Reformulating Small Molecules for Cardiovascular Disease Immune Intervention: Low-Dose Combined Vitamin D/Dexamethasone Promotes IL-10 Production and Atheroprotection in Dyslipidemic Mice

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The targeting of proinflammatory pathways has a prophylactic and therapeutic potential on atherosclerotic cardiovascular diseases (CVD). An alternative/complementary strategy is the promotion of endogenous atheroprotective mechanisms that are impaired during atherosclerosis progression, such as the activity of tolerogenic dendritic cells (tolDC) and regulatory T cells (Treg). There is a need to develop novel low cost, safe and effective tolDC/Treg-inducing formulations that are atheroprotective and that can be of easy translation into clinical settings. We found that apolipoprotein E-deficient (ApoE−/−) mice treated with a low-dose combined formulation of Vitamin D and Dexamethasone (VitD/Dexa), delivered repetitively and subcutaneously (sc) promoted interleukin-10 (IL-10) production by dendritic cells and other antigen presenting cells in the lymph nodes draining the site of injection and the spleens. Expectedly, the treatment also increased the numbers of IL-10-producing CD4+ T cells. Concomitantly, the frequency of IFNγ-producing CD4+ and CD8+ T cells in the spleen, and the IFNγ response of splenocytes to polyclonal stimulation ex vivo were lower after VitD/Dexa treatment, indicating a reduced proatherogenic Th1 response. Interestingly, VitD/Dexa-treated mice had smaller atherosclerotic lesions, with reduced lipid content and lower inflammatory infiltrate of macrophages and T cells in the aortic root. No hypolipidemic or antioxidant effect could be detected, suggesting that a dominantly immunomodulatory mechanism of atheroprotection was engaged under the low-dose sc VitD/Dexa conditions used. Finally, no evidence of clinical, biochemical or immune toxicity was observed in treated ApoE−/− mice and, most importantly, C57BL/6 mice latently infected with Leishmania parasites and treated with an identical VitD/Dexa dose/scheme showed no clinical or microbiological signs of disease reactivation, suggesting the absence of general immunosuppression. Altogether, these results indicate that a non-toxic,
INTRODUCTION

Despite significant advances in the prevention and management of atherosclerotic CVD, they are expected to remain as the major cause of human mortality and disability worldwide (1–3). Whereas the public health situation and economic impact of CVD claims for novel therapies, vascular drug development has paradoxically declined in the last two decades owing to the intrinsic complexities of CVD, as compared to other life-threatening and costly chronic diseases like cancer (3). Although recent advances in the understanding of atherosclerosis highlight the complexity of the pathophysiological events underlying the development of atheromatous plaques and subsequent vascular complications, novel opportunities for therapeutic interventions have also been revealed (see editorial (2) and compendium articles therein). An overwhelming amount of information, mostly derived from animal studies, has implicated the immune system and the inflammatory response as key drivers/ regulators of atherosclerotic CVD. On the one hand, macrophages and T cells play a critical role in atherosclerosis progression by triggering and maintaining proatherogenic innate and adaptive immune responses, involving proinflammatory mediators such as IL-1, IL-6, or IFNγ. On the other hand, concomitant atheroprotective versions of the innate and adaptive responses, such as healing-promoting macrophages and regulatory T (Treg) cells, develop to counterbalance chronic inflammation and promote its resolution via regulatory mediators such as IL-10 (4–6). The translational relevance of this immune hypothesis of atherogenesis remained controverted during decades, since evidence in humans was mostly associative and circumstantial, rather than causal (7). However, the CANTOS study has ended this controversy. It has demonstrated that neutralization of IL-1β, a key proatherogenic/proinflammatory cytokine, significantly reduced the inflammatory burden and protected from secondary CVD events (8), providing a “proof of principle” for targeting of immune pathways, via anti-inflammatory and immunomodulatory agents, as a therapeutic intervention against CVD in humans (9, 10).

Whilst direct blocking of proinflammatory atherogenic pathways has been the obvious focus in the search for a CVD immunotherapy (8), a less explored alternative form of immunomodulation is the reinforcement/restoration of the natural atheroprotective mechanisms that become insufficient as atherosclerosis progresses. Among the immune cell types that promote atheroprotection, Tregs have emerged as dominant players (4–6, 11). Tregs not only exert classical regulatory functions on effector proatherogenic immune cells, but also influence cholesterol metabolism, plaque stability, foam cell formation and endothelial response, participate in plaque regression and tissue repair after ischemic events, and contribute to the athero/cardioprotective effects of statins, vaccination and inflammation resolution inducers (4–6, 12, 13). Thus, from a therapeutic perspective, it is difficult to envision a cell type, other than Treg, that could impact atherogenic responses in a more essential manner. Although a variety of methods have been proposed to restore Treg numbers and functions, including cell therapy, biologicals or particle-based sophisticated approaches (11, 12, 14, 15), their implementation for routine clinical practice is hampered by high costs and/or complexity, making them poorly realistic for widespread clinical use. Small molecules represent an alternative that might offer interesting advantages such as lower cost, which could facilitate preclinical and clinical development, and most importantly, accelerate implementation to clinical settings and its eventual use by the majority of patients. Investigations that explore small molecule route/dose/scheme of administration, delivery strategies and novel combinatory formulations could be of particular interest.

Many synthetic or naturally occurring small molecules have been reported as Treg inducers in vitro and in vivo, including immunosuppressive/anti-inflammatory drugs (such as Dexamethasone, Dexamethasone, Rapamycin and Acetylsalicylic Acid), antioxidants (such as the natural polyphenols Curcumin and Quercetin, or the synthetic N-Acetylcysteine) and vitamins (such as Vitamin D, VitD, Vitamin A, Vitamin C, and Vitamin E) (16–21). Interestingly, but not surprisingly, the Treg-inducing capacity of these compounds is largely explained by their ability to condition antigen presenting cells (APC) toward a tolerogenic form. Many reports have shown that dendritic cells (DC) incubated in vitro with those compounds were potent

**Abbreviations:** APC, antigen-presenting cell; ApoE−/−, apolipoprotein E-deficient; ASA, acetylsalicylic acid; CANTOS, canakinumab anti-inflammatory thrombosis outcomes study; CD, cluster of differentiation; cDC, conventional DC; Curc, curcumin; CVD, cardiovascular diseases; Dexta, dexamethasone; dLN, draining lymph node; FACS, fluorescence-activated cell sorting; FOXP3, forkhead box P3; GR, glucocorticoid receptor; HFD, high-fat diet; IFNγ, interferon-gamma; Ig, immunoglobulin; IL, interleukin; LDL, low-density lipoproteins; MDA, malondialdehyde; MDSC, myeloid derived suppressor cells; MFI, mean fluorescence intensity; MHC, major histocompatibility complex; MLN, mediastinal lymph node; MoDC, monocyte-derived dendritic cells; NAC, N-acetyl cysteine; NK, natural killer; NR, nuclear receptors; pDC, plasmacytoid dendritic cells; PLN, popliteal lymph node; Querc, quercetin; rapa, rapamycin; sc, subcutaneous; SP, spleen; T effector T cells; Th1, type 1 T helper; tolDC, tolerogenic dendritic cells; Tr1, type 1 regulatory T cell; Treg, regulatory T cells; VDR, vitamin D receptor; Vit, vitamin; VitD, vitamin D3; WHO, world health organization.
Treg inducers *in vitro*, or *in vivo* upon adoptive transfer, demonstrating that DC-intrinsic effects are sufficient to explain their tolerogenic capacity. Based on this, we reasoned that a low-dose local (non-systemic) delivery of these compounds could be sufficient to condition local or draining lymph node (dLN) DC to acquire tolerogenic functions and Treg-inducing capacity *in vivo*. A low-dose but repetitive subcutaneous scheme would assure a sustained local/regional activity with little systemic distribution, which appeared particularly appropriate and convenient for agents potentially toxic and/or with a very restricted biodisponibility given their physical-chemical properties. A pioneering work by Kang and collaborators (21) demonstrated that subcutaneous (sc) delivery of low-dose Dexa, in combination with an antigen, efficiently induced tolerogenic DC (tolDC) and antigen-specific Tregs that restored tolerance and impacted disease, providing further support to our rationale. Thus, we initiated the search for a tolDC/Treg-inducing small molecules that exerts atheroprotection under the administration conditions described above in dyslipidemic mice, and found that combined VitD/Dexa is a promising formulation.

**MATERIALS AND METHODS**

For detailed information, see Supplementary Material.

**Mice, Treatment and Samples**

Male ApoE<sup>−/−</sup> mice were injected sc into the footpad with the combination of VitD and Dexa (10 µg each), or vehicle, and fed a chow or a HFD, as shown in Figure 1. This scheme was designed in order to maximize the chances of evidencing a protective effect in the case it existed. After sacrifice (euthanasia with Ketamine/Xylazine 100/10 mg/kg, ip), spleens (SP), popliteal LN (PLN), blood and hearts-aortas were collected. Mediastinal LN (MLN) were also collected in some experiments. Wild-type C57BL/6 and BALB/c mice were also used in some experiments (see Supplementary Material for details) and euthanized as above. The institutional ethical committee (Comité Institucional para el Uso y Cuidado de los Animales de Experimentación, CICUAL) approved all the experiments, and procedures were performed according to the guidelines from Directive 2010/63/EU of the European Parliament.

**Immunological Analysis**

SP and LN cell suspensions were obtained to analyze the frequency of tolerogenic (IL-10-producing) APCs, Tregs (IL-10-producing CD4<sup>+</sup> T cells, called here Tr1-like Tregs and forkhead box P3<sup>+</sup> (FOXP3<sup>+</sup>) CD4<sup>+</sup> T cells, called here FOXP3<sup>+</sup> Tregs), and IFNγ-producing CD4<sup>+</sup> T cells (called here Th1 cells) by flow cytometry. SP cell suspensions were used to assess the amount of IFNγ and IL-10 secreted after stimulation with Concanavalin A (ConA). See Supplementary Material for details.

**Atheroprotection and Serum Analysis**

Cryosections from the aortic root were used to quantify atherosclerotic lesion size, lipid deposition area, and macrophage and T cell infiltration by H/E, Oil red O staining, and immunohistochemistry. Serum samples were obtained to quantify the circulating levels of the lipid peroxidation product malondialdehyde (MDA), cholesterol, triglycerides, calcium, and glucose (22).

**Immunosuppressive Effect of VitD/Dexa**

*C*57BL/6 mice were also used to evaluate the capacity of VitD/Dexa treatment to reactivate a latent infection with *Leishmania* parasites, by following the size of the cutaneous lesions and the amounts of viable parasites at the site of infection. Further details can be found in Supplementary Material.

**Statistical Analysis**

Data were tabulated and analyzed using GraphPad Prism 6.0 software. Normality and equality of variances were tested by D’Agostino-Pearson test and Levene test, respectively. Student’s *t*-test or Mann-Whitney *U*-test were used for comparison between unmatched two groups. Two-way ANOVA followed by Bonferroni’s post-test was conducted in multiple group comparisons. *P* < 0.05 was considered significant.

**RESULTS**

**Combined VitD/Dexa as a Promising Treg-Inducing Formulation *In vivo***

We began by performing a pilot *in vivo* screening in which we tested a collection of nine small molecules with documented Treg-inducing capacity (16–21) using a low-dose repetitive sc injections. In a first experiment we confirmed that all compounds tested were indeed Treg inducers after two rounds of treatment (Supplementary Figure S1A). In a second experiment, in which the number of rounds was increased to three, we found that the combination of VitD and Dexa induced the highest percentage of FOXP3<sup>+</sup> CD4<sup>+</sup> T cells when injected either in absence or presence of the atherosclerosis-relevant antigen LDL (Supplementary Figure S1B).

**In Hypercholesterolemic ApoE<sup>−/−</sup> Mice, Low-Dose sc VitD/Dexa Promotes IL-10 Production by DC and Other APC**

The result of our screening was interesting, since cross-talk between the glucocorticoid receptor (GR) and VitD receptor (VDR), and the synergistic interaction between their ligands have been described (23–25), suggesting that novel combined re-formulations of these agents might offer novel possibilities for the optimization of their immunomodulatory capacities, with better efficacy and safety. In fact, the power of combined VitD/Dexa to induce tolDC/Treg *in vitro* has been well documented (26, 27); however, surprisingly, the prophylactic/therapeutic potential of combined VitD/Dexa has been poorly explored *in vivo*. Based on this, we continued with a more detailed *in vivo* characterization of the immunoregulatory and atheroprotective capacity of VitD/Dexa in dyslipidemic mice. ApoE<sup>−/−</sup> mice were treated with VitD/Dexa (10 µg each) or vehicle and fed a HFD as...
depicted in Figure 1, and the frequency of tolDC in lymphoid organs was determined by flow cytometry at the end of the experiment. Since IL-10 production is a key phenotypic and functional marker of the tolDC induced by VitD/Dexa (26, 27), we evaluated the production of this cytokine by CD11c^{hi} MHC-II^+ cells (conventional DC, cDC) regionally in the PLN, the LN draining the site of drug administration, as well as systemically (in the SP) with a gating strategy that excluded other cell types (including B cells, Supplementary Figure S2A). Our results indicated that VitD/Dexa treatment significantly increases the percentage and absolute number of regional and systemic IL-10-producing cDC (Figure 2A). Because major histocompatibility complex class II (MHC-II) and costimulatory molecules are also important APC functional markers and it is known that VitD and Dexa alter their surface abundance (26, 27), we continued with our ex vivo analysis by determining the expression of MHC-II and CD40 on cDC. The overall density of these maturation markers in cDCs from VitD/Dexa-treated animals was slightly higher than in control mice, with statistical significance confirmed only for CD40 in the spleen (Figure 2B). Thus, in dyslipidemic mice, VitD/Dexa treatment does not appear to substantially alter surface MHC-II and CD40 expression, but it improves the capacity of cDC to produce the regulatory mediator IL-10.

Fluorescence-activated cell sorting analysis also allowed us to assess the production of IL-10 in other non-cDC APC, including B cells (by using the CD19 marker; Supplementary Figure S2B) and the non-B non-CD11c^{hi} MHC-II^+ cells, which comprises mainly monocyte/macrophages and less abundant DC subpopulations (such as plasmacytoid DC and monocyte-derived DC; Supplementary Figure S2A). Interestingly, these two groups of APCs also expressed more IL-10 in the PLN and the spleen when mice were treated in vivo with VitD/Dexa (Figure 3A). Regarding the maturation marker expression, VitD/Dexa induced an overall reduction in the surface density of MHC-II and CD40 on the non-B CD11c^{int-low} MHC-II^+ APCs (Figure 3B, left panel) and a weaker inhibitory effect only on CD40 expression by splenic CD19^+ B cells (Figure 3B, right panel).

Altogether, findings presented so far indicate that although exposure to low-dose VitD/Dexa might have distinct effects on the maturation/activation status of different APC in vivo, all APC subsets become more efficient producers of the regulatory cytokine IL-10.

**VitD/Dexa Treatment Promotes Treg Cells In vivo**

Vitamin D3 and/or Dexa-conditioned DC are potent inducers of the two major Treg subsets, FOXP3^+ and Tr1-like Tregs, both in vitro and in vivo (26, 27). This, together with our observation that combined VitD/Dexa induces IL-10-producing DC/APC in vivo (Figures 2, 3), prompted us to analyze the amounts of IL-10-producing CD4^+ T and FOXP3^+ CD4^+ T cells ex vivo (Supplementary Figure S3). Results confirmed that VitD/Dexa treatment markedly increased the relative and absolute number of IL-10-producing CD4^+ T cells (Tr1-like Tregs) in the SP and PLN (Figure 4A). On the other hand, no changes in the FOXP3^+ CD4^+ T cells (FOXP3^+ Treg) subset were observed in these lymphoid organs (Figures 4B, C). We extended our Treg analysis to MLN, the secondary lymphoid organs draining the heart, and found that animals treated with VitD/Dexa had increased percentages of FOXP3^+ but not of Tr1-like Tregs when compared to control animals (Figures 4A, B). The analysis of the abundance of the FOXP3 marker (as assessed by the mean of fluorescent intensity, MFI) in CD4^+ T cells from MLN indicated a non-statistically significant trend to higher level in VitD/Dexa-treated mice compared to control mice (Figure 4C). In summary, this set of analysis confirmed that VitD/Dexa also induces Tr1-like Tregs in dyslipidemic mice.

**VitD/Dexa-Treated Mice Have Less IFNγ^+ Producing T Cells**

IFNγ^+ -producing T cells are major drivers of local and systemic inflammation and have a pivotal role in atherosclerosis development and in CVD (4–6). Because Tregs inhibit...
Adverse effects such as disturbances in calcium, glucose or lipid modifying or antioxidant systemic effects. VitD/Dexa operates via immune modulation rather than lipid-lowering effect.

VitD/Dexa Protects Against Atherosclerotic Lesion Development in Dyslipidemic Mice

Having demonstrated that VitD/Dexa increases regulatory and promotes suppressive Tregs that might be able to reduce atherosclerosis-related inflammation.

Repetitive Low-Dose VitD/Dexa Treatment Does Not Induce Overall Toxicity or Immunosuppression

Depending on the dose and extension of the treatment, both VitD and Dexamethasone are potentially deleterious and might induce adverse effects such as disturbances in calcium, glucose or lipid.
metabolism, or immunosuppression. It was noteworthy that ApoE<sup>−/−</sup> mice treated with VitD/Dexa gained weight similarly to vehicle-treated mice (Table 1) and had healthy appearance and normal behavior, just like control mice. Furthermore, at the end of the experiment, no changes on serum glucose and lipid levels, or major effects on lymphoid cell viability or total CD4<sup>+</sup> T cell numbers were observed in VitD/Dexa treated mice (Table 1). Although a mild (7%) but significant increase in serum calcium levels was observed in VitD/Dexa-treated mice (Table 1), these levels (9.45 ± 0.95 mg/dl) were comparable to reported reference values in 16-week-old ApoE<sup>−/−</sup> mice (2.28 ± 0.22 mM or 9.12 ± 0.88 mg/dl) (29), as well as to the calcium serum levels reported by Charles River Laboratories in C57BL/6 mice (IC 95% 9.7–12.5 mg/dl). In addition, no clinical signs of hypercalcemia (e.g., dehydration or weight loss, as shown above) were detected. A major safety concern of pharmacological anti-inflammatory agents such as VitD and specially Dexa is their potential immunosuppressive effect, which might impair effector immune mechanisms against tumors and infections. To gain insights into the potential immunosuppressive effects of low-dose VitD/Dexa, we examined its impact on cutaneous leishmaniasis, a chronic and persistent infection in which immunosuppression promotes disease reactivation. In a murine model of cutaneous leishmaniasis by Leishmania panamensis (Lp) established in our laboratory, we have observed that BALB/c mice challenged with infective parasites develop progressive ulcerative cutaneous lesions whereas C57BL/6 mice develop mild self-limited lesions. Although cured C57BL/6 mice become immune to reinfections, they harbor persistent parasites at the site of infection, rendering them suitable models for assessing the reactivation of a latent infection. C57BL/6 mice (and BALB/c mice, as a control for the inoculum infectivity) were inoculated intradermally in the ears to subsequently monitor lesion development (Figure 8A). As expected, BALB/c mice developed a severe disease (Supplementary Figures S5A,B) while C57BL/6 mice showed a mild self-limited clinical form.
that cured within 6–8 weeks postinfection (Figures 8B). Once C57BL/6 mice completely healed the lesions, they were treated with VitD/Dexa using the same route/dose/scheme that resulted in atheroprotection in ApoE<sup>−/−</sup> mice (Figures 1, 8A) and monitored weekly to detect any sign of lesion reactivation. Similarly to vehicle-treated mice, the ears of VitD/Dexa-treated animals presented a normal, healed and healthy appearance, with no redness, swelling, or other clinical signs of disease reactivation.
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during the whole experimentation period (Figure 8B). Both groups of mice also gained weight in a similar manner (Figure 8C), had a healthy overall appearance and normal behavior, and did not exhibit piloerection, lethargy, or signs of dehydration. Most importantly, and consistently with clinical observations, the number of viable parasites in the ears of VitD/Dexa-treated mice did not differ from control mice (Figure 8D), confirming the absence of parasite reactivation. Thus, these results indicate that the effector immune mechanisms that maintained lesion-free parasite latency remained unaffected after chronic low-dose VitD/Dexa treatment.

DISCUSSION

Although significant efforts have been made in the last years to develop useful immunomotherapeutic agents for chronic inflammatory conditions, such as CVD, two major problems remain (30). First, safety issues such as immunosuppression that might increase the susceptibility to infections and tumors (8, 31). And second, the high costs [for example of biological drugs; (32, 33)] and the logistic and technical limitations [for example of cellular therapies; (34)] that make unrealistic the translation of the agents to routine prophylactic or therapeutic clinical use. VitD and Dexa, listed by WHO as essential medicines, are two of the most prescribed drugs by clinicians, are widely available at an exceptionally meager cost, and importantly, there is plenty of scientific information on their pharmacology, toxicology, efficacy and safety. Thus, the repurposing and reformulation of these agents could be inexpensive and rapid, and their clinical use realistic. Here we report that our attempts to find a simple, safe and inexpensive alternative Treg-inducing small molecule led us to test and confirm that the local low-dose repetitive delivery of a tolDC/Treg-inducing formulation that combines VitD and Dexa promotes atheroprotection in ApoE<sup>−/−</sup> mice.

Current therapeutic use of VitD and Dexa in the clinical practice (high dose, oral/muscular delivery) implies systemic distribution with the expected consequences, beneficial and toxic, dictated by the pharmacological response of the ubiquitously expressed VDR and GR (35–37). Given the physical-chemical properties (very low water solubility) we reasoned that local (e.g., footpad, sc route) low-dose (10 µg) treatment using a repetitive but spaced scheme (rounds every other week) would provide biologically relevant concentrations of VitD/Dexa at local/regional level rather than systemically, restricting their effects to the site of injection and dLN. Although a detailed pharmacological analysis is required to formally confirm the absence or the negligible circulation and distribution, all analysis
FIGURE 6 | VitD/Dexa promotes CD4⁺ T cells that are suppressive ex vivo. Wild type C57BL/6 mice were treated with VitD/Dexa or vehicle, and CD3⁺ CD4⁺ T cells were purified by FACS-sorting to be co-cultured with total SP cells from LDL-vaccinated C57BL/6 mice (containing IFNγ-producing effector T cells, Teff), in the presence of the relevant antigen for 72 h (A). The suppressive activity of the CD4⁺ T cells was determined by the amount of IFNγ released to the medium at different Teff:TCD4⁺ ratios (B). CD4⁺ T and Teff were obtained from pooled splenocytes from three mice. OVA was also included as a control of non-relevant antigen stimulation. Data are shown as mean ± SD from pooled triplicate cultures. ND, not detected. Note that whereas in 1:1 Teff:TCD4⁺-Control co-cultures the mean of IFNγ expression was 168 pg/mL, in 1:1 Teff:TCD4⁺-VitD/Dexa co-cultures IFNγ was not detectable (which means less than 3.1 pg/mL).

performed (clinical, biochemical, redox, immunological and immunosuppressive effect; Table 1; Figure 8) indicated the lack of direct systemic effects of VitD/Dexa, including toxicity. In contrast, the desired immunological effects (induction of regulatory cells) on the dLN were evident (Figures 2–4). Moreover, in preliminary experiments, we have detected the induction of apoptotic cells in the dLN but not in the spleen after sc VitD/Dexa injection (data not shown), further supporting our hypothesis of regional VitD/Dexa delivery but negligible systemic exposure. These results confirmed that sc low-dose VitD/Dexa formulation is indeed a safe and non-immunosuppressive tolDC/Treg-inducer in mice.

Despite the hyperlipidemic and proinflammatory conditions characterizing ApoE⁻/⁻/− mice, which are further accentuated by HFD feeding, the treatment with VitD/Dexa was able to efficiently induce IL-10 production in DCs and other APCs (Figure 2, 3), shifting CD4⁺ T cells response toward a less accentuated Th1 profile, and promoting an increased Treg response (Figures 4, 5) and atheroprotection (Figure 7). Our observations are in agreement with an increasing number of reports suggesting that interventions modifying the DC-T cell crosstalk under hyperlipidemic conditions, from the prevailing proimmunogenic form toward a more tolerogenic form, results in atheroprotection (4–6, 9–14, 30). Previous reports have indicated that low-dose sc Dexa promoted tolDC, increased Treg/Teff levels and protected mice from autoimmunity (21) and atherosclerosis (38) when co-injected with a disease related-relevant antigen. Also, that oral delivery of VitD promoted
Our work extends those findings by showing that a combined low-dose formulation of VitD/Dexa, delivered locally in the absence of coinjected antigen, delays atherosclerosis development. Although we did not directly compare the two single versus combined VitD/Dexa within the same trial, two pilot independent experiments revealed the dependency of the co-formulation for inducing significant atheroprotection (Supplementary Figure S6). This result, together with the improved Treg-inducing capability observed for the combined formulation in wild-type mice (Supplementary Figure S1) suggest a potential additive or synergistic atheroprotective effect for these two small molecules that requires further investigation. Considering the well demonstrated synergy between VitD and Dexa to induce tolDC and Treg in vitro (26, 27), it is to be evaluated whether atheroprotective effects of VitD/Dexa can be further optimized by refining the amounts and relation of the agents and the treatment scheme. Similarly, whether coinjection of a relevant atheroantigen such as LDL together with VitD/Dexa will lead to improved atheroprotection is unknown, but is currently being investigated in our laboratory.

Due to technical limitations, we were not able to investigate whether VitD/Dexa-induced Tregs were atheroantigen-specific. Nonetheless, CD4+ T cells from VitD/Dexa-treated mice were more effective suppressors of the LDL-stimulated Th1 effector response ex vivo compared to vehicle-treated mice (Figure 6), and no evidence of immunosuppression was observed after VitD/Dexa administration (Figure 8). Thus, we speculate that oxLDL/LDL present in plasma, lymph, and other fluids from wild-type mice, and more notably from HFD-fed ApoE−/− mice, might have contributed to the tolerogenic conditioning of DCs at the draining site of VitD/Dexa injection, providing relevant evidence for the potential role of these molecules in atheroprotection.

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**FIGURE 7 |** VitD/Dexa attenuates the development of atherosclerosis in dyslipidemic mice. ApoE−/− mice were treated with VitD/Dexa (10 µg) or vehicle and fed a HFD, as shown in Figure 1. Cryosections from the aortic root were stained with H/E (A) and Oil red O (B) or probed with macrophage (CD68)- and T cell (CD3)-specific antibodies for immunostaining (C, D). Representative micrographs for each treatment group are shown (left panel). The areas of the atheromatous lesions, lipid deposition as well as the macrophage and T cell infiltrate were calculated with image analysis software. The black bar on the left panel represents 250 µm. Each point represents the average area per mouse (7–9 sections/mouse), and bars represent the mean (right panel). Representative results of three independent experiments are shown. *P < 0.05; **P < 0.001 (Student’s t-test).
TABLE 1 | Body weight, serum analyses and lymphoid organ cell counts in dyslipidemic ApoE−/− mice.

| Parameters                              | Control               | VitD/Dexa             | P-value  | P-value summary |
|-----------------------------------------|-----------------------|-----------------------|----------|-----------------|
| **Body weight, g (across the treatment)** |                       |                       |          |                 |
| Week 1                                  | 25.38 ± 3.46          | 23.63 ± 3.05          | 0.20     | ns              |
| Week 4                                  | 27.88 ± 2.25          | 26.43 ± 1.91          | 0.10     | ns              |
| Week 8                                  | 30.41 ± 3.42          | 30.82 ± 3.31          | 0.77     | ns              |
| Week 12                                 | 35.38 ± 4.60          | 34.26 ± 6.19          | 0.62     | ns              |
| Week 16                                 | 36.70 ± 4.935         | 33.20 ± 4.50          | 0.08     | ns              |
| **Serum analysis**                      |                       |                       |          |                 |
| Cholesterol Total, mg/dL                | 1301.00 ± 423.10      | 1427.00 ± 384.30      | 0.452    | ns              |
| Triglyceride, mg/dL                     | 142.30 ± 88.33        | 130.10 ± 23.03        | 0.636    | ns              |
| Malondialdehyde (MDA), nmol/mg protein  | 2.75 ± 1.06           | 3.93 ± 2.04           | 0.117    | ns              |
| Glucose, mg/dL                          | 398.10 ± 89.62        | 414.70 ± 105.80       | 0.691    | ns              |
| Calcium, mg/dL                          | 8.77 ± 0.36           | 9.45 ± 0.95           | 0.037    | *               |
| **Lymphoid organ cell counts**          |                       |                       |          |                 |
| SP                                      | 88.35 ± 8.09          | 93.91 ± 1.51          | 0.058    | ns              |
| PLN                                     | 92.05 ± 1.68          | 94.52 ± 3.06          | 0.032    | *               |
| MLN                                     | 85.13 ± 6.45          | 85.65 ± 8.35          | 0.883    | ns              |
| **Number of total leukocytes/organ**    |                       |                       |          |                 |
| SP                                      | 1.43E + 08 ± 4.55E + 07 | 1.59E + 08 ± 6.50E + 07 | 0.900    | ns              |
| PLN                                     | 6.60E + 06 ± 3.09E + 06 | 4.67E + 06 ± 2.13E + 06 | 0.097    | ns              |
| MLN                                     | 1.93E + 06 ± 8.48E + 05 | 1.30 + 06 ± 1.02E + 06 | 0.173    | ns              |
| **Number of CD4+ T cells/organ**        |                       |                       |          |                 |
| SP                                      | 1.35E + 07 ± 1.15E + 07 | 2.62E + 07 ± 2.20E + 07 | 0.114    | ns              |
| PLN                                     | 1.40E + 06 ± 7.20E + 05 | 1.70E + 06 ± 1.39E + 06 | 0.542    | ns              |
| MLN                                     | 1.85E + 05 ± 7.73E + 04 | 2.83E + 05 ± 1.41E + 05 | 0.079    | ns              |

Non-fasted serum samples were collected from animals. Data are presented as the mean ± SD (representative of two independent experiments, 11 to 12 mice per group, Student’s t-test, *P < 0.05).

ApoE−/− mice (Figure 4B). This contrasted with the splenic FOXP3+ Treg response observed in non-dyslipidemic wild-type mice (Supplementary Figure S1B) and opens the possibility that Treg induction might be qualitatively different under dyslipidemic versus non-dyslipidemic conditions and that in VitD/Dexa-treated mice different Treg subsets might play distinct atheroprotective roles.

Similarly, the stimulation of IL-10 production by VitD/Dexa in non-cDC non-B APC (Figure 3A, left) is also interesting, since monocyte-derived DC appear to be important during atherosclerosis development and tolerogenic plasmacytoid DC are atheroprotective (43, 44). IL-10-producing tolerogenic macrophages have been identified in low-dose Dexamethasone-treated mice (45) and, therefore, could also support the atheroprotection observed in our experiments. Also B cells are possible contributors to the atheroprotection reported here via IL-10 production (Figure 3A, right), according to the recent recognition of regulatory B cells (Breg) as potentially atheroprotective cells (46). We could not find differences in the serum levels of LDL-specific total IgG, or the subclasses IgG1 and IgG2c between treated and control mice (data not shown), suggesting that a B cell-dependent atheroprotective role could be more related to IL-10 production or, alternatively, to IgM natural antibodies. Our findings indicate that, in dyslipidemic mice, VitD/Dexa induces a
regulatory network of IL-10-producing innate and adaptive immune cells that are likely mediators of atheroprotection. Moreover, preliminary evidence suggests that other leukocyte types (such as CD8$^+$ T cells or NK cells) also upregulate IL-10 production (not shown), extending the repertoire of immune cells that produce IL-10 in response to the VitD/Dexa treatment in vivo. Accordingly, higher number of cells producing IL-10 in those mice is already evident in total splenocytes, without further gating of subpopulations (Supplementary Figures S7A,B). Taking this in consideration, the known role of IL-10 as inducer of MDSC, and the atheroprotective potential of MDSC (47), we also evaluated the abundance of these cells. However, it was surprising that the numbers of MDSC did not substantially increase with VitD/Dexa treatment (Supplementary Figure S8). Thus, we hypothesize that the induction of an IL-10-producing leukocyte network is a key event underlying the atheroprotection reported here. The wide range of cells in which a regulatory program is triggered by this cytokine enabling the induction anti-inflammatory effects, such as infectious tolerance (48), would be sufficient to explain atheroprotection. More detailed future work will permit to
analyze the kinetics and essentiality of each IL-10 producer for atheroprotection.

Our findings support a mode of action in which tolDC and Tregs are induced in the periphery (skin/dLN), and upon subsequent migration exert distant anti-inflammatory and atheroprotective effects. The presence of tolDC and Tregs in the spleen could be better explained by migration from the periphery rather than by splenic induction, as we don’t expect significant systemic exposure to VitD/Dexa. The spatiotemporal dynamics of VitD/Dexa-induced tolDC and Treg is also to be confirmed. However, according to current knowledge, a possible explanation is that VitD/Dexa-conditioning of DC might occur on skin migratory DC or on resident dLN DC. In both cases, tolDC will be present in dLN to mediate Treg induction. Also, Treg formation in the sc tissue cannot be ruled out, particularly in dyslipidemic mice where skin inflammation and abundant inflammatory lymphocyte infiltrate has been reported (49).

Alternatively, dying cells induced locally by VitD/Dexa could be rapidly removed by efferocytic DC, triggering Treg induction. Both apoptosis induction and effector cytokinin stimulation by these agents, as well as the connection between effectorcytokin, Tregs, inflammation resolution and atheroprotection, have been documented (35, 36, 50, 51). As DC tolerogenic-conditioning via direct effects of the agents or indirectly via apoptosis/efferocytosis are not mutually exclusive, it is possible that a combination of them underlie the increased frequency of Treg observed in VitD/Dexa-treated mice.

How peripherally induced VitD/Dexa-conditioned tolDC and specially Tregs exert atheroprotection is also to be explored. The simplest explanation is that these regulatory cells migrate to the inflamed atheroma. However, our attempts to identify immune cells in the aortic roots by FACS were unsuccessful given the poor yields and limited viability of the cell suspensions obtained. When we analyzed the expression of some immune markers in whole abdominal aortas by qPCR, there was a trend showing reduced levels of expression of inflammatory markers such as IFNγ and MCP-1 in VitD/Dexa-treated mice (Supplementary Figure S9), which is concordant with the reduced inflammatory infiltrate observed in the aortic root (Figures 7C,D). However, mRNA levels of IL-10, and other Treg-related mediators such as TGF-β and FOXP3 exhibited the same tendency (Supplementary Figure S9). Although caution has to be taken with qPCR analysis from whole abdominal aortas, as results reflect the average abundance of mediators in both atheromatous and non-atheromatous areas of the artery, the confirmation of this finding would imply that tolDC and Treg induced by VitD/Dexa could have influenced more systemically in the spleen and relevant LN, rather than in the arteries. This is in line with the recent observation that whereas priming and amplification of effector IFNγ+ T cells occurs in the artery, Treg responses preferentially takes place in secondary lymphoid organs (52) and is concordant with our observations showing more Tregs in the spleen and MLN of VitD/Dexa-treated mice than in controls (Figure 4). Further research is needed to better characterize tolDC and Treg trafficking in VitD/Dexa-treated mice.

An important aspect that deserves further discussion is that the formulation utilized here was composed by cholecalciferol and Dexa, prepared in 2% DMSO. In contrast to Dexa, which directly triggers the GR, cholecalciferol is the VitD prohormone that requires the previous transformation into the active 1,25-hydroxy VitD hormone. VitD production, activation and inactivation have been extensively investigated in the context of its dominant function as a key regulator of mineral homeostasis. Sequential hydroxylation reactions at the positions 25 and 1 of cholecalciferol are known to occur in the liver and kidneys, respectively. In the context of the immunomodulatory actions, it is of great interest that immune cells not only express VDR, making them responsive to the active hormone, but also the key metabolizing enzymes, which enables them to transform cholecalciferol into active VitD (53). Particularly in the skin, where cholecalciferol is bioavailable via endogenous synthesis or diet, dendritic cells have been described as a local source of active VitD (54). Interestingly, under inflammatory conditions and/or through DC-T cell cross-talk, VitD-activating enzymes are upregulated, leading to increased local VitD activation (53, 54). Thus, the inflamed skin and dLN that characterizes dyslipidemic mice (49) are expected to be particularly capacitated to transform inactive VitD into the active form. Although this requires experimental confirmation by measuring inactive and active forms of VitD locally, and VitD-activating enzymes in immune cells, our results of tolDC and Treg induction (Figures 2, 4) and atheroprotection (Figure 7) are consistent with the idea of local activation of cholecalciferol upon sc administration. This is of translational relevance, since obesity associates to skin inflammatory conditions (55) and also to poor VitD status due to liver dysfunction (56). Thus, sc administration of cholecalciferol under inflammatory environments could assure sufficient 25-hydroxy and 1,25-hydroxy VitD local availability for tolDC-conditioning and Treg induction. Moreover, we consider that inflammation-triggered local activation of cholecalciferol is convenient, since the extent of this activation under non-limiting amounts of the prohormone due to its sc repetitive supply would be proportional to the inflammatory burden, leading to a controlled form of active VitD delivery that matches the tolerogenic requirements. Future work comparing the immunoregulatory capacity of sc low-dose cholecalciferol, 25-hydroxy and active VitD, combined to Dexa, will permit a better understanding of the atheroprotective effects reported here and the development of improved Treg-inducing VitD/Dexa-based antiatherogenic formulations.

Metabolism and immunity are two intimately interconnected processes that are considered to play central role in atherosclerosis pathogenesis and CVD (57, 58). Since signaling through the VDR and GR exerts effects on both processes, it was of interest that no evidence of metabolic changes, such as lipid-lowering effects, associated to the VitD/Dexa-mediated atheroprotection were found (Figure 7 and Table 1). As explained before, the dose, via and scheme of treatments utilized here and our findings of no toxicity or immunosuppression, argues to non-systemic relevant distribution of the agents that could have affected directly the liver or other key metabolic organ. However, FOXP3+ Tregs have been shown to influence hepatic metabolism of lipoproteins and control circulating cholesterol levels (59), which raises the question of why our...
VitD/Dexa treatment did not alter blood lipids indirectly via Treg-promotion. We believe that a more complete analysis of lipoprotein distribution could help to clarify this, or that Tr1-like cells (in which VitD/Dexa impact was more pronounced; Figure 4) are weaker at “lipid-lowering” compared to FOXP3+ Treg. Alternatively, taking into account the complexity of this new field of immunometabolism, it is plausible that the magnitude and specificity of the atheroprotective pathways triggered by a particular immunotherapeutic strategy underlies the chances to significantly affect organismal metabolism. Thus, in accordance with our results, a recent work inducing epitope-specific Treg cells via vaccination was atheroprotective without affecting blood lipids (60). Moreover, not only the relevant CANTOS study, but also a number of immunomodulatory interventions in animal models, such as other Treg-inducer (61), a TLR4 inhibitor (62), an antiplatelet agent (63), NLRP3 inflammasome-directed therapies (64, 65) or a M2 macrophage-inducing kinase inhibitor (66) exerted atheroprotection in the absence of lipid-modifying effects, suggesting that lipid-lowering effects of immune-mediated treatments are possibly the exception rather than the rule.

It is tempting to explain the observed VitD/Dexa-induced atheroprotection as a consequence of a better status of VitD and/or direct suppressive effects of Dexa on atheromatous artery. However, we consider this unlikely for the following reasons: (1) this idea implies systemic circulation and distribution of the agents for which as we explained, although not directly proven here, no indirect evidence was found. (2) A better VitD status requires that mice responded to VitD replenishment, which has been demonstrated not to be the case in obese mice (56, 67). (3) Direct atheroprotective activity of glucocorticoids on the atheromatous artery is controversial and not only anti-atherogenic but also pro-atherogenic effects have been reported (68–79). (4) Systemic exposure to Dexa promotes metabolic changes, such as high serum lipids and glucose, and some other alterations, not found here (Table 1). Instead, the vigorous increase in IL-10-producing leukocytes (Supplementary Figure S7) argues in favor of an indirect immunoregulatory-operating process and a rather negligible direct athero-suppressive effect of Dexa or a VitD-improving action in our VitD/Dexa-treated mice as the key contributors to atheroprotection.

One limitation of the present study is that although some immunological insights were presented as likely contributors to the therapeutic effect, definitive experimental evidence of the mechanisms of atheroprotection operating in VitD/Dexa-treated mice was not obtained. Likewise, whether VitD/Dexa treatment inhibits atherosclerosis progression in already established plaques was not investigated here, and both aspects could be of future key value from the translational point of view. This, of course, does not change the value of this preclinical proof of concept for a small molecule formulation that induces Tregs and mitigates chronic inflammation in an extraordinarily relevant pathological scenario like atherosclerosis. Despite the broad clinical interest, no drug is currently available in the clinics specifically intended to induce Tregs. Clinical trials are in course in which the therapeutic potential of low-dose IL-2 is being evaluated as a Treg-inducer for inflammatory diseases, including atherosclerosis (80). Independent of the safety/efficacy outcomes of these trials, it is predictable that the costs of this biological agent will make it unrealistic for the routine medical practice. This contrasts with the low-cost, simplicity and convenience of the VitD/Dexa formulation, which make the two agents excellent candidates for reformulation, repurposing, clinical development and realistic clinical use.

CONCLUSION

In conclusion, we demonstrated that a simple combined formulation of two well-known small molecules, administered locally in the footpad via sc route and following a low-dose scheme with the intention to avoid systemic exposure, was able not only to safely promote regulatory immune cells but also to impact atherosclerosis development in vivo. This alternative immune intervention combining two cheap “old drugs” could be easily reformulated in a clinically compatible preparation for human proof of principle testing and clinical development. Moreover, we propose this approach as an attractive prophylactic or therapeutic option for other medical conditions in which stimulation of Treg have therapeutic potential, such as autoimmune, allergic, and other chronic inflammatory diseases.

DATA AVAILABILITY STATEMENT

The datasets generated for this study are available on request to the corresponding author.

ETHICS STATEMENT

The animal study was reviewed and approved by the Comite Institucional para el Uso y Cuidado de los Animales de Experimentacion, Universidad de Antioquia.

AUTHOR CONTRIBUTIONS

JR-P conceived, designed and directed the study, and wrote the manuscript. LO-Q, JJ, JT-G, and JR-P designed and planned the experiments. LO-Q, JJ, and JT-G conducted the experiments. LO-Q, JJ, and JR-P analyzed the results and interpreted the data. LO-Q and JJ contributed to manuscript writing and preparation. All authors read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2020.00743/full#supplementary-material
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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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