Wheat-derived arabinoyxylans reduced M2-macrophage functional activity, but enhanced monocyte-recruitment capacity†

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The immunomodulatory properties of non-digestible polysaccharides (NDPs) have been recognized in in vitro and in vivo studies. The latter mostly demonstrated altered frequencies and inflammatory status of immune cells as clinical parameters. Most of the NDP activity will be exerted in the intestine where they can directly interact with macrophages. The predominant macrophage phenotype in the intestine is M2-like, with M1-like macrophages arising during inflammation. Here, we investigated transcriptional and functional impact on these macrophage phenotypes by NDP-treatment (i.e. yeast-derived soluble β-glucan (yeast-βG), apple-derived RG-I (apple-RGI), shiitake-derived β-glucan (shiitake-βG) or wheat-derived arabinoyxylan (wheat-AX)). Wheat-AX, and to a lesser extent shiitake-βG and apple-RGI but not yeast-βG, reduced endocytosis and antigen processing capacity of M1- and M2-like macrophages. Moreover, the NDPs, and most notably wheat-AX, strongly induced transcription and secretion of a unique set of cytokines and chemokines. Conditioned medium from wheat-AX-treated M2-like macrophages subsequently demonstrated strongly increased monocyte recruitment capacity. These findings are in line with clinically observed immunomodulatory aspects of NDPs making it tempting to speculate that clinical activity of some NDPs is mediated through enhanced chemoattraction and modifying activity of intestinal immune cells.

Introduction

Non-digestible polysaccharides (NDPs) constitute a large family of different molecules, including β-glucans, pectins, resistant starch, cellulose and arabinoyxylans, and are known for their immune-modulatory properties and beneficial health effects.1,2 These properties were demonstrated in vitro as well as reported in clinical studies using different NDPs. For example, yeast β-glucan intake by obese people lowered pro-inflammatory cytokine levels (i.e. IL-6 and TNF-α) and increased anti-inflammatory IL-10 levels in peripheral blood, accompanied by lowering risk factors for wellbeing such as waist circumference and blood pressure.3 Also, rice bran arabinoyxylan increased NK cell activity and peripheral blood levels of myeloid-derived dendritic cells and T1h1 related cytokines in multiple myeloma patients, unfortunately without reported beneficial effects on disease progress.4 A randomized dietary intervention in healthy young adults demonstrated that consuming shiitake mushrooms containing lentinan reduced CRP levels and therefore lowered peripheral inflammation but also increased immune cell effector potency.5 This coincided with an altered cytokine balance of increased IL-4, IL-10, TNF-α and IL-1α levels and decreased MIP1α/CCL3 level. Other beneficial health effects related to NDP intakes studied in clinical trials include increased NK cell activity linked to a reduction in metastasis in lung and breast cancer patients;6 increased blood leukocytes, neutrophils, IgG and IgM frequencies accompanied by delayed progression of non-small cell lung cancer;7 and transiently increased blood levels of IgG and IgM and NK cell counts linked to a reduction in flu and flu-like symptoms and respiratory tract infections in children.8

The large variation in NDP functionality and structure resulting from source, extraction method, saccharide composition and/or branching hampers identification of the mecha-
nism of action. However, many studies have demonstrated that NDPs are recognized by immune cells through binding to pattern recognition receptors (PRRs). Dectin-1 was the first receptor shown to bind β-glucans to initiate functional responses in macrophages. Increasingly, PRRs are being identified through which NDPs can modulate immune responses including dectin-2, mannose receptor, complement receptor 3, and scavenger receptors.

Orally consumed NDPs will first and foremost exert their immunomodulatory activity in the intestine. NDPs can interact with a variety of intestinal immune cells located in the Peyer's patches through microfold cell-mediated transcytosis across the intestinal epithelium. Also, NDPs can directly be sampled in the lumen by CX3CR1+ antigen presenting cells which were shown to consist for 75–80% of macrophages in mice. Furthermore, studies into macrophage populations in human mucosa of the duodenal-proximal jejunum revealed different subsets, but all positive for the CX3CR1 fractalkine receptor. These CX3CR1+ macrophages were demonstrated to be of an immune suppressive subset (often referred to as M2). In case of intestinal immune activation, a CCR2-dependent influx of circulating CD14+ monocytes is initiated which under local conditions polarize into an inflammatory macrophage subset (often referred to as M1). The immune suppressive and inflammatory macrophage balance is key for maintaining intestinal homeostasis. Furthermore, intestinal macrophages have been shown to endocytose and degrade β-glucans. Beta-glucan fragments were subsequently translocated towards the bone marrow, spleen and peripheral lymph nodes and locally released to exert immunomodulatory effects. We therefore pose that in vitro models based on macrophages are relevant to understand the immunomodulatory potential of NDPs and study the mechanism of action. Given the central position of macrophages in intestinal, and potentially peripheral, immune homeostasis we used an M1/M2 dichotomy model system to further understand the functional impact of NDP intake.

Materials and methods

NDP characteristics

The NDPs and their treatment to remove lipopolysaccharide were described previously. Briefly, Saccharomyces cerevisiae (yeast)-derived soluble 94% pure β-1,3/1,6-glucan preparation (yeast-βG) was provided by BioThera (Eagan, MN, USA); a Lentinula edodes (shiitake)-derived 73% pure β-1,3/1,6-glucan (shiitake-βG) was produced in our lab; an apple-derived 80% pure rhamno-galacturonan-I (apple-RGI) preparation was provided by INRA (Paris, France); and a wheat-derived 77% pure arabinoxylan (wheat-AX) was provided by BioActor (Maastricht, The Netherlands). The NDPs, except yeast-βG, were treated with alkaline-ethanol for 5 hours, neutralized with HCl and lyophilized, resulting in lipopolysaccharide levels below immune stimulatory thresholds.

Primary macrophage differentiation and polarization

Primary macrophages were differentiated as previously described. In brief, human monocytes were obtained from buffy coats from healthy donors (Sanquin, Nijmegen, The Netherlands) using the QuadroMACS system and CD14 microbeads according to the manufacturer's protocol (Miltenyi Biotec, Leiden, The Netherlands). A written informed consent was obtained before sample collection. Monocytes were differentiated into macrophages through a 7 day culture in RPMI 1640 + glutamax medium (Gibco, Bleiswijk, The Netherlands) supplemented with 1% of pen/strep (Sigma, St Louis, MO, USA), 1% of sodium-pyruvate (Lonza, Breda, The Netherlands), 1% of MEM non-essential amino acids (Gibco), 10% of fetal bovine serum (FBS, Hyclone, Eindhoven, The Netherlands) and 50 ng ml⁻¹ M-CSF (R&D systems, Minneapolis, MN, USA) at 1 x 10⁶ cells per 2 ml per well in a 24-well culture plate. After three and five days of culture half of the medium was replaced with fresh culture medium containing 100 ng ml⁻¹ M-CSF (providing a final concentration of 50 ng ml⁻¹). After 7 days of culture all of the medium was replaced with culture medium either without additions to generate M0, with 20 ng ml⁻¹ IFN-γ (R&D systems) and 20 ng ml⁻¹ TNF-α (R&D systems) to generate M1, with 20 ng ml⁻¹ IL-4 (R&D systems) to generate M2 or with 500 μg ml⁻¹ of NDP preparation and incubated for 18 hours. Validation of M0, M1 and M2 macrophage phenotype was performed via Q-PCR and based on gene transcription markers as established previously. Regarding M1 and M2 macrophages, when applicable the culture medium was again completely replaced with either culture medium without additions or 500 μg ml⁻¹ of NDP preparations.

RNA extraction, cDNA synthesis and qPCR

RNA extraction, cDNA synthesis and qPCR was performed as described previously. In brief, RNA was extracted with TRizol (Invitrogen, Bleiswijk, The Netherlands) and RNeasy kits (Qiagen, Venlo, The Netherlands) and quality and quantity controlled. Subsequently, cDNA was synthesized with iScript (Bio-Rad, Veenendaal, The Netherlands). Next, 5 μl cDNA was mixed with SYBRGreen supermix (Bio-Rad), forward and reverse primers, and milliQ water to total 20 μl. Primers were derived from the Harvard Primerbank (http://pga.mgh.harvard.edu/primerbank/) and synthesized by Biologix (Nijmegen, The Netherlands). Primers’ information is available upon request. Amplification was performed with the CFX96 Touch Real-Time PCR Detection System (Bio-Rad) in technical duplicate and all samples were normalized to reference genes and medium control stimulations using the qBase⁺ software (Biogazelle, Gent, Belgium).

Endocytosis assay and antigen processing analysis

Macrophages that were polarized, NDP- or medium-treated in a 24-well plate were washed with PBS and incubated for 1 h in 0.5 ml culture medium containing 2 μg ml⁻¹ FITC-labelled E. coli fragments (excitation at 495 nm, emission at 519 nm;
Molecular Probes, Leiden, The Netherlands) to analyze for phagocytic activity or 10 ng ml\(^{-1}\) DQ-OVA (D12053, Molecular Probes) to analyze antigen processing. OVA is heavily labelled with fluorescent green BDY dyes (excitation at 502 nm, emission at 510 nm) resulting in a quenching effect. Protease mediated hydrolysis of OVA, as expected upon processing, reliefs the quenching effect. Before addition, \(E. \ col i\) fragments were re-suspended in non-heat-inactivated human serum (2 mg per 100 µl) and incubated for 60 minutes at 37 °C. Following incubation with either \(E. \ col i\) or DQ-OVA, macrophages were washed with PBS without calcium and magnesium and detached following 10 minutes incubation with 0.25% Trypsin/EDTA (Gibco, Bleiswijk, The Netherlands) at 37 °C. Subsequently, cells were at least 4 times diluted in PBS with 2% FBS and centrifugation at 300g for 10 minutes. Cells were re-suspended in 200 µl PBS, 5 µl trypan blue was added to cells incubated with \(E. \ col i\) fragments for quenching extracellular fluorophores, and analyzed using flow cytometry (Accuri™ C6, BD Biosciences) with Accuri C6 software. For each individual experiment MFI values were collected of marker positive cells (marker set at 1% on non-stained cells) after which medium stimulated MFI values were set at 100% to correct for donor differences.

**Multiplex cytokine and chemokine analysis**

Secretion of cytokines and chemokines (i.e. CCL1, CCL5, CCL20, CCL24, CXCL8, IL-1β, TNFα, MMP1, CXCL9, CXCL10 and CXCL11) by macrophages was measured using Bio-Plex Pro Reagent Kit (BioRad, Veenendaal, The Netherlands) according to the manufacturer’s instructions. In brief, 50 μL magnetic beads were dispersed in a plate together with 50 μl of undiluted sample or standard and incubated for 1 h at RT. Subsequently, the beads were incubated with 25 μl detection antibodies for 30 min and with 50 μl streptavidin-PE for 10 min at RT. The beads were re-suspended in 125 μl assay buffer and read by Bio-Plex® MAGPIX™ Multiplex Reader (BioRad). Data processing was performed using Bio-Plex Manager 5.0, and concentrations (in pg ml\(^{-1}\)) were interpolated from standard curves. To identify relations and to display our results most effectively, we used lower or upper limit of the standard curve for data points that were below or above the standard curve, respectively.

**Cell migration assay**

Cell migration assays were performed using an xCELLigence RTCA DP system (xCELLigence, ACEA Biosciences, San Diego, USA) and CIM-16 well plates, according to the manufacturer’s instructions. In brief, 160 μl of macrophage-conditioned medium were added to the lower chamber of CIM-16 well plates. Fresh macrophage culture medium was used as negative control and CCL5 (400 ng ml\(^{-1}\)) as positive control. Following placement of the upper chamber the upper chamber wells were filled with 50 µl THP-1 cell culture medium and equilibrated at 37 °C for 1 h which was recorded as background cell index. Next, 50 μl THP-1 cell suspension (8 × 10\(^6\) cells per ml) was added to the upper chamber after which cells settled for 5 minutes at RT. The cell index is a unit-less parameter used to indicate the impedance of electron flow caused by adherent cells and defined as [impedance at time point impedance n – impedance in the absence of cells]/nominal impedance value. The cell index was measured every 5 min over the course of 22 h. Monocyte migration was subsequently calculated as area under the curve using Prism 6 software (Graphpad, La Jolla, USA).

**Statistics**

Statistical analysis and graphing were performed using GraphPad Prism (version 6, Graphpad Software). The data are presented as means ± SD and differences were assessed by one- or two-way ANOVA, followed by multiple comparisons test with the Fisher’s LSD test. Differences with \(p < 0.05\) were considered significant.

**Results**

**Functional characterization of non-polarized macrophages following exposure to NDPs**

Previously, we performed a transcriptional analysis of non-polarized macrophages following their exposure to NDPs. Functionality of these NDP-treated macrophages was assessed here by comparing endocytosis capacity, antigen processing and cytokine and chemokine production to that of M1- or M2-like macrophages. To this end, non-polarized macrophages were treated with yeast-βG (i.e. soluble), shiitake-βG, apple-RGI or wheat-AX or polarized towards M1- or M2-like macrophages.

Macrophage endocytosis capacity following NDP-treatment of polarization was investigated by incubation with fluorescent \(E. \ col i\) fragments and detection using flow cytometry (Fig. 1A and ESI Fig. 1A†). M2-like macrophages demonstrated a trend towards increased \(E. \ col i\) uptake whereas M1-like macrophages significantly reduced \(E. \ col i\) uptake. Similarly, macrophages treated with shiitake-βG or wheat-AX revealed a significant reduced \(E. \ col i\) uptake. Next, to investigate the potential for antigen processing, the NDP-treated or polarized macrophages were incubated with DQ-OVA. Polarization towards M1 or M2 phenotypes did not alter DQ-OVA processing by macrophages (Fig. 1B and ESI Fig. 1B†). In contrast, processing of DQ-OVA was significantly reduced by apple-RGI, shiitake-βG and wheat-AX. Finally, the secretion of chemokines and cytokines by NDP-treated or polarized macrophages was assessed. Tested signaling molecules were selected based on gene transcription levels of M1-like macrophages, with typically and/or significantly increased transcription of CXCL9, CXCL10, CXCL11, and TNFα, M2-like macrophages, with a strongest and significantly increase in CCL24 transcription, and NDP-treated macrophages, with typically and/or significantly increased CCL1, CCL5, CCL20, CXCL8, IL-1β, and MMP-1 transcription for most NDPs (ESI Table 1†). Polarization of macrophages into an M1-like phenotype resulted in significantly increased secretion of TNFα, CXCL9, CXCL10, and CXCL11, whereas macrophage polarization into an M2-phenotype did not significantly affect...
secretion of tested cytokines or chemokines (Table 1). Exposure of macrophages to NDPS, and in particular wheat-AX, affected secretion of signaling molecules. Wheat-AX significantly increased the secretion of CCL1, CCL5, CCL20, CXCL8 and IL1β. These signaling molecules are described to induce migration of monocytes (ESI Table 1†). Therefore, a monocyte migration assay was performed using cell culture medium as negative control, CCL5 as positive control and conditioned medium from non-polarized macrophages, macrophages polarized towards an M1-like phenotype and wheat-AX-treated macrophages. In line with the cytokine and chemokine secretion profiles, conditioned medium from M1-like polarized macrophages or wheat-AX-treated macrophages significantly increased monocyte migration when compared to medium from non-polarized macrophages (Fig. 1C).

**Apple-RGI, shiitake-βG and wheat-AX induced similar transcriptional profiles in M1- and M2-like macrophages**

In a steady state most of the macrophages in the intestine will have an M2-like phenotype.18 Therefore, M2-like macrophages were exposed to medium or NDPS and analyzed for their transcriptional expression of in part previously established gene markers.24 Yeast-βG only demonstrated to significantly increase transcription of CXCL8 and MINCLE (Fig. 2). Apple-RGI, shiitake-βG and wheat-AX in addition also significantly increased transcription of CD80, DECTIN-2, IDO1 and LAMP3 and decreased transcription of TLR7. Shiitake-βG and wheat-AX also significantly increased transcription of IL-1B, TNFA, LOX1 and CCL20 and decreased transcription of MGL. Finally, wheat-AX also significantly increased transcription of IRF1 and decreased transcription of CCL13. These results indicate that NDPS can indeed modify the gene transcription of M2-like macrophages, as they did for non-polarized macrophages.24

To test whether NDPS similarly could impact M1-like macrophages, the transcription of the same set of marker genes was analyzed. Overall, NDPS modified gene transcription similarly as upon treatment of M2-like macrophages, albeit less pronounced (ESI Fig. 2†), with wheat-AX again being most effective. In contrast to modifications to the M2-like macrophage transcriptional profile, however, apple-RGI was more effective than shiitake-βG in inducing gene transcriptional changes in M1-like macrophages.

**NDP-treatment decreased endocytosis and antigen processing capacity in M1- and M2-like macrophages**

Upon identifying transcriptional modifications of M1- and M2-like macrophages by apple-RGI, shiitake-βG and wheat-AX, the functionality of NDP-treated macrophages was investigated regarding endocytosis and antigen processing. Wheat-AX-treatment of M1- and M2-like macrophages resulted in significantly reduced endocytosis of E. coli fragments and DQ-OVA processing (Fig. 3). Although similar to the impact on non-polarized macrophages (Fig. 1A and B), the inhibitory effect of wheat-AX...
was more pronounced in M1- and M2-like macrophages than in non-polarized macrophages.

Apple-RGI and shiitake-βG non-significantly lowered E. coli uptake and significantly lowered DQ-OVA processing by M1-like macrophages which was similar to their effect on non-polarized macrophages. Upon exposure to M2-like macrophages, shiitake-βG also appeared to reduce E. coli uptake and DQ-OVA processing. In contrast, apple-RGI did not affect the M2-like macrophage functionality regarding E. coli uptake and DQ-OVA processing. Finally, yeast-βG did not modulate the endocytosis of E. coli fragments nor processing of DQ-OVA by M1- or M2-like re-polarized macrophages, similar to non-polarized macrophages.

**Wheat-AX-treatment induces signaling molecule secretion in M2-like macrophages with strong chemotactic capacity towards monocytes**

To test whether NDP-mediated altered functionality of M1- and M2-like macrophages was also reflected in alternative cytokine and chemokine production, and as a result monocYTE recruitment, the cytokines and chemokines secreted by M1- and M2-like macrophages from a single donor were analyzed following incubation with apple-RGI, shiitake-βG or wheat-AX for 18 h. Cytokine and chemokine secretion (pg ml⁻¹) was analyzed with Bio-Plex multiplex analysis. Values represent mean values ± SD of n = 3 different donors. Statistically significant differences were analyzed by one-way ANOVA: *p < 0.05.

**Table 1** Polarization of macrophages or NDP-treatment increased cytokine and chemokine secretion

| Proteins | Medium | M1        | M2         | Yeast-βG | Apple-RGI | Shiitake-βG | Wheat-AX |
|----------|--------|-----------|------------|----------|-----------|-------------|----------|
| CCL1     | 22 ± 8 | 394 ± 365 | 8.3 ± 0    | 82 ± 104 | 322 ± 297 | 3295 ± 2463 | 7088 ± 3535 |
| CCL5     | 23 ± 14| 1635 ± 289| 22 ± 13    | 32 ± 18  | 441 ± 194 | 207 ± 53    | 2744 ± 2650 |
| CCL20    | 5.9 ± 8 | 174 ± 173 | 1.2 ± 1    | 5.6 ± 4  | 60 ± 16   | 7.4 ± 3     | 3504 ± 2672 |
| CCL24    | 64 ± 39 | 36 ± 4    | 97 ± 48    | 119 ± 129| 37 ± 21   | 1321 ± 1041 | 775 ± 586  |
| CXCL8    | 2280 ± 1653 | 2764 ± 622 | 668 ± 353  | 3547 ± 1229 | 16581 ± 1840* | 18956 ± 1785* | 19008 ± 1593* |
| IL-1β    | 0.1 ± 0 | 0.3 ± 0   | 0.1 ± 0    | 0.1 ± 0  | 2.3 ± 2   | 0.7 ± 1     | 195 ± 251  |
| TNFR     | 8.1 ± 3 | 31395 ± 16953 | 4.0 ± 1 | 13 ± 6 | 548 ± 477 | 387 ± 366 | 13215 ± 14182 |
| MMP1     | 123 ± 0 | 166 ± 37  | 123 ± 0    | 144 ± 37 | 187 ± 65  | 166 ± 37    | 797 ± 909  |
| CXCL9    | 44 ± 44 | >24199*   | 200 ± 227  | 47 ± 69  | 53 ± 12   | 371 ± 314   | 92 ± 102  |
| CXCL10   | 136 ± 103 | 27617 ± 17822* | 60 ± 31 | 93 ± 30 | 25 ± 29   | 55 ± 19     | 45 ± 39   |
| CXCL11   | 0.5 ± 0 | 3830 ± 3927* | 0.3 ± 0 | 0.4 ± 0 | 0.1 ± 0   | 0.4 ± 0     | 0.5 ± 1   |

Macrophages were non-treated (medium), polarized towards M1-like or M2-like macrophages or treated with 500 μg ml⁻¹ yeast-βG, apple-RGI, shiitake-βG or wheat-AX for 18 h. Cytokine and chemokine secretion (pg ml⁻¹) was analyzed with Bio-Plex multiplex analysis. Values represent mean values ± SD of n = 3 different donors. Statistically significant differences were analyzed by one-way ANOVA: *p < 0.05.

**Discussion**

NDPs are considered immunomodulatory food components with variable bioactive properties. Here we investigated the functional impact of NDP-exposure to different macrophage phenotypes. NDP-treatment of non-polarized macrophages revealed that NDPs in general hampered typical M2-like macrophage functionality such as antigen endocytosis and processing (Fig. 1A and B). These findings are in line with reduced gene transcription in NDP-treated macrophages of NDP-binding lectin-receptors (i.e. CD302, MERTK and dectin-1) and HLA-class II molecules for lysosomal antigen loading (i.e. HLA-DM and HLA-DO (ESI Table 2)). Of note, gene transcription of the receptor for ovalbumin (i.e. mannose receptor), which is likely involved in the uptake of DQ-OVA, was not affected. Results further revealed that NDPs induced secretion of a unique mixture of chemokines and cytokines in non-polarized macrophages which is in line with the M(NDP) transcriptional phenotype. Cytokine and chemokine production patterns can be considered as markers to phenotype macrophages as CXCL9, CXCL10, CXCL11 and TNFα which induce monocyte recruitment are for M1-like macrophages (Table 1 and Fig. 1C). This is in line with literature which indicates that an influx of bone-marrow-derived monocytes replenishes intestinal macrophage populations especially in case of inflammation. NDPs, in particular wheat-AX, induced secretion of a signature mix of CCL1, CCL5, CCL20, CCL24, CXCL8 and IL-1β in the absence of CXCL9, CXCL10 and CXCL11 in non-polarized macrophages. CCL1 and CCL5 were shown in literature to mediate recruitment of CCR8- and CCR5-positive regulatory T cells, respectively, that might limit (chronic) inflammation in the intestine. In contrast, CCL20 is essential for migration of CCR6-positive T₄₁₇ cells towards the intestine, a cell type linked to chronic inflammatory disease, and IL-1β is crucial for Th17 polarization. Moreover, CCL24 was shown to induce inflammatory infiltration of eosinophils, neutrophils and basophils and CXCL8 has been shown to induce migration and activation of a large variety of immune cells. Conditioned medium from
wheat-AX-treated macrophages instigated monocyte migration equal to the conditioned medium from M1-like macrophages (Fig. 1C). Theoretically the wheat-AX-treated macrophages could also induce migration of other immune cell types such as T cells (ESI Table 1†). Unfortunately, our model system appears limited to assessing migration of adherent cell types, preventing analysis of T cell migration as function of conditioned medium from treated macrophages. Previously, exposing non-polarized macrophages to various NDPs induced a general transcriptional phenotype that was termed M(NDP) and was distinct from M1- and M2-like macrophages. 24 Functionally, however, the various NDPs have demonstrated clear differences exemplified by on the one hand soluble yeast-βG that apparently does not affect non-polarized macrophage functionality and on the other hand wheat-AX that strongly impacts non-polarized macrophage functionality. This indicates that the term M(NDP) to classify a phenotypical macrophage subset is limited in scope to transcriptional changes.

In homeostasis intestinal macrophages are in general of an M2-like subset whereas in case of inflammation the macrophages are, locally, predominantly of the M1-like subset.18–20 Therefore M1- and M2-like macrophages were exposed to NDPs to assess the impact on their phenotype. NDPs, and again in particular wheat-AX, indeed induced gene transcription in M2-like macrophages and to some lesser extent M1-like macrophages similar to the M(NDP) transcriptional phenotype (Fig. 2 and ESI Fig. 2†). In addition, NDP-treatment similarly altered endocytosis and antigen processing of M1- and M2-like macrophages.

**Fig. 2** NDPs induced an M (NDP) transcriptional phenotype in M2-like macrophages. M2-like macrophages were stimulated with medium, yeast-βG, apple-RGI, shiitake-βG or wheat-AX at 500 μg ml⁻¹ for 24 h and analyzed for gene transcription using qPCR. Bars represent mean fold change ± SD of n = 2–3 different donors. Statistically significant differences compared to medium control were analyzed by one-way ANOVA: * p < 0.05, ** p < 0.01, *** p < 0.001.
**Fig. 3** NDPs affected endocytosis and antigen procession activity in M1-like and M2-like macrophages. M1-like macrophages (A and C) and M2-like macrophages (B and D) were stimulated with medium, yeast-βG, apple-RGI, shiitake-βG or wheat-AX at 500 μg ml\(^{-1}\) for 24 h followed by 1 h incubation with *E. coli*-fragments (A and B) or DQ-OVA (C and D). *E. coli*-fragment and DQ-OVA uptake was analyzed by measuring MFI using flow cytometry. Bars represent mean values normalized to medium control (100%) ± SD of *n* = 3 different donors. Statistically significant differences compared to medium control were analyzed by one-way ANOVA: *p* < 0.05, **p** < 0.01, ***p** < 0.001.

**Table 2** NDP-treatment of M1- and M2-like macrophages increased cytokine and chemokine secretion

| Macrophage polarization status | M1 + medium | M1 + Apple-RGI | M1 + Shiitake-βG | M1 + Wheat-AX | M2 + medium | M2 + Apple-RGI | M2 + Shiitake-βG | M2 + Wheat-AX |
|-------------------------------|-------------|----------------|------------------|---------------|-------------|----------------|------------------|--------------|
| CCL5                          | 916         | 1898           | 705              | 4961          | 9.9         | 375            | 890              | 1929         |
| CCL20                         | 17.8        | 110            | 13               | 1784          | 0.8         | 18             | 223              | 859          |
| CCL24                         | 14.9        | 54             | 16               | 648           | 352         | 272            | 1970             | 1784         |
| CXCL8                         | 921         | 15 865         | 4336             | 19 590        | 298         | 21 968         | 23 592           | 23 213       |
| IL-1β                         | 0.2         | 0.3            | 0.2              | 3.1           | 0.1         | 0.2            | 1.3              | 4.1          |
| TNFa                          | 203         | 3035           | 284              | 30 331        | 2.3         | 549            | 10 012           | 23 877       |
| CXCL9                         | >24 199     | >24 199        | >24 199          | >24 199       | 25.2        | 37.7           | 37.7             | 126          |
| CXCL10                        | 224 944     | 2603           | 699 198          | 17 327        | 13.5        | 18.7           | 28.8             | 64.8         |
| CXCL11                        | 1188        | 36.6           | 410              | 698           | 0.1         | 0.1            | 0.2              | 0.4          |

M1- or M2-like macrophages were incubated with medium or treated with 500 μg ml\(^{-1}\) apple-RGI, shiitake-βG or wheat-AX for 24 h. Cytokine and chemokine secretion (pg ml\(^{-1}\)) was analyzed with Bio-Plex multiplex analysis. Values represent average of a technical duplicate measurement of a single donor.
macrophages (Fig. 3). These changes in functional responses depend on both the NDP as well as the polarization state of the cells. When comparing apple-RGI and shiitake-βG, apple-RGI demonstrated more potency to alter M1-like macrophage functionality whereas shiitake-βG was more effective in changing M2-like macrophage functionality (Fig. 3 and Table 2). This suggests that the macrophage polarization status dictates towards which NDP it is most responsive, which in turn might relate to the expressed PRRs to recognize and bind NDPs. Indeed, PRRs dectin-1, LacCer, scavenger-, mannose-, and complement receptor were shown to bind to the shiitake-derived β-glucan termed lentitin of which transcription was higher in M2-like macrophages, whereas toll-like receptors 1 and 2 were shown to bind RG-I of which transcription was higher in M1-like macrophages.

Cytokine and chemokine secretion of M1- and M2-like macrophages following NDP-treatment (Table 2) revealed a similar unique mixture as NDP-treated non-polarized macrophages. The cumulative effect of wheat-AX treatment of M2-like macrophages was the strongly increased recruitment of monocytes (Fig. 4). Results shown here on the induction of monocyte migration need to be extended to other immune cell types, but also investigated at lower NDP concentrations, to show that NDPs modify immune responses towards a lowered activation state, but with enhanced vigilance. Our findings, albeit in vitro and with suboptimal macrophage phenotypes, are in line with these results. NDPs reduced M2-macrophage, but also M1-macrophage, capacity to endocytose and process antigen but increased the macrophage potential to recruit monocytes providing a rationale for the observed effects in clinical studies.

Conflicts of interest
There are no conflicts to declare.

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