21 anaesthesia (intraperitoneal injection of 150 mg/kg ketamine-HCL and 30 mg/kg xylazine). 22 Before the exsanguination, a cell suspension from the peritoneum was obtained by peritoneal 23 lavage with 5 mL of ice-cold PBS. Blood was withdrawn from the right ventricle of the heart 24 and collected in EDTA tubes for blood cell analysis. The heart and aorta were dissected, 25 photographed and mounted on cryomolds for further histomorphological analysis based on

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cryosections. The experiment was repeated twice with similar results. Pooled number of mice
 from both experiments who survived and developed aneurysms in each group are summarized
 in Table 2.

Of note, in RMB mice, DT induces both MC and basophil depletion. While basophils fully repopulate the blood within 12 days, MCs repopulation of peripheral tissues such as skin or peritoneum is much slower (Dahdah et al., 2014; Ngo Nyekel et al., 2018). Consistently, two weeks after DT administration (Day 28 of AngII infusion), MCs were totally absent from the peritoneal cavity whereas the basophil population was totally replenished in the blood of ApoE-RMB mice (Figure 4–figure supplement 1). DT injection had no effect on blood basophils, peritoneal MCs, and aortic tissue MCs in control ApoE^{-/-} mice (data not shown).

11 Histology on mouse tissues

12 MC depletion was analysed on cryosections of mouse aortic roots stained with Toluidine Blue. 13 Images were captured on a Zeiss Axio Observer Z7 inverted microscope, and MCs were 14 counted manually. A detailed blind analysis of pseudoaneurysms was performed for mouse 15 abdominal aortas displaying macroscopic evidence of the occurrence of a pseudeoaneurysm. 16 Cross-cryosections (10 µm) of the aortic segments taken at different levels of the 17 pseudoaneurysms (every 300 µm) were stained for collagen with picrosirius red (Figure 4-18 figure supplement 2A). Images were acquired under polarised light. The size of the different 19 layers (media, haematoma, adventitial fibrous cap) and the extent of collagen deposition were 20 quantitatively assessed using Image J.

21 Image acquisition

Images were digitally captured using an AxioObserver epi-fluorescent microscope equipped
with a Colibri 7 LED generator (Zeiss) and an ApoTome system and running Zen Software

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(Zeiss). Macroscopic images of the human samples were acquired using the NanoZoomer
 Digital slide scanner (Hamamatsu Photonics).

3 Flow cytometry analysis

4 Mouse blood samples collected in EDTA were incubated in Ammonium-Chloride-Potassium 5 lysis buffer for 5 minutes at room temperature to lyse red blood cells. Cells from blood samples 6 or 1 mL of peritoneal lavage were stained for dead cells with Live/Dead fixable far red Dead 7 cell stain kit (Invitrogen) for 30 minutes at 4°C, then incubated with a purified rat anti-mouse 8 CD16/32 (FcBlock, BD Biosciences) for 15 minutes at 4°C. Cells were then incubated for 20 9 minutes at 4°C with the following antibodies (from BD Biosciences unless stated otherwise): 10 BV605 rat anti-mouse CD19 (clone 1D3), FITC rat anti-mouse CD117 (clone 2B8), PerCP rat 11 anti-mouse CD45 (clone 30-F11), APC hamster anti-mouse FccRIa (clone MAR-1, 12 Biolegend), AF700 rat anti-mouse CD3 (clone 172A). References for antibodies are shown in 13 Supplementary File 1.

14 For human tissue analysis, fresh adventitial layer samples were weighed, cut into small pieces (<1 mm) and digested using a previously described protocol (Fletcher et al., 2011). After a wash 15 16 step, the cells were stained for dead cells with Live/Dead fixable far red Dead cell stain kit 17 (Invitrogen) for 30 minutes at 4°C, then incubated for 20 minutes at 4°C with a combination of 18 the following mouse anti-human antibodies: AF488 anti-CD3 (clone UCHT1, BD Biosciences), PE anti-CD203c (clone NP4D6, Biolegend), PerCP anti-FcεRIα (clone AER-37, Biolegend), 19 20 BV421 anti-CD117 (clone 104D2, BD Biosciences), BV500 anti-CD45 (clone HI30, BD 21 Biosciences), PE-Cy7 anti-CD63 (clone H5C6, BD Biosciences), AF700 anti-CD19 (clone HIB19, BD Biosciences), AF647 anti-CD107a (clone H4A3, BD Biosciences), BV421 anti-22 23 CD27 (clone O323, Biolegend), BV570 anti-HLA-DR (clone L243, Biolegend), BV785 anti-24 CD19 (clone HIB19, Biolegend), APC anti-IgD (clone IA6-2, BD Biosciences), FITC anti-

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CD38 (clone HIT2, BD Biosciences), PE anti-CD45 (clone HI30, BD Biosciences), BV605
 anti-CD45 (clone HI30, BD Biosciences).

For mouse peritoneal cells and human adventitial cells, flow-count fluorospheres (Beckman Coulter) were added to the samples before the acquisition on the cytometer in order to calculate absolute count of cells. Data were acquired on a LSRII flow cytometer or an ARIA III cell sorter (BD Biosciences) and analysed using FACSDiva (BD Biosciences) and FlowJo (TreeStar) softwares.

8 Statistical analysis

9 Statistical analysis was performed using the JMP9 and Prism software. To compare two 10 populations, we used Student t-tests or non-parametric Mann-Whitney tests when sample size 11 was too small (n<10) or when values were not normally distributed. Paired values were 12 compared using Wilcoxon matched-pairs signed rank test. Pearson correlation coefficients or 13 p-values of chi-square contingency tests were calculated to assess correlation between 14 continuous or categorical variables, respectively. Quantitative data are expressed as mean +/-15 standard error. Source data with summary statistics can be found in source data files associated 16 to each figure.

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13

14 **Related interests**

A Eggel is a cofounder and scientific advisor of Excellergy, INC. and ATANIS Biotech AG.
M. Arock is on DSMB for AB Science and advisory board for Blueprint Medicines; receives
consulting fees and/or honoraries from AB Science, Blueprint Nedicines and Novartis; and
declares patent #WO2013064639A1 'Human mastocyte lines, preparation and uses. P Launay
is the CEO of Inatherys. Details can be found in ICMJE forms.

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1 Materials availability statement

- 2 Use of DARPin[®] protein bi53_79 and RMB mice requires a MTA. Figure 1-source data 1,
- 3 Figure 2-source data 1, Figure 3-source data 1, and Figure 4-source data 1, contain the
- 4 numerical data used to generate the figures.
- 5
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1 Figures

2 Figure 1. IgE-producing TLOs are present in the adventitia of human AAAs.

3 (A) Cross-sections of AAAs were stained for CD20 (magenta) and IgE (green). Dense DAPI⁺ 4 lymphoid aggregates highly enriched in CD20+ B cells correspond to TLOs. Several TLOs 5 displayed a strong IgE staining projecting in the CD20+ B cells comprised in the DAPI "light 6 zone" of TLOs (white arrowheads). The magnified inset shows 5 adventitial TLOs (cluster of 7 DAPI+ CD20+ B cells), 3 of which are IgE+. adv: adventitia, m: media, thr; thrombus. 8 Representative image of 9 different samples. Scale bar: 2.5 mm. (B) Maximum diameter size 9 of AAAs depending on the absence (no) or the presence of TLOs. p value (Mann-Whitney test) 10 is indicated on the plots. (C) Serial cross-sections of AAAs were coloured with Carstairs' 11 staining, where fibrin appears in bright red, collagen in blue, red blood cells in yellow and 12 nuclei in black; Perl's+DAB staining allows to visualise redox-active iron; Orcein staining 13 permits to detect elastin fibres. The lumen is at the bottom of each picture and the media (m) is indicated by the blue bar on the side of the pictures. Black arrowheads point at red blood cells 14 15 outside blood vessels, reminiscent of recent intramural haematomas. Scale bar: 1 mm. Right 16 panels: magnified insets. (D) Contrast-enhanced tomography angiograms of an AAA 17 displaying blood disruption from the aortic lumen to the aortic wall through the intraluminal 18 thrombus (white arrow).

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1 Figure 1-figure supplement 1. IgEs are elevated in the plasma and adventitia of

2 AAA patients.

IgEs were titrated in the plasma (A) and conditioned medium from adventitia (B) of NAA organ
donors and AAA patients. Mann-Whitney tests were used to compare groups. **: p<0.01; ***:
p < 0.001.

6

Figure 1-figure supplement 2. GC B cells and plasma cells are elevated in AAA adventitia.

9 Adventitia from NAA organ donors and AAA patients were digested and analysed by flow 10 cytometry after the addition of fluorescent count beads. (A) B cells were identified as singlet, autofluorescent⁻, live CD45⁺ CD19⁺ cells, and their number was calculated in each sample, 11 showing a statistically significant increase of B cells in AAA samples. ***, p < 0.001, Mann-12 13 Whitney test. (B) B cells were identified as in (A), and subsets were defined as follows: naïve B cells, IgD⁺ CD38⁻; plasma cells (PC), IgD⁻ CD38^{hi}; germinal centre B cells (GC), IgD⁻ CD38⁺ 14 HLA-DR^{hi}; memory B cells, IgD⁻ CD38⁻ HLA-DR⁺. Representative samples show that GC B 15 16 cells and plasma cells were present in the adventitia of AAA, while they were barely detected 17 in the matched blood of the AAA patient, or in the adventitia of NAA.

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19 Figure 1–figure supplement 3. AAAs with ancient or no intramural haematomas.

Histological staining of NAAs (A) and AAAs (B-C). Serial sections were stained with Carstairs's stain, Perl's+DAB stain and Orcein stain as in Figure 1C. The squares frame the localisation of the details shown in the insets. The lumen is at the bottom of each picture, and the media (m) is indicated by the blue bar on the side of the pictures. In A, intact elastin fibers

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can be seen in the media on the Orcein stain. In B, black arrows indicate iron deposits in the
absence of red blood cells, suggestive of an ancient intramural haematoma. An intraluminal
thrombus (*) can be seen in C, where no signs of intramural haematoma were seen. Scale bar:
1 mm. (D) Contrast-enhanced tomography angiograms of AAAs. White arrows: blood
disruption from the aortic lumen to the aortic wall through the intraluminal thrombus.

6

Figure 1-figure supplement 4. Proximity of micro-fissures and MC degranulation in human AAAs.

9 Serial sections of an AAA sample with a micro-fissure (A-C) and from a NAA sample (D-F) 10 were stained with: Carstair's stain (A and D); orcein to reveal elastin fibres (B and E); and 11 tryptase (red) and glycophorin A (gray) to detect MCs/tryptase⁺ granules and red blood cells, 12 respectively (C and F; the elastin fibres were autofluorescent: green). The pictures in B and C, 13 and E and F, correspond to the green insets in A and D, respectively. The entry site of blood 14 into the aortic wall in the AAA sample can be observed on the Carstair's stain (A, red blood 15 cells appear in yellow). The blood entry site visible on the Carstair's stain (A) and with the 16 glycophorin A stain (C) was associated with degraded elastin fibres (black arrow on orcein stain 17 in B) and the presence of MCs (white arrowhead) and tryptase⁺ granules (C). Note the normal 18 aspect of the elastin fibers (E) as well as the absence of tryptase and red blood cells (F) in the 19 control. Scale bars: A and D 2.5 mm; others: 200 µm.

20

21 Figure 1–source data 1.

This file contains the numerical data and summary statistics used to generate the graphspresented in Figure 1 and its supplements.

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Figure 2. IgE-producing TLOs, activated MCs and intramural haematomas are in close proximity in human AAAs.

3 (A) Cross-sections of AAAs were coloured with Perl's+DAB staining. TLOs (circled with 4 dashed line) are found at proximity of haematomas (arrowheads). Scale bar: 500 µm. (B) Consecutive sections were stained for CD20 (magenta), IgE (green), CD117 (red) and DAPI 5 6 (grey). B cells (CD20⁺ and DAPI⁺) appear in light pink, autofluorescent red blood cells (low 7 intensity in the CD20 channel and DAPI⁻) appear in magenta, and MCs (CD117⁺ IgE⁺) appear 8 in yellow. Scale bar: 250 µm. (C) Magnified inset highlighting the close proximity of IgE+ 9 TLOs (dashed circle), red blood cells (white arrowheads) and MCs (blue arrows), (D) 10 Magnified inset highlighting MCs (blue arrows) at proximity to intramural haematoma. (E-F) 11 Adventitia from NAA organ donors and AAA patients were digested and analysed by flow cytometry after the addition of fluorescent count beads. Singlet, autofluorescent, live CD45⁺ 12 13 cells were selected, and MCs were identified as CD3⁻ CD19⁻ FcsRIa⁺ CD117⁺ cells (E). The number of MCs (F, left) and activated (CD107a^{hi}) MCs (F, right) was calculated in each sample. 14 Error bars represent mean +/- standard error. **, p < 0.01; ***, p < 0.001, Mann-Whitney test. 15 16 (G-H) Cross-sections of AAAs were stained for CD20 (magenta), IgE (green), and tryptase 17 (red). MCs (tryptase⁺ IgE⁺) were detected around TLOs. Scale bar: 100 µm. The magnified inset (H) shows a degranulating tryptase⁺ MC at proximity of a TLO. 18

19

20 Figure 2–figure supplement 1. MCs accumulate in the adventitia of human AAAs.

Carstairs' stain (A) and toluidine blue stain (B-C) on serial sections of a representative microfissured AAA sample. MCs appear purple on toluidine blue stain (indicated with black circles).
TLO and indicated with black circles).

- 23 TLOs are circled with a dotted line. Scale bar: 500 $\mu m.$
- 24

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1 Figure 2–source data 1.

- 2 This file contains the numerical data and summary statistics used to generate the graphs
- 3 presented in Figure 2.
- 4

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Figure 3. MC degranulation and IgE-dependent IL-4 production in response to conditioned medium from AAA adventitia.

(A-D) ROSA MCs were cultured in the presence of conditioned medium from adventitia of 3 4 control organ donors (NAA) and AAA patients. DARPin® protein bi53 79 (5 µM) was added (+D) or not (-D) to inhibit IgE binding to MCs' FccRI (C-D). (A, C) After 1 hr, degranulation 5 6 (MFI CD63) was assessed by flow cytometry. The dotted line indicates CD63 MFI in nonstimulated cells. (B, D) After 4 hrs. mRNA IL-4 level was analysed by RT-PCR ($2^{-\Delta\Delta Ct}$. 7 8 normalised to RP18S and non-stimulated cells). *, p < 0.05 (A,B: Mann-Whitney test; C,D: Wilcoxon matched-pairs signed rank test). (E, F) Concentrations of IL-4, tryptase, and IgE were 9 measured in conditioned medium from AAA adventitia. *, p < 0.05; ***, p < 0.001 (Pearson 10 11 correlation analysis).

12

Figure 3-figure supplement 1. No correlation between tryptase, IgE and IL-4 in the plasma of AAA patients.

IL-4, tryptase, and IgE concentrations were measured in the serum of AAA patients (same
 patients as in Fig. 4D). r² and p-values from Pearson correlation analysis are indicated.

18 Figure 3–source data 1.

This file contains the numerical data and summary statistics used to generate the graphspresented in Figure 3 and its supplement.

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Figure 4. MCs depletion decreases aneurysmal growth after dissections in ApoE RMB mice.

(A) Experimental protocol, showing the normal course of pseudoaneurysm development in the 3 4 Ang II infusion model, and the presence or absence of MCs depending on the treatment (PBS 5 in blue or DT in red). (B) Toluidine blue stain on aortic root cross-sections of DT- and PBS-6 treated mice. The arrow points to one of the MCs detected in the aortic root adventitia of a PBS control mouse. Scale bar: 250 μ m. (C). Quantification of MCs in a rtic roots. ***, p < 0.001, 7 8 Mann-Whitney test. (D) Macroscopic images of the hearts and aortas of DT- and PBS-treated 9 mice displaying aneurysms (indicated by arrows). (E) Consecutive cross-sections stained with 10 Masson Trichrome (top, scale bar: 500 µm) or with Sirius red (bottom; polarised light). Aorta 11 outer diameter (F) and collagen density of the aortic wall (G) for mice presenting 12 pseudoaneurysms (PBS: n=7; DT; n=6) were calculated by computer-assisted morphometry on 13 cross-sections stained with Sirius red. Data from sections taken at different levels (every 300 14 µm) from each pseudoaneurysm were used for calculations (Supplemental Figure 8A), and 15 aligned to the layer with the largest haematoma. Mean +/- standard error; Mann-Whitney test, **: p < 0.01; ***: p < 0.001. 16

17

Figure 4–figure supplement 1. Repopulation of basophils in the blood and MCs in the peritoneum after DT depletion in ApoE RMB mice.

ApoE-RMB mice were treated as in Figure 4A. At day 14 after DT (n=9) or PBS (n=10) injection (day 28 of Ang II infusion), mice were sacrificed. (A) Blood basophils were identified among singlet cells, as CD45⁺ Live/Dead⁻ CD3⁻ CD19⁻ FccRIa⁺ Tomato⁺ cells (19). (B) Percentage of basophils within singlet cells, CD45⁺ Live/Dead⁻ cells. (C) Peritoneal MCs were identified among singlet cells as CD45⁺ Live/Dead⁻ CD3⁻ CD19⁻ FccRIa⁺ Tomato⁺ cells

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- CD117⁺ cells ²⁰. (D) Number of MCs per peritoneal lavage. ***: p<0.0001, Mann-Whitney test.
 Data are representative of two experiments.
- 3

4 Figure 4-figure supplement 2. MCs promote pseudoaneurysm expansion after

5 dissection in ApoE-RMB mice.

- 6 (A) Sections taken at different levels (every 300 μ m) from each pseudoaneurysm were used for 7 analysis. (B) Vascular wall layers were defined by computer-assisted morphometry on Sirius 8 red-stained cross-sections (Figure 4E) of pseudoaneurysm (scale bar: 500 μ m). Size (C) and 9 collagen density (D) of the different layers were calculated by computer-assisted morphometry, 10 and plotted relatively to the distance from the layer with the largest haematoma. Mann-Whitney 11 test, **: p < 0.01; ***: p < 0.001.
- 12

13 Figure 4–source data 1.

This file contains the numerical data and summary statistics used to generate the graphspresented in Figure 4 and its supplements.

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1 Figure 5. Schematic overview of the IgE/MC/B cell self-sustained loop in dissecting

2 **AAAs.**

Locally produced IgEs (and may be other isotype Igs) complexed with lesion-specific antigens activate MCs. Additional triggers from the arterial wall, such as pro-inflammatory cytokines and DAMPs, may contribute to MC activation/degranulation. Proteases and inflammatory mediators hence released from MCs increase the risk of micro-fissures, and MC-derived IL-4 promotes the synthesis of more IgEs by TLO B cells. These events lead to an amplification loop of the pathology.

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1 Tables

2 Table 1: Patients' characteristics according to the presence of intramural

3

haematomas.

	None	Ancient	Recent
	(n=2, 8%)	(n=2, 8%)	(n=21, 82%)
Age (years)	70 +/- 7	67+/- 1	71 +/- 2
Male (%)	100%	100%	90%
Max. aortic diameter (cm)	63 +/- 3	66 +/- 14	61 +/- 4
Diabetes (%)	0%	0%	7%
Hypertension (%)	100%	100%	87%
Hyperlipidaemia (%)	0%	50%	53%
Smoking (%)	50%	50%	93%

4

5

Values are mean +/- SEM or %. Ancient intramural haematomas are characterised by ferrous
iron deposits with few red blood cells in the media (Figure 1–figure supplement 3B) and recent
intramural haematomas are characterized by abundant red blood cells in the media (Figure 1C).

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Table 2: ApoE-RMB survival and pseudoaneurysm occurrence.

	PBS	DT	p-value	
	(n = 17)	(n = 18)		
Pseudoaneurysm	n = 7 (47%)	n = 8 (44%)	0.85	
Death [*]	$n = 0 (0\%^{\dagger})$	$n = 2 (25\%^{\dagger})$	0.15	

2

- 3 *, dead mice presented pseudoaneurysms and died 21 days after the beginning of AngII infusion
- 4 (7 days after DT treatment). † , % of mice with AAA.

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Table 3: Patients' characteristics.

	All	Histology	FC	СМ
	(n=57)	(n=25)	(n=25)	(n=37)
Age (yrs)	68 +/- 2	70 +/- 2	73 +/- 2	68 +/- 2
Male	86%	92%	92%	84%
Aneurysm location				
Suprarenal	2.5%	5%	0%	4%
Juxtarenal	2.5%	5%	0%	0%
Subrenal	59%	58%	83%	57%
Subrenal+iliac	36%	32%	17%	39%
Maximum aortic diameter (cm)	61 +/- 2	61 +/- 3	67 +/- 4	63 +/- 3
Clinical Features				
Diabetes	5%	5%	0%	7%
Hypertension	81%	89%	86%	87%
Hyperlipidaemia	40%	47%	58%	37%
Smoking	83%	84%	86%	80%
Anti-coagulants	15%	17%	27%	15%

2

Values are mean +/- SEM or %, for all AAA samples, or samples according to processing (some
samples were used for several kind of analysis). FC: flow cytometry analysis (MCs and/or B
cells); CM: conditioned medium (IgE concentration and/or MC stimulation).

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	All	Histology	FC	СМ
	(n=39)	(n=7)	(n=16)	(n=24)
Age (yrs)	55 +/- 3	53 +/- 9	55 +/- 5	56 +/- 4
Male	61%	80%	56%	61%
Atherosclerotic lesion				
None	28%	29%	31%	25%
Fatty streak	44%	43%	31%	54%
Fibrolipidic	26%	29%	31%	17%
Intraplaque hemorrhage	3%	0%	6%	4%

Table 4: NAA healthy donors' characteristics.

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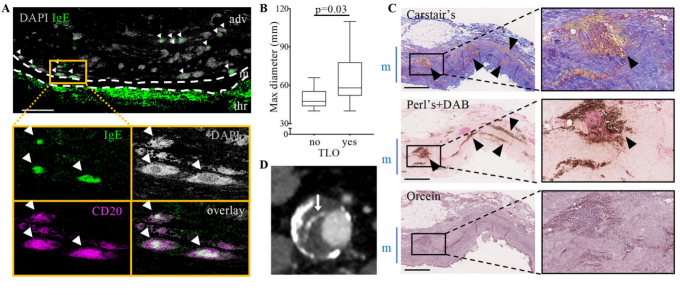
Values are mean +/- SEM or %, for all NAA samples, or samples according to processing (some
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cells); CM: conditioned medium (IgE concentration and/or mast cell stimulation).

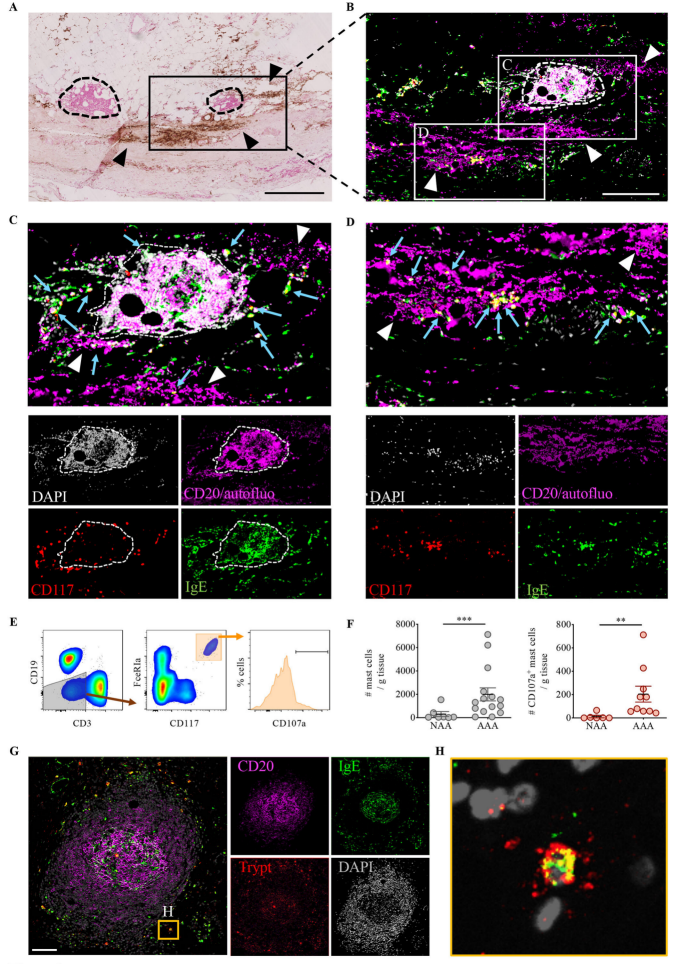
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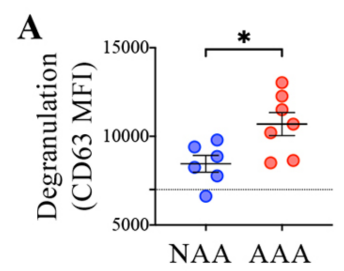
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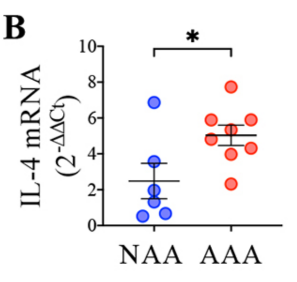
1 Additional files

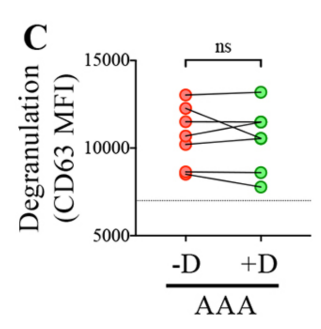
2 Supplementary file 1: References for antibodies used in this study.

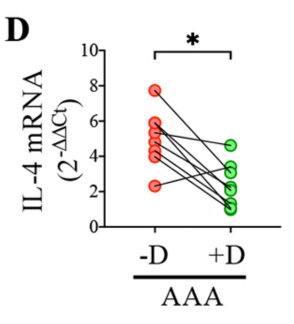


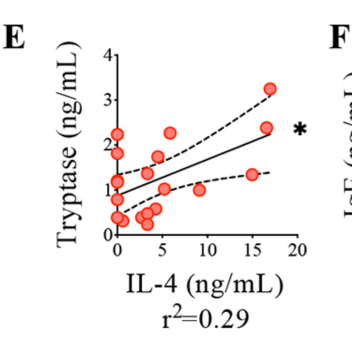


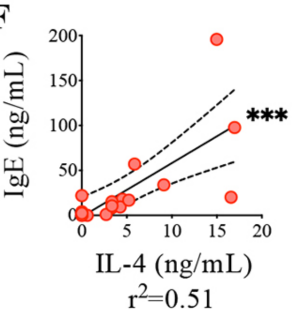


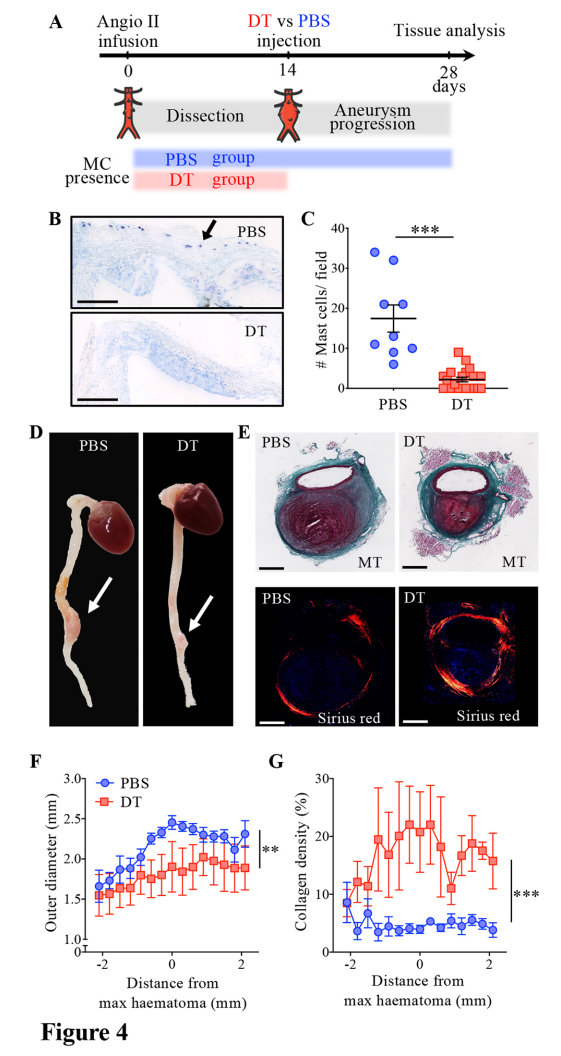


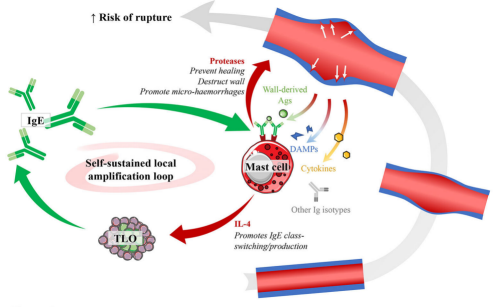












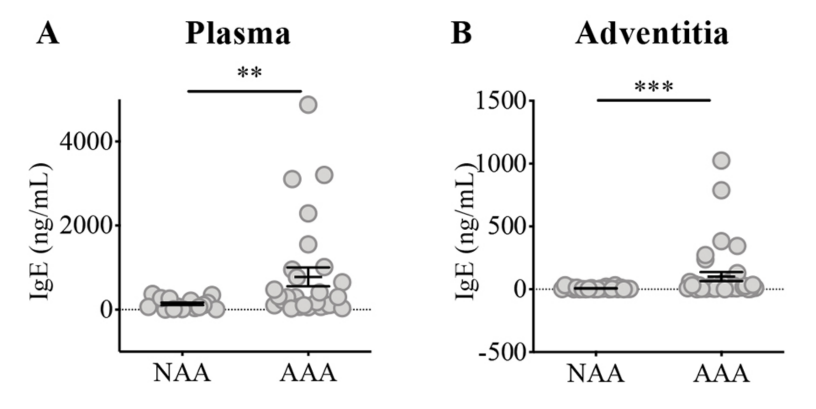
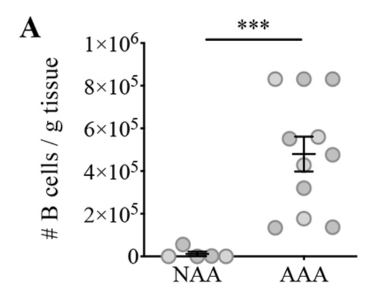


Figure 1-figure supplement 1



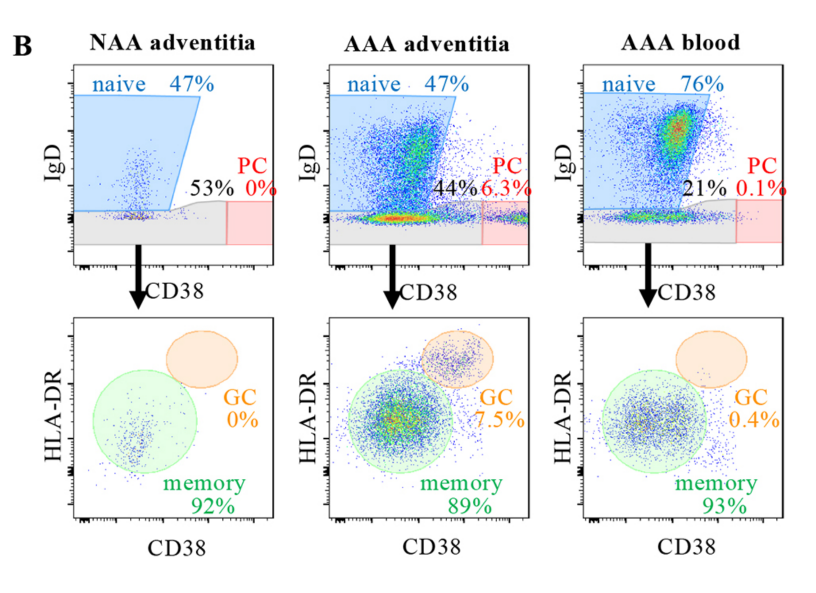


Figure 1-figure supplement 2

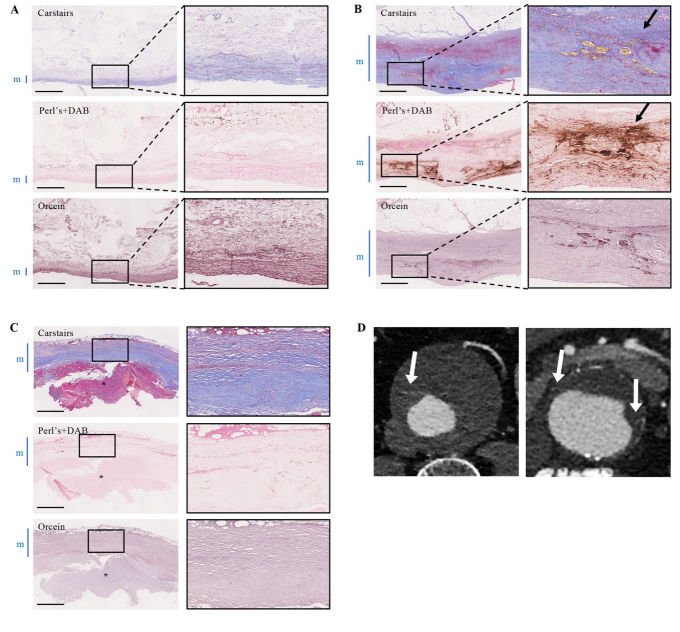


Figure 1-figure supplement 3

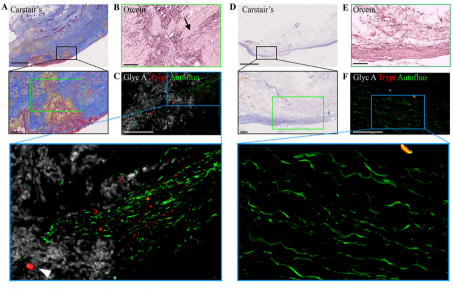


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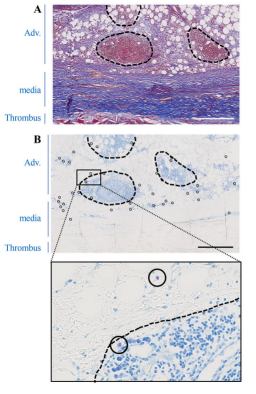


Figure 2-figure supplement 1

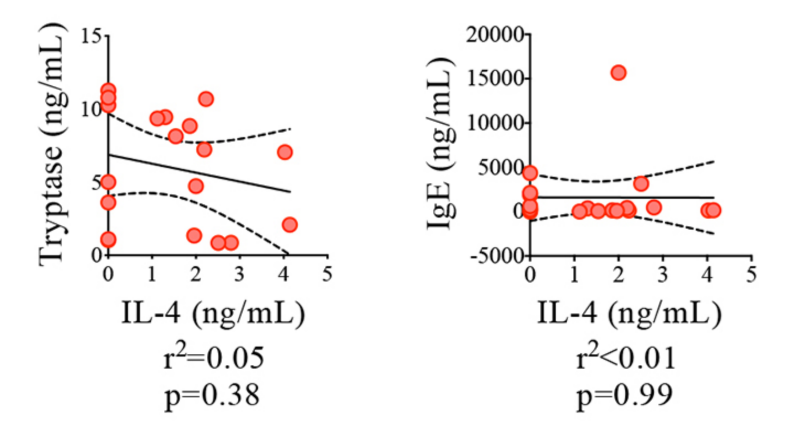


Figure 3-figure supplement 1

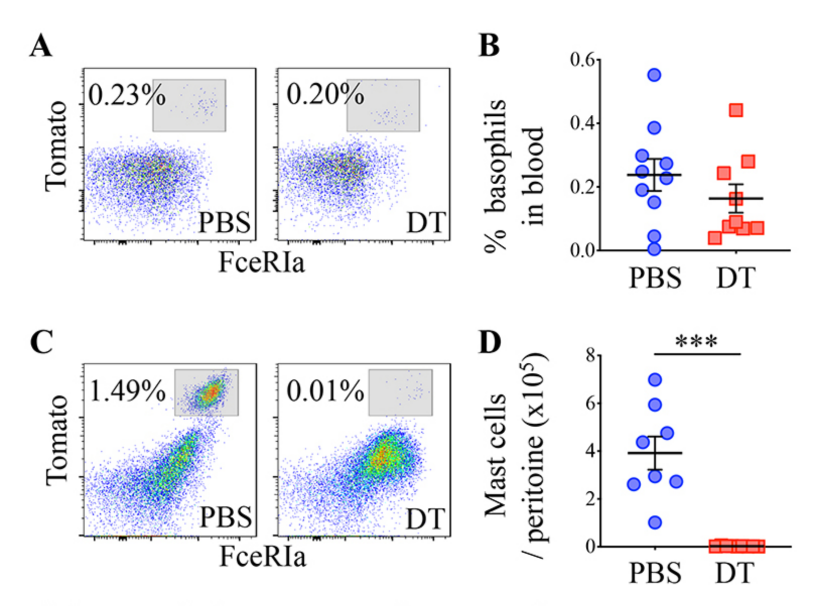
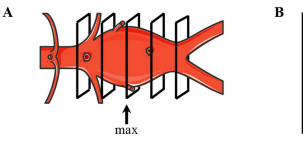
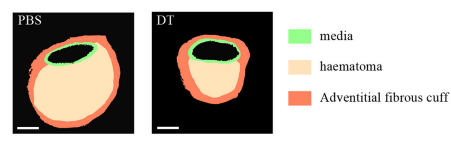


Figure 4-figure supplement 1





Adventitial fibrous cuff

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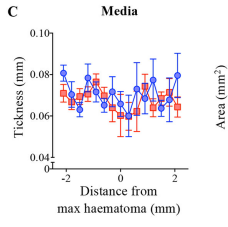
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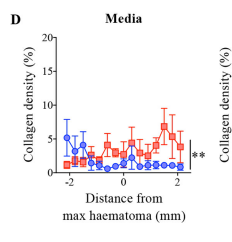
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Tickness (mm)





Haematoma

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Distance from

max haematoma (mm)

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Haematoma

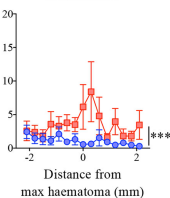
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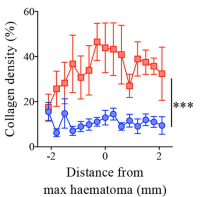


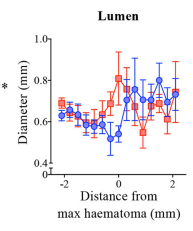
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Distance from

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PBS

DT

Figure 4-figure supplement 2