Introduction

Cyclooxygenase (COX) is expressed in two isoforms: constitutive COX-1 and inducible COX-2 [1]. COX-1 is associated with homeostatic processes including protection of the gastro-intestinal tract and thrombosis. COX-2 is induced by cytokines and mitogens, and contributes to inflammation, pain, angiogenesis and cancer. As such, COX-2 is the therapeutic target for non-steroidal anti-inflammatory drugs (NSAIDs) including diclofenac (Voltaren), and COX-2 selective NSAIDs including celecoxib (Celebrex) and rofecoxib (Vioxx). Drug development in this area, including study of the potentially promising chemopreventative benefits of COX-2 inhibitors in cancer, was dramatically halted in 2005 with the world-wide withdrawal of rofecoxib and the continued controversy surrounding the cardiovascular safety issues of these drugs.

In line with the protective roles of COX isoforms in the gut, traditional NSAIDs, which inhibit both isoforms of COX, are associated with serious and potentially fatal gastro-intestinal side effects [2,3]. We now know that gastro-intestinal side effects are a result of the combined inhibition of COX-1 and COX-2 in the gastric mucosa [4] and, as such, highly selective COX-2 NSAIDs, such as rofecoxib spare the gut [5]. Despite this proven advantage, rofecoxib was voluntarily withdrawn by its manufacturers when results of a placebo controlled clinical trial (the APPROVe study) suggested an increased risk of atherothrombotic events [6]. We now know that, in general, all drugs that inhibit COX-2, including traditional NSAIDs, which also inhibit COX-1, are associated with increased risk of cardiovascular events. This particularly manifests as an increase in rates of myocardial infarction [1,7,8,9,10], which is consistent with data from animal models [1,11,12,13]. Similarly, some studies show a pro-atherogenic phenotype of COX-2 inhibition [14,15] or deletion [13,16], although other pharmacological studies have suggested no change [17,18,19,20] or reduced atherosclerosis [21] with pharmacological blockade of COX-2. Taken together, we have no unifying explanation for the association between COX-2 inhibitor use and cardiovascular events in man.

Inhibition of urinary prostacyclin metabolites by drugs such as rofecoxib [22] and celecoxib [22] has been used as the main evidence to support the idea that COX-2 drives prostacyclin
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locally within the vessel wall [8,23]. Accordingly, inhibition of COX-2 derived prostacyclin release by NSAIDs has been suggested to remove a local protective block on platelet reactivity, vascular inflammation and atherosclerosis [8]. However, our group [24,25], and others [26,27,28], have demonstrated that it is COX-1 and not COX-2 that drives vascular prostacyclin production in a healthy cardiovascular system and that urinary metabolites do not reflect levels in the circulation [24,29]. We have gone on to use an unbiased systems analysis of COX-2 gene expression and found that, whilst COX-2 is not in large vessels, it is expressed in discrete hot spots including within the thymus, kidney and brain [30]; all locations previously shown to be sites of COX-2 activity. These findings leave us with no clear explanation of how COX-2 inhibitors might influence cardiovascular health [24], especially in those without overt cardiovascular disease such as those participating in the APPROVe study [6] or in healthy mice [11].

Data from animal models and human tissue demonstrate that COX-2 can be induced by inflammation in atherosclerotic vessels [18,31,32,33,34]. In these conditions, therefore, COX-2 may provide an additional pathway for local production of protective prostacyclin. Atherosclerotic disease is likely to be prevalent in NSAID target populations, such as arthritis patients, and subclinical atherosclerosis is present in even apparently healthy individuals. As such, it is essential that the role of COX-2 in the prostanoid generation in atherosclerotic vessels be understood in order for us to move the field of COX-2 biology and NSAID development forward. Previous studies have provided mixed results as to how total prostacyclin formation is altered in animal models of atherosclerosis [35,36,37] but to our knowledge only one has addressed the role of particular COX isoforms in this process. Using a cholesterol-fed rabbit model, Wong et al. [38] demonstrated aortic prostacyclin and PGE2 formation to be unaltered by selective COX-2 inhibitors, but partially inhibited by a non-selective COX-1/COX-2 inhibitor. Although this was a model of mild fatty streak formation with only modest COX-2 induction and data using pharmacological tools must be interpreted cautiously, these results suggest that COX-1 remains the dominant isoform for prostacyclin generation in mildly atherosclerotic vessels. By contrast, recently published data from Tang et al. [39] show endothelial-specific deletion of COX-2 exacerbates atherosclerosis in an LDL receptor knockout model, suggesting a protective role for vascular COX-2-derived prostacyclin.

Here we have used a fat-fed apoE−/− mouse model to investigate the effect of genetic deletion of COX-2 on local prostacyclin release in profoundly atherosclerotic vessels. We have gone on to establish the extent to which global COX-2 protects against atherosclerosis and used microarray and pathway analysis techniques to identify COX-2 regulated gene networks, which may provide new targets for understanding how COX-2 regulates cardiovascular health. Whilst these experiments do not yet deliver definitive answers, this study is the first to demonstrate that COX-2 protects the cardiovascular system independently of vascular prostacyclin and shows that systems biology approaches are valid in this area for the identification of new targets for NSAID-induced cardiovascular risk. This will, in turn, allow for the development of safe, efficacious, next generation NSAIDs.

Methods

Animals and induction of atherosclerosis

ApoE−/−/COX-2−/− double knockout mice were generated by cross of COX-2−/− mice [40] with apoE−/− mice (C57Bl/6j background; Charles River, UK) for 2 generations to produce apoE−/−/COX-2−/− mice, which were then inter-crossed. The resulting apoE−/−/COX-2−/− progeny were identified by genomic PCR, with apoE−/−/COX-2+/− littersmates serving as controls. To accelerate atherosclerosis in these animals, 12 week old male and female apoE−/−/COX-2−/− and apoE−/−/COX-2+/− animals were fed a high fat, high cholesterol diet (21% fat, 1.25% cholesterol, cholate-free; D12108; Research Diets, USA) for 10 weeks. The reported effects of COX-2 deletion on atherosclerosis and vascular prostanoid release were apparent in both male and female mice.

At the termination of all studies, animals were killed by CO2 narcosis, and the vasculature immediately perfused with phosphate buffered saline (PBS) across the heart. For some studies, mice were additional perfused with 10% neutral buffered formalin at physiological pressure.

This study was carried out in strict accordance with the guidelines set out under Animals (Scientific Procedures) Act 1986. All procedures have been approved by Imperial College Ethical Review Panel and under a designated Home Office License (PPL 70/7013) and were carried out by Home Office approved personnel. All efforts were made to minimise suffering.

Optical projection tomography

Perfusion fixed aortic arches from apoE−/−/COX-2+/+ and apoE−/−/COX-2−/− mice were cleaned of peri-adventitial material and embedded in agarose (Invitrogen, UK); before dehydration in methanol and optical clearing in 66% benzyl alcohol, 34% benzyl benzoate (Acrros Organics, UK). Tissue morphology and plaque distribution was recorded by optical projection tomographic imaging of auto-fluorescent emission (405–445 nm excitation) using a Bioptons 3001 tomograph as previously described [41]. Briefly, 400 projection images were recorded at 0.9° increments of rotation, and the three-dimensional volume reconstructed by Hamming-filtered back-projection (NRecon; Skyscan, Belgium). Plaque volume was quantified using CTAn software (Skyscan, Belgium) by a blinded observer.

Histological analysis and immunohistochemistry

Brachiocephalic arteries were collected from perfusion fixed apoE−/−/COX-2+/+ and apoE−/−/COX-2−/− mice and post-fixed in 10% neutral buffered formalin (Sigma, UK) for 48 hrs, before dehydration and processing to paraffin wax. 5 μm thick serial sections were cut perpendicular to the vessel axis, and stained using Verhoeff’s van Gieson method (Elastic Stain Kit; Sigma, UK). For all stains, non-specific binding was blocked with 10% normal goat serum (Vector Labs, UK), 2.5% bovine serum albumin (Sigma, UK). Before staining for B220, heat induced epitope retrieval was applied and incubated as indicated- anti-Mac2, B220 and CD3 was performed according to the labeled streptavidin-biotin method using rehydrated paraffin sections. Lipidated intra-plaque voids, as previously described [42].

Immunohistochemistry for α-smooth muscle actin (αSMA), Mac2, B220 and CD3 was performed according to the labeled streptavidin-biotin method using rehydrated paraffin sections. Before staining for B220, heat induced epitope retrieval was performed using pH 10 Tris-HCl antigen retrieval solution (Sigma, UK). For all stains, non-specific binding was blocked with 10% normal goat serum (Vector Labs, UK), 2.5% bovine serum albumin (Sigma, UK) and 2.5% non-fat milk powder (Marvel, UK) in PBS (Sigma, UK). Unconjugated primary antibodies were applied and incubated as indicated- anti-αSMA (Sigma, UK): 1/400 dilution; 30 min incubation; anti-Mac2 (Cederlane Labs, USA): 1/6000 dilution, overnight incubation; anti-B220 (Biole-
Sudan IV lipid staining

The aortic tree, including its major branches, was dissected from apoE<sup>−/−</sup>/COX-2<sup>+/+</sup> and apoE<sup>−/−</sup>/COX-2<sup>−/−</sup> mice perfused with PBS. Vessels were cut open to reveal their luminal surface, and pinned en face onto Sylgard coated plates (Dow Corning, USA). After brief fixation (10 mins; 10% neutral buffered formalin; Sigma, UK), vessels were rinsed in 70% ethanol before incubation (20 mins, room temperature) with filtered sudan IV solution (0.5% in 50% ethanol, 50% acetone; Sigma, UK). The stain was differentiated in 80% ethanol (5 mins), and vessels rinsed in running water. Aortas were photographed and the sudan IV-stained area quantified by image analysis (Photoshop CS5 Extended; Adobe Systems, USA) by a blinded observer.

En face confocal imaging of the aortic arch

Perfusion fixed aortic arches were carefully dissected of peri-adventitial material and 2 mm rings of the proximal (between aortic root and base of the brachiocephalic artery) and mid-arch (between the base of the brachiocephalic artery and left carotid artery) removed for immunostaining. Briefly, aortic arch rings were treated with 0.1% Triton X-100 (Sigma, UK) and 20% normal goat serum (Vector Labs, UK; 12 hrs, room temperature) to permeabilize membranes and block non-specific binding, then incubated with primary antibodies raised against COX-1 (1:50; 4 hrs, room temperature; Cayman Chemical, USA) or COX-2 (1:50; 4 hrs, room temperature; Cayman Chemical, USA). Aortic rings were then treated with AlexaFluor-594-conjugated goat anti-rabbit IgG secondary antibodies (Invitrogen, UK) before incubation with AlexaFluor-488-conjugated anti-CD31 (1:100; 12 hrs; 4°C; Biolegend, UK) and DAPI (25 μg/ml; 5min room temperature; Invitrogen, UK) to label endothelial cells and cell nuclei, respectively. Tissues were washed thoroughly with PBS between incubations. After staining, aortic rings were cut open to reveal the luminal surface, mounted flat between a glass slide and coverslip with aqueous hard-set media (Vector Labs, UK) and pressed until the media had firmly set. The luminal surface of the arch was visualized with a Leica SP5 inverted confocal microscope at 40X objective magnification. Laser and gain settings were fixed at the beginning of each imaging protocol. Areas corresponding to lesser and greater curvature of the aortic arch were determined by tissue orientation and confirmed by the cell morphology in the CD31<sup>+</sup> endothelial cell layer. In these regions, Z-stacks were recorded which included the endothelial cell, intra-plaque and medial layers. These layers were identified based cell and nuclear morphology and orientation, and CD31 immunoreactivity. Images presented and analysed represent a single plane from each Z-stack which best corresponds to the layer of vessel under study (i.e. endothelial layer, intra-plaque layer or medial layer). Due to the uneven morphology of the atherosclerotic vessel, in some cases individual Z-stack planes may have cut across more than one vessel layer. COX-1 and COX-2 was quantified as mean fluorescence intensity using ImageJ software (National Institutes of Health, USA).

Aortic COX activity bioassays

Aortic arch, brachiocephalic artery and thoracic aorta (regions both with and without grossly visible plaque formation) were isolated from animals perfused with PBS. Vessels were carefully dissected into 2 mm rings and placed immediately into individual wells of a 96 well microtitre plate containing DMEM (Sigma, UK) containing Ca<sup>2+</sup><sup>2+</sup> ionophore A23187 (50 μM; Sigma, UK). Vessels were incubated at 37°C for 30 mins then conditioned media was removed and subsequently analysed by competitive immunoassay for 6-keto-PGF1α (a stable breakdown product of prostacyclin; Cayman Chemical, USA) and PGE2 (Gibso, France).

qPCR

Tissues (heart, lung, liver, spleen, thymus, kidney) were removed from fat-fed apoE<sup>−/−</sup>/COX-2<sup>+/+</sup> and apoE<sup>−/−</sup>/COX-2<sup>−/−</sup> mice and snap frozen, homogenised, and total RNA extracted using a silica column-based kit (Nucleospin RNA II; Machery-Nagal, UK). RNA was converted to cDNA by reverse transcriptase (Superscript II; Invitrogen, UK) with oligo(dT) primers (Invitrogen, UK) and qPCR performed using a Rotor-Gene Q instrument (Qiagen, UK). Ptg2 (probe ID: Mm00477214_m1), ptg2 (probe ID: Mm00478374_m1) and rgl1 (probe ID: Mm00444088_m1) gene expression levels were determined using TaqMan expression assays and quantified relative to levels of 18S rRNA (probe ID: Mm03928990_g1) using the comparative Ct method.

Microarray analysis

Total RNA extracted from apoE<sup>−/−</sup>/COX-2<sup>+/+</sup> and apoE<sup>−/−</sup>/COX-2<sup>−/−</sup> mouse lung, liver and thymus homogenates as above were subject to standard microarray procedures. Samples were converted to cDNA, fragmented, labelled, and hybridized to Illumina MouseRef-8v3 BeadChip arrays (Illumina, UK). Quality control and basic interpretation of data was performed at Bart’s and The London Genome Centre, Queen Mary, University of London before datasets were received for further analysis. Data were subsequently analysed for differential expression levels and gene pathways using the Ingenuity iReport service (Ingenuity Systems, USA) and statistical significance determined using the linear models for microarray data (LIMMA) modified t-test method. Full methodological details can be found at http://www.ingenuity.com/getireport/pdf/ingenuity_robust_unattended_microarray_analysis.

Blood cell and cytokine analyses

Venous blood was collected from the inferior vena cava. Plasma was separated from heparinized blood (10 UD/ml; Leo Laborato- ries, UK) and levels of IL-1β, IL-2, IL-4, IL-10, IL-12 (total), TNFα, KC and TNFα were measured using a multiplex immunoassay system (Meso Scale Diagnostics, USA). Plasma cholesterol and triglyceride levels and whole blood leucocyte counts were determined by a commercial veterinary diagnostics service (IDEXX Laboratories, UK).
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(a) COX2+/+  
(b) COX2+/+  
(c) COX2+/+  
(d) COX2+/+
The role of COX-2 enzyme activity on prostacyclin production by atherosclerotic vessels

It is generally accepted that the inflammation associated with atherosclerosis leads to the induction of COX-2 [18,31,32,33,34]. However, COX-1 remains detectable in atherosclerotic tissue and the relative contribution of COX-2 to prostacyclin production in atherosclerotic vessels remains unclear [38] [39]. This is a critical question since the prevailing dogma dictates that, in patients who are likely to have pre-existing atherosclerosis, COX-2 drives local vascular prostacyclin production [13], an idea that has been adopted by the American Heart Association and has influenced prescribing advice [46]. We found evidence of some COX-2-like immunoreactivity in atherosclerotic vessels alongside abundant COX-1-like immunoreactivity (Figure 2). This COX-2 staining was specific since it was lost in COX-2-deficient mouse vessels (Figure S2). Sequential (Z) scanning through vessels suggested COX-2-like immunoreactivity was most abundant in the endothelial and intraplaque layers and somewhat less present in the media (Figure 2; Figure S2), although the heterogeneous composition and uneven morphology of the diseased vessels complicates precise localisation. Further, immunohistochemical analysis does not allow quantitative comparison of different protein levels, and protein levels, in turn, may poorly reflect bioactivity. Consequently, in this study we have systematically assessed the ability of isolated atherosclerotic vessels from apoE−/− mice lacking COX-2 to release prostacyclin (measured as its stable breakdown product, 6-keto-PGF1α) and PGE2, and how this is altered by deletion of the COX-2 gene. In order to ensure that COX expression rather than the level of phospholipase A2 activation was the rate-limiting step in prostaglandin production we measured release after stimulation with Ca2+ ionophore A23187. Our analysis determined the role of COX-2 in prostacyclin release from segments of the carotid artery (Figure 2a), aortic arch (Figure 2b) and thoracic aorta, which were each heavily burdened with atherosclerosis (Figure 2c), as well as plaque-free areas of the thoracic aorta (Figure 2d).

Previous studies have provided mixed results as to how total prostacyclin formation is altered in animal models of atherosclerosis [35,36,37]. Here we observed similar levels of prostacyclin release between vessels burdened with atherosclerosis and those free of disease by gross examination (Figure 2c,d). Moreover, regardless of the level of atherosclerosis, COX-2 did not detectably contribute to prostacyclin production by these isolated vessels (Figure 2). Similar results were seen when PGE2 release by the same vessels was measured (Figure 2). The physical trauma of removal of the vessel from the animal, ex vivo assay conditions and A23187 stimulation result in maximum activation of the vessel, conditions which may not reflect those in vivo and which may confound interpretation of these results. In order to address this we have previously measured plasma levels of 6-keto-PGF1α, a marker we have previously found to reflect systemic vascular prostacyclin production [24]. 6-keto-PGF1α was readily detectable in plasma and the levels did not differ between apoE−/−/COX-2+/− (419±106 pg/ml) and apoE−/−/COX-2−/− mice (436±66 pg/ml; p = 0.90). These findings are consistent with previous reports from hyperlipidemic rabbits [38] and our previous observations in healthy mouse vessels where COX-2 depletion did not alter vascular prostacyclin release, unless vessels are treated with inflammatory or mitogenic stimuli to induce COX-2 [24] or plasma prostacyclin unless animals were treated with high dose of LPS [47]. Whereas COX-1 deletion removed prostacyclin release in vitro and in vivo by >90% [24,30]. Taken together, our data demonstrate that, despite modest enzyme induction, COX-2 does not meaningfully drive local vascular prostacyclin production in diseased plaque burdened atherosclerotic vessels or, as in healthy animals [24,30] contribute to global circulatory prostacyclin release in animals with atherosclerosis.
Figure 2. COX-2 deletion does not alter prostacyclin production by atherosclerotic vessels. Production of Prostacyclin and PGE$_2$ by isolated rings of brachiocephalic artery (a) and aortic arch (b), and from plaque-bearing (c) and plaque-free regions of thoracic aorta (d) from fat-fed apoE$^{-/-}$/COX-2$^{++;}$ and apoE$^{-/-}$/COX-2$^{++;}$ mice demonstrated little role for COX-2 in the production of either prostaglandin in atherosclerotic vessels. In agreement, en face confocal immunofluorescence imaging of the lesser curvature of the aortic arch (b) suggests that whilst both COX-1 and COX-2-like immunoreactivity was present in atherosclerotic vessels, the COX-1 isoform was the predominant isoform. Sequential (Z) scanning through en face vessel preparations suggested COX-2 was more abundant in the endothelial and intraplaque layers than the media. Images are merges of red (COX1 or COX2), green (CD31 – endothelial cells) and blue (DAPI – cell nuclei) channels. n = 4–6.
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shown to protect against atherogenesis [39], despite our findings that vascular COX-2 has no role for prostacyclin production in affected vessels. In order to explore these hypotheses, we profiled COX-1 and COX-2 gene expression in a panel of tissues from atherosclerotic apoE−/− mice to find sites where loss of COX-2 may influence pathophysiology. COX-2 message was differentially expressed across tissues with highest levels present in the thymus, and lowest in the liver (Figure 3a), results that are consistent with our recent observations using a COX-2 luciferase mouse revealing COX-2 gene hot spots in these tissues [30]. COX-1 expression was more consistent between tissues (Figure 3b) and in agreement with our previous observations of COX activity in tissues from healthy animals, was the dominant isoform expressed in all tissues examined. Interestingly, the thymus, the site of T-lymphocyte maturation, possessed not only the highest level of COX-2 expression, but also the lowest level of COX-1 expression, suggesting prostanooid production in this tissue may be relatively dependent on COX-2 in this model (Figure 3c).

Effect of COX-2 deletion on the transcriptome in atherosclerotic mice; identification of T-lymphocyte pathways

To generate new ideas about how COX-2 at distant sites might regulate atherosclerotic disease in the vessel wall we took a systems biology approach and measured the transcriptome of tissues from apoE−/−/COX-2−/+ and apoE−/−/COX-2−/− mice. We examined the thymus, the highest COX-2 expressing tissue found, the lung, which is highly vascularised and sensitive to inflammation, and the liver, the major site of lipid metabolism. We then performed pathway analysis of microarray data to understand how changes in gene expression might influence biological processes (Figure 4, Figures S3-S5). In all three tissues, gene pathways involved in lymphocyte function and antigen presentation were amongst those most consistently and significantly altered (Figure 4, Figures S3-S5). To validate the importance of these pathways, we measured inflammatory cell counts in blood and found increased numbers of circulating lymphocytes in atherosclerotic mice lacking COX-2, but normal levels of monocytes, neutrophils, and eosinophils (Figure 5a). Similarly, deletion of COX-2 was associated with an increase in plasma levels of the lymphocyte cytokines IL-2 and IL-12, as well as the neutrophil chemottractant, KC (Figure 5b). To determine if these changes extended to the atherosclerotic vessels themselves we examined T- (CD3+ and B-lymphocyte (B220+)) accumulation in histological sections. B-lymphocytes were rarely observed in atherosclerotic vessels (Figure 6a), but T-lymphocytes were present as focal accumulations in the adventitial layer (Figure 6b). Crucially, the number of T-lymphocytes present was increased in atherosclerotic vessels from apoE−/− lacking COX-2 (Figure 6b).

These findings suggest that T-lymphocytes may mediate enhanced atherosclerosis in COX-2-deficient apoE−/− mice. Mature T-lymphocytes themselves, however, do not normally express COX-2 [48,49,50], and indeed, it has been reported that T-lymphocyte-specific COX-2-deletion does not alter atherosclerosis in an apoE-deficient model [50]. By contrast, it is known that deletion or inhibition of COX-2 alters the development of T-lymphocytes in the thymus [51], a site we have identified as highly expressing COX-2 in these mice. Further, prostanooids are reported to regulate a range of T-lymphocyte functions including proliferation, apoptosis and Th1/Th2 balance [48]. This latter effect may be particularly relevant to our study because we observed that deletion of COX-2 produced an increase in pro-atherogenic Th1-type cytokines (IL-2 and IL-12) and genes (STAT1, IRF1) but not Th2-type cytokines (IL-4, IL-10). A full

Figure 3. COX gene expression in tissue from atherosclerotic apoE−/− mice. COX-2 (ptgs2) (a) and COX-1 (ptgs1) (b) gene expression levels measured by quantitative RT-PCR in a panel of tissues from atherosclerotic apoE−/− mice indicated that COX-1 was expressed at a similar level and was the dominant isoform across all studied tissues. COX-2 expression showed more variation across tissues, with the highest levels present in the thymus, which also had the greatest COX-2:COX-1 expression ratio of any tissue examined (c). n = 4-6. doi:10.1371/journal.pone.0098165.g003

These observations however leave us now with no explanation of how COX-2 inhibition or gene deletion exacerbates atherosclerosis. Thus, with the idea that COX-2 mediates prostacyclin release in healthy [24] and atherosclerotic vessels (this study) removed, we must now address the possibility that COX-2 in sites remote to the affected vessel can protect indirectly. When doing so, it must be remembered that COX-2 may be expressed not only in parenchyma, but also in organ specialized microvasculature or other stromal cells. Indeed, endothelial specific COX-2 has been
Figure 4. Deletion of COX-2 is associated with up-regulation of lymphocyte-related gene expression patterns, and down-regulation of rgl1 in tissues from fat-fed apoE^{-/-} mice. Microarray analysis indicated 82, 87 and 90 differentially expressed genes (DEGs), respectively, in thymus (a), liver (b) and lung (c) from fat-fed apoE^{-/-} and apoE^{-/-}/COX-2^{+/+} mice. Volcano plot summaries of these data sets illustrate the distribution of effect size and statistical significance. Gene expression differing between genotypes 1.2-fold with \( p < 0.05 \) highlighted in red, were considered for pathway analysis. Differential gene expression patterns demonstrated only weak overlap between tissues, but one gene, rgl1, was...
gene validation and interpretation of how the thymus and associated lymphocyte pathways may be linked to COX-2, NSAIDs and atherosclerosis is beyond the scope of this study and remains the subject of investigation. Nevertheless, these ‘proof of concept’ studies provide a direction by which the cardioprotective properties of COX-2 at sites distant to the vessel may be identified.

Figure 5. COX-2 deletion increases circulating levels of lymphocytes and lymphocyte-related cytokines in fat-fed apoE<sup>−/−</sup> mice.

Measurement of plasma levels of cytokines by multiplex immunoassay (a) demonstrated an increased in the lymphocyte-related cytokines IL-2 and IL-12 (total) and the neutrophil chemoattractant KC, but no change in levels of TNFα, IL-4, IL-10 or IFNγ. Cell counts in whole blood (b), measured in parallel, indicated an increase in circulating lymphocytes, but not monocytes, neutrophils or eosinophils. *; p<0.05 vs apoE<sup>−/−</sup>/COX-2<sup>−/−</sup> by unpaired t-test; n = 10–25.

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tissues (thymus, lung, liver), and the only common altered gene between them (Figure 4d; Figures S3-S5). This was confirmed in each tissue by qPCR (Figure 4e). The mechanism of rgl1 down-regulation is unclear since it was observed in tissues with both high (thymus, lung) and low (liver) COX-2 expression. This suggests the link between rgl1 and COX-2 must be mediated either (a) by a single COX-2 expressing cell type present in all tissues such as leucocytes or nerves, or (b) by a circulating factor or other systemic influence triggered by loss of COX-2 at a distant site.

There is little literature describing the function and regulation of rgl1 in mammalian systems. Rgl1 is a Ral guanine nucleotide exchange factor (RalGEF) closely related to RalGDS (Ral guanine nucleotide dissociation stimulator). RalGEFs are activated by Ras (and related molecules) and catalyse the activation of small G-proteins of the Ral family. The Ras/RalGEF/Ral pathway has been linked to many biological effects (reviewed in [52]) including proliferation, migration and inflammation (NFκB activation). If and how down-regulation of rgl1 by COX-2 deletion might regulate atherogenesis is unclear since reduced Ral signaling might be expected to reduce inflammation and proliferation. It should also be noted that Ras/RalGEF/Ral is a key pathway in cancer [52] and as such our findings may be of equal or greater relevance to understanding how NSAIDs exert their chemo-preventative effects. Clearly the mechanism(s) by which COX-2 regulates rgl1 expression and the biological consequences of this warrant further investigation.

**Summary and conclusions**

Our previous work showed that COX-1, and not COX-2, mediates prostacyclin release in healthy vessels, and that urinary metabolites of prostacyclin do not reflect those in the circulation [24]. Here, we extend these observations, making the critical observation that COX-2 protects against atherosclerosis, but that this protection is mediated at some distant site and independent of local prostacyclin by the vessel. With the idea that COX-2 mediates vascular prostacyclin release removed, we must now revisit the question of where COX-2 is expressed and how this might influence vascular disease at distant sites. In this study we have performed microarray analysis of gene expression and used a systems biology approach to identify lymphocyte pathway signatures. This leads us to propose a role for the thymus and T-lymphocytes in the increased atherosclerosis observed in mice lacking COX-2. In addition, we have identified a novel target gene, rgl1, which is strongly linked to COX-2 in tissue from atherosclerotic mice. The importance of this study is to show that in a model where COX-2 protects against atherosclerosis, this is independent of prostacyclin at the site of vascular lesions. Our study does not provide definitive answers but offers important new hypotheses for investigation and illustrates approaches that will, in the future, lead to a full understanding of how COX-2 might protect the cardiovascular system. NSAIDs remain the most common over the counter medications taken worldwide and there is a clear
requirement for the continued development of new drugs in this class. These should spare the cardiovascular system as well as the gut, to fulfill their potential as anti-inflammatory drugs as well as in new clinical indications such as cancer.

Supporting Information

Figure S1 COX-2 deletion increases plaque macrophage but not lipid and smooth muscle content. Atherosclerotic lesions in the brachiocephalic artery of fat-fed apoE
−/−/COX-2+/+ and apoE
−/−/COX-2−/− mice were examined for estimated lipid content, smooth muscle/ myofibroblast content and macrophage content respectively by measuring intra-plaque voids in elastic Van Gieson (EVG) stained histological sections (a), or by immunohistochemistry for α-smooth muscle actin (αSMA; b) and Mac2 (c). Intra-plaque voids reminiscent of extracellular cholesterol crystals and Mac2 immunoreactivity were abundant in the intimal layer of all student vessels but rarer in the medial and adventitial layers, whereas αSMA was present in both the intimal and medial layers. The relative abundance of lipid-like voids and αSMA immunoreactivity was not altered by COX-2 deletion. By contrast, vessels from apoE
−/− mice lacking COX-2 exhibited significantly more Mac2 immunoreactivity than vessels from control mice. *; p < 0.05 vs apoE
−/−/COX-2+/+ by unpaired t-test; n = 8–10.

Figure S2 COX-2 is expressed in the atherosclerotic vessel and absent in COX-2−/− tissue. Atherosclerotic vessels from the lesser curvature of the aortic arch exhibited some COX-2-like immunoreactivity when examined by en face confocal immunofluorescence imaging. Sequential (Z) scanning through the vessel suggested this was present primarily in the endothelial and intraplaque layers with less present in the medial layers. In each case this was lost in COX-2-deficient mice demonstrating the antibody specificity. *; p < 0.05 by two-way ANOVA. n = 4–6.

Author Contributions

Conceived and designed the experiments: JAM NSK. Performed the experiments: NSK ML WRW. Analyzed the data: NSK WRW MJ. Contributed reagents/materials/analysis tools: TDW. Wrote the paper: JAM NSK.

References

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Figure S3 Effect of COX-2 deletion on the transcriptome of the thymus in apoE
−/−/COX-2+/+ mice. Whole thymus from fat-fed apoE
−/−/COX-2+/+ and apoE
−/−/COX-2−/− mice was examined for differential gene expression by microarray analysis. Genes exhibiting >1.25-fold expression level between genotypes are displayed. Data was analysed by the linear models for microarray analysis method. n = 4.

Figure S4 Effect of COX-2 deletion on the transcriptome of the lung in apoE
−/−/COX-2+/+ mice. Whole lung from fat-fed apoE
−/−/COX-2+/+ and apoE
−/−/COX-2−/− mice was examined for differential gene expression by microarray analysis. Genes exhibiting >1.25-fold expression level between genotypes are displayed. Data was analysed by the linear models for microarray analysis method. n = 4.

Figure S5 Effect of COX-2 deletion on the transcriptome of the liver in apoE
−/−/COX-2+/+ mice. Whole liver from fat-fed apoE
−/−/COX-2+/+ and apoE
−/−/COX-2−/− mice was examined for differential gene expression by microarray analysis. Genes exhibiting >1.25-fold expression level between genotypes are displayed. Data was analysed by the linear models for microarray analysis method. n = 4.

Author Contributions

Conceived and designed the experiments: JAM NSK. Performed the experiments: NSK ML WRW. Analyzed the data: NSK WRW MJ. Contributed reagents/materials/analysis tools: TDW. Wrote the paper: JAM NSK.


