Hypersensitivity pneumonitis (HP) is an immunological disease characterized by a prominent interstitial infiltrate composed predominantly of lymphocytes secreting inflammatory cytokines. Dendritic cells (DCs) are known to play a pivotal role in the lymphocytic response. However, their cross talk with microorganisms that cause HP has yet to be elucidated.

This study aimed to investigate the initial interactions between human monocyte-derived DCs (MoDCs) and four microorganisms that are different in nature (Saccharopolyspora rectivirgula [actinomycetes], Mycobacterium immunogenum [mycobacteria], and Wallisma sebi and Eurotium amstelodami [filamentous fungi]) and are involved in HP. Our objectives were to determine the cross talk between MoDCs and HP-causing agents and to determine whether the resulting immune response varied according to the microbial extract tested. The phenotypic activation of MoDCs was measured by the increased expression of costimulatory molecules and levels of cytokines in supernatants. The functional activation of MoDCs was measured by the ability of MoDCs to induce lymphocytic proliferation and differentiation in a mixed lymphocytic reaction (MLR). E. amstelodami-exposed (EA) MoDCs expressed higher percentages of costimulatory molecules than did W. sebi-exposed (WS), S. rectivirgula-exposed (SR), or M. immunogenum-exposed (MI) MoDCs ($P < 0.05$, Wilcoxon signed-rank test). EA-MoDCs, WS-MoDCs, SR-MoDCs, and MI-MoDCs induced $CD4^{+}$ T cell proliferation and a Th1-polarized immune response. The present study provides evidence that, although differences were initially observed between MoDCs exposed to filamentous fungi and MoDCs exposed to bacteria, a Th1 response was ultimately promoted by DCs regardless of the microbial extract tested.

Dendritic cells (DCs) are professional antigen-presenting cells that play a crucial role in initiation of the adaptive immune system. They act as sentinels that quickly respond to foreign antigens; when activated, DCs transfer important information about the invading pathogens and the innate response in the periphery to T cells, thus inducing an appropriate adaptive immune response (14). Mature DCs also trigger NK cell effector functions, which promote Th1 polarization (15, 16). On the basis of their ability to take up antigens, present the antigens to naive T cells, and activate those T cells, DCs constitute useful tools for assessing lymphocytic induction and polarization after exposure to microorganisms involved in HP.

Our study aimed to investigate the initial interactions between human monocyte-derived DCs (MoDCs) and four microorganisms that are involved in HP and are different in nature (actino-
In order to determine the cross talk between MoDCs and HP causative agents and to determine whether the resulting immune responses differ according to the microbial extract tested, human MoDCs were exposed to microbial extracts of three main microorganisms involved in FLD, i.e., *S. rectivirgula* (SR), *M. immunogenum* (MI), *W. sebi* (WS), or *E. amstelodami* (EA). The phenotypic activation of MoDCs was evaluated by measuring costimulatory molecule levels and cytokine/chemokine levels in supernatants. The functional activation of MoDCs was measured by assessment of their ability to induce lymphocytic proliferation and determination of the lymphocytic polarization.

### FIG 1
Expression of the costimulatory molecules CD80, CD86, and CD40 by immature MoDCs (iMoDCs) and MoDCs exposed to LPS (positive control), *S. rectivirgula* (SR), *M. immunogenum* (MI), *W. sebi* (WS), or *E. amstelodami* (EA). White peaks, isotypes; gray peaks, corresponding antibodies. The mean fluorescence intensity (MFI) is indicated for each condition.

### TABLE 1
Percentages of costimulatory molecules CD80, CD86, and CD40 measured on human MoDCs before (iMoDCs) and after (positive control) exposure to LPS, *E. amstelodami*, *W. sebi*, *S. rectivirgula*, or *M. immunogenum*.

| Cells tested   | % of expression (mean ± SD) |
|---------------|-----------------------------|
|               | CD80 | CD86 | CD40 |
| iMoDCs        | 12 ± 10 | 10 ± 5 | 9 ± 5 |
| LPS-exposed MoDCs | 94 ± 10 | 97 ± 13 | 98 ± 12 |
| EA-MoDCs      | 60 ± 18 | 66 ± 16 | 78 ± 16 |
| WS-MoDCs      | 36 ± 13 | 32 ± 15 | 41 ± 12 |
| SR-MoDCs      | 51 ± 19 | 45 ± 18 | 41 ± 15 |
| MI-MoDCs      | 26 ± 16 | 25 ± 13 | 31 ± 12 |

mycetes, mycobacteria, and filamentous fungi), in order (i) to determine the cross talk between MoDCs and HP causative agents and (ii) to determine whether the resulting immune responses differ according to the microbial extract tested. Human MoDCs were exposed to microbial extracts of three main microorganisms involved in FLD, i.e., *S. rectivirgula*, *E. amstelodami*, and *W. sebi*, and to a microbial extract of *M. immunogenum*, the main causative agent of MWF-HP. The phenotypic activation of MoDCs was evaluated by measuring costimulatory molecule levels and cytokine/chemokine levels in supernatants. The functional activation of MoDCs was measured by assessment of their ability to induce lymphocytic proliferation and determination of the lymphocytic polarization.
TABLE 2 Statistical analysis of differences in the expression of costimulatory molecules among MoDCs pulsed with E. amstelodami, W. sebi, S. rectivirgula, or M. immunogenenum

| Comparison          | P for expression of: |
|---------------------|----------------------|
|                     | CD80 | CD86 | CD40 |
| EA-MoDCs vs WS-MoDCs| 0.002 | 0.002 | 0.008 |
| EA-MoDCs vs SR-MoDCs| 0.002 | 0.002 | 0.008 |
| EA-MoDCs vs MI-MoDCs| 0.001 | 0.001 | 0.001 |
| WS-MoDCs vs SR-MoDCs| NS*  | NS   | NS   |
| WS-MoDCs vs MI-MoDCs| NS   | NS   | NS   |
| SR-MoDCs vs MI-MoDCs| 0.03  | 0.04 | NS   |

*NS, not significant.

MATERIALS AND METHODS

Strains. We used Lacey’s strain of S. rectivirgula (DSMZ 43113), two fungal strains isolated from FLD patient hay from Franche-Comté (a region in eastern France), namely, E. amstelodami (BCCM/IHEM 16286) and W. sebi (BCCM/IHEM 16284), and a strain of M. immunogenenum (DSMZ 45496) isolated in MWF taken from a factory where cases of MWF-HP were diagnosed (17). All of the strains were cultured for 1 week under the following conditions: at 44°C on R8 medium for S. rectivirgula, at 20°C on Sabouraud agar (Becton, Dickinson and Company, Le Pont de Claix, France) for W. sebi, at 30°C on DG18 agar (Oxoid Unipath, Basingstoke, England) for E. amstelodami, and at 30°C on Muller-Hinton culture medium for M. immunogenenum. Temperatures and media were chosen to reflect the optimal growth conditions for each species.

Preparation of total extracts. For each species, 4 culture plates were gently brushed with a swab and the fungal structures were harvested in 2 ml of sterile water, resulting in milk-like suspensions (McFarland Scale values of >7). Each suspension was frozen at −20°C overnight. The following day, lyophilization was carried out in a Labconco lyophilizer (Labconco, Kansas City, MO). The microbial extracts were weighed and then suspended in sterile water at 1 mg/ml. The presence of endotoxins was measured in the total extracts using the Chromo-LAL method (Biogenic, Perols, France).

Generation of MoDCs. Peripheral blood mononuclear cells (PBMCs) were isolated by density gradient centrifugation, using Ficoll-Paque Premium (Dutscher, Brumath, France), from healthy donor blood obtained from the French Blood Institute of the Bourgogne and Franche-Comté regions. A total of 1.1 × 10^9 PBMCs were incubated in a 75-cm² culture flask (Fischer Scientific, Illkirch, France) containing culture medium, i.e., RPMI 1640 (Dutscher) supplemented with 10% fetal calf serum (Dutscher) and 5% penicillin-streptomycin (Dutscher). Two hours later, nonadherent cells (peripheral blood lymphocytes [PBLs]) were removed; adherent cells were gently washed three times with phosphate-buffered saline (Dutscher). In order to generate MoDCs, adherent cells were cultured for 5 days in culture medium supplemented with interleukin 4 (IL-4) (1,000 IU/ml; R&D Systems, Lille, France) and granulocyte-macrophage colony-stimulating factor (GM-CSF) (1,000 IU/ml; R&D Systems). Incubations were conducted in a humidified 5% CO₂ atmosphere at 37°C for all experiments. No cytokines and no fresh medium were added during the 5 days of differentiation.

Exposure of MoDCs to microbial extracts. The cocultivation of MoDCs (5 × 10^6 cells) and each of the HP causative agents tested (S. rectivirgula, E. amstelodami, W. sebi, and M. immunogenenum), at 10 μg/ml, was carried out for 24 h. Lipopolysaccharide (LPS) (Sigma-Aldrich) was used as a positive control (also at 10 μg/ml). The exposure concentration of 10 μg/ml was selected after concentration assays using 0.1, 1, 10, and 100 μg/ml for each of the microbial extracts. The quantity of IL-8 secreted in the supernatants by MoDCs after 24 h of exposure and their expression of the costimulatory molecule CD40 were used to select the concentration of 10 μg/ml for further assays (see Fig. S1 in the supplemental material).

Phenotypic activation of MoDCs. (i) Costimulatory molecule expression. Immature MoDCs (iMoDCs) (unexposed) and MoDCs 24 h after exposure were stained with labeled antibodies (BD Biosciences Pharmingen, Pont de Clairaix, France) against CD11c (phycoerythrin [PE]-labeled), CD80 (fluorescein isothiocyanate [FITC]-labeled), CD86 (PE-
and Cy7-labeled), and CD40 (allophycocyanin [APC]-labeled). MoDCs were analyzed by flow cytometry (BD Canto II) using FACSDiva software (BD Biosciences).

(ii) Cytokine/chemokine production. The secretion of IL-10, IL-12, IL-23, and IL-8 was analyzed by enzyme-linked immunosorbent assays (ELISAs) (DiaciOne, Besançon, France) according to the manufacturer’s instructions (detection thresholds of the ELISAs were 29 pg/ml for IL-8, 20 pg/ml for IL-23, 5 pg/ml for IL-10, and 2.2 pg/ml for IL-12).

Functional activation of MoDCs. (i) Mixed lymphocytic reaction with total allogeneic PBLs. The ability of exposed MoDCs to induce the proliferation of both CD4+ and CD8+ T lymphocytes was investigated using total allogeneic PBLs. MoDCs (5 × 10^5) that had been previously exposed for 24 h to each of the microbial extracts tested were incubated for 5 days with 1 × 10^6 PBLs in round-bottomed wells (Sarstedt, Marnay, France). T cell proliferation was assessed by flow cytometry with cytarbino-fluorescein succinimidyl ester (CFSE) labeling, as recommended by the manufacturer (eBioscience, Paris, France).

(ii) Mixed lymphocytic reaction with purified naive CD4+ T lymphocytes. (a) Cells. The lymphocytic CD4+ polarization was investigated using naive CD4+ T lymphocytes (CD3+CD4+CD45RA-CD25-), that had been previously purified from PBMCs by immunomagnetic depletion using the Naive CD4+ T Cell Isolation Kit II (Milteny Biotec, Paris, France), according to the manufacturer’s instructions. MoDCs (5 × 10^5) that had been previously exposed to each of the microbial extracts tested were incubated for 5 days with 1 × 10^5 CD4+ T lymphocytes in round-bottomed wells (Sarstedt). A MoDC/T cell ratio of 1:2 was selected due to the results of mixed lymphocytic reaction (MLR) testing with different MoDC/T cell ratios (1:2, 1:5, 1:10, and 1:100), using MoDCs that had been previously exposed to LPS for 24 h (see Fig. S2 in the supplemental material).

(b) Transcription factors. After 3 days of cocultivation, the RNA of naive CD4+ T lymphocytes was extracted using an RNAeasy minikit and an RNase-free DNase set (Qiagen, Courtaboeuf, France), according to the manufacturer’s instructions. Reverse transcription was carried out in a LightCycler LC 480 system (Roche Diagnostics, Meylan, France) as follows: 20 µl RNA was added to 8 µl of RNA-to-cDNA Master Mix (Applied Biosystems, Courtaboeuf, France). Real-time PCR was carried out in a LightCycler LC 480 system (Roche Diagnostics, Meylan, France), in a 20-µl final volume containing 10 µl TaqMan Universal PCR Master Mix (Applied Biosystems), 1 µl TaqMan Gene Expression Assays-on-Demand primer solution (Applied Biosystems), and 2 µl cDNA. The expression of four transcription factors (target genes), each representative of a polarization pathway, i.e., T-bet (Th1), Gata-3 (Th2), FoxP3 (Treg), and RORc (Th17), was measured. Values for each sample were normalized on the basis of the contents in comparison with a reference gene, GAPDH. The results were expressed as the N-fold difference in target gene expression, relative to GAPDH gene expression (termed N_{target}), and were obtained using the following formula: N_{target} = 2^{ΔCt_{sample}}

(c) Intracellular markers. After 5 days of cocultivation, CD4+ T lymphocytes were stimulated for 4 h with phorbol myristate acetate (PMA) (50 ng/ml; Sigma) and ionomycin (500 ng/ml; Sigma). T cells were fixed and permeabilized with a BD Cytofix/Cytoperm Plus kit with GolgiPlug (BD Biosciences) and then were incubated with labeled antibodies (BD Biosciences) against IL-17 (PE-labeled), IL-4 (FITC-labeled), tumor necrosis factor alpha (TNF-α) (PE- and Cy7-labeled), and gamma interferon (IFN-γ) (APC-labeled). Cells were washed, and cytokine profiles were analyzed.

(d) Cytokine/chemokine levels. After 5 days of cocultivation, secreted chemokines and cytokines were quantified using Luminex multiplex bead technology. Human cytokine/chemokine panel 1 (MPXHYCITO-60K, Milliplex Map; Millipore, Billerica, MA) was used to quantify IL-6, IL-8, IFN-γ, IL-17, IL-4, IL-10, IL-12, and TNF-α ( assay sensitivity of 0.4 pg/ml for all). Each sample was measured in duplicate, and the assay was performed following the manufacturer’s instructions.

Th1, Th2, Treg, and Th17 polarization controls. Naive CD4+ T lymphocytes were incubated in round-bottomed wells with Dynabeads Human T-Activator CD3/CD28 (Life Technologies, Saint Aubin, France) plus IL-12 (50 ng/ml) and anti-IL-4 (10 µg/ml) for Th1 polarization; IL-4 (50 ng/ml) and anti-IFN-γ (10 µg/ml) for Th2 polarization; transforming growth factor β1 (TGFB1) (50 ng/ml), anti-IFN-γ (10 µg/ml), and anti-IL-12 (50 ng/ml) for Treg polarization; or TGFB1 (50 ng/ml), IL-6 (50 ng/ml), IL-23 (25 ng/ml), IL-1β (25 ng/ml), anti-IFN-γ (10 µg/ml), and anti-IL-4 (10 µg/ml) for Th17 polarization.

Transcription factor expression, intracellular marker levels, and cytokine secretion were measured as described above. In the case of the Treg polarization control, Human Regulatory T Cell Staining Kit 2 (eBioscience, San Diego, CA) was used to measure the intracellular markers FoxP3 and CD25.

Statistical analysis. Data are presented as means ± standard deviations (SD) from at least six separate experiments. For comparisons of continuous paired data (related samples), the Friedman test and the Wilcoxon signed-rank test were performed using the software package Stata v10 (StataCorp LP, College Station, TX). All tests were two-tailed, and P values of <0.05 were considered statistically significant.

RESULTS
Phenotypic activation of MoDCs. (i) Costimulatory molecule expression. We observed slight T lymphocyte contamination in each preparation (3.26% ± 1.4% T lymphocytes). MoDCs were selected on the basis of their CD11c expression. Exposure to microbial extracts may lead to changes in the expression of costimulatory molecules present on the surface of MoDCs. The percentages of CD80, CD86, and CD40 obtained for immature MoDCs (iMoDCs) and MoDCs exposed to either LPS, E. amastelodami (EA-MoDCs), W. sebi (WS-MoDCs), S. rectivirga (SR-MoDCs), or M. immunogenenum (MI-MoDCs) were measured by flow cytometry, and an example of the results observed is presented in Fig. 1. The percentages of activation of the costimulatory molecules are detailed in Table 1 and the corresponding statistical analysis in Table 2. The analysis revealed that EA-MoDCs expressed higher percentages of costimulatory molecules (CD80, CD86, and CD40) than did WS-MoDCs, SR-MoDCs, and MI-MoDCs (P < 0.01, Wilcoxon signed-rank test).

(ii) Cytokine/chemokine production. Exposure to microbial extracts involved in HP may induce the secretion of chemokines and cytokines such as IL-8, IL-23, IL-12, and IL-10. A basal level of

| Condition tested | IL-12 level (pg/ml [mean ± SD]) |
|------------------|----------------------------------|
| Phenotypic assays |
| iMoDCs           | 0 ± 0                            |
| LPS-exposed MoDCs| 246 ± 80                         |
| EA-MoDCs         | 21 ± 4.5                         |
| WS-MoDCs         | 7.2 ± 2.9                        |
| SR-MoDCs         | 6.7 ± 2.2                        |
| MI-MoDCs         | 0 ± 0                            |
| Functional assays |
| Allogeneic T lymphocytes + iMoDCs | 1.4 ± 0.1 |
| Naive CD4+ T lymphocytes + iMoDCs | 1.5 ± 0.2 |
| Naive CD4+ T lymphocytes + LPS-exposed MoDCs | 10.8 ± 4.1 |
| Naive CD4+ T lymphocytes + EA-MoDCs | 11.5 ± 4.7 |
| Naive CD4+ T lymphocytes + WS-MoDCs | 11.9 ± 3.7 |
| Naive CD4+ T lymphocytes + SR-MoDCs | 9.5 ± 3.6 |
| Naive CD4+ T lymphocytes + MI-MoDCs | 9.8 ± 2.5 |

Table 3: Levels of IL-12 (pg/ml) in supernatants in assays investigating phenotypic and functional activation of MoDCs.
IL-8 (122 ± 41 pg/ml) was detected in the supernatants of iMoDCs, but IL-23, IL-12, and IL-10 were undetectable. LPS-exposed MoDCs produced 10,060 ± 651 pg/ml IL-8, 1,170 ± 350 pg/ml IL-23, 246 ± 80 pg/ml IL-12, and 491 ± 80 pg/ml IL-10. EA-MoDCs, WS-MoDCs, SR-MoDCs, and MI-MoDCs all produced high levels of IL-8 (means, 2,076 pg/ml to 2,931 pg/ml). MoDCs exposed to filamentous fungi (EA-MoDCs and WS-MoDCs) produced higher levels of IL-10 and IL-23 than did MoDCs exposed to bacteria (SR-MoDCs and MI-MoDCs) (P < 0.05, Wilcoxon signed-rank test). These results are presented in Fig. 2. The levels of IL-12 detected were low for EA-MoDCs, WS-MoDCs, and SR-MoDCs, and undetectable for MI-MoDCs (Table 3).

Functional activation of MoDCs. (i) MLR with total allogeneic PBLs. The capacity of exposed MoDCs to stimulate T cell proliferation was initially investigated using total allogeneic PBLs, in order to determine whether the lymphocytic response involved mainly CD4+ or CD8+ lymphocytes. The results are presented in Fig. 3. Cocultivation of total allogeneic PBLs with LPS-exposed MoDCs, EA-MoDCs, WS-MoDCs, SR-MoDCs, or MI-MoDCs for 5 days, in three separate experiments, induced greater proliferation of CD4+ cells (means, 34% ± 13% to 58% ± 19%) than CD8+ cells (means, 11% ± 6% to 21% ± 9%). Moreover, greater proliferation of PBLs was observed in the case of stimulation with bacteria (S. rectivirgula and M. immunogenenum) (P = 0.028, Wilcoxon signed-rank test). FSC, forward scatter; SSC, side scatter.

Functional activation of MoDCs. (ii) MLR with purified naive CD4+ T lymphocytes. (a) Transcription factors. In order to determine the polarization of the proliferating CD4+ lymphocytes, the MLR was repeated with only CD4+ T cells and the expression of four transcription factors (target genes), each representative of a polarization pathway, i.e., T-
The upregulation of the MoDCs for 3 days, in three separate experiments, induced marked exposure of MoDCs, EA-MoDCs, WS-MoDCs, SR-MoDCs, or MI-MoDCs were significantly higher (P < 0.05, Wilcoxon signed-rank test) than those produced by T lymphocytes alone and also showed that all microbial extracts promoted Th1 polarization, as measured by T-bet.

(b) Intracellular markers. In order to determine the polarization of the proliferating CD4 T lymphocytes, the MLR was repeated with only CD4 T cells and the presence of intracellular markers such as IL-17a, IL-4, TNF-α, and IFN-γ in CD4 T cells was investigated by flow cytometry. After 5 days of cocultivation of naive CD4 T lymphocytes with LPS-exposed MoDCs, EA-MoDCs, WS-MoDCs, SR-MoDCs, or MI-MoDCs, the percent-

### TABLE 4 Intracellular staining for IL-17a, IL-4, IFN-γ, and TNF-α for allogeneic T lymphocytes and naive CD4 T lymphocytes cocultured with MoDCs previously exposed to LPS, E. amstelodami, W. sebi, S. rectivirgula, or M. immunogenen

| Cells tested                                      | % (mean ± SD) | CD4 CD25 FoxP3 cells* (% [mean ± SD] of parents) |
|--------------------------------------------------|---------------|-----------------------------------------------|
| T lymphocytes cocultured with MoDCs exposed to  |               |                                               |
| microbial extracts                               |               |                                               |
| Naive CD4 T lymphocytes                          | 0.1 ± 0.08    | 12 ± 0.7                                      |
| Naive CD4 T lymphocytes + LPS-exposed MoDCs      | 0.17 ± 0.1    | 13.5 ± 0.5                                    |
| Naive CD4 T lymphocytes + EA-MoDCs               | 0.1 ± 0       | 6.8 ± 1.9                                     |
| Naive CD4 T lymphocytes + WS-MoDCs               | 0.13 ± 0.06   | 7.9 ± 4.9                                     |
| Naive CD4 T lymphocytes + SR-MoDCs               | 0.1 ± 0       | 6.5 ± 1.1                                     |
| Naive CD4 T lymphocytes + MI-MoDCs               | 0.1 ± 0       | 8.9 ± 0.4                                     |
| T lymphocytes with LPS-exposed MoDCs             | 0.3 ± 0.06    | 25 ± 7.3                                      |
| T lymphocytes with EA-MoDCs                      | 0.3 ± 0       | 0.3 ± 0.1                                     |
| T lymphocytes with WS-MoDCs                      | 0 ± 0         | 0.4 ± 0.08                                    |
| T lymphocytes with SR-MoDCs                      | 0 ± 0         | 0 ± 0                                          |
| T lymphocytes with MI-MoDCs                      | 0 ± 0         | 0 ± 0                                          |
| T lymphocytes with MI-MoDCs                      | 0 ± 0         | 55 ± 9.8                                       |

* Specific CD25/FoxP3 staining was added for the Treg polarization control. NR, no response.

FIG 4 Standardized mRNA expression of the FoxP3, T-bet, Gata-3, and RORc genes after exposure of MoDCs to LPS, E. amstelodami, W. sebi, S. rectivirgula, or M. immunogenen. Calculated with the ΔΔCt method, with GAPDH as the reference gene. The data are presented as means ± SEMs from three separate experiments. In three separate experiments, levels of T-bet produced by T lymphocytes (L1) after cocultivation with LPS-exposed MoDCs, EA-MoDCs, WS-MoDCs, SR-MoDCs, or MI-MoDCs were significantly higher (P < 0.05, Wilcoxon signed-rank test) than those produced by T lymphocytes alone and also showed that all microbial extracts promoted Th1 polarization, as measured by T-bet.
Cytokine/chemokine levels were measured in supernatants using the Luminex technique after 6 days of incubation.
cytokines according to the microbial extracts tested (E. amstelodami, W. sebi, S. rectivirgula, or M. immunogenum) showed no significant differences in IL-8, IL-6, and IFN-γ production, while lower production of TNF-α was observed in the case of coculture of CD4^+ T lymphocytes with SR-MoDCs (P < 0.05, Wilcoxon signed-rank test) and higher production of IL-10 and IL-17a was observed in the case of coculture of CD4^+ T lymphocytes with EA-MoDCs (P < 0.05, Wilcoxon signed-rank test).

All of the results are detailed in Tables 3 and 5, including those obtained for the negative control (CD4^+ T cells alone) and polarization controls. These results support the fact that all microbial extracts promoted Th1 polarization, as measured by IFN-γ, TNF-α, IL-8, and IL-6 levels.

### Endotoxin levels

The presence of endotoxins was measured in the total extracts using the Chromo-LAL method (Biogenic, Perols, France). Microbial extracts were used at 10 μg/ml for MoDC exposure. Endotoxin levels for the W. sebi and M. immunogenum extracts were 0.007 endotoxin units (EU)/ml and 0.0006 EU/ml, respectively. A first set of experiments was performed using extracts of S. rectivirgula and E. amstelodami with endotoxin levels higher than 0.01 EU/ml (18). A second set of experiments was performed using S. rectivirgula and E. amstelodami extracts with levels under the 0.01 EU/ml limit (0.008 EU/ml and 0.0005 EU/ml, respectively). Significant differences were observed between the two sets of experiments in two cases, i.e., (i) higher production of IL-8 in cocultures with the S. rectivirgula extract with <0.01 EU/ml and (ii) higher production of IFN-γ in cocultures with the E. amstelodami extract with >0.01 EU/ml (Wilcoxon signed-rank test) (Table 6). The results were considered similar for phenotypic and functional assays using either set of S. rectivirgula and E. amstelodami total extracts, and we concluded that the presence of endotoxin did not affect the Th1 polarization (Table 6).

### DISCUSSION

The present study, which aimed to determine the cross talk between MoDCs and microbial extracts from microorganisms involved in HP, provides evidence that, although differences initially were observed between MoDCs exposed to filamentous fungi and MoDCs exposed to bacteria (high levels of IL-23 and IL-10 for MoDCs stimulated by E. amstelodami or W. sebi), a Th1 response ultimately was promoted by MoDCs regardless of the microbial extract tested. The results of our study are consistent with the hypothesis that upregulation of Th1 signaling plays a critical role in the pathogenesis of HP.

Th17 cells are a recently described subset of Th cells, and they play a pivotal role in immunity against fungi (19, 20). IL-23, a member of the IL-12 cytokine family produced by MoDCs, is involved in the Th17 lineage and plays a critical role in the development of inflammation. A previous study demonstrated that the interaction between MoDCs and the polysaccharide β-glucan, which is present in the cell walls of fungi, triggered the syk-CARD9 signaling pathway and thus the production of IL-23 (21). E. amstelodami and W. sebi are both filamentous fungi with β-glucan in their cell walls, and high levels of IL-23 were detected in supernatants of EA-MoDCs and WS-MoDCs. This result suggested that differentiation of naïve CD4^+ T cells into Th17 cells could occur. In MLR assays, however, a Th1-polarized immune response was observed for the four microbial extracts tested, including E. amstelodami and W. sebi. This observation is consistent with the recent demonstration that IL-23, while able to maintain a Th17 phenotype, does not mediate commitment to this phenotype (22).

Previous studies, focusing mainly on Aspergillus fumigatus, a filamentous fungi involved in invasive aspergillosis, demonstrated that surface components of fungi stimulate the Toll-like receptors (TLRs) TLR2 and TLR4 (23, 24). Ligands that are present on nearly all fungi and might be responsible for stimulating the TLRs include β-glucans, manannans, and chitin (25). Several studies involving either the genus Mycobacterium (mostly Mycobacterium tuberculosis) or Gram-positive bacteria (but not specifically actinomycetes) also demonstrated interactions between bacterial components and TLRs (also mostly TLR2 and TLR4) (26, 27).

Thus, it seems most probable that the mechanism involves interactions with TLR2 and TLR4.

Previous studies have established that HP is caused by a Th1-type immune response; bronchoalveolar lavage fluid (BALF)

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**TABLE 6 Comparison of results obtained in phenotypic and functional assays using different batches of S. rectivirgula and E. amstelodami total extracts, either below or above the 0.01 EU/ml limit**

| Polarization result | Extracts with >0.01 EU/mla | Extracts with <0.01 EU/mla | Differenceb |
|---------------------|----------------------------|----------------------------|-------------|
| Phenotypic assays   | SR-MoDCs | EA-MoDCs | SR-MoDCs | EA-MoDCs | SR-MoDCs | EA-MoDCs |
| % with CD80         | 51 ± 19   | 60 ± 18   | 61.2 ± 15.2  | 58.3 ± 15.2  | NS        | NS        |
| % with CD86         | 45 ± 18   | 66 ± 16   | 38.8 ± 19   | 58.4 ± 13.1  | NS        | NS        |
| IL-23 level (pg/ml) | 72 ± 38   | 2,416 ± 875 | 122 ± 46   | 2,590 ± 422  | NS        | NS        |
| IL-8 level (pg/ml)  | 1,987 ± 358 | 2,708 ± 296 | 3,876 ± 823 | 2,189 ± 752  | 0.03      | NS        |
| IL-10 level (pg/ml) | 79 ± 52   | 475 ± 89  | 86 ± 48     | 546 ± 66     | NS        | NS        |
| Functional assays   | % with CD4 (CFSE) | 48 ± 19.6  | 36.4 ± 11.2 | 73 ± 27 | 30.2 ± 7.9 | NS        | NS        |
| T-bet mRNA expression (fold) | 4.16 ± 1.8 | 2.9 ± 1.2 | 4.78 ± 1.3 | 3.3 ± 1.6 | NS        | NS        |
| % with intracellular TNF-α | 7.6 ± 0.2 | 7 ± 0.5 | 8.2 ± 2.4 | 6.1 ± 0.9 | NS        | NS        |
| % with intracellular IFN-γ | 6.1 ± 0.2 | 8 ± 2.3 | 6.8 ± 1.8 | 5.6 ± 1.6 | NS        | NS        |
| IL-8 level (pg/ml) | 3,522 ± 243 | 4,356 ± 487 | 4,200 ± 386 | 3,987 ± 612  | NS        | NS        |
| IL-6 level (pg/ml) | 1,423 ± 253 | 2,289 ± 423 | 1,923 ± 358 | 1,678 ± 269  | NS        | NS        |
| IFN-γ level (pg/ml) | 53,620 ± 1.245 | 75,283 ± 2,641 | 48,888 ± 3,200 | 60,069 ± 2,823 | NS | 0.044 |

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a Values are means ± SD.

b Differences were tested using the Wilcoxon signed-rank test. NS, not significant.
specimens obtained from individuals with HP displayed abundant IFN-γ, TNF-α, IL-8, and IL-12 (28–30). The low levels of IL-12 detected in our assays, in both phentypic and functional assays, are probably due to the use of MoDCs instead of macrophages. In patients’ BALF, macrophages are well represented and may be responsible for IL-12 production. Data showed that MoDCs were able to induce marked Th1 responses, with strong IFN-γ production, but failed to produce high levels of IL-12 (31).

The results of our study are consistent with the hypothesis that upregulation of Th1 signaling plays a critical role in the pathogenesis of HP. The fact that the nature of the microorganisms (filamentous fungi, actinomycetes, or mycobacteria) tested did not influence the outcome of the final immune response gives weight to the hypothesis that the pathophysiology of HP and the consequent clinical form are due more to the type of exposure (32) and to the hypothesis that the pathophysiology of HP and the consequent clinical form are due more to the type of exposure (32) and individual genetic susceptibility (33–35) than to the nature of the microbial environment of the patient.

In conclusion, this study provides evidence that, although initial differences are observed between MoDCs exposed to filamentous fungi and MoDCs exposed to bacteria, all of the microbial extracts ultimately lead to a Th1-type immune response. This study also clarifies the interactions between MoDCs and microbial extracts from microorganisms involved in HP, highlighting the fact that MoDCs are useful for acquiring a better understanding of antimicrobial immunity.

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