Dectin-1/2–induced autocrine PGE$_2$ signaling licenses dendritic cells to prime Th2 responses

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Abstract

The molecular mechanisms through which dendritic cells (DCs) prime T helper 2 (Th2) responses, including those elicited by parasitic helminths, remain incompletely understood. Here, we report that soluble egg antigen (SEA) from *Schistosoma mansoni*, which is well known to drive potent Th2 responses, triggers DCs to produce prostaglandin E$_2$ (PGE$_2$), which subsequently—in an autocrine manner—induces OX40 ligand (OX40L) expression to license these DCs to drive Th2 responses. Mechanistically, SEA was found to promote PGE$_2$ synthesis through Dectin-1 and Dectin-2, and via a downstream signaling cascade involving spleen tyrosine kinase (Syk), extracellular signal-regulated kinase (ERK), cytosolic phospholipase A$_2$ (cPLA$_2$), and cyclooxygenase 1 and 2 (COX-1 and COX-2). In addition, this pathway was activated independently of the actions of omega-1 (ω-1), a previously described Th2-priming glycoprotein present in SEA. These findings were supported by in vivo murine data showing that ω-1–independent Th2 priming by SEA was mediated by Dectin-2 and Syk signaling in DCs. Finally, we found that Dectin-2$^{−/−}$ mice, displayed impaired Th2 responses and reduced egg-driven granuloma formation following *S. mansoni* infection, highlighting the physiological importance of this pathway in Th2 polarization during a helminth infection. In summary, we identified a novel pathway in DCs involving Dectin-1/2-Syk-PGE$_2$-OX40L through which Th2 immune responses are induced.
Author summary

T helper 2 (Th2) responses, which are initiated by dendritic cells (DCs), can cause allergic diseases, but they can also provide protection against metabolic disorders and parasitic helminth infections. As such, there is great interest in better understanding how their activity is induced and regulated by DCs. Parasitic helminths can potently induce Th2 responses. However, how helminths condition DCs for priming of Th2 responses remains incompletely understood. Here, we find that egg antigens from the parasitic helminth Schistosoma mansoni bind to pattern-recognition receptors (PRRs) Dectin-1 and Dectin-2 on DCs. This binding triggers a signaling cascade in DCs that results in synthesis of eicosanoid prostaglandin E2 (PGE_2). PGE_2 is sensed by the DCs themselves, resulting in expression of OX40 ligand (OX40L), which subsequently enables the DCs to promote Th2 differentiation. We show that this pathway is activated independently of omega-1 (ω-1), which is a glycoprotein secreted by the eggs and previously shown to condition DCs for priming of Th2 responses. Moreover, we demonstrate that this ω-1–independent pathway is crucial for Th2 induction and egg-driven immunopathology following S. mansoni infection in vivo. In summary, we identified a novel pathway in DCs involving Dectin-1/2–induced autocrine PGE_2 signaling through which Th2 responses are induced.

Introduction

Dendritic cells (DCs) are key players in the immune system because of their unique capacity to prime antigen–specific T helper 1 (Th1), Th2, Th17, or regulatory T cell (Treg) responses tailored against the pathogen they encounter [1–3]. It is well known that allergens and parasitic helminths can evoke strong type 2 immune responses, which largely depend on DCs that prime Th2 responses [4–8]. However, the molecular mechanisms through which DCs prime Th2 responses are still not fully defined.

Soluble egg antigen (SEA) from Schistosoma mansoni is a widely used antigen mixture to study Th2 responses to helminths. SEA is well recognized for its ability to condition DCs for priming of Th2 responses [9–13]. Omega-1 (ω-1), a glycosylated T2 RNase [14] present in SEA, was found to be a major Th2-polarizing molecule [9,15–18]. Mechanistic studies revealed that ω-1 is bound and internalized via its mannose receptor (MR) and that, following uptake, ω-1 impairs protein synthesis in an RNase-dependent manner that is essential for conditioning of DCs for Th2 polarization [9,10]. However, while ω-1 by itself was sufficient to condition DCs for Th2 polarization, SEA from which ω-1 was depleted still retained most of its Th2 priming potential both in vitro and in vivo. Moreover, eggs in which ω-1 expression was silenced [19] retained most of their Th2-polarizing potential, suggesting that there are additional mechanisms through which DCs become conditioned by schistosome eggs to prime Th2 responses [20].

Lipid mediators (LMs)—which arise from the enzymatic oxidation of polyunsaturated fatty acids (PUFAs), such as arachidonic acid (AA), docosahexaenoic acid, or linoleic acid—play an important role in immunological responses. In particular, prostanoids such as tromboxanes and prostaglandins (PGs)—which are derivatives of AA and are primarily released by myeloid cells, including macrophages and DCs—have been shown to have to capacity to influence immune cells by affecting their migration, differentiation, effector function, and/or polarization [21–24]. We recently performed a detailed analysis of the lipidome of the different life stages of S. mansoni and found that the eggs and egg-derived antigen preparations had a unique lipid profile characterized by the presence of various PGs and leukotrienes [25]. Thus
far, efforts to identify molecules responsible for Th2 polarization by helminths have primarily focused on glycans and (glyco)proteins. Whether LMs directly derived from schistosomes, or derived from immune cells in response to infection by this parasite, may additionally affect immune polarization remains unknown.

To identify potential novel pathways through which Th2 responses are induced by S. mansoni, we set out to assess the role of PUFAs and LMs in S. mansoni egg-driven Th2 polarization. We here report that SEA, apart from containing various LMs, induces DCs to generate several PGs and leukotrienes, including prostaglandin E2 (PGE$_2$), independently of ω-1. We show that this de novo synthesis of PGE$_2$ by SEA-stimulated DCs is driven by signaling through Dectin-1 and Dectin-2 and is crucial for Th2 priming. Mechanistically, we provide evidence that this PGE$_2$ through autocrine signaling induces OX40 ligand (OX40L) expression, to license DCs to prime Th2 responses. Finally, we show that this pathway is also crucial for Th2 priming by S. mansoni in vivo.

Results

SEA contains PGE$_2$ and promotes its synthesis by human monocyte-derived DCs independently of ω-1

In a recent lipidome analysis, we identified the presence of various LMs in SEA [25]. To explore the immunological significance of these findings, we first quantified these LMs in SEA using a sensitive liquid chromatography tandem mass spectrometry (LC-MS/MS)–based platform. We discovered that SEA contains 22 out of 55 monitored analytes (S1 Table), including docosahexaenoic acid, linoleic acid, and AA in relative high abundance, but also PGs such as PGE$_2$ and PGD$_2$ (S1 Fig, panel A). To assess potential consumption or uptake of such lipids by DCs, we tested supernatants of monocyte-derived DCs (moDCs) at 0, 6, 12, and 24 h after stimulation with lipopolysaccharide (LPS), LPS plus SEA, and LPS plus ω-1. We observed that levels of the majority of the LMs present in SEA decreased over time in DC culture supernatants following SEA stimulation, indicative of consumption/uptake or degradation (S1 Fig, panel B). We also observed that some of these lipids (i.e., 13-Hydroxyoctadecadienoic acid [13-HODE], lipoxin A$_4$ [LXA$_4$], PGD$_2$, and PGE$_2$) were accumulating over time in DC culture supernatants in response to SEA stimulation both in the presence (S1B Fig and quantitated for PGE$_2$ in Fig 1A) or absence of LPS (S1 Fig, panel C,D), suggestive of active production by moDCs in response to SEA. Stimulation of moDCs with LPS alone or LPS plus ω-1 did not drive accumulation of any of these compounds in the supernatants (S1 Fig, panel B). These results show that SEA contains a wide range of LMs as well as induces the release of particular LMs by DCs, including PGE$_2$.

Th2 polarization by SEA is dependent on PGE$_2$ synthesis by moDCs in absence of ω-1

To test whether LMs present in SEA or generated by moDCs upon stimulation with SEA contribute to Th2 polarization by SEA, we stimulated moDCs with several of these LMs—in concentrations similar to those found in SEA or in supernatants of SEA-stimulated moDCs—and analyzed their ability to condition DCs to induce Th2 polarization. Amongst all tested lipids, we identified PGE$_2$ as the only lipid capable of inducing Th2 polarization (Fig 1B). Based on this observation and given that ω-1 did not promote PGE$_2$ synthesis by moDCs (Fig 1A), we wondered whether PGE$_2$ may play a role in the previously observed ω-1-independent ability of SEA to prime Th2 responses [20]. To test this, we depleted ω-1 from SEA. Alongside ω-1, IPSE/α-1, which is another glycoprotein present in SEA but without Th2-priming capacity
[20], was also depleted from this preparation. We found that treatment of moDCs with SEAΔα-1/ω-1 reduced expression of the Th1-polarizing cytokine interleukin 12 (IL-12) induced by LPS (Fig 1C) and promoted Th2 polarization as determined by increased IL-4, IL-5, and IL-13 secretion by T cells (Fig 1D). The Th2-polarizing effect of SEAΔα-1/ω-1 occurred both in the presence and absence of LPS (S2 Fig, panel A) but was less potent than Th2 polarization by total SEA (Fig 1D and S2 Fig, panel A). In contrast to complete SEA, we could hardly detect PGE2 in SEAΔα-1/ω-1 itself (time point 0 h in Fig 1E versus in Fig 1A), which suggests that during the depletion step of ω-1 and α-1 from SEA, PGE2 was removed from SEA as well. We observed that SEAΔα-1/ω-1 still promoted PGE2 synthesis in moDCs both with (Fig 1E)
and without the presence of LPS (S1 Fig, panel D). This shows that SEA, in addition to containing PGE$_2$ itself, stimulates PGE$_2$ secretion by moDCs in an $\omega$–1–independent fashion.

Next, we investigated the contribution of the synthesized PGE$_2$ by SEA-stimulated moDCs to $\omega$–1–independent Th2 induction. Strikingly, when PGE$_2$ was neutralized using a specific anti-PGE$_2$ antibody during stimulation of moDCs with SEAD$\alpha$–1/$\omega$–1, the ability of SEAD$\alpha$–1/$\omega$–1–stimulated moDCs to drive Th2 polarization was totally lost (Fig 2A). In contrast, neutralization of PGE$_2$ in cultures of moDCs stimulated with $\omega$–1 or complete SEA had no effect on the Th2-priming potential of these cells, which is consistent with our recently published study showing that $\omega$–1—either alone or in the context of SEA—can prime Th2 responses via other mechanisms [9,10]. Moreover, we found that later neutralization of PGE$_2$ limited to the coculture of SEAD$\alpha$–1/$\omega$–1–stimulated moDCs with T cells did not impair Th2 polarization (Fig 2B), indicating that PGE$_2$ synthesized by moDCs does not act as a polarizing signal on T cells but rather directly conditions moDCs in an autocrine manner to acquire a Th2-priming phenotype. In line with this observation, we found that simultaneous inhibition of the 2 major receptors of PGE$_2$—prostaglandin E$_2$ receptor 2 (EP2) and EP4—on moDCs reduced the ability of SEAD$\alpha$–1/$\omega$–1–stimulated moDCs to induce a Th2 response (Fig 2C). These results collectively demonstrate that SEA, independently of $\omega$–1, promotes PGE$_2$ synthesis by moDCs.

Fig 2. $\omega$–1–independent Th2 polarization by SEA is dependent on PGE$_2$ synthesis by moDCs. (A–C) T-cell polarization assay as described in Fig 1B. (A) Neutralizing anti-PGE$_2$ antibody was added during stimulation of moDCs with indicated reagents or (B) during DC–T cell coculture. (C) EP2 and EP4 receptor inhibitors (EP2-I and EP4-I) were added during stimulation of moDCs with indicated stimuli. (A–C) Left: representative flow cytometry plots are shown of intracellular staining of CD4$^+$ T cells for IL-4 and IFN-$\gamma$. Numbers in plots represent frequencies of cells in indicated quadrants. Right: these data were used to calculate the fold change in frequency of IL-4$^+$ and IFN-$\gamma$ T cells polarized by moDCs stimulated with indicated stimuli relative to the cytokine production by T cells polarized by LPS-stimulated moDCs, for which the values were set to 1. Bars represent mean ± SEM of at least 4 independent experiments. Significance was calculated based on the ratio of IL-4 over IFN-$\gamma$ between conditions. *$P < 0.05$ and **$P < 0.01$ for significantly different from control conditions based on paired analysis (paired Student t test). Underlying data can be found in S1 Data. $\omega$–1, omega-1; CD4, cluster of differentiation 4; EP2, prostaglandin E$_2$ receptor 2; IL-4, interleukin 4; IFN$\gamma$, interferon $\gamma$; LPS, lipopolysaccharide; moDC, monocyte-derived DC; PGE$_2$, prostaglandin E$_2$; SEA, soluble egg antigen; Th2, T helper 2.
which subsequently, in an autocrine manner, conditions these cells to acquire a Th2-polarizing phenotype.

OX40L is induced by SEA via PGE$_2$ signaling and is required for Th2 induction

moDCs matured in the presence of PGE$_2$ are characterized by the expression of OX40L, a costimulatory molecule linked to Th2 polarization [26–28]. Moreover, an earlier study showed that moDCs stimulated with SEA express OX40L [3]. Indeed, we observed that stimulation of moDCs with PGE$_2$, SEA, or SEA$\Delta \alpha -1/\omega -1$ induced expression of OX40L on moDCs either in the presence (Fig 3A) or absence of LPS (S2 Fig, panel B), whereas $\omega -1$ did not induce OX40L expression (Fig 3A). While both SEA$\Delta \alpha -1/\omega -1$ and SEA promote PGE$_2$ synthesis, OX40L induction by SEA$\Delta \alpha -1/\omega -1$ was lower than the levels induced by SEA. This might be explained by the fact that in contrast to SEA$\omega -1/\alpha -1$, SEA additionally contains pre-existing PGE$_2$ itself, resulting in higher overall concentrations of PGE$_2$ that SEA-stimulated DCs are exposed to compared to SEA$\omega -1/\alpha -1$–primed DCs. Neutralizing PGE$_2$ prevented the induction of OX40L expression by SEA$\Delta \alpha -1/\omega -1$ (Fig 3B and S2 Fig, panel C). Finally, neutralizing OX40L

![Fig 3. OX40L is induced by SEA via PGE$_2$ signaling and is required for Th2 induction.](https://doi.org/10.1371/journal.pbio.2005504.g003)
during the coculture with T cells significantly reduced the Th2-polarizing capacity of SEAΔα-1/ω-1–primed moDCs (Fig 3C and S2 Fig, panel D). These data demonstrate that OX40L expression by SEAΔα-1/ω-1–conditioned moDCs is dependent on PGE₂ and that subsequently, induction of OX40L is important for Th2 polarization.

**SEA promotes PGE₂ synthesis and drives Th2 polarization via Dectin-1 and Dectin-2 in human moDCs**

Classically, central to the synthesis of PGE₂ is the release of AA from membrane phospholipids by cytosolic phospholipase A₂ (cPLA₂), which can then be converted into PGs, including PGE₂, through constitutively expressed cyclooxygenase 1 (COX-1) and stimulus-induced COX-2. We observed that SEA induced a small but significant increase in cPLA₂ activity (Fig 4A). We found that SEA did not change protein expression of COX-1, which was consistently expressed in all conditions. Moreover, SEA did not appear to promote COX-2 expression nor to alter LPS-driven COX-2 expression (Fig 4B), suggesting that SEA primarily promotes PGE₂ synthesis through induction of cPLA₂ activation. Indeed, selective inhibition of cPLA₂ activity using pyrrophenone attenuated SEAΔα-1/ω-1–induced PGE₂ synthesis (Fig 4C). For these experiments in which we analyzed the signaling events leading to PGE₂ synthesis, we used PGE₂-free SEAΔα-1/ω-1 and not PGE₂-containing complete SEA in order to be able to selectively assess de novo synthesis of PGE₂ by DCs. In addition, both COX-1 and COX-2 were important for PGE₂ synthesis by SEAΔα-1/ω-1, as treatment of moDCs with COX-1 and COX-2 inhibitors indomethacin and SC236 abrogated SEAΔα-1/ω-1–driven PGE₂ release (Fig 4C). IL-4 is known to suppress cPLA₂ expression via IL-4Rα/γc signaling, while IL-13–driven signaling through IL-4Rα/IL-13Rα has been described to promote cPLA₂ activity [29,30]. Because we cultured moDCs in the presence of IL-4, we wondered whether SEA might increase cPLA₂ expression/activity by interfering with IL-4Rα/γc and/or enhancing IL-4Rα/IL-13Rα signaling to promote PGE₂ synthesis. However, when we generated moDCs using IL-13 (S3 Fig, panel A)—which only signals through IL-4Rα/IL-13Rα—SEAΔα-1/ω-1 still promoted PGE₂ production to similar levels as by IL-4–cultured moDCs (S3 Fig, panel B), suggesting that the SEA-induced increase in cPLA₂ activity occurs through a different mechanism. Given that SEA has previously been reported to promote phosphorylation of extracellular-signal-regulated kinase (ERK) [31] and that ERK can drive activation of cPLA₂ [32], we evaluated the role of ERK in SEA-driven PGE₂ synthesis. We found that SEA, in contrast to ω-1–induced phosphorylation of ERK (Fig 4D) and that inhibition of ERK signaling, using U0126, abrogated PGE₂ synthesis induced by SEAΔα-1/ω-1 (Fig 4E).

Next, we aimed to identify the receptors through which SEA activates this pathway leading to PGE₂ synthesis in moDCs. Previous studies have identified various c-type lectin receptors (CLRs) through which SEA can be recognized by antigen-presenting cells (APCs). For human moDCs primarily fucose- and mannose-binding dendritic cell-specific intercellular adhesion molecule-3-grabbing non-integrin (DC-SIGN) and the MR have been implicated in this process [13,33–35]. Moreover, studies with murine APCs have also pointed to a possible role in recognition of components in SEA for Dectin-1 and Dectin-2, which are classically known for their ability to bind and respond to β-glucans and α-mannans from fungal origins, respectively [36]. Yet whether components within SEA can be recognized and induce signaling via human Dectin-1 and/or Dectin-2 expressed by DCs remains to be determined. As a first step towards the identification through which receptor(s) SEA induces PGE₂, we determined which of these receptors are involved in binding of SEA by moDCs. In line with earlier observations [9,13], blocking of the MR reduced binding of fluorescently labelled SEA, which could be further reduced when DC-SIGN binding was neutralized simultaneously. Blocking of both Dectin-1
Fig 4. SEA promotes PGE₂ synthesis and drives Th2 polarization via signaling through Dectin-1 and Dectin-2 in human moDCs. (A) cPLA₂ activity 8 h after stimulation. Zymosan was taken along as a positive control for cPLA₂ activation. (B) Protein expression of COX-1 and COX-2 were assessed by western blot; β-actin was used as housekeeping protein. One of 3 experiments is shown. (C) Following 1 h pre-incubation with specific inhibitors for cPLA₂ (Pyr.) or COX-1 and COX-2 (SC and ind., respectively), moDCs were stimulated for 12 h with LPS plus SEAΔα/ω-1, and supernatants were collected for PGE₂ determination by LC-MS/MS. (D) At the indicated time points after stimulation with depicted stimuli, phosphorylation of ERK was...
and Dectin-2 in conjunction with MR plus DC-SIGN neutralization further reduced binding relative to blocking of just MR plus DC-SIGN, suggesting that all 4 receptors contribute to recognition of glycans or glycoproteins present in SEA (Fig 4F). Next, we aimed to identify through which of these CLRs SEA promotes PGE\(_2\) synthesis. We found that blocking of either Dectin-1 or Dectin-2, but not DC-SIGN, strongly attenuated PGE\(_2\) synthesis induced by SEAΔα-1/ω-1, while blocking the MR also resulted in reduced PGE\(_2\) synthesis albeit to a lesser extent. This suggests a major role for Dectin-1 and Dectin-2 in SEAΔα-1/ω-1–driven PGE\(_2\) synthesis (Fig 4G).

Dectins phosphorylate and activate spleen tyrosine kinase (Syk) through the immunoreceptor tyrosine-based activation motifs (ITAMs) present in their cytoplasmic domain (Dectin-1) or in the recruited Fc gamma receptor (FcRG) chain (Dectin-2). Syk in turn can promote ERK phosphorylation [37]. Indeed, we observed that SEA stimulation resulted in phosphorylation of Syk, which was dependent on both Dectin-1 and Dectin-2 but not the MR (Fig 4H), and found that SEA-driven ERK phosphorylation was dependent on Dectin-1, Dectin-2, and Syk signaling (Fig 4I and 4J). Of note, blocking either Dectin-1 or Dectin-2 alone only had minor effects on Syk and ERK phosphorylation, while these signals were totally blunted when both Dectin-1 and Dectin-2 signaling were blocked simultaneously (Fig 4H and 4I), suggesting that SEA depends on both receptors to activate this pathway in human moDCs. In line with these findings, inhibition of Syk signaling blunted SEAΔα-1/ω-1–induced PGE\(_2\) synthesis (Fig 4K). Finally, blocking either Dectin-1 or Dectin-2 attenuated OX40L expression (Fig 4L) as well as the Th2 response induced by SEAΔα-1/ω-1 (Fig 4M). Because signaling through Dectin-1 and Dectin-2 by fungal pathogen associated molecular patterns (PAMPs) is known to lead to IL-23 production and priming of Th17 responses [38,39], we analyzed IL-23 secretion and Th17 induction by SEA–primed DCs. Although we found that SEA promoted IL-23 release by DCs (S4 Fig, panel A), this did not result in any discernible Th17 polarization (S4 Fig, panel B). In conclusion, these data suggest that components in SEA promote PGE\(_2\) synthesis by moDCs through the MR, Dectin-1, and Dectin-2 and via a signaling cascade involving Syk, ERK, cPLA\(_2\), COX-1, and COX-2 that is required for ω-1–independent Th2 induction by SEA.

**PGE\(_2\) isomer generation via autoxidation contributes to Th2 induction by SEA**

The observations that PGE\(_2\) promoted Th2 induction by SEA and that this PGE\(_2\) synthesis was dependent on COX activity led us to hypothesize that blocking of COX activity in SEAΔα-1/ω-1–driven PGE\(_2\) synthesis via autoxidation contributes to Th2 induction by SEA.
1–stimulated moDCs would abrogate their ability to prime Th2 responses. However, inhibition of COX-1 and COX-2 only partly reduced the Th2 response induced by these SEAΔα/ω–conditioned DCs (Fig 5A). A possible explanation for this unexpected result came from a careful reanalysis of the extracted ion chromatogram of the PGE$_2$ trace in which we noted that alongside PGE$_2$, moDCs stimulated with SEA produced PGE$_2$ isomers, also known as isoprostanes (IsoPs) (Fig 5B). Several studies have suggested that IsoPs may have similar properties as PGE$_2$ but that, in contrast to the latter, they are generated by an autoxidation process directly from AA fueled by reactive oxygen species (ROS), independently of COX activity [40]. We found IsoP isomers “1” and “2” (Fig 5B) to have a fragment ion m/z 189, characteristic for 15-series IsoPs. (D) Showing the MS/MS spectrum of IsoP 3, possibly identifying this IsoP as a 15-series IsoP, based on the fragment ions m/z 115, 217, and 191. (E) ROS generation was determined by flow cytometry (H$_2$-DCFDA) of moDCs pulsed for 6 h with indicated reagents. On the left, representative histograms for ROS induction are shown. On the right, the geometric mean fluorescence of these signals is enumerated and shown as fold change relative to unstimulated moDCs (dashed line set to 1). (F) ROS production was quantified as in panel E following pretreatment with general ROS scavenger NAC or R406 for 1 h. (G) moDCs were stimulated with indicated PGs with or without anti-PGE$_2$ after which a T-cell polarization assay was performed as described in Fig 2A. (H) moDCs were stimulated with indicated reagents in the presence following 1 h pre-incubation with COX inhibitors (S and I) and NAC after which a T-cell polarization assay was performed as described in Fig 2A. Bars represent mean ± SEM of at least 3 independent experiments. *** and #*: P < 0.005; ** and #: P < 0.01 for significant differences with the LPS control (*) or between-test conditions (#) based on paired analysis (paired Student t test). Underlying data can be found in S1 Data. COX, cyclooxygenase; H$_2$-DCFDA, 2',7'-dichlorodihydrofluorescein diacetate; I, indomethacin; IsoP, isoprostane; LC-MS/MS, liquid chromatography tandem mass spectrometry; LPS, lipopolysaccharide; moDC, monocyte-derived DC; NAC, N-acetyl-L-cysteine; PGE$_2$, prostaglandin E$_2$; ROS, reactive oxygen species; S, SC236; SEA, soluble egg antigen; Th2, T helper 2.

Fig 5. SEA-induced ROS production by moDCs results in PGE$_2$ isomer synthesis that contributes to Th2 polarization.

1–stimulated moDCs would abrogate their ability to prime Th2 responses. However, inhibition of COX-1 and COX-2 only partly reduced the Th2 response induced by these SEAΔα/ω–conditioned DCs (Fig 5A). A possible explanation for this unexpected result came from a careful reanalysis of the extracted ion chromatogram of the PGE$_2$ trace (m/z 351 → 271) in which we noted that alongside PGE$_2$, moDCs stimulated with SEA produced PGE$_2$ isomers, also known as isoprostanes (IsoPs) (Fig 5B). Several studies have suggested that IsoPs may have similar properties as PGE$_2$ but that, in contrast to the latter, they are generated by an autoxidation process directly from AA fueled by reactive oxygen species (ROS), independently of COX activity [40]. We found IsoP isomers “1” and “2” (Fig 5B) to have a fragment ion m/z 189 that is characteristic for 15-series IsoPs with identical relative retention times to commercially available 15β-PGD$_2$ and 11β-PGE$_2$, respectively, suggesting these isomers are 15β-PGD$_2$ and 11β-PGE$_2$ (Fig 5C) [41]. IsoP “3” showed a somewhat different tandem MS spectrum indicating it belongs to the class of 5-series IsoPs, but this could not be confirmed due to a lack of standard material (Fig 5D) [42].
Syk and Dectin-2 responses were suppressed by SEA BMDCs was not significantly affected (Fig 6E and S5 Fig, panel A,B). Furthermore, Th17 induction by DCs in response to egg antigen challenge in vivo, we made use of We next aimed to assess the importance of this Dectin–Syk signaling in mediating Th2 polarization by DCs in response to egg antigen mixture without Syk signaling in DCs play a key role in Th2 priming by SEA in vivo following adoptive transfer into wild-type (WT) recipient mice. We found that, following subcutaneous immunization with SEA, cluster of differentiation 4 (CD4)$^+$ T cells from draining lymph nodes (LNs) from CD11c$^{ΔSyk}$ mice, compared to CD11c$^{WT}$ controls, produced less Th2 cytokines ex vivo in response to both polyclonal (Fig 6A) and antigen-specific restimulation (Fig 6B), while interferon-γ (IFN-γ) production was not different between the 2 groups of mice. To explore the contribution of ω-1 in driving this Syk-dependent Th2 polarization by SEA in vivo, we additionally immunized these mice with SEAΔα-1/ω-1 and observed that CD11c$^{ΔSyk}$ mice, also in response to this egg antigen mixture without ω-1, failed to significantly drive Th2 polarization (Fig 6C). Skewing towards Th2 following immunization with ω-1, however, was unaffected by loss of Syk in the DCs (Fig 6D), showing that ω-1 drives a Th2 response through a Syk-independent route as well as ruling out a role for ω-1 in driving Syk-dependent Th2 polarization by SEA in vivo. This is in line with our in vitro findings with human moDCs and provides evidence that Syk signaling in DCs plays a key role in Th2 priming by S. mansoni egg antigens in vivo.

Next, to establish the role of Dectin-1 and/or Dectin-2 in driving this Syk-dependent response in vivo, we assessed the ability of SEAΔα-1/ω-1–pulsed bone marrow–derived DCs (BMDCs) derived from Dectin-1–deficient (Clec7a$^{-/-}$) and Dectin-2–deficient (Clec4n$^{-/-}$) mice to prime Th2 responses in vivo following adoptive transfer into wild-type (WT) recipient mice. We found that SEAΔα-1/ω-1–pulsed Dectin-2$^{-/-}$ BMDCs—just like SEAΔα-1/ω-1–pulsed CD11c$^{ΔSyk}$ BMDCs—failed to prime a Th2 response, whereas Th2 induction by Dectin-1$^{-/-}$ BMDCs was not significantly affected (Fig 6E and S5 Fig, panel A,B). Furthermore, Th17 responses were suppressed by SEAΔα-1/ω-1–pulsed WT BMDCs, which was even further reduced in draining LNs of mice immunized with SEAΔα-1/ω-1–pulsed Dectin-1$^{-/-}$ BMDCs (S6 Fig). Finally, in accordance with the impaired ability of Dectin-2$^{-/-}$ and CD11c$^{ΔSyk}$ BMDCs to prime Th2 responses, these cells—but not Dectin-1$^{-/-}$ BMDCs—displayed a reduced uptake of SEA (Fig 6F), ROS production (Fig 6G), and OX40L expression in response to SEAΔα-1/ω-1 stimulation (Fig 6H and S5 Fig, panel C-E). Together, these data reveal that Dectin-2 and Syk signaling in DCs play a key role in Th2 priming by S. mansoni egg antigens in vivo independently of ω-1.

Finally, we set out to assess the importance of this signaling axis in Th2 differentiation and Th2-driven immunopathology during a natural infection with S. mansoni, using Dectin-1$^{-/-}$ and Dectin-2$^{-/-}$ mice. During S. mansoni infection, adult worms residing in the portal vasculature release eggs that get trapped in the liver, where they induce strong Th2 responses that...
Th2 via Dectin-induced PGE2

Fig 6. Th2 polarization induced by SEA is mediated via Dectin-2 and Syk signaling in vivo. (A–D) WT or CD11cΔSyk mice were injected with SEA (panel A, B), SEAΔα-1/ω-1 (panel C), or ω-1 (panel D) in the hind footpad, and draining pLNs were analyzed 7 d later. (A, C, D) pLN cells were restimulated with PMA/Ionomycin in the presence of brefeldin A, and CD4+ T cells were stained for indicated intracellular cytokines and percentage cytokine-positive CD4+ T cells enumerated. Based on these data, the ratio between the percentage IL-4 over IFN-γ CD4+ T cells was determined as a measure for overall skewing towards Th2. (B) pLN cells were restimulated with SEA for 72 h, and cytokine levels in culture supernatants were determined. (A–D) Bar graphs represent mean ± SEM of 5 to 6 mice per group and are representative of 2 (panel A–C) or 1 (panel D) experiment. (E) BMDCs cultured from BM from WT, Dectin-1−/−, or Dectin-2−/− mice were pulsed overnight with SEAΔα-1/ω-1 and injected into hind footpads after which CD4+ T-cell responses were analyzed as in panel A. Representative flow cytometry plots of intracellular staining of CD4+ T cells are depicted, of which the data are enumerated in bar graphs representing mean ± SEM of 2 independent experiments with 3 to 6 mice per group. (F) SEA binding and uptake was determined as in Fig 4F. (G) ROS production by indicated BMDCs was determined as described in Fig 5E, 1 h after stimulation.
orchestrate the development of granulomatous lesions surrounding the eggs [8]. The intensity of the Th2 response and associated granulomatous inflammation peaks at 8 wk after infection. To compare the Th2 response induced by this infection in WT and Dectin-1<sup>−/−</sup> and Dectin-2<sup>−/−</sup> mice, cells from mesenteric LNs and spleens from 8-wk–infected mice were restimulated with SEA or anti-CD3/CD28. We found that infected Dectin-2<sup>−/−</sup> mice displayed lower production of Th2 cytokines IL-4 and IL-5 in both lymphoid organs in comparison to their infected WT counterparts, while IFN-γ production was not different between the 2 groups (Fig 7A and 7B). In infected Dectin-1<sup>−/−</sup> mice, only IL-5 production by splenocytes was reduced (S7 Fig, panel A,B). Of note, no SEA-specific Th17 responses, as determined by IL-17A production, were detected in spleen and mesenteric LNs of infected mice (Fig 7A and 7B and S7 Fig, panel A,B). In line with the reduced Th2 responses found in the infected Dectin-2<sup>−/−</sup> mice, granuloma size around the eggs trapped in the liver was smaller relative to infected WT mice (Fig 7C). This difference in Th2 response was not due to differences in infection load because both Dectin-2<sup>−/−</sup> and WT mice were found to harbor similar numbers of eggs and adult worms (Fig 7D and 7E). Dectin-1<sup>−/−</sup> mice did not show an altered granulomatous response towards liver eggs (S7 Fig, panel C), nor did they show differences in numbers of adult worms and eggs (S7 Fig, panel D,E). We previously found that inflammasome activation, which can be triggered by SEA via Dectin-2 signaling, can alter T-cell polarization and contribute to granuloma formation during <i>S. mansoni</i> infection [33]. However, levels in IL-1β protein, as readout for inflammasome activity, in liver were similar between Dectin-2<sup>−/−</sup> and WT mice (Fig 7F). Altogether, these findings highlight an important role for Dectin-2 in promoting Th2 differentiation and immunopathological outcome of this response during <i>S. mansoni</i> infection.

**Discussion**

The molecular mechanisms through which DCs prime Th2 responses, including those elicited by helminths, are still incompletely understood. We here explored the role of PUFAs and LMs in Th2 induction by <i>S. mansoni</i> eggs, which are well known for their potent ability to elicit strong Th2 responses. This enabled us to identify a novel signaling axis in DCs involving Dectin-Syk-PGE<sub>2</sub>-OX40L through which Th2 responses are induced in vitro and in vivo.

Some studies have documented that different life stages of <i>S. mansoni</i> are able to produce LMs from both COX products (e.g., PGE<sub>1</sub>, PGE<sub>2</sub>, PGD<sub>2</sub>, and PGA<sub>2</sub>) and Lipoxygenase products (e.g., Leukotriene B<sub>4</sub>, 5-Hydroxyeicosatetraenoic acid [5-HETE], 12-HETE) [40,43]. However, the existence of LMs in eggs or SEA had not been examined before. Here, we uncovered that SEA itself contains various PUFAs and LMs with potential immunomodulatory properties. In particular, the presence of the well-studied immunomodulatory eicosanoid PGE<sub>2</sub> caught our attention because, among the pleiotropic properties that have been attributed to this lipid, it has been associated with promoting Th2 polarization by functional modulation of DCs [3,27]. Moreover, we found PGE<sub>2</sub> not only to be present in SEA but also to be synthesized by DCs themselves in response to SEA stimulation. While other life cycle stages of <i>S. mansoni</i>
have been shown to promote PGE$_2$ synthesis in host cells such as cercariae in keratinocytes [44], we are the first to report and mechanistically investigate the ability of egg-derived antigens to promote PGE$_2$ synthesis in immune cells. We found that this PG, in contrast to several other LMs secreted by SEA-stimulated DCs, was not only sufficient to condition moDCs for Th2 polarization but also crucial for mediating ω-1--independent Th2 polarization by SEA. This identifies PGE$_2$ as a key factor through which SEA, independently of ω-1, primes Th2 responses. A question that remains to be answered is what the relative contribution is of the pre-existing PGE$_2$ (present in SEA) versus PGE$_2$ synthesized by moDCs in mediating the Th2-polarizing effect. The observation that SEA from which ω-1 was depleted was fully dependent on de novo--synthesized PGE$_2$ by the moDCs for Th2 polarization at least shows that moDC-derived PGE$_2$ can be sufficient for promoting a Th2 response. Moreover, the fact that SEA requires Syk signaling in DCs to prime a Th2 response in vivo would argue that PGE$_2$ derived from SEA itself is insufficient to condition DCs for Th2 priming and that SEA instead rather depends on Syk-driven de novo PGE$_2$ synthesis for this response.

Fig 7. Dectin-2 signaling is required for induction of a Th2 response during *S. mansoni* infection. WT and Dectin-2$^{-/-}$ mice were infected with *S. mansoni*. After 8 wk of infection, cells from spleens (A) or mLN (B) were restimulated with SEA or anti-CD3/CD28 for 72 h, and cytokine levels were analyzed in supernatants by Luminex or ELISA. Bars represent mean ± SEM of combined data of at least 2 or 3 independent experiments with 5 to 10 mice per group. (C) Granuloma sizes around eggs trapped in the liver of 8-wk–infected mice were assessed in Masson blue–stained liver sections. Data are based on 10 mice per group. Number of worms (D) and liver and intestinal eggs (E) in mice infected with *S. mansoni* for 8 wk. (F) IL-1β protein levels in livers of mice infected with *S. mansoni* for 8 wk. $^\ast P < 0.01$ and $^{***} P < 0.001$ for significant differences relative to the control mice based on unpaired analysis (unpaired Student t test). Underlying data can be found in S1 Data. CD3, cluster of differentiation 3; IL-1β, interleukin 1β; mLN, mesenteric lymph node; SEA, soluble egg antigen; Th2, T helper 2; WT, wild-type.

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Mechanistically, we found that PGE$_2$ derived from moDCs acts in an autocrine manner to promote Th2 polarization by promoting the expression of OX40L in moDCs. OX40L expression has been shown to be important for Th2 polarization by DCs stimulated with various other Th2-priming stimuli, such as allergens and thymic stromal lymphopoietin (TSLP) [28,45,46]. Congruent with these studies, we found that OX40L expression was crucial for Th2 polarization by SEA from which $\omega$-1 was depleted. While expression of OX40L in response to SEA has been documented before [3,12], we now provide evidence that SEA-induced OX40L expression in moDCs is secondary to its ability to induce PGE$_2$ synthesis by these cells.

Moreover, we found that SEA from which $\omega$-1 was depleted was dependent on signaling through both Dectin-1 and Dectin-2 to condition moDCs to drive Th2 polarization. Dectin-2 has been linked to Th2 polarization before in the context of allergic responses induced by house dust mite [47,48]. In these studies, Dectin-2 was found to mediate Th2 induction through the generation of cysteinyl leukotrienes by murine DCs. However, we did not observe induction of cysteinyl leukotrienes by SEA. Instead, we found that SEA interacts with Dectin-1, Dectin-2, and the MR to promote PGE$_2$ synthesis. Downstream of these receptors, we identified a pathway involving Syk, ERK, and cPLA$_2$, that leads to the release of AA, which subsequently acts as a substrate for COX to produce PGE$_2$. We currently do not have a clear explanation for the observations that PGE$_2$ synthesis and Th2 induction by human moDCs can be blocked by either anti–Dectin-1 or Dectin-2 antibody while full inhibition of Syk phosphorylation is only seen when both receptors are blocked. Possibly, this suggests that Syk phosphorylation needs to reach a certain threshold level before it can start relaying signals leading to PGE$_2$ synthesis and Th2 polarization. In this scenario, reducing Syk phosphorylation below that threshold (seen with single CLR blockade) would be sufficient to block PGE$_2$ synthesis/Th2 induction, and further inhibition of Syk phosphorylation would not have additional functional consequences.

The observation that the MR also seems to play a role in SEA-driven PGE$_2$ synthesis—despite the fact that the MR itself, in contrast to Dectin-1 and Dectin-2, does not harbor an intracellular signaling motif—leads us to speculate that the MR may collaborate with Dectin-1 and/or Dectin-2 to form heterodimers or multimers that effectively bind glycans or glycoproteins in SEA that allow for efficient activation of the signaling cascade downstream of Dectins resulting in PGE$_2$ synthesis. Associations of different CLRs to potentiate glycan-induced signaling have been described before for Dectin-2 and Dectin-3 [49]. Glycans derived from the cell wall of fungi such as *Candida albicans* [50] are well known to promote PGE$_2$ synthesis through this pathway via activation of Dectin-1, Dectin-2, and the MR [39,51]. However, in this context, the production of PGE$_2$ seems to contribute to Th17 priming by APCs and not Th2 [39]. This difference might be explained by differences in glycan repertoire between fungi and schistosome eggs. For instance, classical $\beta$-glucans expressed by fungi are not present in SEA [52]. Therefore, the carbohydrates in SEA that mediate Dectin binding may interact differently with, or have a lower affinity for, these receptors than fungal carbohydrates do. This may induce a qualitatively and/or quantitatively different signaling cascade that could trigger sufficient PGE$_2$ synthesis and OX40L expression to allow for Th2 induction to occur, without promoting the expression of high levels of cytokines required for Th17 polarization. Secondly, fungal Dectin agonists may trigger additional pattern-recognition receptors (PRRs) that are not activated by SEA to induce pro-inflammatory cytokine expression [53]. Finally, the immunological outcome of Dectin engagement can also be cell type dependent. For instance, Dectin-1 ligand curdlan was found to promote Th2 responses via plasmacytoid DCs, while this same ligand conditioned myeloid DCs to inhibit Th2 responses [54] or to promote Th9 responses [55]. Similar to our observations with SEA, the conditioning of plasmacytoid DCs by curdlan to promote Th2 responses was dependent on induction of OX40L expression [54].
Currently, studies are underway to identify the glycoproteins or glycan moieties present in SEA that bind to Dectin-1, Dectin-2, and the MR to promote this response and identify how these receptors collaborate to promote Syk activation and PGE\(_2\) synthesis.

We provide evidence that COX-independent generation of several PGE\(_2\) isomers (IsoPs) by SEA, independently of enzymatically synthesized PGE\(_2\), is capable of conditioning moDCs for Th2 priming. While COX-independent generation of these isomers has been described before, as a result of auto-oxidation of AA by free radicals [40], we now here show a role for these IsoPs in regulation of an immune response. In line with the free radical–dependent synthesis of IsoPs, we found that SEA could drive ROS production in a Syk-dependent manner, which corroborates a recent study showing that SEA can induce ROS in murine DCs [33]. However, the latter study focused on the role of ROS in SEA-driven inflammasome activation and did not report on other ROS-mediated effects. Our data suggest that enzymatically synthesized PGE\(_2\) and its ROS-induced isomers act in concert in Dectin-mediated conditioning of moDCs for Th2 priming by SEA. Moreover, we found that the widely used neutralizing anti-PGE\(_2\) antibody [56] that we have used in this study not only neutralizes PGE\(_2\) but also harbors cross-reactivity towards 2 of the main IsoPs that we found to be generated by SEA-stimulated DCs. This can explain our observation that PGE\(_2\) neutralization, in contrast to COX inhibition, did fully block Th2 polarization.

Our findings that murine DCs that are deficient for Dectin-2 or Syk display reduced OX40L expression following exposure to SEA and fail to mount a Th2 response in vivo provide strong support for a key role of the Dectin-Syk-PGE\(_2\)-OX40L axis in Th2 polarization by Schistosoma egg–derived antigens in vivo. Our studies with S. mansoni–infected Dectin-2\(^{-/-}\) mice suggest that also during natural infection, this signaling axis seems to be crucial for induction of Th2 responses. Nonetheless, our in vivo work can currently not formally exclude the possibility that apart from PGE\(_2\), there are other factors downstream of Dectin–Syk signaling that contribute to Th2 priming by S. mansoni. For instance, SEA has previously been reported to activate the Nlrp3-inflammasome in a Dectin-2–dependent manner, and inflammasome-deficient mice were shown to have an altered T-cell polarization profile and a reduction in granuloma size during S. mansoni infection similar to our observations reported here in infected Dectin-2\(^{-/-}\) mice. However, the fact that—in contrast to what was observed in inflammasome-deficient mice [33]—there was no reduction in total IL-1\(\beta\) levels in livers of Dectin-2\(^{-/-}\) mice during S. mansoni infection would suggest that, downstream of Dectin-2, the PGE\(_2\)-OX40L axis rather than inflammasome activation plays a key role in Th2 priming during, and in the immunopathological outcome of, this infection. Nonetheless, additional studies will be needed to definitively determine the individual contribution of each pathway to the immunopathology in vivo. Somewhat surprising was the observation that in contrast to the in vitro data with human moDCs, Dectin-1 appears to be less important for Th2 polarization in vivo in mice. This may suggest that murine Dectin-1, as opposed to its human counterpart, does not play a major role in recognition of glycans present in SEA, which, although currently still speculation, might be due to differences in glycan specificity or in expression of Dectin-1 isoforms between murine and human DCs [38,57]. More detailed comparative studies between the SEA-binding characteristics of human and murine Dectin-1 could provide more molecular insight in the mechanisms underlying the difference in requirement for Dectin-1 in Th2 polarization by SEA between the human in vitro and murine in vivo models.

In summary, we propose a model (Fig 8) in which SEA can condition DCs for Th2 polarization independently of \(\omega-1\) by triggering Dectin-1, Dectin-2, and the MR to induce in a Syk-dependent fashion the synthesis of PGE\(_2\) and IsoPs, which subsequently promote OX40L expression in an autocrine manner. OX40L then enables the SEA-stimulated moDCs to prime a Th2 response. The fact that neutralization of PGE\(_2\) and its isomers completely blunted the
Th2-priming ability of moDCs that had been stimulated with SEA from which ω-1 had been depleted provides strong support for the notion that this pathway can fully account for “residual” ability of SEA to prime Th2 polarization in the absence of ω-1. However, the fact that this same intervention had little or no effect on the Th2-priming capacity of complete SEA in our in vitro moDC assay suggests that the Dectin-PGE2-OX40L signaling axis can be compensated for by the presence of ω-1, which employs distinct mechanisms to condition moDCs for Th2 polarization, and that this novel axis is only unmasked in vitro when ω-1 is removed from SEA. In vivo, however, the contribution of Dectin-PGE2-OX40L signaling axis in egg antigen-driven Th2 polarization appears to be much more dominant, given that interference with Syk or Dectin signaling did result in an impaired Th2 response induced by complete SEA or by a natural infection with S. mansoni, respectively. This would be corroborated by our previous work showing that SEA from which ω-1 was depleted was still as potent in inducing a Th2 response as complete SEA in vivo [20] and that ω-1 knockdown in S. mansoni eggs by lentiviral transduction did not reduce Th2 responses induced by the eggs in vivo [19]. Together, this leads us to speculate that in vivo, the Dectin-PGE2-OX40L signaling axis plays a more important role in schistosome egg-driven Th2 polarization than the one that is induced by ω-1 in which the MR and suppression of protein synthesis play a central role [9].

In conclusion, we have delineated a previously unrecognized pathway involving Dectin-1/Dectin-2, PGE2, and OX40L through which Th2 immunity is induced. It is interesting to note that antigens from liver fluke Fasciola hepatica as well as from house dust mite have been shown to be recognized by Dectin-1 [58] and Dectin-2 [59], respectively, which makes it tantalizing to speculate that this Dectin-PGE2-dependent signaling axis in DCs is not only triggered by S. mansoni but possibly also by other helminths or allergens to promote Th2 responses. As...
such, targeting this axis may hold promise as an approach to regulate type 2 immune responses for therapeutic purposes, not only in the context of schistosomiasis but possibly also in other helminth infections and in major diseases of the Western world, such as allergies and type 2 diabetes, that are caused by overzealous and defective type 2 immune responses, respectively.

**Materials and methods**

**Ethics statement**

All animal experiments were performed in accordance with local government regulations and the EU Directive 2010/63/EU and Recommendation 2007/526/EC regarding the protection of animals used for experimental and other scientific purposes as well as approved by the Regierung von Oberbayern (animal license number 55.2-1-54-2532-28-1), the animal ethics committee at CNIC (animal license number CNIC10-16-PROEX 240/16), and the Dutch Central Authority for Scientific Procedures on Animals (CCD) (animal license number AVD116002015253). For infection and immunization experiments as described in this study, animals were euthanized using cervical dislocation or an overdose of ketamine/xylazine.

**Mice**

Clec7a<sup>−/−</sup> and Clec4n<sup>−/−</sup> mice (C57BL/6) were housed and bred at the MIH, TUM, Germany, under SPF conditions. Itgax<sup>cre</sup> Syk<sup>fl/fl</sup> mice [60] were housed and bred at the CNIC, Madrid, Spain, under SPF conditions.

**Preparation and purification of S. mansoni egg–derived antigens**

SEA, IPSE/α-1, ω-1, and SEAΔα-1/ω-1 from *S. mansoni* eggs were prepared and isolated as described previously [3,20].

**Human DC culture, stimulation, and analysis**

Peripheral blood mononuclear cells were isolated from venous blood of healthy volunteers by density centrifugation of Ficoll as described before [10]. Monocytes were isolated by positive magnetic cell sorting using CD14-microbeads (Miltenyi Biotech, Bergisch Gladbach, Germany) and cultured in 10% FCS RPMI medium supplemented with 20 ng/mL rGM-CSF (BioSource/Invitrogen, Carlsbad, CA) and 0.86 ng/mL of rIL-4 (R&D Systems, Minneapolis, MN). On day 2, medium including supplements was replaced. In the presence or absence (if indicated) of 100 ng/mL ultrapure LPS (*Escherichia coli* 0111 B4 strain; InvivoGen, San Diego, CA), immature moDCs were stimulated on d 6 with indicated reagents: SEA (50 μg/mL), ω-1 (500 ng/mL), IPSE (500 ng/mL), SEAΔα-1/ω-1 (50 μg/mL), PGE<sub>2</sub> (2.5 ng/mL; Cayman Chemical, Ann Arbor, MI), IFN-γ (1,000 U/mL) as Th1 control, or 0.1 mg/mL Zymosan (Z4250; Sigma-Aldrich, St. Louis, MO). Alternatively, moDCs were stimulated with 2.5 ng/mL LXA<sub>4</sub>, 2.5 μg/mL PGD<sub>2</sub>, 12.5 μg/mL 5-HETE, 12.5 μg/mL 8-HETE, 12.5 μg/mL 11-HETE, 25 μg/mL 9-HODE, or 25 μg/mL 13-HODE (all Cayman Chemical). For blocking experiments, cells were pre-incubated for 60 min at 37˚C with 20 μg/mL anti–DC-SIGN (clone AZN-D1; Beckman-Coulter, Fullerton, CA), anti-MR (clone 15.2; Biolegend, San Diego, CA), anti–Dectin-1 (clone #259931; R&D Systems), anti–Dectin-2 (clone Q7-4B5, InvivoGen), 20 μg/mL IgG1 control antibody for both anti–DC-SIGN and anti-MR, IgG2a (clone RTK2758; Biolegend) control antibody for anti–Dectin-1 and anti–Dectin-2, 1 μM R406, 4 μM U0126 (Merck, Amsterdam, the Netherlands), 1 μM Pyrrophenone (Merck), 10 μM SC-236 (Sigma-Aldrich) in combination with 10 μM Indometachine (Sigma-Aldrich), 10 μg/mL neutralizing anti-PGE<sub>2</sub> antibody (2B5; Cayman Chemical), 10 μg/mL IgG1 antibody (clone MOPC-21; Biolegend) as control for anti-
PGE_2 antibody, 10 μM EP2 (AH6809; Cayman Chemical), or 10 μM EP4 (AH23848; Cayman Chemical) receptor antagonist. After 24 or 48 h of stimulation, surface expression of costimulatory molecules was determined by flow cytometry (FACS-Canto; BD Biosciences, Breda, the Netherlands) using the following antibodies: CD14 (clone MφP9), CD86 (clone 2331 FUN-1), CD40 (clone 5C3), and CD80 (clone L307.4) (all BD Biosciences); HLA-DR (clone LN3) (eBioscience, San Diego, CA); CD83 (clone HB15e) and CD1a (clone BL6) (both Beckman-Coulter); and CD252/OX40L (clone ANC10G1; Ancell, Bayport, MN). Only live cells that were negative for 7-AAD (eBioscience) were included in the analysis. Routinely, in all tested culture conditions, at least 85% of the cells were alive.

Cytokine detection
A total of 1 × 10^4 moDCs matured for 48 h were cocultured with 1 × 10^4 CD40L-expressing J558 cells for 24 h, and supernatants were collected to determine IL-12p70 levels, using mouse anti-human IL-12 (Clone 20C2) as capture antibody and biotinylated mouse anti-human IL-12 (Clone C8.6) (both BD Biosciences) in a sandwich ELISA. IL-23 levels were quantified in supernatants of moDCs stimulated for 40 h by ELISA.

Human DC and T cell coculture and determination of T-cell polarization
For analysis of T-cell polarization, 5 × 10^3 moDCs pulsed for 48 h were cultured with 2 × 10^4 allogenic naive CD4^+ T cells for 11 d in the presence of Staphylococcal enterotoxin B (10 pg/mL). On d 6 and 8, rhuL-2 (10 U/mL; R&D Systems) was added to expand the T cells. Intracellular cytokine production was analyzed after restimulation with 100 ng/mL phorbol myristate acetate plus 2 μg/mL ionomycin for 6 h; 10 μg/mL brefeldin A was added during the last 4 h. Subsequently, the cells were fixed with 1.9% paraformaldehyde (all Sigma-Aldrich). The cells were permeabilized with 0.5% saponin (Sigma-Aldrich) and stained with antibodies against IL-4 and IFN-γ, respectively (BD Biosciences). For analysis of Th17 polarization, 5 × 10^3 moDCs pulsed for 24 h were cultured with 2 × 10^4 allogenic memory CD4^+ T cells for 6 d in the presence of Staphylococcal enterotoxin B (10 pg/mL). On d 6, supernatants were harvested and analyzed for IL-17A by ELISA (eBioscience). For blocking experiments, moDC–T cell cocultures were pre-incubated for 15 min with 10 μg/mL neutralizing PGE_2 antibody (2B5; Cayman Chemical), 10 μg/mL EP2 (AH6809; Cayman Chemical), 10 μg/mL EP4 (AH23848; Cayman Chemical) receptor antagonist, 10 μg/mL anti-OX40L antibody (Clone 159403; R&D Systems), or IgG1 control antibody (clone P3.6.2.8.1; eBioscience).

Detection of ROS
Detection of ROS was performed according to a published protocol (http://www.bio-protocol.org/e313) with minor modifications. In brief, after 6 h or 1 h of stimulation of moDCs or BMDCs, respectively, the cells were harvested, washed using 1% FCS RPMI, and resuspended in 50 μl containing 10 μM CM-H_2DCFDA (C6827; Invitrogen), followed by an incubation at 37°C for 30 min. Prior to sample measurement, 7AAD was added. ROS levels were quantified by flow cytometry.

Western blot
moDCs were harvested after 8 h of stimulation. Then, cells were washed twice with PBS before being lysed in EBSB buffer (8% [w/v] glycerol, 3% [w/v] SDS, and 100 mM Tris–HCl [pH 6.8]). Lysates were immediately boiled for 5 min, and their protein content was determined using a bicinchoninic acid protein assay kit (Thermo-Scientific, Waltham, MA). Proteins were
separated by SDS-PAGE, followed by transfer to a PVDF membrane. Membranes were blocked for 1 h at room temperature in TTBS buffer (20 mM Tris–HCl [pH 7.6], 137 mM NaCl, and 0.25% [v/v] Tween 20) containing 5% (w/v) fat-free milk and incubated overnight with primary antibodies. The primary antibodies used were COX-1 (Cell Signalling Technology, Danvers, MA), COX-2 (Cell Signalling Technology), and actin (Merck). The membranes were then washed in TTBS buffer and incubated with horseradish peroxidase-conjugated secondary antibodies for 1 h at room temperature. After washing, blots were developed using enhanced chemiluminescence.

**Syk and ERK phosphorylation**

For detection of phosphorylation of Syk (pSyk) and ERK (pERK), 2.5 × 10⁴ immature moDCs were seeded overnight in a 96-well flat-bottom plate. moDCs were stimulated with SEA (50 μg/ml), SEAΔα-1/ω-1 (50 μg/mL), or ω-1 (500 ng/mL) in the presence or absence of blocking antibodies or inhibitors (R406, anti-MR, anti–Dectin-1, anti–Dectin-2, combination of anti–Dectin-1 and anti–Dectin-2 or IgG1 and IgG2 control antibodies) for indicated periods, and the moDCs were fixed for 15 min with 4% ultrapure formaldehyde (Polysciences, War- rington, PA) directly in the plate. The cells were harvested and washed first with PBS and then with 0.5% of saponin for permeabilization. Cell were intracellularly stained with anti–phospo-Try525/526 Syk (clone C87C1) and anti–phospo-p44/42 MAPK (Erk1/2) (both Cell Signalling Technology). Following 2-h incubation at room temperature, cells were washed with 0.5% of saponin, and Syk and ERK phosphorylation was determined by flow cytometry.

**cPLA₂ activity**

cPLA₂ activity was determined according to the manufacturer’s recommendation (Cayman Chemical). Briefly, 1 × 10⁶ moDCs stimulated for 8 h with indicated reagents were harvested. moDCs were lysed with lysis buffer (containing 50 mM Hepes, pH 7.4, 1 mM EDTA, NP-40, protease and phosphatase inhibitors) followed by 4 rounds of sonication for 10 s. The cells were then concentrated using a 30 KDa Amicon filter (Merck). To 10 μl cell lysate, 200 μl substrate solution was added to initiate the reaction, and the plate was briefly shaken and incubated for 1 h at room temperature. To stop the reaction, 10 μl DTNB/EGTA was added and the plate was briefly shaken, followed by 5-min incubation at room temperature. The cPLA₂ activity was measured using a plate reader with absorbance of 405 nm.

**Antigen binding and uptake by DCs**

SEA was fluorescently labelled with PF-647 using promofluor labelling kit (Promokine, Heidelberg, Germany) according to the manufacturer’s recommendations. Approximately 2 × 10⁴ immature moDCs or BMDCs per well were seeded in a flat-bottom 96-well plate. Where indicated, cells were pre-incubated with 20 μg/mL of anti-MR, anti–DC-SIGN, anti–Dectin-1, anti–Dectin-2, or control antibodies at 37˚C for 45 min. Subsequently, cells were incubated with 2 μg/mL PF-647–labelled SEA at 37˚C for 45 min for testing both binding and uptake of the antigen. After 45 min, cells were washed with PBS followed by flow cytometry measurement.

**LC-MS/MS analysis of PUFAs and LMs**

Twenty μl of SEA or supernatants from each condition were collected at indicated time points after stimulation and stored at −80˚C until analysis. A volume of 10 μl sample was mixed with 28.4 μL methanol (MeOH) and 1.6 μL of internal standard (containing Leukotriene B4-d4, Leukotriene D4-d4, and Leukotriene E4-d4).
15-HETE-d8, PGE₂-d4, and DHA-d₅ at a concentration of 50 ng/mL in MeOH). The samples were subsequently kept at −20°C for 10 min for completion of protein precipitation, followed by centrifugation for 10 min, 16,000 × g at 4°C. Subsequently, samples were diluted 1:1 with water and transferred into auto-sampler vials. LC-MS/MS analysis using a QTrap 6500 (Sciex, the Netherlands) was carried out as described previously [61,62].

**BMDC culture and adoptive transfer**

BMDCs were generated from indicated WT and KO mice as described previously [63]. Bone marrow cells were differentiated for 8 d in the presence of rGM-CSF (20 ng/ml) (PeproTech, Rocky Hill, NJ) in RPMI-1640 medium containing 5% FCS, 100 U/ml penicillin-streptomycin, and 2 mM glutamine. On d 3 and 6, culture medium was refreshed. Nonadherent cells, containing routinely >80% CD11c⁺MHC-II⁺ DCs were used for various assays, including ROS production, SEA binding/uptake, OX40L expression, and Th2-priming experiments in vivo. OX40L expression was assessed 18 h after stimulation with SEAΔα⁻¹/ω⁻¹ (50 μg/mL) by flow cytometry following surface staining with anti-OX40L (clone RM134L; Biolegend). For in vivo experiments, BMDCs were stimulated with SEAΔα⁻¹/ω⁻¹ (50 μg/mL) for 18 h, washed 3 times in HBSS, and injected into WT recipient mice (400,000/footpad). Analysis of T-cell responses in draining popliteal LNs 7 d later was performed as described below.

**SEA, BMDC immunization, and S. mansoni infection**

Mice were injected subcutaneously with SEA (20 μg), SEAΔα⁻¹/ω⁻¹ (20 μg), ω⁻¹ (2 μg), or 400,000 SEAΔα⁻¹/ω⁻¹–pulsed BMDCs in the hind footpad. Seven d later, cells from both draining and nondraining lymph nodes were isolated and analyzed as described below. For S. mansoni infection, mice were infected with 100 cercariae from a Brazilian strain of S. mansoni obtained from our in-house cycle of infected Biomphalaria glabrata snails (also of Brazilian origin). Mice were killed after 8 wk of infection. Liver samples were fixed in 4% buffered formalin and embedded in paraffin. Sections (4 μm) were stained with Masson blue and examined microscopically (Axioskop; Zeiss, Oberkochen, Germany) for measuring the diameters to calculate the size of spherical granulomas. To determine the parasite burden, pieces of weighed liver and intestine samples from individual mice were digested in 4% KOH at 37°C for 4 h. After centrifugation, the released eggs were microscopically counted. The absolute number of eggs in the liver and intestine was then calculated in accordance with the total organ weight. Worm burden was calculated as adult worm recovery after portal perfusion and microscopic examination of livers and intestines.

**Analyses of murine T-cell responses**

Antigen-specific recall responses were determined by culturing 3 × 10⁵ LN or spleen cells per well in 96-well round-bottom plates in 200 μl complete medium (RPMI containing 10% fetal calf serum, 100 U/ml penicillin/streptomycin, and 2 mM l-glutamine) in the presence of 20 μg/ml SEA or 1 μg/ml anti-CD3/CD28 antibody (eBioscience). 2.5 μg/ml IL-4R blocking antibody (M1) was added to the cultures to retain IL-4 in culture supernatants. After 72 h, culture supernatants were stored for cytokine determination. Cell culture supernatants were analyzed for cytokines using the Cytokine Bead Array (BD) or the mouse Ready-Set-Go ELISA kits (eBioscience) according to the manufacturer’s recommendation. Samples were analyzed on a BD Canto II Flow Cytometer and Sunrise ELISA microplate reader (Tecan, Morrisville, NC), respectively. Alternatively, assessment of cytokine production by intracellular staining of T cells from LNs was determined after polyclonal restimulation in 96-well round-bottom plates for 5 h with phorbol 12-myristate 13-acetate (PMA; 50 ng/ml) and ionomycin (1 μg/ml).
in the presence of brefeldin A (10 μg/ml; all from Sigma-Aldrich) for the last 3 h. Afterwards, cells were fixed with 4% PFA and subsequently stained in 0.5% saponin with antibodies against the following antigens: CD44 (IM7), IL-4 (11B11), IFN-γ (XMG1.2), IL-13 (eBio13A), IL-17A (TC11-18H10.1), and CD4 (RM4-5) (all BD Bioscience or Biolegend). Samples were analyzed on a BD Canto II Flow Cytometer.

**Statistical analysis**

The heatmap was generated using Microsoft Excel. Data were analyzed for normal distribution (Shapiro-Wilk normality test) and statistical significance using two-way ANOVA test, two-sided paired Student t test, or unpaired Student t test. Statistical analysis was performed using GraphPad Prism version 6.00 (GraphPad Software, La Jolla, CA) for Windows.

**Supporting information**

**S1 Fig.** LM composition of SEA and in supernatants of SEA-conditioned moDCs. (A) Concentration of 22 LMs, out of 55 potentially detectable LMs, that are present in SEA from *S. mansoni* as determined by LC-MS/MS. LMs are ordered according to abundance, and concentrations are determined based on internal standards. (B) moDCs were pulsed with SEA or ω-1 in combination with LPS, after which supernatants were collected at 0, 6, 12, and 24 h after stimulation. Relative amounts of PUFAs and LMs detected by LC-MS/MS in supernatants are shown in a heat map. Data represent an average of 3 independent experiments. Color coding is based on relative abundance of each lipid in comparison to other time points or stimulations. (C) As in panel B but without LPS. Data represent 1 of 2 independent experiments. (D) moDCs were pulsed with indicated reagents after which supernatants were collected at 24 h after stimulation. Relative amounts of PGE₂ detected by LC-MS/MS in supernatants are shown. Bar graphs represent means ± SEM of 3 independent experiments. *P < 0.05, for significant differences with the control conditions based on unpaired Student t test. Underlying data can be found in S1 Data. ω-1, omega-1; LC-MS/MS, liquid chromatography tandem mass spectrometry; LM, lipid mediator; LPS, lipopolysaccharide; moDC, monocyte-derived DC; PGE₂, prostaglandin E₂; PUFA, polyunsaturated fatty acid; SEA, soluble egg antigen.

**S2 Fig.** OX40L is induced by SEA via PGE₂ signaling and is required for Th2 induction independently of LPS. (A) T-cell polarization assay as described in main figures. (B, C) moDCs were stimulated as indicated for 48 h in the presence or absence of neutralizing anti-PGE₂ antibody after which expression of OX40L was analyzed by flow cytometry. The fold change based on geometric mean fluorescence is shown relative to LPS, which is set to 1 (dashed line). (D) T-cell polarization assay as described in main figures. Neutralizing OX40L antibody was added during the DC–T cell coculture. Bar graphs represent means ± SEM of at least 3 independent experiments. ** and #: P < 0.05 for significant differences with the control conditions (* ) or between-test conditions (#) based on unpaired analysis (unpaired Student t test). Underlying data can be found in S1 Data. LPS, lipopolysaccharide; moDC, monocyte-derived DC; OX40L, OX40 ligand; PGE₂, prostaglandin E₂; SEA, soluble egg antigen; Th2, T helper 2.

**S3 Fig.** SEA-driven PGE2 synthesis is similar between IL-4– and IL-13–cultured human moDCs. (A) CD1a expression was assessed as a marker for moDC differentiation of monocytes that were differentiated for 6 d in the presence of GM-CSF plus IL-4 or GM-CSF plus IL-13. Representative graphs of 2 independent experiments are shown. (B) PGE₂ production by
IL-4– or IL-13–cultured moDCs in response to SEAΔα-1/ω-1 16 h after stimulation. Bar graphs represent means ± SEM of 2 independent experiments. Underlying data can be found in S1 Data. ω-1, omega-1; GM-CSF, granulocyte-macrophage colony-stimulating factor; IL-4, interleukin 4; LPS, lipopolysaccharide; moDC, monocyte-derived DC; PGE₂, prostaglandin E₂; SEA, soluble egg antigen.

**S4 Fig. SEA does not condition human moDCs for priming of Th17 responses.** (A) IL-23 levels were determined in supernatants of moDC cultures that were stimulated with indicated reagents for 40 h. (B) IL-17 production was assessed in culture supernatants of T cells that were cultured with moDCs that were stimulated with indicated reagents. Zymosan was taken along as positive control stimulus for Th17 induction. Bar graphs represent means ± SEM of at least 4 independent experiments. Underlying data can be found in S1 Data. IL-23, interleukin 23; moDC, monocyte-derived DC; O.p., outpositive; SEA, soluble egg antigen; Th17, T helper 17.

**S5 Fig. BMDCs require Syk expression for Th2 priming by S. mansoni egg antigens in vivo.** (A) BMDCs cultured from BM from CD11cWT or CD11cΔSyk mice were pulsed overnight with SEAΔα-1/ω-1, injected into hind footpads after which CD4⁺ T-cell responses were analyzed as in Fig 6A. Representative flow cytometry plots of intracellular staining of CD4⁺ T cells are depicted, of which the data are enumerated in bar graphs representing mean ± SEM of 3 to 4 mice per group. (B) Ratio between percent IL-4– and IFN-γ–producing T cells as described in panel A. (C) SEA binding and uptake by indicated BMDCs was determined as in Fig 4F. (D) ROS production by indicated BMDCs was determined as described in Fig 5E 1 h after stimulation with SEAΔα-1/ω-1. (E) BMDCs were stimulated as indicated for 18 h after which expression of OX40L was analyzed by flow cytometry. Representative plots are depicted, of which the data are enumerated in bar graphs and shown as fold change relative to control condition, which is set to 1. (C–E) Bar graphs represent duplicates ± SEM of 2 independent experiments. “*” and “#”: P < 0.05; “**” and “##”: P < 0.01; “***” and “###”: P < 0.001 for significant differences with the control conditions (*) between-test conditions (#) based on unpaired analysis (unpaired Student t test). Underlying data can be found in S1 Data. ω-1, omega-1; BMDC, bone marrow–derived DC; CD4, cluster of differentiation 4; H2-DCFDA, 2',7'-dichlorodihydrofluorescein diacetate; IFN-γ, interferon γ; IL-4, interleukin 4; ndLN, non-draining lymph node; OX40L, OX40 ligand; ROS, reactive oxygen species; SEA, soluble egg antigen; Syk, spleen tyrosine kinase; Th2, T helper2; WT, wild-type.

**S6 Fig. Analysis of Th17 responses by SEAΔα-1/ω-1–pulsed BMDCs induced in vivo.** BMDCs cultured from BM from WT, Dectin-1−/− or Dectin-2−/− mice were pulsed overnight with SEAΔα-1/ω-1 and injected into hind footpads after which Th17 responses were analyzed as in Fig 6A. Representative flow cytometry plots of intracellular IL-17A staining of CD4⁺ T cells are depicted, of which the data are enumerated in bar graphs representing mean ± SEM of 2 independent experiments with 4 mice per group. “*” and “#”: P < 0.05 for significant differences with the control conditions (*) or between-test conditions (#) based on unpaired analysis (unpaired Student t test). Underlying data can be found in S1 Data. ω-1, omega-1; BMDC, bone marrow–derived DC; CD4, cluster of differentiation 4; H2-DCFDA, 2',7'-dichlorodihydrofluorescein diacetate; IFN-γ, interferon γ; IL-4, interleukin 4; ndLN, non-draining lymph node; OX40L, OX40 ligand; ROS, reactive oxygen species; SEA, soluble egg antigen; Syk, spleen tyrosine kinase; Th2, T helper2; WT, wild-type.

**S7 Fig. Dectin-1 signaling plays a minor role in Th2 priming during S. mansoni infection.** WT and Dectin-1−/− mice were infected with S. mansoni. After 8 wk of infection, cells from
spleens (A) or mLNs (B) were restimulated with SEA or anti-CD3/CD28 for 72 h, and cytokine levels were analyzed in supernatants by ELISA. Bars represent mean ± SEM of combined data of 3 independent experiments with 3 to 4 mice per group. (c) Granuloma sizes around eggs trapped in the liver of 8-week–infected mice were assessed in Masson blue–stained liver sections. Data are based on 10 mice per group. Number of worms (D) and liver and intestinal eggs (E) in mice infected with S. mansoni for 8 wk. *P < 0.05 for significant differences relative to the control mice based on unpaired analysis (unpaired Student t test). Underlying data can be found in S1 Data. CD3, cluster of differentiation 3; mLN, mesenteric lymph node; SEA, soluble egg antigen; Th2, T helper 2; WT, wild-type.

S1 Table. LMs including PUFAs that were measured using LC-MS/MS. LC-MS/MS, liquid chromatography tandem mass spectrometry; LM, lipid mediator; PUFA, polyunsaturated fatty acid.

S1 Data.

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