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Veld LGM, van Eck van der

# Saponin-based adjuvants enhance antigen cross-presentation in human CD11c<sup>+</sup> CD1c<sup>+</sup> CD5<sup>-</sup> CD163<sup>+</sup> conventional type 2 dendritic cells

Nataschja I Ho <sup>(b)</sup>, <sup>1</sup> Lisa G M Huis in 't Veld <sup>(b)</sup>, <sup>1</sup> Jesper van Eck van der Sluijs <sup>(b)</sup>, <sup>1,2</sup> Branco M H Heuts <sup>(b)</sup>, <sup>3</sup> Maaike W G Looman, <sup>1</sup> Esther D Kers-Rebel, <sup>1</sup> Koen van den Dries, <sup>4</sup> Harry Dolstra, <sup>2</sup> Joost H A Martens, <sup>3</sup> Willemijn Hobo, <sup>2</sup> Gosse J Adema <sup>(b)</sup>

### ABSTRACT

**Background** Adjuvants are key for effective vaccination against cancer and chronic infectious diseases. Saponinbased adjuvants (SBAs) are unique among adjuvants in their ability to induce robust cell-mediated immune responses in addition to antibody responses. Recent preclinical studies revealed that SBAs induced crosspresentation and lipid bodies in otherwise poorly crosspresenting CD11b<sup>+</sup> murine dendritic cells (DCs).

**Method** Here, we investigated the response of human DC subsets to SBAs with RNA sequencing and pathway analyses, lipid body induction visualized by laser scanning microscopy, antigen translocation to the cytosol, and antigen cross-presentation to CD8<sup>+</sup> T cells.

**Results** RNA sequencing of SBA-treated conventional type 1 DC (cDC1) and type 2 DC (cDC2) subsets uncovered that SBAs upregulated lipid-related pathways in CD11c<sup>+</sup> CD1c<sup>+</sup> cDC2s, especially in the CD5<sup>-</sup> CD163<sup>+</sup> CD14<sup>+</sup> cDC2 subset. Moreover, SBAs induced lipid bodies and enhanced endosomal antigen translocation into the cytosol in this particular cDC2 subset. Finally, SBAs enhanced cross-presentation only in cDC2s, which requires the CD163<sup>+</sup> CD14<sup>+</sup> cDC2 subset.

**Conclusions** These data thus identify the CD163<sup>+</sup> CD14<sup>+</sup> cDC2 subset as the main SBA-responsive DC subset in humans and imply new strategies to optimize the application of saponin-based adjuvants in a potent cancer vaccine.

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For numbered affiliations see end of article.

#### **Correspondence to**

Professor Gosse J Adema; gosse.adema@radboudumc.nl

### INTRODUCTION

The development of therapeutic vaccines against cancer and persistent infections has received much attention in recent decades. These vaccines are usually accompanied by an adjuvant to enhance the magnitude and durability of the immune response to the vaccine antigen.<sup>1</sup> Saponin-based adjuvants (SBAs) belong to the group of new-generation of adjuvants. They consist of triterpene glycosides found in a variety of plants and trees including the South American soapbark tree *Quillaja saponaria*. Specific saponin fractions

### WHAT IS ALREADY KNOWN ON THIS TOPIC

⇒ Saponin-based adjuvants (SBAs) have been used in the clinic against infectious diseases. They have the advantage to induce both CD4<sup>+</sup> T helper and CD8<sup>+</sup> T cells in addition to antibody responses. We have shown previously that only the CD11b<sup>+</sup> dendritic cell (DC) subset in the mouse responds to SBAs by cross-presentation of exogenous antigens to CD8<sup>+</sup> T cells in a lipid-body dependent manner. Here, we expand our research to human DC subsets, including the recently discovered subpopulations within the cDC2s.

### WHAT THIS STUDY ADDS

⇒ We defined that only the cDC2 subset is sensitive to SBAs, and not the cDC1 subset. Especially the recently identified subpopulation of the cDC2 subset, CD5<sup>-</sup> CD163<sup>+</sup> CD14<sup>+</sup> cDC2s, showed enhanced antigen translocation and upregulation of several lipidrelated pathways, which correlated with increased lipid body presence in the cells. Most importantly, our study revealed that SBAs can specifically enhance antigen cross-presentation in CDC2s, which requires the CD5<sup>-</sup> CD163<sup>+</sup> CD14<sup>+</sup> cDC2s.

### HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE OR POLICY

⇒ Understanding the DC subset-specificity and mechanisms of SBAs in humans is important for further development and application of SBA based vaccines against infectious diseases and cancer.

derived from this particular tree have been shown to exhibit potent immunostimulatory effects and are used for SBAs.<sup>2</sup> The first approved human vaccines containing SBAs are against the herpes zoster virus (RZV, Shingrix, GlaxoSmithKline) and malaria (RTS, S/AS01, Mosquirix, GlaxoSmithKline). The SBA-containing COVID-19 vaccine (NVX-COV2373, Novavax) has also successfully

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been used in the clinic. SBAs are extensively investigated as an adjuvant since they can induce robust cell-mediated immune responses activating both CD4<sup>+</sup> T helper and CD8<sup>+</sup> T cells in addition to antibody responses.<sup>3–6</sup> Especially for cancer, a strong CD8<sup>+</sup> T- cell response is necessary to eradicate the tumor.<sup>7</sup>

Dendritic cells (DCs) are well known for their ability to capture exogenous (tumor) antigens, and present them on major histocompatibility complex class I (MHCI) molecules to CD8<sup>+</sup> T cells, a process called antigen crosspresentation.<sup>8</sup> Different subsets can be distinguished within the murine DCs, namely the  $CD8\alpha^+$   $CD103^+$  DCs (conventional type 1 DCs (cDC1s)) and the CD8a<sup>-</sup> CD11b<sup>+</sup> DCs (conventional type 2 DCs (cDC2s)). The human BDCA-3<sup>+</sup> Clec9a<sup>+</sup> DCs (cDC1s) from human peripheral blood resemble murine  $CD8\alpha^+$   $CD103^+$  DCs, whereas the BDCA-1<sup>+</sup> DCs (cDC2s) display a gene expression profile resembling that of murine  $CD8\alpha^{-}$   $CD11b^{+}$ DCs.<sup>9 10</sup> The general consensus for murine DCs is that cDC1s are superior in cross-presenting exogenous antigens to CD8<sup>+</sup> T cells compared with cDC2s, whereas cDC2s mainly present antigens to CD4<sup>+</sup> T cells.<sup>11-13</sup> However, this distinction between cDC1s and cDC2s in humans is less pronounced.14-18

We and others have shown that an important characteristic of SBAs is their potency to enhance cross-presentation in murine DC and thereby induce efficient cytotoxic CD8<sup>+</sup> T- cell immunity.<sup>19-21</sup> Importantly, we have demonstrated using an in situ tumor ablation model that SBA injection induced tumor-specific CD8<sup>+</sup>T- cell responses, resulting in superior synergistic antitumor immunity compared with other adjuvants.<sup>22</sup> Interestingly, we also demonstrated that in vitro generated murine MHCII<sup>lo</sup> CD11b<sup>hi</sup> DCs and the corresponding *in vivo* CD11b<sup>+</sup> DCs from lymph nodes were most responsive to SBAs and exhibited high crosspresentation capacity.<sup>20</sup> We further reported that SBAs enhance endosomal antigen translocation to the cytosol, and that the endoplasmic reticulum (ER) stress-related PKR-like ER kinase (PERK) pathway, and proteasomal antigen degradation are crucial for SBA-induced crosspresentation.<sup>20 21</sup> Moreover, only the CD11b<sup>+</sup> DCs showed induction of lipid bodies (LBs) after treatment with SBAs. LBs, also called lipid droplets, are cytoplasmic organelles that play a major role in the storage of neutral lipids and energy metabolism, but also play a role in immune regulation.<sup>23 24</sup> Importantly, the presence of LBs is essential for SBA-induced increase in antigen cross-presentation and subsequent CD8<sup>+</sup> T- cell activation.<sup>20</sup>

The SBA-responsiveness of human DC subsets is still largely unknown, especially concerning their capacity to cross-present exogenous antigens. Recent RNA sequencing studies uncovered new members in the human cDC2 subset family.<sup>25–27</sup> Initially, it had been reported that cDC2s comprised two subsets, one is named DC2s which is genetically and functionally closer related to cDC1s, and the other is called CD14<sup>+</sup> DC3s that are more related to monocytes.<sup>25–26</sup> Now, cDC2s are proposed to be delineated and independent of monocytes and macrophages, and

comprise four subsets where the CD5<sup>+</sup> subset corresponds to DC2s, and CD5<sup>-</sup> subsets to DC3s.<sup>27 28</sup> The CD5<sup>-</sup> cDC2s could further be refined with the additional expression of CD163<sup>+</sup> or CD14<sup>+</sup>. The CD163<sup>+</sup> CD14<sup>+</sup> cDC2 subset has been described as being inflammatory.<sup>27</sup> There is not much known yet about their function, although a recent study showed that CD1c<sup>+</sup> CD163<sup>+</sup> cDC2 can prime CD8<sup>+</sup> CD103<sup>+</sup> T cells.<sup>28</sup>

In the current study, we investigated SBA-mediated cross-presentation in human cDC1s, the total cDC2 fraction, and the four cDC2 subsets using SBA, formulated with immunoactive saponin Fraction C in combination with cholesterol and phospholipid to generate so-called immune stimulatory complexes (ISCOMs) Matrix C.<sup>29</sup> We show that ISCOMs Matrix C can specifically induce crosspresentation in cDC2s, not cDC1s, and that this depends on the CD163<sup>+</sup> CD14<sup>+</sup> cDC2 subset. RNA sequencing analysis revealed that mainly lipid-related pathways were upregulated in the CD163<sup>+</sup> CD14<sup>+</sup> cDC2 subset. Additional data demonstrate enhanced antigen translocation and LB formation in this particular subset after treatment with ISCOMs. The discovery that specifically the human CD163<sup>+</sup> CD14<sup>+</sup> cDC2 subset can benefit from ISCOMs is an important step toward the design and development of new vaccines for diseases that require T cell-mediated immunity, including a potent cancer vaccine.

#### RESULTS

#### ISCOMs induce lipid-related pathways specifically in cDC2s

In order to identify the human DC subset that is responsive to ISCOMs, CD141<sup>+</sup> Clec9a<sup>+</sup> cDC1s and CD11c<sup>+</sup>  $CD1c^+$  cDC2s were isolated from blood (figure 1A), stimulated with ISCOMs or mock-treated, and subjected to RNA sequencing. The top 25 expressed genes in the cDC1 (figure 1B, left) and cDC2 subsets (figure 1B, right) confirmed the purity of the isolated subsets, as most of these genes correspond with genes that were previously found to be associated with these subsets (gene names in red, figure 1B).<sup>26 27 30</sup> Principal component analysis (PCA) indicated that cDC1s and cDC2s were separated along the PC2 axis, accounting for 33% variances, confirming substantial difference in gene expression between the subsets (figure 1C). Stimulation with ISCOMs showed 52% variances along the PC1 axis, indicating that ISCOMs treatment led to a substantial change in gene expression in both subsets (figure 1C). Analysis of differentially expressed genes (DEGs) indicated that in cDC1s 1327 and 1047 genes were significantly upregulated and downregulated upon ISCOMs treatment, respectively, whereas 1379 and 1174 genes were significantly upregulated and downregulated in cDC2s. Gene ontology (GO) analysis of upregulated genes was performed to indicate enrichment of Biological Processes upon ISCOMs treatment. Immunologically relevant GO pathways were selected and displayed as a radar plot showing the gene ratio. Upon ISCOMs treatment, mostly T-cell activation-related pathways were upregulated in cDC1s (figure 1D, orange),



Figure 1 ISCOMs upregulate distinct pathways in cDC2 and cDC1 subsets. (A) Purity of isolated cDC1s and cDC2s after MACS isolation analyzed by flow cytometry. (B) Top 25 highest expressed genes for cDC1s (left heatmap) and cDC2s (right heatmap) without ISCOMs stimulation based on transformed normalized counts from RNA sequencing. Known genes that are specific for each subset according to literature are depicted in red. (C) Principal component analysis (PCA) for bulk-sequenced cDC1s and cDC2s before and after 8 hours of ISCOMs stimulation. (D) Gene ontology (GO) enrichment for Biological Processes based on DEGs with p<0.05 and fold change ≥2 after stimulation with ISCOMs for 8 hours. Immunologically relevant GO pathways were selected and displayed as a radar plot showing the gene ratio. RNA sequencing for bulk subsets was performed on two to three healthy donors. cDC1, conventional type 1 DC; cDC2, conventional type 2 DC; DEGs, differentially expressed genes; IFN, interferon; IL, interleukin; ISCOMs, immune stimulatory complexes; JAK-STAT, Janus kinases-signal transducer and activator of transcription proteins; MACS, magnetic-activated cell sorting; Th, T helper; VEGF, vascular endothelial growth factor

whereas lipid-related pathways, and leukocyte proliferation and differentiation pathways were upregulated in cDC2s (figure 1D, blue). Previously, we showed that induction of PERK, activated upon ER stress, is crucial for ISCOMs-induced cross-presentation in murine DCs.<sup>21</sup> Here, both DC subsets induced "Regulation of response to stress" upon ISCOMs treatment, indicating that stressrelated pathways could also play a role in human DC subsets. Interestingly, "Lipid storage" is specifically upregulated in the cDC2s upon ISCOMs treatment. Since LBs are the main lipid storage organelles in the cell, this data encouraged us to further investigate LB induction upon ISCOMs treatment in the cDC2 subset. Moreover, "Cholesterol biosynthetic process" and "Regulation of lipid localization" further point towards involvement of LBs. Overall, ISCOMs induce a variety of immune



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**Figure 2** ISCOMs induce lipid bodies in cDC2s. (A) LB staining in cDC1s and cDC2s with Bodipy 493/503 (green) after 18 hours stimulation with ISCOMs or OA, analyzed by laser scanning microscopy. A magnification of a selected cell in the overview picture is shown in the upper right corner. Nucleus staining is done with DAPI (blue). (B) Quantification of the frequency of LB-positive cells was done with Fiji. Statistical analyses were done using one-way ANOVA using post hoc Tukey. Data are shown here from two healthy donors. cDC1 \*\*\*\*p<0.0001 (DF=2; *F* (2, 24) = 560.3), cDC2 \*\*\*\*p<0.0001 (DF=2; *F* (2, 37) = 366.2). (C) The number of LBs within the LB-positive cells was quantified with Fiji and divided into three categories, 1–3 LBs, 4–8 LBs, and 9–20 LBs per cell, and shown as frequency per subset. At least five overview pictures per sample were taken for quantification. ANOVA, analysis of variance; cDC1, conventional type 1 DC; cDC2, conventional type 2 DC; ISCOMs, immune stimulatory complexes; LBs, lipid bodies; OA, oleic acid.

signaling and stress pathways in both human DC subsets, and specifically lipid-related pathways in the cDC2s.

### ISCOMs induce lipid bodies in cDC2s

Since lipid-related pathways were upregulated in human cDC2s after ISCOMs stimulation, we investigated its impact on LB induction via staining with the neutral lipid dye Bodipy 493/503. Feeding the fatty acid oleic acid (OA) was used as a positive control for the overall capacity of the DC subsets to induce LBs. As expected, treatment with OA was able to induce LBs in both DC subsets (figure 2A,B). In contrast, treatment with ISCOMs for 18hours showed more than 80% LB-positive cDC2s,

with no detectable increase in cDC1s (figure 2A,B). Quantification of the number of ISCOMs-induced LBs within the LB-positive cells showed that the majority of cDC2s contained 4–8 LBs per cell, whereas ~40% of the cDC2 cells had even 9–20 LBs per cell after stimulation with ISCOMs (figure 2C). In the cDC1s, a very low number of LBs was observed upon ISCOMs treatment, and the number of LBs per cell was similar to untreated cells. Shorter stimulations of cDC2s with ISCOMs showed that LBs were still mostly absent after 5 hours, appearing in ~50% of the cells after 10 hours, reaching ~90% after 18 hours (figure 2 and online supplemental extended

data figure 1A,B). OA addition induced LB formation in ~30% of the cells already after 5 hours incubation, ~70% LB positive cells were present after 10 hours incubation, reaching ~90% after 18 hours (figure 2 and online supplemental extended data figure 1A,B). The cDC2s showed a similar increase in the number of LBs per cell in time following ISCOMs or OA stimulation (online supplemental extended data figure 1C). These data indicate that only cDC2s are capable of inducing LBs after ISCOMs stimulation, consistent with the upregulation of genes in lipid-related pathways.

### Enhanced antigen cross-presentation in cDC2s upon ISCOMs stimulation

Next, we investigated the ability of ISCOMs to enhance antigen cross-presentation in human cDC1s and cDC2s. Here, purified cDC1s and cDC2s were either loaded

with minor histocompatibility antigen (MiHA) HA-1 long peptide (VARFAEGLEKLKECVLHDDLLEARRPRA-HEZL) in the presence or absence of ISCOMs or with the short nominal peptide epitope (VLHDDLLEA) and co-cultured with peripheral blood mononuclear cells (PBMCs) from patients containing HA-1-specific CD8<sup>+</sup> T cells. Both cDC1s and cDC2s induced a comparable increase in frequencies of HA-1-specific CD8<sup>+</sup> T cells after stimulation with the positive control HA-1 short peptide as detected by dual-color HA-1 tetramer staining (figure 3A). Loading with the HA-1 long peptide also led to HA-1-specific CD8<sup>+</sup> T-cells expansion by both subsets, although to a lesser extent as compared with the nominal peptide, which is consistent with the requirement of prior antigen processing and cross-presentation. As expected, long peptide pulsed cDC1 (mean 19.7±12SD)



**Figure 3** Enhanced antigen cross-presentation in cDC2s upon ISCOMs. (A) HA-1- specific CD8<sup>+</sup> T-cell expansion indicated by double-positive HA-1 tetramer staining (in red) for cDC1s and cDC2s after stimulation with short peptide, long peptide, or ISCOMS+ long peptide. DCs (100,000 cells) were incubated with PBMCs (750,000 cells) in a 1:7.5 ratio. Representative flow cytometry data are shown from one healthy DC donor. (B) Fold expansion of HA-1-specific CD8<sup>+</sup> T cells in cDC2s and cDC1s. The numbers of different healthy DC donors used are depicted in the graphs. Statistical analyses were done using paired non-parametric t-test. \*p<0.05. cDC1, conventional type 1 DC; cDC2, conventional type 2 DC; ISCOMs, immune stimulatory complexes; PBMCs, peripheral blood mononuclear cells.

were much more effective than cDC2 (mean  $8.7\pm4.6$  SD) in cross-presenting antigen without adjuvant treatment (figure 3B). Strikingly, ISCOMs-treated long peptide loaded cDC2s increased frequencies and expansion of HA-1-specific CD8<sup>+</sup> T cells in six independent donors consistent with an enhanced antigen cross-presentation capacity, while ISCOMs-treated cDC1s did not show any enhancement (figure 3A,B). Taken together, these data demonstrate that ISCOMs specifically enhance antigen cross-presentation in cDC2s.

### ISCOMs upregulate lipid-related pathways in CD163<sup>+</sup> CD14<sup>+</sup> cDC2s

Since our data so far indicated that especially the cDC2s were responsive to ISCOMs and that ISCOMs induced higher numbers of LBs (9-20 per cell) in ~40% of the total cDC2 fraction, it was imperative to further investigate the impact of SBAs on the cDC2 subsets. Dutertre et al proposed the presence of at least four subsets within the cDC2s based on the expression of CD5, CD163, and CD14.<sup>27</sup> We have isolated these four cDC2 subsets according to the following markers: subset 1 (P1, CD11c<sup>+</sup> CD1c<sup>+</sup> CD5<sup>+</sup>), subset 2 (P2, CD11c<sup>+</sup> CD1c<sup>+</sup> CD5<sup>-</sup> CD163<sup>-</sup> CD14<sup>-</sup>), subset 3 (P3, CD11c<sup>+</sup> CD1c<sup>+</sup> CD5<sup>-</sup> CD163<sup>+</sup> CD14<sup>-</sup>), and subset 4 (P4, CD11c<sup>+</sup> CD1c<sup>+</sup> CD5<sup>-</sup> CD163<sup>+</sup>  $CD14^+$ ) (figure 4A). Since P1 and P2 were described to be closely related<sup>28</sup> and due to the low amounts of P1 and P2 that could be isolated per donor, we combined these two subsets for subsequent RNA sequencing analysis.

The top 25 expressed genes were identified in untreated cDC2 subsets and compared with the genes found by Dutertre *et al* (figure 4B. P1+2, left; P4 right). Similar genes (gene names in red) were identified for P1+2 and P4. Despite the fact that none of the genes identified by Dutertre *et al* were expressed in the top 25 of P3, no P1+2 nor P4 genes were found in P3 either (figure 4B. middle). This confirms that all cDC2 subsets exhibit distinct marker and gene expression profiles. Interestingly, one of the highest expressed genes within P4 is ITGAM, which encodes CD11b, the marker that distinguishes the SBA-responsive DC subset in mice.

PCA indicated that untreated P1+2 and P3 were clustered more closely together, and that P4 was clearly separated from P1+2 and P3 based on gene expression on the PC2 axis (figure 4C). After ISCOMs stimulation, all cDC2 subsets responded with a change in gene expression on the PC1 axis. P4 was still the most separated from the other subsets on the PC2 axis (figure 4C). The shift induced by OA stimulation in the subsets appears closely related to the shift observed for ISCOMs treatment, while their activation using the Toll-like receptor 4 (TLR4) ligand lipopolysaccharide (LPS) induced a gene expression profile distinct from the other treatments (figure 4C). Interestingly, PCA of the different human and murine DC subset after ISCOMs stimulation showed a closer similarity between the human total cDC2s, P3 cDC2s, P4 cDC2s, and the SBA-responsive murine MHCII<sup>lo</sup> CD11b<sup>hi</sup> DC subset based on the PC2 axis, whereas human P1+2

cDC2s were more closely related to the non-responsive murine MHCII<sup>hi</sup> CD11b<sup>int</sup> DC subset (figure 4D). Analvsis of DEGs in the cDC2 subsets induced upon ISCOMs treatment indicated that in P1+2 cDC2s 1721 and 1529 genes were significantly upregulated and downregulated, respectively, whereas 1346 and 1472 genes were significantly upregulated and downregulated in P3 cDC2s, respectively. In P4 cDC2s 1288 were significantly upregulated and 1468 genes were downregulated upon ISCOMs treatment. GO enrichment analysis and subsequent selection of immunologically relevant pathways of upregulated genes upon ISCOMs treatment revealed distinctive pathways, but also some overlap in upregulated pathways in the different cDC2 subsets (figure 4E). One of the pathways upregulated in all subsets is "Regulation of response to stress", which is in line with what was described earlier for the total cDC2s. While different immune pathways were induced in P1+2 and P3, the pathways induced in P4 were mostly lipid-related. Interestingly, lipid-related pathways were also observed earlier in total cDC2s upon ISCOMs stimulation. The "Regulation of lipid storage", "Regulation of lipid transport", "Unsaturated fatty acid metabolic process" and "Lipid biosynthetic process" pathways all strongly suggest the induction of LBs in this subset. Upon LPS treatment, a different set of induced pathways was observed in P4 cDC2s showing upregulation of inflammatory response pathways including "Positive regulation of acute inflammatory response" and "Inflammatory response to antigenic stimulus" (online supplemental extended data figure 2A). This is in line with the description of P4 as the inflammatory subset in literature.<sup>27</sup> By using a combinatorial approach filtering out ISCOMs-induced genes in the other subsets and OA/ LPS-induced genes in P4, we identified 65 genes that are specifically upregulated by ISCOMs in P4, but not upregulated upon ISCOMs treatment in cDC1s, P1+2, P3, nor upon OA or LPS treatment in P4 (online supplemental extended data figure 2B and table 1). Strikingly, this included genes such as TRIB3 (~4-fold induction), which is an ER stress-inducible gene, downstream of the PERK pathway. We have published before that TRIB3 and other genes downstream of PERK are upregulated in the murine MHCII<sup>lo</sup> CD11b<sup>hi</sup> DC subset upon ISCOMs stimulation and that PERK activation is crucial for ISCOMs-induced cross-presentation in this DC subset.<sup>21</sup> Moreover, lipid-related genes APOE (~5-fold induction), CDS1 (~5-fold induction), GBA (~4fold induction), and HSD17B14 (~14-fold induction) are specifically upregulated in P4. APOE is best known for its role in lipoprotein metabolism,<sup>31</sup> whereas CDS1, also known as CPG-diacylglycerol synthase 1, is crucial for the regulation of lipid droplet growth.<sup>32</sup> Mutations in the GBA gene have been linked to lipid alterations that are associated with Parkinson's disease. HSD17B14 is a 17-beta-hydroxysteroid dehydrogenase that is involved in peroxisomal  $\beta$ -oxidation of fatty acids, and the breakdown of fatty acid chain elongation in the ER.<sup>33 34</sup> In conclusion, these data indicate that ISCOMs upregulate





**Figure 4** ISCOMs upregulate lipid-metabolism-mediated pathways in CD163<sup>+</sup> CD14<sup>+</sup> cDC2s. (A) Gating strategy of the CD11c<sup>+</sup> cDC2 subsets: P1 (CD5<sup>+</sup>), P2 (CD5<sup>-</sup> CD163<sup>-</sup>), P3 (CD5<sup>-</sup> CD163<sup>+</sup> CD14<sup>-</sup>), and P4 (CD5<sup>-</sup> CD163<sup>+</sup> CD14<sup>+</sup>) by flow cytometry. (B) Top 25 highest expressed genes for cDC2 subsets P1+P2 (left heatmap), P3 (middle heatmap), and P4 (right heatmap) without ISCOMs stimulation based on transformed normalized counts from RNA sequencing. Known genes for each subset according to literature are depicted in red. (C) PCA for bulk-sequenced P1+2, P3, and P4 cDC2s before and after 8 hours of ISCOMs, LPS, or OA stimulation. (D) PCA of bulk-sequenced human cDC1, cDC2, P1+2, P3, P4 after 8 hour ISCOM stimulation and murine MHCII<sup>hi</sup> CD11b<sup>int</sup> and MHCII<sup>lo</sup> CD11b<sup>hi</sup> DCs after 5 hours of ISCOMs stimulation. (E) GO enrichment for Biological Processes based on DEGs with p<0.05 and fold change  $\ge 2$  after stimulation with ISCOMs for 8 hours. Immunologically relevant GO pathways were selected and displayed as a radar plot showing the gene ratio. RNA sequencing for bulk cDC1s and cDC2s was performed on two healthy donors, for cDC2 subsets on three healthy donors, and for the murine subsets from two mice. cDC1, conventional type 1 DC; cDC2, conventional type 2 DC; DEGs, differentially expressed genes; GO, gene ontology; IL, interleukin; ISCOMs, immune stimulatory complexes; LPS, lipopolysaccharide; MHC, major histocompatibility complex; OA, oleic acid; PCA, principal component analysis; ROS, reactive oxygen species; TGF, transforming growth factor; VEGF, vascular endothelial growth factor.

Table 1 Genes specific for P4 I	SCOMs
Genes	Fold change
ABCB1	7.713929243
APOE	4.975581838
CAPNS2	9.979767603
CCAT1	28.39131712
CDKN3	6.914625195
CDS1	5.098935519
CHST1	27.72334149
CLEC4A	4.008313234
COL22A1	33.00824066
COL4A2-AS1	14.13743764
FBXO10	4.190391801
FBXO2	26.89625517
FIGF	33.40963375
GAL	44.38457189
GBA	4.231682052
GIN1	4.366585996
GNAO1	36.76779046
HEBP1	4.068944088
HSD17B14	14.44046381
KHDRBS3	4.952637264
LANCL3	25.65677356
LINC00659	15.24906197
LOC101929371	26.25785865
LOC102723524	6.926784041
LOC102723569	11.42477861
LOC102724153	4.150846501
LOC102724316	6.632749517
LOC102724369	4.265184456
LOC102724571	20.1669791
LOC285638	4.432657248
MAS1	117.7526355
MICA	6.128474378
MIR4257	4.978114229
MIR499B	31.61367763
MIR591	6.12076253
MKNK1	4.082959086
MPP4	9.526311288
NDUFAF4P1	18.4677358
NRBF2	4.098259785
OR10P1	50.69324162
PCDHGA11	21.87064488
PCDHGA12	26.06845056
PCDHGB5	25.65677356
PHF23	4.04618254
PITX1	25.15984089

Continued

Table 1 Continued	
Genes	Fold change
POM121L9P	34.77758985
RBM47	4.153873052
RBP1	18.80462484
RP11-265P11.2	28.33723066
RP11-290F20.2	47.3134694
SCARNA20	7.661874874
SDS	5.593669543
SLC4A2	4.11043159
SPOCD1	92.05931005
TACSTD2	34.8083394
TMEM52B	5.153274705
TNFSF18	43.01283957
TRIB3	4.091024102
TRIM47	12.67510751
TRNAR22	9.36268282
TRNQ	6.691803218
UNC5B	4.492870072
YPEL4	6.67719103
ZBTB7C	27.06881449
ZNF827	6.877323214

ISCOMs, immune stimulatory complexes.

lipid-related pathways specifically in the P4 (CD163<sup>+</sup> CD14<sup>+</sup>) cDC2 subset.

### ISCOMs induce lipid bodies mainly in CD163<sup>+</sup> CD14<sup>+</sup> cDC2s

Since ISCOMs upregulate lipid-related pathways in the CD163<sup>+</sup> CD14<sup>+</sup> (P4) cDC2 subset, we further investigated their ability to induce LBs. No LBs were detected in all isolated cDC2 subsets directly after cell sorting (figure 5A). Analysis of the P1 and P2 subsets revealed that there were already LBs present in ~70% of the cells after 18 hours in medium (figure 5B,C). However, this was not further enhanced by ISCOMs stimulation. LBs were also present in populations P3 and P4 (~30% and 20%, respectively) when cultured with medium (figure 5B,C). Interestingly, the frequency of LB-containing cells in P3 and P4 was significantly increased after ISCOMs stimulation to ~80% and 90%, respectively. Further quantification of the number of LBs per cell revealed that  $\sim 15\%$ of P4 contained ~11-20 LBs per cell compared with ~3% in the P1, P2, and P3 subsets when treated with ISCOMs (figure 5D). Shorter stimulation with ISCOMs showed comparable results, where significant induction of LBs was observed in P3 and P4 (online supplemental extended data figure 3A-C). To further confirm the identity of the lipid aggregates, we co-stained the cells with Bodipy and the LB marker ADRP (also known as Perilipin 2) after stimulation with ISCOMs or OA. By performing a fluorescence intensity profile analysis, we observed that





the Bodipy signals are highly correlated with the localization of ADRP in all four cDC2 subsets, indicating that the lipid aggregates were indeed LBs (figure 6, online supplemental extended data figure 4). Our results indicate that ISCOMs induce LBs mostly in the P4 (CD163<sup>+</sup> CD14<sup>+</sup>) cDC2s, both with respect to the frequency of LB positive cells and the number of LBs per cell.

## Enhanced antigen translocation by ISCOMs in CD163<sup>+</sup> CD14<sup>+</sup> cDC2s

We and others have proposed that enhanced endosomal antigen translocation to the cytosol, thereby allowing more antigens to be available for processing, could play a role in SBA-induced cross-presentation.<sup>20 35</sup> To investigate antigen translocation, the cDC2 subsets received mitochondrial protein cvtochrome c in addition to ISCOMs. The translocation and subsequent exposure of cytochrome c from endosomes to the cytosol induces cell apoptosis, which can be measured by reduced cell metabolic activity and viability. The translocation of cytochrome c in non-stimulated conditions and subsequent decrease in cell metabolic activity is higher in P3 and P4 (~60%) compared with P1+2 (~10–15%) (figure 7A). This baseline difference might be caused by differences in the uptake of cytochrome c or by the differential capacity of translocation between the subsets. Strikingly, P4 was the only cDC2 subset that showed significantly reduced cell metabolism activity in the presence of ISCOMs in combination with different concentrations of cytochrome c (figure 7A). Antigen translocation of cytochrome c was observed in cDC1s, however, this was not further enhanced by ISCOMs (online supplemental extended data figure 5A). These data demonstrate that ISCOMs specifically enhance antigen translocation in the CD163<sup>+</sup> CD14<sup>+</sup> (P4) cDC2s subset.

### Enhanced antigen cross-presentation by ISCOMs requires CD163<sup>+</sup> CD14<sup>+</sup> cDC2s

Since we demonstrated earlier that ISCOMs could only augment antigen cross-presentation to HA-1<sup>+</sup> T cells in cDC2s, we further zoomed in on the cDC2 subsets. The subsets P1+2, P3, and P4 were individually pulsed with HA-1 long peptide in the presence or absence of ISCOMs stimulation and then incubated with PBMCs containing HA-1 specific CD8<sup>+</sup> T cells. HA-1 long peptide loading onto P1+2 and P3 cDC2 subsets resulted in an increase in absolute numbers of HA-1 specific CD8<sup>+</sup> T cells, however, there was no further enhancement by addition of ISCOMs (figure 7B). Unexpectedly, co-culture of subset P4 and PBMCs showed a remarkable decrease in cell viability of the PBMCs containing HA-1 specific CD8<sup>+</sup> T cells regardless of the stimuli that were added (online supplemental extended data figure 5B). Upon co-culture with P4 there is a decrease in CD3<sup>+</sup> T cells, especially in the amount of CD4<sup>+</sup> T cells (online supplemental extended data figure 5B). Further experiments using PBMCs from healthy donors showed similar results, ruling out that this is a specific effect of the PBMCs from patients containing

HA-1 CD8<sup>+</sup> T cells (online supplemental extended data figure 5C). As this effect was not observed in the PBMCs co-cultured with the total cDC2 population, we combined P4 with either P1+2 or P3. Combining P4 with either of the other populations prevented the loss of CD3<sup>+</sup> CD4<sup>+</sup> T cells induced by population P4 only (online supplemental extended data figure 5C). Therefore, we loaded different subset combinations with HA-1 long peptide with or without ISCOMs. Combining the subsets P1+2 with P3 or P4 showed no enhanced expansion of HA-1 specific CD8<sup>+</sup> T cells after ISCOMs stimulus (figure 7D). However, the combination of P3 with P4 showed a superior increase in the frequencies and amount of HA-1 specific CD8<sup>+</sup> T cells in the presence of ISCOMs for multiple donors (figure 7C,D). Titrating the amounts of P4 when combined with P3 implies that there is a ratio optimum for the enhancement of ISCOM-mediated antigen cross-presentation, too high or too low amounts of P4 will cause diminished effects (online supplemental extended data figure 6A). Taken together, our results indicate that enhanced antigen cross-presentation is induced by ISCOMs in cultures consisting of P4 cDC2s supplemented with P3 cDC2s.

### DISCUSSION

SBAs are new-generation adjuvants that are recently implemented in vaccination strategies against diseases such as COVID-19. The most outstanding advantage of SBAs over older adjuvant types, for example, aluminum salts or oil-in-water emulsion, is their ability to induce both CD4<sup>+</sup> T helper and CD8<sup>+</sup> T-cell responses.<sup>3–6</sup> CD8<sup>+</sup> T cells play a prominent role in battling cancer and in infectious diseases such as HIV and COVID-19.<sup>7 36 37</sup> Although the efficiency of SBA-containing vaccines has clearly been demonstrated, its effect on inducing cross-presentation in human DCs remains largely unknown. In the current study, we demonstrate that SBAs can enhance antigen cross-presentation in a specific human cDC2 subset, identified as CD11c<sup>+</sup> CD1c<sup>+</sup> CD5<sup>-</sup> CD163<sup>+</sup> cDC2s, leading to antigen-specific CD8<sup>+</sup> T-cell expansion.

It has been suggested that human cDC2s are the homologue of murine CD8α<sup>-</sup> CD11b<sup>+</sup> DCs.<sup>9 10</sup> Recent studies have reported the heterogenicity of the human cDC2s.<sup>25–27</sup> One out of the four newly identified cDC2 subsets is the CD163<sup>+</sup> CD14<sup>+</sup> cDC2s.<sup>27</sup> We have previously shown that SBAs could only induce cross-presentation in murine MHCII<sup>lo</sup> CD11b<sup>hi</sup> DCs.<sup>20</sup> With the current study, we discovered a similar specificity of SBAs for their human counterpart, most specifically in CD163<sup>+</sup> CD14<sup>+</sup> (P4) cDC2s. Interestingly, the CD163<sup>+</sup> CD14<sup>+</sup> (P4) cDC2s highly expressed ITGAM (CD11b), which corresponds to the murine SBA-responsive DC subset. The CD163<sup>+</sup>  $CD14^+$  (P4) cDC2s are the cells most likely responsible for the enhanced cross-presentation ability upon SBAtreatment. However, as the addition of the CD163<sup>+</sup> CD14<sup>-</sup> (P3) subset to the  $CD163^+$   $CD14^+$  (P4) co-culture with PBMCs was required to sustain the viability of PBMCs,



**Figure 6** Co-localization between separate markers for LBs in cDC2 subsets. LB staining with Bodipy 493/503 (green) and ADRP (magenta) in P3 and P4 cDC2 subsets after 18 hours stimulation with ISCOMs or OA. Overlay pictures between Bodipy and ADRP are shown as Merge. Co-localization histograms for each fluorophore were created for a selected area in Fiji (indicated by a line on the Merge+DAPI picture). Co-localizations were depicted by arrows in the histograms. At least five pictures were taken per sample, and representative data are shown here for two healthy donors. ISCOMs, immune stimulatory complexes; LBs, lipid bodies; OA, oleic acid.



Figure 7 Enhanced antigen translocation and cross-presentation by CD14<sup>+</sup> CD163<sup>+</sup> cDC2s upon ISCOMs. (A) Antigen translocation measured by CCK8 assay after stimulation with ISCOMs and titration of cytochrome c, indicated as relative metabolic activity and viability to medium control. Statistical analyses were done comparing using one-way ANOVA using post hoc Tukey. Representative data from one donor with two technical replicates are shown for two healthy donors. \*p<0.05, \*p<0.01, \*\*\*p<0.001 (DF=10; F (10, 11) = 444.0). (B) Fold expansion of HA-1-specific CD8<sup>+</sup> T cells in P1+2 and P3. P1+2 (25,000+25,000 cells) and P3 (50,000 cells) were incubated with PBMCs (500,000 cells) in a 1:10 ratio. Data are shown for four healthy DC donors. (C) HA-1-specific CD8<sup>+</sup> T-cell expansion indicated by double-positive HA-1 tetramer staining (in red) for P3+4 (87,000+12,500 cells) after stimulation with short peptide, long peptide, or ISCOMS+long peptide. DCs (100,000 cells) were incubated with PBMCs (1×10<sup>6</sup> cells) in a 1:10 ratio. Representative flow cytometry data are shown from one healthy DC donor. (D) Fold expansion of HA-1-specific CD8<sup>+</sup> T cells in different combinations of cDC2 subsets for the number of donors indicated per graph. P1+2 (12,500+12,500 cells, respectively), P3 (25,000 cells), and P4 (25,000 cells) were combined into P1+2+3, and P1+2+4 (50,000 cells total). For P3+4 the following amounts were used: Donor 1 and 2 (P3 25,000+P4 25,000), donor 3 (P3 75.000+P4 25.000 cells), donor 4 (P3 87.000+P4 12.500 cells), and donor 5 (P3 93.750+P4 6.250 cells). The DCs were incubated with PBMCs containing HA-1-specific CD8<sup>+</sup> T cells in a 1:10 ratio. Statistical analyses were done using paired non-parametric t-test. N.s. non significant, \*p<0.05, \*\*p<0.01. ANOVA, analysis of variance; DCs, dendritic cells; DMSO, dimethyl sulfoxide; ISCOMs, immune stimulatory complexes; PBMCs, peripheral blood mononuclear cells.

particularly the CD4<sup>+</sup> T cells, we cannot formally rule out a contribution of P3. It has been suggested that splenic  $CD8\alpha^+$  DCs from mice can kill  $CD4^+$  T cells through Fas/Fas-ligand interaction as part of regulating T-cell responses.<sup>38</sup> However, we could not detect high expression of tumor necrosis factor superfamily ligands (such as TRAIL, FasL, or OX40L) or PD-L1 on the CD163<sup>+</sup> CD14<sup>+</sup> (P4) cDC2s based on our RNA sequencing data. Our preliminary data showed that overnight cDC2 stimulation with TLR ligands LPS, R848, or Poly(I:C) did not prevent the decrease in T-cell numbers after 7 days, but rather resulted in a further reduction in T-cell viability in case of P3 and P4. This suggests that in our co-culture system with purified DCs the T cells are sensitive for (over)stimulation and require further studies. Since these cDC2 subsets have been discovered only recently, there is still much unknown about their individual functions and their relation to each other. It has been suggested that the four cDC2 subsets reflect their differentiation stages starting from CD5<sup>+</sup> cells (P1), to CD5<sup>-</sup> CD163<sup>-</sup> cells (P2), to CD5<sup>-</sup> CD163<sup>+</sup> CD14<sup>-</sup> cells (P3), and finishing with CD163<sup>+</sup> CD14<sup>+</sup> (p4) cells, proposing a differentiation and/or activation continuum.<sup>27</sup> An important note is that CD163<sup>+</sup> CD14<sup>-</sup> (P3) cells are the largest subset of cDC2s followed by  $CD163^+$   $CD14^+$  (P4) cells. We demonstrated that there was an optimum ratio between the two subsets for crosspresentation and T-cell expansion, where lower amounts of CD163<sup>+</sup> CD14<sup>+</sup> (P4) cells were required compared with CD163<sup>+</sup> CD14<sup>-</sup> (P3) cells. This might be a reflection of how the subsets are naturally occurring in the body with vet unidentified mechanisms of interplay between the subsets. Interestingly, a recent study showed that CD163<sup>+</sup> cDC2s, encompassing both CD14<sup>+</sup> and CD14<sup>-</sup> cDC2s (comparable with our P3 and P4 combined), could prime CD8<sup>+</sup> CD103<sup>+</sup> tissue-resident memory T cells.<sup>28</sup> They investigated whether CD163<sup>+</sup> cDC2s could differentiate from the CD163<sup>-</sup> cDC2s (comparable with our P1 and P2 combined) or from monocytes in the presence of granulocyte-macrophage colony-stimulating factor (GM-CSF)-expressing stromal cells. They showed that after 2 days neither CD163<sup>-</sup> cDC2s nor monocytes differentiated into CD163<sup>+</sup> cDC2s, whereas CD163<sup>+</sup> cDC2s did upregulate CD14 expression. We observed similar results when we incubated the four different cDC2 subsets individually for 1 day in the presence of GM-CSF. Interestingly, the authors also showed that CD163<sup>+</sup> cDC2s could only be differentiated from human cord blood-derived CD34<sup>+</sup> hematopoietic stem and progenitor cells in the presence of GM-CSF but not FLT3L, while FLT3L was already sufficient to induce the differentiation of CD163<sup>-</sup> cDC2s.<sup>28</sup> Taken together, these results indicate that CD163<sup>+</sup>  $CD14^-$  (P3) and  $CD163^+$   $CD14^+$  (P4) cDC2s are probably more closely related compared with  $CD5^+$  (P1) and CD5<sup>-</sup> CD163<sup>-</sup> (P2) cDC2s. The reason why only cDC2s, and specifically the  $CD163^+$   $CD14^+$  (P4) subset, are sensitive to SBAs remains not fully understood. There could be a difference in SBAs and antigen uptake between the different DC subsets. Another possibility is that there are

distinct pathways activated in the different DC subsets on SBA uptake. We have recently published that the PERK pathway of the unfolded protein response is selectively upregulated in murine MHCII<sup>lo</sup> CD11b<sup>hi</sup> DCs upon SBA stimulation.<sup>21</sup> One of the genes that is related to the PERK pathway, TRIB3, was specifically upregulated in both the murine CD11b<sup>+</sup> DCs and the human CD163<sup>+</sup> CD14<sup>+</sup> (P4) subset upon ISCOMs stimulation. It would be of interest to investigate whether PERK also plays a dominant role in SBA-induced cross-presentation in CD163<sup>+</sup> CD14<sup>+</sup> (P4) cDC2s.

In addition to SBA-mediated cross-presentation, we showed an increase in antigen translocation in CD163<sup>+</sup>  $CD14^+$  (P4) cDC2s, indicating that SBAs facilitate antigen escape from endosomes. Others have shown comparable results in human monocyte-derived DCs, where SBAs could induce efficient cross-presentation and antigen translocation of the cancer testis antigen NY-ESO-1 based on interferon (IFN)- $\gamma$  production by CD8<sup>+</sup> T cells.<sup>35</sup> We have published before that SBA-induced cross-presentation is proteasome-dependent, suggesting that antigens are processed and loaded on MHCI molecules through the cytosolic pathway.<sup>20</sup> Therefore, antigen translocation is a critical step to transport antigens that are initially stored in endosomal compartments into the cytosol for proteasomal degradation and subsequent antigen crosspresentation. It is still not fully understood how SBAs facilitate antigen translocation. It has been proposed that the SBA QS-21 mediates pore formation in lysosomes, resulting in the release of macromolecules to the cvtosol.<sup>39</sup> However, others argue that antigen translocation is a highly regulated process that involves dislocation through the endoplasmic-reticulum-associated protein degradation (ERAD) machinery or a transmembrane pore complex instead of a simple lysosome leakage.<sup>40-42</sup> Their main argument is that uncontrolled lysosome leakage would lead to the cytosolic release of cathepsins and activate the NLRP3 inflammasome resulting in cell death.<sup>43 44</sup> Recent work by Gros *et al* showed that the repair of endosomes is a highly regulated process, which means that endosomal leakage can still be controlled within the cell.<sup>45</sup> Since our previous studies in mice demonstrated that not all pore-forming saponins (which mostly interact with the cell membrane) showed adjuvant activity in DCs, SBA probably induce endosomal translocation in a more controlled and localized manner.<sup>20</sup>

One of the characteristics of SBA sensitivity in DCs is their ability to induce LBs. We have demonstrated before that LB induction was observed only in SBA-stimulated murine MHCII<sup>lo</sup> CD11b<sup>hi</sup> DCs and that this is correlated with their enhanced cross-presentation ability.<sup>20</sup> Importantly, we showed in the same study that blocking LB formation with inhibitors such as 5-tetradecyloxy-2-furoic acid, diacylglycerol acyltransferase 1 and 2, or by inhibiting acyl-CoA synthetase or acetyl-CoA carboxylase, was correlated with impaired antigen cross-presentation by DCs. Moreover, IGTP (an IFN-related GTPase associated with lipid bodies) and ADRP knockout mice showed reduced lipid body accumulation and levels of crosspresentation after ISCOM treatment.<sup>20</sup> These results indicate the vital role of LB formation for SBA-induced cross-presentation by DCs. We now show that SBAs could only augment LBs in cDC2s, especially in the CD163<sup>+</sup> CD14<sup>-</sup> (P3) and CD163<sup>+</sup> CD14<sup>+</sup> (P4) cDC2 subsets, and not in cDC1s. Interestingly, we already found many  $CD5^+$  (P1) and CD5<sup>-</sup> CD163<sup>-</sup> cDC2s (P2) containing LBs after incubation with medium for 18 hours, although the frequency or number of LBs were not further enhanced upon SBA stimulation. Importantly, we show with RNA sequencing that lipid-associated pathways and genes were specifically upregulated in the CD163<sup>+</sup> CD14<sup>+</sup> (P4) cDC2s by SBAs, the same subset where we found the highest induction of LBs and enhanced cross-presentation capacity. Although ISCOMs could also induce LBs in the CD163<sup>+</sup>CD14<sup>-</sup> (P3) subset, it was to a much lower extent compared with P4. Ouantification of the data from confocal microscopy analysis showed that the amount of LBs per cell is much higher in P4 than P3. Moreover, RNA sequencing analysis showed that the lipid metabolic pathways that were upregulated in P3 were less dominant than in P4 after ISCOMs treatment, and therefore did not fall into the selection of top upregulated pathways in P3. It is known that different stimuli or metabolic states in cells can affect their protein and/or lipid compositions, and even different LB populations could be observed within the same cell.<sup>46–48</sup> While some studies have pointed out the essential role of LBs in inducing DC cross-presentation,<sup>20 49-51</sup> contradictory studies showed the opposite where LBs hamper crosspresentation in tumor-associated DCs.<sup>52–54</sup> This suggests that the type of LBs and their content is crucial for the desired functional outcome. Further analysis of lipid content in the different DC subsets, for example, by lipidomics, would give better insight into the different types of LBs and their distinct functions.

A recent study demonstrated that the infiltration of CD163<sup>+</sup> cDC2s in luminal breast cancer primary tumors, containing both CD14<sup>+</sup> and CD14<sup>-</sup> cells, was associated with the abundance of CD8<sup>+</sup> CD103<sup>+</sup> T cells,<sup>28</sup> which can contribute to improved breast cancer prognosis.<sup>55 56</sup> In contrast, a study reported that CD1c<sup>+</sup> CD14<sup>+</sup> cells were expanded in patients with melanoma and that cDC1 vaccines with high amounts of CD1c<sup>+</sup> CD14<sup>+</sup> cells were less effective in inducing antigen-specific CD4<sup>+</sup> T-cell responses in patients with melanoma.<sup>57</sup> Another study described that CD14<sup>+</sup> cDC2s have a higher capacity to prime naïve CD4<sup>+</sup> T cells towards Th2 and Th17 cells, which might correspond to disease activity in patients with systemic lupus erythematosus.<sup>27</sup> It would be interesting to investigate whether the use of SBAs could skew the immune responses of CD14<sup>+</sup> cDC2s into a more favorable outcome in those patients. Moreover, it is important to further investigate the functions of CD14<sup>+</sup> cDC2s in various tissues and tumor microenvironments besides blood circulation. Interestingly, Bosteels et al recently showed that the activity of the adjuvant AS01, a combination of saponin and monophosphoryl lipid A (MPL),

is associated with the recruitment of cDC1s and cDC2, including the inflammatory cDC2s.<sup>58</sup> They showed that these (inflammatory) cDC2s are equally efficient at priming both OVA-specific CD4<sup>+</sup> and CD8<sup>+</sup> T cells. Mice lacking CCR2-dependent and Flt3-dependent inflammatory cDC2s failed to raise proper antibody and T-cell responses. All in all, these and our data support the finding that the inflammatory cDC2 subset could be a key target for adjuvant systems like saponin to induce adaptive immunity in humans.

In conclusion, we show that SBAs enhance antigen cross-presentation in recently identified human  $\text{CD11c}^+$   $\text{CD5}^ \text{CD163}^+$  cDC2s. Since SBAs are currently gaining ground in vaccination strategies, it is important to understand the mechanisms of SBA adjuvant activity and their specificity for DC subsets. Further work would be essential to understand the roles of these subsets in SBA-mediated immune responses and contribute to improving cancer vaccination development.

### **METHODS**

### **Cell isolation**

PBMCs were isolated from buffy coats or apheresis from healthy donors (Sanquin, Nijmegen, The Netherlands) using Lymphoprep (STEMCELL Technologies). Total CD141<sup>+</sup> CLEC9A<sup>+</sup> DCs (cDC1s) were MACS isolated from PBMCs using the human CD141 (BDCA-3) MicroBead kit (Miltenyi Biotec), followed by isolation of total CD1c<sup>+</sup> DCs (cDC2s) using the human CD1c (BDCA-1)<sup>+</sup> dendritic cell isolation kit (Miltenyi Biotec). A purity check of the subsets after isolation was performed by additional FACS staining and measured flow cytometry (BD FACSCanto II). The cDC1s were stained with anti-Clec9a (SONY Biotechnology) and anti-CD141 antibodies (BioLegend), cDC2s were stained with anti-CD11c (BD Biosciences) and anti-CD1c antibodies (Miltenyi Biotec). Further cDC2 subset isolation was performed by FACS sorting (BD FACS Aria II SORP) using anti-CD11c (BC Biosciences), anti-CD1c (Miltenyi Biotec), anti-CD5 (BioLegend), anti-CD163 (eBioscience), and anti-CD14 (BD Biosciences) antibodies. The four cDC2 subsets were gated according to the following markers: P1 (CD11c<sup>+</sup> CD1c<sup>+</sup> CD5<sup>+</sup>), P2 (CD11c<sup>+</sup> CD1c<sup>+</sup> CD5<sup>-</sup> CD163<sup>-</sup>), P3 (CD11c<sup>+</sup> CD1c<sup>+</sup> CD5<sup>-</sup> CD163<sup>+</sup> CD14<sup>-</sup>), and P4 (CD11c<sup>+</sup> CD1c<sup>+</sup> CD5<sup>-</sup> CD163<sup>+</sup> CD14<sup>+</sup>). Cells were cultured in X-VIVO 15 serum-free hematopoietic cell medium (Lonza) with an addition of 10% fetal bovine serum (Gibco), and 40U/mL human granulocyte-macrophage colony-stimulating factor (hGM-CSF) (Immunotools).

### **RNA sequencing**

The different cell populations were isolated, as described above, from healthy apheresis donors. Cells were stimulated with Matrix C ISCOMs (400 ng/mL, MSD Animal Health, Boxmeer, the Netherlands), LPS (1  $\mu$ g/mL, Sigma-Aldrich), or OA (50  $\mu$ M, Sigma-Aldrich) for 8 hours. RNA isolation for each subset was performed with TRIzol

Reagent (Invitrogen) and DNAse treatment (DNA-free DNA Removal Kit, Invitrogen). RNA sequencing was performed and Fastq files were mapped to the reference human genome GRCh38 using the Seq2science pipeline (https://github.com/vanheeringen-lab/seq2science) (STAR as default aligner). Tag count normalization, dimension reduction, and differential expression analysis (twofold change, p value<0.05) were performed using DESeq2 in R.<sup>59</sup> To investigate the subset purity, the top 25 highest expressed genes of the transformed normalized counts were selected per subset and compared with the other subsets and genes known from literature. Subsequently, enriched gene sets were determined by Generally Applicable Gene-set Enrichment.<sup>60</sup> L2F change and go.bp.gs set were used to determine the enriched biological processes (GO-terms). Immunologically relevant GO pathways that were upregulated with p<0.05 were selected and displayed as a radar plot showing the gene ratio (observed gene count/ total gene count per pathway). Murine RNA sequencing data for MHCII<sup>hi</sup> CD11b<sup>int</sup> and MHCII<sup>lo</sup> CD11b<sup>hi</sup> were from our previous work.<sup>21</sup>

### Lipid body staining

Isolated DC subsets were incubated with Matrix C ISCOMs (400 ng/mL) or OA  $(50 \mu \text{M})$  for 5, 10, or 18 hours. Cells (100,000) were then washed and seeded on chamber slides (Nunc Lab-Tek II) precoated with Poly-L-Lysine (100µg/mL, Sigma-Aldrich). After 1 hour attachment, cells were fixated with 4% formaldehyde (Merck) and stained with Bodipy 493/503 (7 µg/ml in phophatebuffered saline (PBS), Invitrogen) for 10 min at room temperature. For the co-staining studies, cells were first permeabilized with saponin (0.1% in PBS, Sigma) for 5 min, followed by 5% human blocking serum (30 min, 37°C). Cells were then incubated with anti-ADRP Alexa-647 (in 0.1% saponin in PBA, 30 min at 37°C, Abcam), washed, and incubated with Bodipy 493/503 as described earlier. DAPI (Santa Cruz Biotechnology) was used for nucleus staining and the chamber slides were mounted with Prolong Diamond Antifade Mountant (Invitrogen) and analyzed with laser scanning microscopy (Leica SP8 SMD, x63 water objective). At least five pictures were taken per sample for quantification. The LB frequency and number of LBs per cell were quantified with Fiji and a script developed by Paul Rijken (Department of Radiation Oncology, Radboud University Medical Center, Nijmegen, The Netherlands). Co-localization histograms for each fluorophore were created for a selected area (indicated by a line on the image) in Fiji.

### **Antigen translocation**

Isolated DC subsets (50,000 cells) were seeded in a flatbottom 96-wells plate (Costar) and incubated with Matrix C ISCOMs (400 ng/mL) with or without cytochrome c (2.5 mg/mL unless indicated otherwise, Sigma-Aldrich) for 18 hours. As a control, 10% DMSO (Merck) was added to the cells to induce cell death. Cell metabolic activity and viability were measured by Cell Counting Kit-8 (CCK8, Sigma-Aldrich) following the manufacturer's protocol. The absorbance was measured by a spectrophotometer at 450 nm. Relative metabolic activity was calculated as (treatment-blank)/(control-blank)×100%.

### **DC cross-presentation**

For MiHA-specific T-cell expansion assays, cryopreserved PBMCs containing HA-1-specific CD8<sup>+</sup> memory T cells from allogeneic stem cell transplantation patients were used.<sup>61</sup> All patient material was obtained in accordance with the Declaration of Helsinki and institutional guidelines and regulations (CMO 2012/064). The different DC populations were isolated, as described above, from HLA-A\*02:01<sup>+</sup> buffy coats from healthy donors (Sanquin, Nijmegen, The Netherlands) and typed HA-1<sup>-</sup> with PCR to prevent endogenous expression of HA-1 by the DCs. The different DC subsets were incubated with Matrix C ISCOMs (400 ng/mL) in combination with or without HA-1 long peptide (5µM, VARFAEGLEKLKECVLHDDL LEARRPRAHEZL) in Iscove's modified Dulbecco's medium (IMDM) (Gibco) containing 2% human serum, 40 U/mL hGM-CSF (Immunotools), and 1% penicillinstreptomycin (PS) (Gibco) for 3 hours. After 3 hours of pulse, additional human serum was added until 10% end concentration and cells were further incubated overnight. Positive control samples were incubated with HA-1 short peptide (5µM, VLHDDLLEA) for 2 hours in the presence of 2% human serum. After washing the DCs, PBMCs containing HA-1 specific CD8<sup>+</sup> T cells were added and incubated for 7 days in IMDM containing 10% human serum, 40U/mL hGM-CSF, and 1% PS. On day 4, medium was refreshed with IMDM containing 10% human serum, 1% PS, 50U/mL human interleukin (hIL)-2 (Immunotools), and 40U/mL hIL-15 (Immunotools). Cells were harvested and counted after 7 days and HA-1-specific CD8<sup>+</sup> T-cell expansion was measured by flow cytometry (Cytoflex LX 21-color, Beckman Coulter). Cells were stained with anti-CD3 (BioLegend), anti-CD8 (Invitrogen), HA-1 tetramer PE, and HA-1 tetramer APC. Fold expansions of tetramer<sup>+</sup> HA-1 specific T cells were quantified by dividing the absolute numbers of double tetramer<sup>+</sup> T cells after treatment by the absolute numbers of double tetramer<sup>+</sup> T cells in the no peptide sample. The HA-1 tetramers, were kindly gifted from Mirjam Heemskerk and Frederik Falkenburg (Department of Hematology, Leiden University Medical Center, Leiden, The Netherlands).

### **PBMC viability assay**

The isolated DC subsets (50,000 cells) were incubated with freshly isolated PBMCs (500,000 cells) from healthy donors in IMDM, 10% human serum, 40 U/mL hGM-CSF, and 1% PS. On day 4, medium was refreshed with IMDM with 10% human serum, 1% PS, 50 U/mL hIL-2, and 40 U/mL hIL-15. Cells were harvested and counted after 7 days, and cell viability was measured by flow cytometry (BD FACSCanto II). Cells were stained with anti-CD8 (BioLegend), anti-CD3 (BD Biosciences),

anti-CD4 (BioLegend), and Fixable Viability Dye eFluor 780 (eBioscience).

### **Statistical analysis**

One-way analysis of variance with post hoc Tukey, or paired non-parametric t-test was performed as indicated in the figure legends. All results are expressed as mean values with SD. The following indications are used in all figures: n.s., non significant; \*p<0.05, \*\*p<0.01, \*\*\*p<0.001, and \*\*\*\*p<0.0001.

#### Author affiliations

<sup>1</sup>Radiotherapy and Oncolmmunology Laboratory, Department of Radiation Oncology, Radboud University Nijmegen Radboud Institute for Molecular Life Sciences, Nijmegen, The Netherlands

<sup>2</sup>Department of Laboratory Medicine, Laboratory of Hematology, Radboud University Nijmegen Radboud Institute for Molecular Life Sciences, Nijmegen, The Netherlands <sup>3</sup>Department of Molecular Biology, Faculty of Science, Radboud University Nijmegen Radboud Institute for Molecular Life Sciences, Nijmegen, The Netherlands <sup>4</sup>Radboud Technology Center Microscopy, Radboud University Nijmegen Radboud Institute for Molecular Life Sciences, Nijmegen, The Netherlands

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### ORCID iDs

Nataschja I Ho http://orcid.org/0000-0003-4071-9370 Lisa G M Huis in 't Veld http://orcid.org/0000-0002-8538-8408 Jesper van Eck van der Sluijs http://orcid.org/0000-0003-3136-7799 Branco M H Heuts http://orcid.org/0000-0002-7258-8312 Gosse J Adema http://orcid.org/0000-0002-6750-1665

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