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Citation

Published Version
doi:10.1371/journal.pone.0045628

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Accessibility
Heterozygosity for Fibrinogen Results in Efficient Resolution of Kidney Ischemia Reperfusion Injury

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Abstract

Fibrinogen (Fg) has been recognized to play a central role in coagulation, inflammation and tissue regeneration. Several studies have used Fg deficient mice (Fg⁺⁻) in comparison with heterozygous mice (Fg⁺⁻) to point the proinflammatory role of Fg in diverse pathological conditions and disease states. Although Fg⁺⁻ mice are considered ‘normal’, plasma Fg is reduced to ~75% of the normal circulating levels present in wild type mice (Fg⁺⁺). We report that this reduction in Fg protein production in the Fg⁻⁄⁻ mice is enough to protect them from kidney ischemia reperfusion injury (IRI) as assessed by tubular injury, kidney dysfunction, necrosis, apoptosis and inflammatory immune cell infiltration. Mechanistically, we observed binding of Fg to ICAM-1 in kidney tissues of Fg⁺⁻ mice at 24 h following IRI as compared to a complete absence of binding observed in the Fg⁻⁻ and Fg⁺⁻ mice. Raf-1 and ERK were highly activated as evident by significantly higher phosphorylation in the Fg⁻⁻ kidneys at 24 h following IRI as compared to Fg⁻⁻ and Fg⁻⁻ mice kidneys. On the other hand Cyclin D1 and pRb, indicating higher cell proliferation, were significantly increased in the Fg⁻⁻ and Fg⁻⁻ as compared to Fg⁻⁻ kidneys. These data suggest that Fg heterozygosity allows maintenance of a critical balance of Fg that enables regression of initial injury and promotes faster resolution of kidney damage.

Introduction

Fibrinogen (Fg) is a 340 kDa glycoprotein, a homodimer linked by disulfide bonds with each unit comprising of 3 distinct polypeptide chains (αA, βB and γγ) that are encoded by 3 separate genes (FGA, FGB and FGG) [1]. Apart from its prominent role in the coagulation cascade, Fg serves as an acute phase response protein by acting as a ligand for receptors expressed on cells recruited to the site of inflammation [2]. In humans, several polymorphisms have been described most of them clustered in the FgB gene [3] resulting in chronically elevated levels of Fg [4,5]. Hyperfibrinogenemia (characterized by high circulating plasma levels of Fg) is consistently associated with an increased risk of cardiovascular diseases [6]; conversely, afibrinogenemia causes severe hemorrhagic risks in affected patients [7]. This suggests the necessity to maintain a critical balance in the levels of Fg that is high enough to maintain adequate clot formation yet low enough to reduce its interactions with cellular receptors along with reducing the availability of fibrin matrices that act as centers of migration and proliferation of immune and endothelial cells in instances of acute and chronic inflammation.

The contribution of Fg in disease pathophysiology of various organs has been studied using Fg deficient mice (Fg⁻⁻) that lack the αA chain, which precludes assembly of functional circulating protein [8,9]. Fg deficient mice (Fg⁻⁻) that lack the αA chain [8] have been shown to be protected from variety of injury/disease states such as atherosclerosis [10], colitis [11], crescentic glomerulonephritis [12], Duchenne muscular dystrophy [13], endotoxemia [14], fibrosis [15,16], multiple sclerosis [17], myocardial ischemia-reperfusion injury [18], ischemic neurodegeneration [19] and rheumatoid arthritis [20]. It should be noted that all of these studies used littermate Fg⁻⁻ mice as experimental controls and not as experimental groups for comparison. Although the Fg⁻⁻ mice do not show any symptoms of abnormal clotting and are for all purposes ‘normal’ when compared to Fg⁻⁻ mice, the plasma level of the Fg protein is reduced to ~75% of the normal circulating levels present in wild type mice (Fg⁺⁺) [9]. The reduction in Fg may not be significant enough to impair the coagulation cascade but could still suffice to alter the binding response to various cellular receptors thereby transforming the immune system’s inflammatory response.

The objective of our study, therefore, was to evaluate the expression profile of Fg following kidney ischemia reperfusion injury (IRI) and to characterize the phenotype of the Fg⁻⁻ and Fg⁻⁻ mice against animals homozygous for the αA gene (Fg⁺⁺) in the context of kidney IRI.

Methods

Ethics

All animal maintenance and treatment protocols were in compliance with the Guide for Care and Use of Laboratory

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Editor: Carlo Gaetano, Goethe University, Germany

Received May 31, 2012; Accepted August 21, 2012; Published September 19, 2012

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Funding: This work in Vaidya laboratory was supported by National Institutes of Health/National Institute of Environmental Health Sciences Outstanding New Environmental Scientist Award (ES017543). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

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animals as adopted and promulgated by the National Institutes of Health and were approved by the Harvard Medical School Animal Care and Use Committees (IACUC).

Animals

Littermate male wild type (Fg+/+), heterozygous (Fg+/-) and knock out (Fg-/-) mice for fibrinogen on Balb/c background (25–29 g) were used for the experiment [9]. Dr. Jay L. Degen at Children’s Hospital Research Foundation, Cincinnati, Ohio, kindly provided breeding pairs of genetically modified Fg mice. Neonate mice experience spontaneous bleeding events, which proves fatal only in 30–50% of cases (depending on strain) and those who survive display otherwise normal organ physiology [9].

Experimental Design

In the first set of experiments twenty male Balb/c mice were anesthetized using pentobarbital sodium (30 mg/kg, ip) and subjected to 25 min bilateral IRI [21] for characterization of fibrinogen expression and excretion. Mice were sacrificed at 24, 48 and 72 hours after reperfusion (n=5/timepoint). In the next set of experiments genetically manipulated mice (54 male wild type, heterozygous and knockout mice) were anesthetized as mentioned above and subjected to 29 min of bilateral renal I/R surgery by the retroperitoneal approach. Sham surgery was performed with exposure of both kidneys but without induction of ischemia. Mice (n=6/group/timepoint) in the respective groups (sham or I/R) were injected with BrdU (50 mg/kg, ip) 3 hr prior to sacrifice. Mice were sacrificed at 12 and 24 h following reperfusion using overdose of pentobarbital (180 mg/kg, ip).

Serum creatinine (Scr) concentrations and blood urea nitrogen (BUN) were measured using a VetScan VS2 (Abaxis, Union City, CA). Plasma Fg (D-Dimer) test was performed by Asserachrom D-Di ELISA kit from Diagnostica Stago, Inc. (Parsippany, NJ) as per manufacturer’s instruction. Urinary Fg levels were measured using commercially available Luminox assay based kit from Millipore (Billerica, MA). Urinary creatinine concentration was used to normalize fibrinogen in order to account for the influence of urinary dilution.

Real Time PCR

Total RNA was extracted by TRIzol reagent (Invitrogen Corporation) as per manufacturer’s protocol. Forward and reverse primer sequences for mouse specific genes were designed using MacVector software (MacVector Inc., Cary, NC) and are listed in Table S1.

In situ Hybridization

Kidney, liver and heart cryosections were washed with TBS and subjected to in situ hybridization was performed with universal digoxigenin-based in situ hybridization and detection kit as per manufacturer’s instructions (Invitrogen, Carlsbad, CA). The probe sequences used are as follows: Fgα-AAT ATG CAA AGA TAG GCC TCA CCC AGA TTA AAG TAG CCTA CTG CCG ACC TGC CTG T; Fgβ-AGT ATA CTC TGT AGC GCT GTA TGG AGG TGT CAG GCT GGA TGA GAT ACG TTT CCG A; Fgγ-CTC AGT GCA TAT GGA ATT GTG GAC TGC ATG CTT ATC AAA TGA ATC TTC TCA TTT C.

Immunofluorescence Staining

Immunostainings in the frozen kidney sections was performed using rat monoclonal anti-F4/80 a kind gift from Dr. Bonventre’s laboratory, BWH, Boston, MA and rat monoclonal anti-BrdU (Cell Signaling Technology, Danvers, MA). The primary antibody was detected using donkey anti rat Cy3 labeled and donkey anti rat FITC labeled secondary antibodies respectively (Jackson ImmunoResearch Laboratories, West Grove, PA). DAPI (Sigma Aldrich, St. Louis, MO) was used for nuclear staining. Apoptosis was measured in kidney tissues by TUNEL assay using the In Situ Cell Death detection kit (Roche Applied Science) according to manufacturer’s instructions [22]. The images were captured by Nikon DS-Qi1Me camera attached to Nikon eclipse 90i fluorescence microscope using oil immersion objective 60/1.4 NA by Nikon NIS elements AR ver 3.2 software.

Immunoblotting and Immunoprecipitation

Kidney tissues were homogenized in RIPA buffer [50 mM Tris-HCl pH 7.4, 150 mM NaCl, 1% NP40, 1 mM PMSE, 1 mM NaF, 20 mM Na4P2O7, 2 mM Na3VO4, 1X protease inhibitor cocktail (Roche Applied Science, Indianapolis, IN)] and equal protein (30 μg) was resolved by polyacrylamide gel electrophoresis. For plasma 0.02 μl was loaded on the gel. Proteins were transferred onto nitrocellulose membrane and western blotting was performed with rabbit polyclonal anti-fibrinogen (Dako), mouse monoclonal anti-pERK, anti-pERK2 (BD Biosciences San Diego, CA), anti-Cyclin D1, anti-pRb, anti-β -Actin (Cell Signaling Technology), anti-i-Tubulin (Sigma) and goat polyclonal HRP conjugated anti-mouse albumin (Abcam, Cambridge, MA). HRP conjugated secondary antibodies against mouse, rabbit and goat was purchased from Jackson Immunoresearch (West Grove, PA). For Immunoprecipitation (IP) tissues were lysed in IP buffer (20 mM Tris-HCl pH 8.0, 137 mM NaCl, 10% glycerol, 1% NP40, 2 mM EDTA) containing protease inhibitor cocktail and 300 μg protein was incubated overnight at 4 °C with 4 μg of rabbit polyclonal anti-fibrinogen antibody (Dako). Fifty microlitre of protein A/G agarose was added and incubated for 2 h at room temperature. Beads were washed thrice with IP buffer. Immune complex was eluted by adding 1X SDS loading dye and heating at 100°C and western blot was performed to detect ICAM-1 (goat polyclonal, R&D Systems) and goat polyclonal anti-fibrinogen (Nordic lab, The Netherlands).

Statistics

Data are expressed as average ± standard error. Statistical difference (p<0.05) as calculated by one-way ANOVA or student’s t-test. P<0.05 was considered significant and represented by * as compared to shams, ** as compared to wild type at similar time points, † as compared to heterozygous at similar time points where applicable. All graphs were generated by GraphPad Prism (GraphPad, Inc., La Jolla, CA).

Results

We found a significant increase in the mRNA (Fig. 1A, 1B) and protein expression (Fig. 1C) of Fgα, Fgβ and Fgγ in the kidney and urinary excretion of Fg (Fig. 1D) in mice following IRI corresponding to the kidney dysfunction and proximal tubular necrosis (Fig. S1). In situ hybridization (ISH) with Fgα, Fgβ and Fgγ in the liver tissue revealed strong diffuse cytoplasmic staining in the hepatocytes (Fig. 1B, biological positive control). In the kidney, the staining varied in intensity and distribution between the chains and with respect to the presence or absence of injury (Fig. 1B, first two columns). Fgα in uninjured (sham) kidney revealed diffuse cytoplasmic staining that was less intense than the reactivity in liver under the same conditions. The staining was more intense and perinuclear in distribution 24 h after IRI. The staining was more intense and perinuclear in distribution 24 h after IRI. ISH for Fgγ in uninjured kidney was as intense as in liver, but revealed
more intense diffuse cytoplasmic distribution at 24 h. ISH for Fgγ was of similar reactivity to Fgα in distribution and intensity, in uninjured and injured kidney; there was more diffuse cytoplasmic staining in uninjured tissue, with less intense reactivity when compared to the liver tissue, but in IRI, the reactivity intensified and distributed around the nuclei, similar to the Fgα.

Phenotypically the Fgγ+/−, and mutant mice for FgA- α chain [heterozygous (FgA−/−) or knockout (FgA−/−)] demonstrated following features (Fig. S2A): i) loss of FgA chain with, albeit modestly decreased, but detectable transcription and translation of Fgβ and Fgγ chains in the liver of FgA−/− mice (Fig. S2B) as reported previously [9]; ii) significant decrease in transcription and translation of FgA and Fgβ with no alterations in Fgγ chain in the kidney of Fg−/− mice (Fig. S2C) as compared to Fg−/− mice; iii) approximately 50% decrease in plasma Fg D-Dimer levels in Fg−/− mice and undetectable Fg protein in the circulation in Fg−/− mice as compared to Fg+/+(Fig. S2D); iv) unaltered urinary Fg protein excretion (Fig. S2E).

Following IRI, Fg−/− and Fg−/− mice showed greater kidney dysfunction as assessed by BUN and SCr (Fig. 2A) at 12 h as compared to Fg+/* mice (Fig. 2A). However, by 24 h the kidney dysfunction in the Fg+/* and Fg−/− mice further escalated whereas, the Fg−/− mice appeared to show rapid functional restoration with BUN and SCr levels ~2-3 fold lower than the Fg+/* and Fg−/− mice (Fig. 2A). Urinary Fg, serving as a sensitive indicator of kidney tubular injury [21], also showed a massive increase at 12 h in all three groups but significantly reduced levels in Fg−/− (~5-fold) and Fg−/− (~10-fold) mice at 24 h as compared to Fg−/− mice (Fig. 2A). Kidney histology showed severe necrosis in the cortico-medullary junction of all post ischemic kidneys particularly in the S3 segments at 12 and 24 h (Fig. 2B). Although differences between the Fg−/−/+/*, Fg−/− and Fg−/− mice in the extent of necrosis were subtle, there was a marked reduction in the number of apoptotic cells at 24 h in the Fg−/− and Fg−/− mice as compared to Fg−/− mice (Fig. 2C).

We next evaluated the contribution of inflammatory immune cell invasion in progression of kidney injury by immunostaining for markers of macrophage (F4/80) and neutrophil (Ly-6G) infiltration. The number of F4/80 positive cells was similar at 12 h in all three groups but at 24 h the Fg−/− and Fg−/− mice showed a statistically significant decrease in the number of macrophages (Fig. 3A) as well as Ly-6G positive neutrophils (Fig. S3) as compared to Fg−/− mice.

To evaluate the kidney tissue repair we quantitated the number of proliferating epithelial cells by BrdU immunostaining and found that Fg−/− and Fg−/− mice exhibited significantly greater number of BrdU positive cells as compared to Fg−/− mice at 24 h (Fig. 3B). This suggests that although the initiation and early phase of injury was similar in all three groups there was a timely and efficient tissue repair response in the Fg−/− and Fg−/− mice, which curbed inflammation and apoptosis resulting in regression of injury. We hypothesized that in the Fg−/− mice there is progression of the initial injury because of Fg binding to Intercellular Adhesion Molecule-1 (ICAM-1) that promotes apoptosis and cell death through ERK phosphorylation.

We found that there was significant binding of Fg to ICAM-1 in kidney tissues of Fg−/− mice at 24 h following IRI as compared to a complete absence of binding observed in the Fg−/− and Fg−/− mice (Fig. 4A). Consistently there was a significant decrease in expression of Fg protein at 24 h in Fg−/− and Fg−/− mice kidneys as compared to Fg+/* mice following IRI (Fig. 4B). We further confirmed Fg binding to ICAM-1 in tubular epithelial cells by co-immunostaining that showed a Pearson co-localization coefficient between Fg and ICAM-1 to be 0.6 in Fg−/− as compared to 0.4 and 0.05 in Fg−/− and Fg−/− respectively (Fig. S4). Raf1 and ERK were highly activated as evident by significantly higher phosphorylation in the Fg−/− kidneys at 24 h following IRI as compared to Fg−/− and Fg−/− mice kidneys. pERK was upregulated at 12 h in all three groups confirming the earlier observations (Fig. 2) about similar level of necrosis, apoptosis and

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**Figure 1.** Transcription and translation of Fgs, Fgβ and Fgγ in the kidney as urinary Fg excretion is significantly increased following IRI in mice. A) To characterize de novo expression of Fgs at the mRNA and protein level in the kidney, male BALB/c were subjected to IRI and kidneys, blood and urines were collected over time (n = 5/time point). A) Real time PCR analysis in kidney for Fgs, Fgβ and Fgγ chains, normalized to GAPDH, and fold change determined over sham. B) In situ hybridization for Fgs, Fgβ and Fgγ mRNA in kidney, 1:25 represents competition with unlabeled probe, liver (positive control) and heart (negative control). C) Western blot analysis for Fg protein in the kidney. β-Actin served as loading control. D) Urinary Fg levels measured by Luminex assay. *represents p<0.05 as determined by student’s t-test as compared to sham. Bar represents 10 μm. doi:10.1371/journal.pone.0045628.g001
inflammation at 12 h in the three groups of mice. On the other hand Cyclin D1 and pRb, indicating higher cell proliferation, were significantly greater at 24 h in the \(\text{Fg}^+/-\) and \(\text{Fg}^-/-\) as compared to \(\text{Fg}^+/+\) kidneys (Fig. 4B).

**Discussion**

Our results support the hypothesis that a reduction in availability of excess Fg from interacting with ICAM-1 prevents progression of injury thereby allowing timely induction of Cyclin D1 and pRb mediating efficient kidney tissue repair and resolution of injury (Fig. 4C). This hypothesis is consistent with the previous reports demonstrating therapeutic potential of Fg-derived peptides (B15-42 and γ377-385) by competing with native Fg for binding to vascular endothelial cadherin (VE-cadherin), ICAM-1, CD11b/CD18 which in turn inhibits infiltration of leukocytes at the site of injury and prevents exacerbation of injury [18,23]. Furthermore, others and we have previously shown that B15-42 peptide administration protected from kidney IRI by increasing tissue repair and decreasing apoptosis [21,24] confirming that pharmacological reduction in excess Fg paves the way for faster structural and functional recovery.
Figure 3. Heterozygous and knockout Fg mice exhibit efficient immune cell clearance coupled with robust tubular epithelial cell proliferation. Fixed frozen sections following IRI were stained for A) macrophage F4/80 (red). Number of F4/80 cells per 60X field is represented graphically on the right of the photomicrograph. B) BrdU positive cells (green) by immunofluorescence. Percentage of positive staining for BrdU positive nuclei is represented graphically on the right of photomicrographs. Arrowheads indicate positive cells/nucleus and bar represents 10 μm. *represents p<0.05 in comparison to sham; †represents p<0.05 as compared to wild type within the time point and ‡represents p<0.05 as compared to heterozygous within the time point as determined by one-way ANOVA. Bar represents 10 μm.

doi:10.1371/journal.pone.0045628.g003
Figure 4. Fibrinogen binds to ICAM-1 in the kidney leading to sustained tissue injury. A) Kidney tissue lysates were immunoprecipitated with anti-fibrinogen antibody and immunoblot was performed for ICAM-1. IgG light chain served as loading control for IP and β-Actin served as loading control for input. B) Following 29 min IRI kidney tissue lysates were prepared and equal protein was resolved on SDS-PAGE for western blot analysis for pERK, ERK, Cyclin D1 and pRb. α-Tubulin served as loading control. C) Schematic showing that in the Fg\textsuperscript{+/–} mice a reduction in availability of excess Fg from interacting with ICAM-1 prevents progression of injury thereby allowing timely induction of Cyclin D1 and pRb mediating efficient kidney tissue repair and resolution of injury.
Thus, the new findings of this study are i) Fg (Fgα, Fgβ, and Fgγ) is transcribed in the kidney and its mRNA levels, protein expression and urinary excretion significantly increase following IRI; ii) Heterozygosity of mouse Fgα chain results in global reduction of Fg production to a moderate level that protects the Fg−/− mice from IRI-induced kidney tubular injury, kidney dysfunction, inflammation and apoptosis by launching an efficient tissue regeneration response. Although Fg−/− mice showed a reduction in apoptosis, reduced immune cell infiltration and increased regeneration, the functional and structural restoration of the kidney after IRI was not as rapid as Fg+/− mice potentially due to the impedance with clotting.

Fibrinogen binding to ICAM-1 through its γ157-159 domain has been well documented on endothelial cells [25,26] and has been shown to promote leukocyte transmigration by acting as an intermediary molecule that can bind both ICAM-1 and leukocytes through the Mac-1 receptor [27]. We extended these studies and show that Fg binds to ICAM-1 in the kidney following IRI thereby potentially activating Raf-1, triggering the Raf-MEK-ERK pathway, which in turn can activate an apoptotic response. Experiments using anti-ICAM-1 antibodies as well as ICAM-1 deficient mice have shown ICAM-1 to be a key mediator of acute IRI injury via potentiation of neutrophil–endothelial interactions [28]. ICAM-1 expression significantly increases on proximal tubular epithelial cells in patients with acute renal allograft rejection [29] and here we found significant co-localization of ICAM-1 with Fg (Pearson’s co-localization coefficient of 0.6, Fig. S3) predominantly on the proximal tubular epithelial cells emphasizing the paradigm that tubular epithelium is not merely a passive victim of injury but also an active participant in the inflammatory response in kidney IRI [30].

In summary, our experiment shows that kidney expresses Fgα, Fgβ and Fgγ transcripts and genetic manipulation resulting in decreased availability of Fg protein to interact with cellular receptors diminishes the molecular response cascade and dampens the inflammatory response leading to faster resolution of injury.

Supporting Information

**Figure S1** Characterization of kidney dysfunction and tubular injury following bilateral renal ischemia/reperfusion injury (IRI). Male BALB/c were subjected to IRI and kidneys, blood and urines were collected over time (n = 5/time point). A) Serum creatinine (Scr) and Blood Urea Nitrogen (BUN) measurements. B) Representative histological H&E stained images following IRI at 24, 48 and 72 h showing proximal tubular necrosis as compared to sham. Bar represent 100 μm. (TIF)

**Figure S2** Genotype and phenotype characterization of Fg wild type, heterozygous and knockout mice. A) Genotyping results from a representative group (n = 7) of Fg wild type, heterozygous and knockout mice as described in methods section. B) Real time PCR and Western Blot analysis for Fg (Fgα, Fgβ and Fgγ) in the liver and C) Kidney of Fg wild type, heterozygous and knockout mice. D) Plasma levels of Fg in wild type, heterozygous and knockout mice were measured by D-Dimer ELISA test and by western blot analysis. E) Urinary levels of Fg were measured using a Luminex based assay in wild type, heterozygous and knockout mice. *represents p<0.05 as determined by student’s t-test in comparison to wild type. (TIF)

**Figure S3** Heterozygous and knockout Fg mice show significantly decreased neutrophil infiltration following IRI. Fixed frozen section following IRI at 12 and 24 h were stained for Ly-6G (green). Number of Ly-6G positive nuclei was represented graphically on the right of photomicrographs. Arrowheads indicate neutrophils and bar represent 10 μm. *represents p<0.05 in comparison to sham; #represents p<0.05 as compared to wild type within the time point and ‘represents p<0.05 as compared to heterozygous within the time point as determined by one-way ANOVA. Bar represent 10 μm. (TIF)

**Figure S4** Significant colocalization of Fg and ICAM in the kidney following IRI. Fixed frozen section following IRI at 12 and 24 h were co-stained for Fg (red) and ICAM-1 (green). Pearson’s coefficient was plotted as a measure of co-localization on the right of photomicrographs. *represents p<0.05 in comparison to sham; #represents p<0.05 as compared to wild type within the time point and ‘represents p<0.05 as compared to heterozygous within the time point as determined by one-way ANOVA. Bar represent 10 μm. (TIF)

**Table S1** Primer sequences for genotyping and Real Time PCR analysis for candidate genes.

**Acknowledgments**

We thank Dr. Jay L. Degen, Professor, Department of Pediatrics, Children’s Hospital Research Foundation, Cincinnati, Ohio, for providing breeding pairs of genetically modified mice for fibrinogen α chain. We thank Dr. Joseph V. Bouventre, Director, Renal Division, Brigham and Women’s Hospital, Harvard Medical School, Boston, MA for kindly providing the F4/80 antibody.

**Author Contributions**

Conceived and designed the experiments: AKA VSV. Performed the experiments: AKA JS. Contributed reagents/materials/analysis tools:. Wrote the paper: AKA JS VSV. Provided unbiased pathological report in a blinded manner: VB. Analyzed the data as well as contributed to the interpretation of the data: AKA JS VB VSV.

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