Recoupling of eNOS with Folic Acid Prevents Abdominal Aortic Aneurysm Formation in Angiotensin II-Infused Apolipoprotein E Null Mice

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Abstract

We have previously shown that eNOS uncoupling mediates abdominal aortic aneurysm (AAA) formation in hph-1 mice. In the current study we examined whether recoupling of eNOS prevents AAA formation in a well-established model of Angiotensin II-infused apolipoprotein E (apoE) null mice by targeting some common pathologies of AAA. Infusion of Ang II resulted in a 92% incidence rate of AAA in the apoE null animals. In a separate group, animals were treated orally with folic acid (FA), which is known to recouple eNOS through augmentation of dihydrofolate reductase (DHFR) function. This resulted in a reduction of AAA rate to 19.5%. Imaging with ultrasound showed that FA markedly inhibited expansion of abdominal aorta. FA also abolished elastin breakdown and macrophage infiltration in the AAA animals. The eNOS uncoupling activity, assessed by L-NAME-sensitive superoxide production, was minimal at baseline but greatly exaggerated with Ang II infusion, which was completely attenuated by FA. This was accompanied by markedly improved tetrahydrobiopterin and nitric oxide bioavailability. Furthermore, the expression and activity of DHFR was decreased in animals derived from activated vascular smooth muscle cells (VSMCs) and studies have shown that oxidative stress implicated in AAA is often regulated to some degree by reactive oxygen species (ROS), degradation, and remodeling [3–6]. As all of these processes are new treatment options.

Studies into the underlying mechanisms of this condition show several important processes, such as inflammation, matrix degradation, and remodeling [3–6]. As all of these processes are regulated to some degree by reactive oxygen species (ROS), oxidative stress plays an important role in the pathogenesis of AAA in both animal models [7–10] and humans [11,12]. Previous studies have shown that oxidative stress implicated in AAA is often derived from activated vascular smooth muscle cells (VSMCs) and infiltrating macrophages [13]. However, there are some evidences that endothelial cells or dysfunctional eNOS plays in the development of AAA warrants further investigation.

In a previous study from our laboratory, we showed that in a strain of mice with a modest deficiency in the eNOS cofactor tetrahydrobiopterin (H4B), infusion with angiotensin II (Ang II) for two weeks induces AAA that appears to be among the most severe of available murine models [18]. We further observed that the augmented H4B deficiency to restore eNOS function, with oral treatment of folic acid (FA) that restores the expression and activity of the H4B salvage enzyme dihydrofolate reductase (DHFR) in the endothelium, eliminated the incidence of AAA [18]. While FA was extremely potent in the prevention of AAA in this model, it is important to examine whether FA targets common pathways of AAA to prevent the disease in other established models of AAA.

In the current study, we examined whether the preventive effect of FA extends to other models of AAA. We chose to use the Ang II infused apoE null model of AAA, as it is a well-characterized murine model for AAA [7,19,20] that is different from our previous model of hph-1 mice. The results show that oral treatment of FA significantly reduced the incidence of AAA from 92% to 19.5%. Furthermore, FA treatment significantly improved endothelial specific DHFR expression and activity, restored H4B...
bioavailability, recoupled eNOS, and increased nitric oxide production. Taken together, these data reveal a novel role of eNOS uncoupling in the development of AAA in Ang II-infused apoE null mice, which is exploited by oral treatment of FA that improves eNOS function to attenuate AAA. These data further confirm a significant role of uncoupled eNOS in mediating AAA formation, and a universal efficacy of FA in preventing AAA formation via restoration of DHFR to restore eNOS function.

Materials and Methods

1) Reagents

Unless otherwise noted, all chemicals were purchased from Sigma-Aldrich at the highest possible purity. Collagenase was obtained from Gibco. Isoflurane was purchased from Piramal Healthcare.

2) Animals

All animals and experimental procedures were approved by the Institutional Animal Care and Usage Committee at the University of California, Los Angeles (UCLA). Breeders of apoE null mice were purchased from Jackson Labs (Bar Harbor, ME, Strain B6.129P2-Apoemtm1Tur /J), then bred in house for experimental use. Animals were kept in ventilated cages, with free access to water and standard chow, and cared for by the DLAM staff until experiments. Male animals were kept until 6–8 months old before experiments.

3) Ang II infusion by osmotic pump

Animals were anesthetized in an isoflurane chamber with 5% isoflurane, and then moved to a nose cone supplying 1.5–2% isoflurane to maintain the anesthetic state. The area on the back between the shoulder blades was cleaned of hair and disinfected. A small incision was made at the cleaned site, and the osmotic pump (Alzet, model 2004) containing Ang II (1000 ng/kg/min) in a 5% solution was inserted into the animal under the skin. The wound site was closed using surgical staples, and the animal was allowed to recover in a heated cage.

4) Folic acid treatment

For animal groups treated with folic acid, standard chow was replaced with customized food tablets containing FA (15 mg/kg/day) [18] two days prior to the osmotic pump implantation, and throughout the study period of 4 weeks of Ang II infusion.

5) Tissue collection

Animals were euthanized with CO2 4 weeks after the implantation of osmotic pump containing Ang II. The aorta was rapidly removed from the body, rinsed with ice cold PBS, and centrifuged at 12,000 g for 3 min. The tissue was then minced with a razor blade and incubated with freshly prepared nitric oxide specific spin trap Fe2(CN)6(ETC)4 (0.5 mmol/L) in modified Krebs/HEPES buffer at 37°C for 60 min, in the presence of potassium ferri cyanide (50 μmol/L). After the incubation, the aorta was snap frozen in liquid nitrogen and loaded into a finger Dewar for analysis with ESR spectrophotometer (eScan, Bruker). The instrument settings were as the followings: bio-field, 3494.50; field sweep, 9 G; microwave frequency, 9.75 GHz; microwave power, 21.02 mW; modulation amplitude, 2.47 G; 4,096 points of resolution; receiver gain, 1000; and kinetic time, 10 min.

6) Ultrasound imaging of abdominal aorta

Animals were anesthetized with isoflurane, and then placed onto a temperature-controlled table, which also measures ECG for the determination of heart rate. Isoflurane levels were adjusted to maintain heart rate between 400–500 bpm while keeping the animal sufficiently anesthetized (~1.5–2%). Hair from the abdomen was removed using a hair removal lotion (Veet), and pre-heated ultrasound transmission gel was applied over the cleaned abdomen area. An ultrasound probe (Visualsonics 2100, MS400, 30 MHz) was placed onto the gel to visualize the abdominal aorta transversely. The aorta was confirmed using Doppler mode to detect the presence of pulsatile flow. Consistent localization of image acquisition was insured by visualizing the aorta immediately superior to the branch of the left renal artery in all animals. In the cases of abdominal aneurysms, the measurements were done at the site of maximal aortic diameter.

7) Electron spin resonance determination of aortic superoxide production and eNOS uncoupling activity

Aortic superoxide production was determined by electron spin resonance (ESR) as previously described [18]. Briefly, entire freshly isolated aortas were homogenized in lysis buffer (Tris-HCl, NaCl, EDTA, EGTA, Sodium pyrophosphate, β-glyceroophosphate, sodium orthovanadate, triton-X) containing 1:100 protease inhibitor cocktail (Sigma, P8340), centrifuged at 12,000 g for 15 min, and protein supernatant collected. After determination of protein concentration using a protein assay kit (Bio-Rad), 5 μg of protein was loaded into ice-cold and nitrogen bubbled modified Krebs/Hepes buffer (KHB, in mmol/L: NaCl 99; KCl 4.7; MgSO4.1.2; KH2PO4 1.0; CaCl2 1.9; NaHCO3 25; glucose 11.1, NaHEPES 20) containing diethylthiodiethiocarbamic acid (3 μmol/L), delferoxamine (25 μmol/L), and the superoxide specific spin trap methoxycarbonyl-2,5,5-tetramethylpyrroline (CMH, 500 μmol/L, Alexis). The mixture was loaded into a glass capillary (Kimble, 71900-50), and assayed using electron spin resonance (ESR) spectrophotometer (eScan, Bruker) for superoxide production by taking the difference in the presence or absence of SOD (100 U/mL). To determine eNOS uncoupling activity, measurements were made with the addition of L-NAME (100 μmol/L). The ESR settings used were: bio-field, 3494.50; field sweep, 9 G; microwave frequency, 9.75 GHz; microwave power, 21.02 mW; modulation amplitude, 2.47 G; 4,096 points of resolution; receiver gain, 1000; and kinetic time, 10 min.

8) Electron spin resonance determination of aortic nitric oxide production

Freshly isolated aortic rings from non-aneurismatic sections were incubated with freshly prepared nitric oxide specific spin trap Fe2(CN)6(ETC)4 (0.5 mmol/L) in modified Krebs/HEPES buffer at 37°C for 60 min, in the presence of calcium ionophore A23187 (10 μmol/L). After the incubation, the aorta was snap frozen in liquid nitrogen and loaded into a finger Dewar for analysis with ESR spectrophotometer (eScan, Bruker). The instrument settings were as the followings: bio-field, 3,280; field sweep, 77.54 G (1 G = 0.1 mT); microwave frequency, 9.78 GHz; microwave power 40 mW (4 dB); modulation amplitude, 10 G; 4,096 points of resolution; and receiver gain 900.

9) HPLC determination of aortic H4B content

Entire freshly isolated aortas were lysed in H4B lysis buffer (0.1 mol/L, phosphoric acid, 1 mmol/L EDTA, 10 mmol/L DL-Dithiothreitol), and then centrifuged at 12,000 g for 3 min. Lysates were then subjected to differential oxidation in acidic (0.2 mol/L, trichloroacetic acid with 2.5% I2 and 10% KI) and alkaline (0.1 mol/L, NaOH with 0.9% I2 and 1.5% KI) solutions as described previously [18]. After centrifugation, 10 μL of the supernatant was injected into a HPLC system equipped with a fluorescent detector (Shimazu). Excitation and emission wavelengths
Figure 1. Folic acid greatly reduces the incidence of AAA in Ang II-infused apoE null mice. A) Representative full length aortas from sham operated, Ang II-infused, and Ang II-infused, folic acid (FA) treated apoE null animals. B) The top panel shows the percentage of AAA development in Ang II-infused apoE null mice fed regular chow or chow mixed with FA. The data show that with regular chow, the AAA incidence rate is 92%, while with FA it reduces to 19.5% (*p < 0.01). The bottom panel shows the actual number of animals used across all groups: apoE n = 27, apoE+Ang II n = 38, apoE+Ang II+FA n = 41.
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Figure 2. Time course of AAA development and efficacy of folic acid in Ang II-infused apoE null mice measured with ultrasound. A) Representative images of ultrasound taken from sham operated apoE null mice (top row), Ang II-infused apoE null mice with (bottom row) and without (middle row) folic acid (FA) treatment. B) Summarized data from ultrasound measurements of abdominal aorta area. FA significantly reduced abdominal aortic size compared with Ang II alone starting at 2 weeks (n = 4–5 each). * p < 0.05 vs. control, # p < 0.01 vs. all, $ p < 0.001 vs. all.
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of 350 nm and 450 nm were used to detect H$_2$B and its oxidized species.

10) Verhoeff-Van Gieson (VVG) staining

Paraffin embedded tissue sections were deparaffinized by sequential washes in xylene (2x), descending alcohol from 100% to 50%, then into distilled water. Sections were then stained in Verhoeff's solution for 70 min, followed by differentiation in 2% ferric chloride for 70 seconds. Sections were then incubated with 5% sodium thiosulfate for 75 seconds, followed by counterstaining with Van Gieson's solution and dehydration with 95% and 100% alcohol, and finally washed in xylene. After drying, the tissues were mounted with Permount (Fisher Scientific).

11) Macrophage staining

Tissue sections were deparaffinized and hydrated as described above. After dehydration, sections were incubated in methanol containing 1% H$_2$O$_2$ for 30 min. Sections were then washed with PBST, and blocked in 2% goat serum at RT for 3 hrs prior to being incubated with primary antibody (Mac-3, BD Pharmingen, 550292, 2% in PBS-T) overnight at 4°C. After washing in PBST, sections were incubated with secondary antibody for 1 hr, followed by another wash. Sections were then incubated with the ABC system (goat IgG, Vectastain, Vector Laboratories, Burlingame, CA, 1.5% reagent A and B in PBST) for 30 min, after which the sections were washed in PBST. The Mac-3 staining was visualized by developing with 3,3'-diaminobenzidine (Sigma-Aldrich) for 3 min to achieve the brown coloration. The final sections were dehydrated in ascending grades of ethanol and xylene, and mounted with Permount.

The degree of macrophage infiltration was quantified using Image J by measuring the brightness of mac-3 signals (dark brown) from each vessel segment was measured and normalized by the tissue area.

12) Endothelial cell isolation

Endothelial cells were isolated from entire freshly isolated mouse aortas as per our previous study [21]. Briefly, freshly isolated aortas were cut into small pieces (~2 mm each), and digested in PBS containing collagenase (0.6 mg/mL) for 20 min at 37°C. After digestion, the aortic rings were gently shaken in the buffer to remove endothelial cells from denuded aortas. The endothelial cells in the buffer were pelleted by centrifugation at 1,000 g for 5 min, prior to being subjected to analyses of DHFR expression and activity.

13) Western blotting

Western blotting was performed as per standard protocols, using 10% SDS/PAGE gel and transferring to nitrocellulose membrane. Proteins were probed with primary antibodies for DHFR (1:500, Novus Biologicals, Littleton, CO, H00001719-M01), actin (1:3000, Sigma-aldrich, A2066), and eNOS (1:2000, BD transduction laboratories, 610297).

14) DHFR Activity assay

DHFR activity was measured via HPLC as previously described [21]. Briefly, cell lysates were incubated with dihydrofolate (50 μmol/L) and NADPH (200 μmol/L) at 37°C for 20 min in a potassium phosphate assay buffer (0.1 mol/L) containing 1 mmol/L DTT, 0.5 mmol/L KCl, 1 mmol/L EDTA, and 20 mmol/L sodium ascorbate at pH 7.4. The reaction was stopped by addition of 0.2 mol/L trichloroacetic acid, and stabilized with a stabilization solution (200 mg sodium ascorbate and 30 mg DTT in 1 mL water, 1:10 dilution in final solution).

Figure 3. Folic acid reduces adventitial hypertrophy in Ang II-infused apoE null mice. Abdominal aortas were collected from sham operated (left column), Ang II-infused (center column), and Ang II-infused and folic acid (FA, right column) treated apoE null mice 4 weeks after infusion. Tissues were then sectioned and stained for H&E; note partially ruptured aneurysm and attenuation of adventitial hypertrophy by FA. doi:10.1371/journal.pone.0088899.g003
The product of this reaction, tetrahydrofolate, was measured using a Shimadzu HPLC system with a C-18 column (Alltech, Deerfield) and a fluorescent detection at 295 nm excitation and 365 nm emission. The mobile phase of the HPLC was a 7% acetonitrile with 5 mmol/L KH₂PO₄ at pH 2.3.

15) Statistical analysis
Differences among different groups of data were compared using unpaired t-test for two groups, and ANOVA for multiple groups which was followed by Newman-Keuls post-hoc test. Statistical significance was set at p<0.05. All grouped data are presented as Mean±SEM.

Results
1) Oral FA treatment reduces the incidence of AAA in Ang II-infused apoE null mice
The total incidence of AAA across all groups for this study is shown in Figure 1. Data show that after 4 weeks of Ang II-infusion, 92% of the animals developed AAA (n = 38), which is consistent with previous observations from other independent laboratories. Oral FA treatment significantly reduced this incidence to 19.5% (p<0.001, n = 41), indicating that FA is effective in the prevention of AAA in the Ang II-infused apoE null model. None of the animals in this study died from aortic rupture within the study period of 4 weeks.

Figure 4. Folic acid reduces matrix degradation in Ang II-infused apoE null mice. Abdominal aortas were collected from sham operated (left column), Ang II-infused (center column), and Ang II-infused and folic acid (FA, right column) treated apoE null mice 4 weeks after infusion. Tissues were then sectioned and stained for VVG; the right arrow in the VVG sections points to a breakdown of elastin fiber, while the left arrow points to a flattening of elastin fiber.

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2) Oral FA treatment prevents enlargement of abdominal aorta measured via ultrasound
To assess the time course of AAA development in the Ang II-infused apoE null animals, and the efficacy of FA during this time, we used non-invasive ultrasound to measure the size of the abdominal aorta weekly from a subset of the animals. Figure 2A shows the representative set of images across the 4 weeks, while Figure 2B shows the summarized data (n = 4–5 each). The data show that the abdominal aortas of Ang II-infused apoE null animals were significantly enlarged compared to those of vehicle-infused control animals starting one week after Ang II infusion. FA treatment resulted in marked reduction in aortic size starting from week 2 (Figure 2B).

3) Oral FA treatment reduces maladaptive vascular remodeling
To further analyze the extent of the vascular remodeling that occurred during AAA, we harvested and stained tissue sections of abdominal aortas. Representative images of H&E, VVG, and macrophage staining are shown in Figure 3, 4, and 5. Of note, Ang II infusion induced a marked adventitial hypertrophy (Figure 3). The representative picture also indicated a partially ruptured aneurysm. Furthermore, VVG staining reveals that the elastin fibers were flattened and broken in the Ang II-infused animals (Figure 4), implicating matrix degradation. Treatment with FA largely abolished these responses. Macrophage staining (Figure 5) shows that Ang II infusion significantly increased
4) Oral FA treatment reduces total aortic superoxide production, recouples eNOS, improves NO levels, and restores aortic H4B bioavailability

A previous study by our group showed that eNOS recoupling mediates FA’s protective effect against development of AAA [18]. Here, we examined whether the key role that eNOS plays in that model also extends to the model of Ang II-infused apoE null mice. Figure 6A shows the production of superoxide measured from aortic lysates of apoE null animals (n = 6), with and without the presence of the NOS inhibitor L-NAME. The measurements made without L-NAME, shown in white bars, indicate that Ang II infusion significantly increased production of superoxide, while FA treatment reduced this enhanced level.

The measurements made with L-NAME, as shown in the black bars of Figure 4A, are done to assess the coupling state of eNOS. Under normal conditions when eNOS is coupled, the addition of the eNOS inhibitor will increase the measured superoxide production, as eNOS is producing NO to scavenge superoxide. However, when eNOS is uncoupled and producing superoxide, its inhibition will lead to a decrease in measured superoxide. The data show that in the sham operated apoE null animals, there was slight eNOS uncoupling, which was however greatly exacerbated with the infusion of Ang II. FA treatment resulted in re-coupling of eNOS, as shown by the increase in superoxide production with the addition of L-NAME. Since FA restored eNOS function, we next measured NO levels in isolated aortas from treated apoE null mice. The results, shown in Figure 6B, demonstrate that Ang II infusion significantly decreased the level of NO in the abdominal aortas, while FA treatment significantly increased it (n = 5-6 each).

H4B is the essential cofactor of eNOS, while its deficiency is indicative of eNOS uncoupling. Here, we measured H4B content from aortas of treated apoE null mice to examine whether FA treatment affects aortic H4B content. The data, summarized in Figure 6C (n = 5), show that Ang II infusion significantly reduced aortic H4B bioavailability (p < 0.01), which was markedly restored by oral FA treatment (p < 0.05). Taken together with the previous data, these data show that FA not only reduces superoxide production in the aortas of Ang II-infused apoE null mice, but also recouples eNOS to improve NO levels. This improvement in NO is likely due to an increase in H4B levels.

5) Oral FA treatment improves DHFR expression and activity in endothelial cells in Ang II-infused apoE aortas

The above data show that restoration of eNOS coupling, which is tied to the bioavailability of H4B, may be important in FA’s protection against Ang II induced AAA development in the apoE null animals. Previous studies have shown that FA treatment can recouple eNOS through the improvement of endothelial DHFR function, which is essential in salvaging H4B [18,21]. Here, we examined endothelial DHFR activity and expression in Ang II-infused apoE null mice to test whether DHFR is also improved during FA prevention of AAA in this model. Endothelial cells were

macrophage infiltration (n = 4 each), while FA treatment was able to attenuate this back to near baseline levels.

Figure 5. Folic acid reduces inflammation in Ang II-infused apoE null mice. Abdominal aortas were collected from sham operated (left column), Ang II-infused (center column), and Ang II-infused and folic acid (FA, right column) treated apoE null mice 4 weeks after infusion. Tissues were then sectioned and stained for Mac-3 for infiltrating macrophage. Areas within the box were enlarged to better show details of the macrophage infiltration. The degree of macrophage infiltration was measured and quantified in Image J (*p < 0.01 vs. all, n = 4 each).

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isolated from freshly prepared aortas, and Western blots and HPLC were performed to assess DHFR expression and activity respectively on both the isolated ECs and EC-denuded aortas. The top panels of Figure 7A shows the representative Western blots for eNOS (144 kD), actin control (42 kD), and DHFR (21 kD) (n = 8–11 each). eNOS was stained for quality control to examine the effectiveness of the EC isolation/removal. The increase in eNOS expression in the Ang II-infused groups was expected and well documented in previous works [22–24]. The summarized data indicate that DHFR was significantly decreased in aortic ECs of Ang II infused apoE null animals, while FA treatment restored this level back to baseline. Similarly, DHFR activity (Figure 7B, n = 5–7) was also markedly reduced in ECs specifically in Ang II infused apoE null mice, but then fully recovered to above baseline levels with oral FA treatment. Of note, DHFR expression and activity remained unchanged in EC-denuded aortas.

Discussion

In this work, we identified an essential role of endothelial DHFR deficiency and eNOS uncoupling in mediating AAA formation in a well-established AAA model of Ang II infused apoE null mice. This shares similarity with our previous findings establishing causal roles of endothelial DHFR deficiency and eNOS uncoupling in Ang II infused hph-1 mice [18]. Further, we demonstrated that an oral treatment of FA potently attenuates AAA formation in this model from 92% to 19.5%. In addition, Ang II infusion resulted in a large increase in superoxide production, reduced NO production, eNOS uncoupling, and reduction in DHFR expression and activity, all of which were restored by FA treatment. Taken together, these data suggest that FA’s protective effects against AAA development are tied to its ability to restore eNOS function. These data further confirm a significant role of uncoupled eNOS in mediating AAA formation and a universal efficacy of FA in preventing AAA formation via restoration of DHFR to restore eNOS function.

In a previous study in our laboratory, we identified a new model for AAA which is characterized by extensive uncoupling of eNOS [18]. In that study, FA was effective in eliminating the incidence of AAA in the animals. Hence, it is important to examine whether eNOS uncoupling plays a role in the development of AAA in other models. In the current study we used the Ang II infused apoE null model of AAA that does not begin with any deficiencies in eNOS cofactor pathways. While it has been reported that apoE null mice
do exhibit mild eNOS uncoupling activity at baseline, the general model is a lipid deficiency based model, which is different from our previous study. The results clearly show that FA treatment was protective against AAA, which suggests that FA is likely protective against AAAs in general. In additional experiments we found that FA did not affect lipid profiles during the treatment period.

Oxidative stress has been implicated in the development of AAA. In humans, it was reported that antioxidant systems are inhibited in aneurismal tissue when compared with non-aneurismal sections [11,12]. However, clinical trials using antioxidants such as vitamin E did not result in any significant benefit in terms of AAA prevention [25,26]. In this study, while FA did reduce oxidative stress as measured by superoxide production, it is important to note that it also resulted in the restoration of eNOS function. The restoration of eNOS not only reduces superoxide through inhibition of eNOS-derived superoxide production, but also increases NO, which is generally protective in attenuating vascular remodeling. These results match well with our previous study, where the restoration of eNOS was the key in the prevention of AAA development in the hph-1 mice [18]. Taken together, these studies suggest that the restoration of eNOS function could be a possible therapeutic target for the prevention of AAA.

Previous studies using apoE/eNOS double knockout mice demonstrated AAA development when exposed to a Western-type diet for 16 weeks [15,27], which suggests that eNOS is protective against AAA. In the current study, we used Ang II to induce AAA, which has been shown to result in eNOS uncoupling in this study as well as previous works [21,24,28,29]. Taken together, this suggests that depending upon the coupling state of eNOS, this enzyme can be protective while coupled and producing NO, or causal to the development of AAA while uncoupled and producing superoxide.

One limitation of this study is that for several of the assays, the entire aorta was used, while AAA is localized in the suprarenal region of the aorta. The reason for this is that these assays require more tissue than the size of the AAA. Further, the aneurismal sections sometimes included blood clots in the tissue that could not be removed. The presence of this blood will interfere with the nitric oxide and superoxide measurements. On the other hand, these data seem to suggest that adjacent aortic tissues share similar mechanisms of remodeling, and are indicative of AAA formation in these animals.

In conclusion, this study shows that oral treatment with FA is effective in reducing the incidence of AAA in the Ang II infused apoE null model of AAA. This protective effect can be attributed to an EC specific restoration of the enzyme DHFR, which improves the bioavailability of H4B to result in recoupling of eNOS. Taken together with our previous findings from the hph-1 model of AAA, these studies demonstrate that eNOS uncoupling plays an important role in the development of AAA that can be generally observed cross different model systems, and that FA can
and/or treatment of AAA via restoration of DHFR to restore serve as a universally effective treatment option for prevention and/or treatment of AAA via restoration of DHFR to restore}

**Author Contributions**

Conceived and designed the experiments: HC. Performed the experiments: KLS XNM. Analyzed the data: HC KLS XNM. Wrote the paper: HC KLS.

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