Activation of inflammatory responses in human U937 macrophages by particulate matter collected from dairy farms: an in vitro expression analysis of pro-inflammatory markers

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Abstract

Background: The purpose of the present study was to investigate activation of inflammatory markers in human macrophages derived from the U937 cell line after exposure to particulate matter (PM) collected on dairy farms in California and to identify the most potent components of the PM.

Methods: PM from different dairies were collected and tested to induce an inflammatory response determined by the expression of various pro-inflammatory genes, such as Interleukin (IL)-8, in U937 derived macrophages. Gel shift and luciferase reporter assays were performed to examine the activation of nuclear factor kappa-light-chain-enhancer of activated B cells (NF-κB) and Toll-like-receptor 4 (TLR4).

Results: Macrophage exposure to PM derived from dairy farms significantly activated expression of pro-inflammatory genes, including IL-8, cyclooxygenase 2 and Tumor necrosis factor-alpha, which are hallmarks of inflammation. Acute phase proteins, such as serum amyloid A and IL-6, were also significantly upregulated in macrophages treated with PM from dairies. Coarse PM fractions demonstrated more pro-inflammatory activity on an equal-dose basis than fine PM. Urban PM collected from the same region as the dairy farms was associated with a lower concentration of endotoxin and produced significantly less IL-8 expression compared to PM collected on the dairy farms.

Conclusion: The present study provides evidence that the endotoxin components of the particles collected on dairies play a major role in mediating an inflammatory response through activation of TLR4 and NF-κB signaling.

Keywords: AhR, IL-8, LPS, NF-κB, PM, TLR, Dairy Farms, PM, Inflammation

Background

Inhalation of particulate matter (PM) and bioaerosol exposure, specifically from agricultural settings, has been shown to have a negative impact on the respiratory system of individuals and animals. Dairies are a large contributor to agriculture revenues in California, with little known about worker exposure to PM and bioaerosol. Exposure to dust on dairy farms may induce systemic reactions, increased bronchial responsiveness and chronic respiratory symptoms; all of which are frequently observed in farm workers [1].

Dairy farmers are exposed to organic dusts of a complex nature, and chronic respiratory symptoms are frequently observed in dairy farm workers. PM from dairies contain toxic and immunogenic constituents including histamine, endotoxins, mite antigen, cow urine antigen and microorganisms [2]. The bacterial content includes whole bacteria and cell wall components, such as endotoxin, lipopolysaccharide (LPS) derived from Gram negative bacteria and peptidoglycan, which is the
main cell wall constituent of Gram positive bacteria. These substances are known to be biologically active, and some can induce chronic airway inflammation [3].

Previous studies have elucidated the initiation of inflammatory events following exposure to PM from dairies or organic dust [4-6]. Exposure to endotoxin has been associated with increased respiratory symptoms in occupational settings and described as a risk factor for organic dust toxic syndrome [7], but studies also suggest that endotoxin exposure may protect agricultural workers from allergic disease [8]. Recent reports indicate that workers without a farm childhood had an increased risk of allergic sensitization (defined by a positive response to allergens) and the development asthma (a chronic inflammatory disorder of the airways) compared to a population with an early life exposure to farming [8,9]. Furthermore, two cross-sectional studies have shown, that children who lived on farms had a lower risk to develop asthma than the children in the reference group [10]. From their results, the authors concluded that the exposure to a wider range of microbes by growing up on a farm could explain part of the protective effect against asthma [10].

The inflammatory response is thought to be caused by bacteria and fungi present in the PM [11]. Along with being exposed to organic dust on dairies, agricultural workers in dry climate regions are also exposed to substantial concentrations of inorganic dusts from agricultural soils. This inorganic component is associated with increased small airway disease among California farm workers [12]. Although an association between chronic bronchitis and dust exposure has been found, asthma was only associated with keeping livestock but not with dust exposure [1]. This supports the inflammatory potential of PM from dairies.

The purpose of the present study was to explore the inflammatory effects of macrophage exposure to dairy PM that may be of relevance for the generation of health effects, a type of study that is difficult to perform in vivo. Our main aim was to investigate the activation of inflammatory markers in human U937 macrophages after exposure to PM collected on various dairy farms in California.

Methods
Reagents and PM collection
National Institute of Standards and Technology (NIST) Standard Reference Material (SRM) 1649, an atmospheric particulate material collected in an urban area, and a diesel exhaust particulate sample, NIST SRM 2975, were purchased from NIST (Gaithersburg, MD). [γ-32P] ATP (6000 Ci/mmol) was purchased from ICN (Costa Mesa, CA). PM from dairy farms were collected with a high-volume air sampler (model GS2310; Andersen Instruments Inc., Smyrna, GA) equipped with a four-stage cascade impactor (series 230, Andersen Instruments, Inc.) in the summer months of the year 2008 from various dairy farms and an urban area (Fresno, CA) of the San Joaquin Valley in California. The dairies included in the study were modern style dairies that included freestall barns and flush lanes, with at least 2,000 lactating cows (dairy #51: 2620 total cows, #54: 3200 total cows, #56: 4950 total cows, #57: 9550 total cows, #60: 2400 total cows).

Slotted aluminum substrates (Tisch Environmental, Cleves, OH) were used for PM collection. The nominal flow rate used for collection was 20 ft³/min, with particle size cutoffs of 10.2, 4.2, 2.1, and 1.3 μm, and these four fractions were named fraction A, B, C and D, respectively. Coarse PM is referred to as particles with a mass median aerodynamic diameter range of 10.2-2.1 μm and fine PM as particles within 2.1-1.3 μm. After collection, substrates from each stage were weighed; particles were removed by scraping with a spatula and stored at -80°C in vials. Stock solutions of particles were prepared before use by suspending them in autoclaved distilled water and by ultrasonication for 2 min at maximum power (100 W). Particles were used at 1, 5, or 10 μg/ml culture medium. 2,3,7,8-Tetrachlorodibenzo-p-dioxin (TCDD, > 99% purity) was originally obtained from Dow Chemicals Co. (Midland, MI). Dimethylsulfoxide (Me₂SO), Phorbol-12-myristate-13-acetate (TPA), polymyxin B (PMB), Tumor Necrosis Factor (TNF)-α and lipopolysaccharide (LPS, Escherichia coli O55:B5) were obtained from SIGMA (St. Louis, MO). Other molecular biological reagents were purchased from Qiagen (Valencia, CA) and Roche (Indianapolis, IN).

Endotoxin analysis
Samples were extracted by vortexing the filter in a TWEEN (10 ml at 0.05%) solution with pyrogen free water for 1 hour at 20-22°C and analyzed for biologically active endotoxin using the recombinant factor C assay (rFC) according to the manufacturer (Lonza Inc., Walkersville, MD), which detects activation of Factor C utilizing a fluorogenic according to Saito et al. [13]. The samples, 100 μl of blank, and an endotoxin standard (Escherichia coli O55:B5) were added to a 96 well plate. The plates were then pre-incubated for 10 minutes at 37°C. A mixture of 100 μl of rFC enzyme solution, buffer, and fluorogenic substrate at a 1:4:5 ratio were then added. The plates were incubated for 1 hour at 37°C. Fluorescence was read at time 0 and 1 hour in a fluoroscence microtiter plate reader (Biotek Instruments, Winooski, VT). Excitation was read at 380 nm and emission at 440 nm. To obtain the endotoxin concentration, the log difference in fluorescence (RFU) was plotted against the log endotoxin concentration (EU/ml) in a linear regression curve. The standards used linear
axis and polynomial regression curve, degree 2. Blanks taken in the field and plate well blanks along with spiking assays were used for quality control in order to account for any field contaminates and lab factors affecting fluorescence, such as pyrogen free water, reagent water centrifuge tubes pipette tips and microplates. Dilution of some samples was necessary; in these cases, a 50-fold dilution was performed.

Cell culture and transient transfection
We obtained human U937 monocytic cells from the American Tissue Culture Collection (Manassas, VA) and maintained them in RPMI 1640 medium containing 10% fetal bovine serum (Gemini, Woodland, CA), 100 U penicillin, and 100 μg/ml streptomycin supplemented with 4.5 g/L glucose, 1 mM sodium pyruvate, and 10 mM HEPES. Cell culture was maintained at a cell concentration between 2 × 10⁵ and 2 × 10⁶ cells/ml. For differentiation into macrophages, U937 cells were treated with TPA (5 μg/ml) and allowed to adhere for 48 hr in a 5% CO₂ tissue culture incubator at 37°C, after which they were fed with TPA-free medium.

For transient transfection of U937 macrophages, luciferase reporter constructs were transfected via Nucleofector technology as described previously [14]. Briefly, 10⁶ U937 macrophages were resuspended in 100 μl Nucleofector Solution V (Amaza GmbH, Köln, Germany) and nucleofected with 1.0 μg plasmid DNA using program V-001, which is preprogrammed into the Nucleofector device (Amaza GmbH). Following nucleofection, the cells were immediately mixed with 500 μl of prewarmed RPMI 1640 medium and transferred into six-well plates containing 1.5 ml RPMI 1640 medium per well. After 24 h transfection, macrophages were treated with PM or LPS (control) for 4 h. Luciferase activities were measured with the Luciferase Reporter Assay System (Promega, Madison, WI) using a luminometer (Berthold Lumat LB 9501/16, Pittsburgh, PA). Relative light units are normalized to β-galactosidase activity and to protein concentration using Bradford dye assay (Bio-Rad, Hercules, CA).

Cell viability assay
To assess the effect of PM on the viability of U937 macrophages, a trypan blue exclusion test was used [15]. A 10-μl portion of re-suspended cell pellet was placed in 190 μL PBS with 200 μl trypan blue (0.5% dilution in 0.85% NaCl). After a 5 min, 10 μl of the cell suspension was loaded into a hemocytometer, and the proportion of nonviable to viable cells was determined.

Quantitative real-time reverse transcription-PCR
Total RNA was isolated from U937 cells using a high-purity RNA isolation kit (Roche), and cDNA synthesis was done as previously described [16]. Quantitative detection of β-actin and differentially expressed genes was performed with a LightCycler Instrument (Roche Diagnostics, Mannheim, Germany) using the QuantiTect SYBR Green PCR Kit (Qiagen) according to the manufacturer’s instructions. DNA-free total RNA (1.0 μg) was reverse-transcribed using 4 U Omniscript reverse transcriptase (RT; Qiagen) and 1 μg oligo(dT)₁₅ in a final volume of 40 μl. The primers for each gene (Table 1) were designed on the basis of the respective cDNA or mRNA sequences using OLIGO primer analysis software provided by Steve Rozen and the Whitehead Institute/MIT Center for Genome Research [17] so that the targets were 100-200 bp in length. PCR amplification was carried out in a total volume of 20 μl containing 2 μl cDNA, 10 μl 2 × QuantiTect SYBR Green PCR Master Mix, and 0.2 μM of each primer. The PCR cycling conditions were 95°C for 15 min followed by 40 cycles of 94°C for 15 sec, 60°C for 20 sec, and 72°C for 10 sec. Detection of the fluorescent product was performed at the end of the 72°C extension period. Negative controls were concomitantly run to confirm that the samples were not cross-contaminated. A sample with DNase- and RNase-free water instead of RNA was concomitantly examined for each of the reaction units described above. To confirm the amplification specificity, the PCR products were subjected to melting curve analysis. All PCR assays were performed in triplicate. The intra-assay variability was < 7%. Data was analyzed with the LightCycler analysis software.

Table 1 Primers for quantitative real time PCR analyses

| Gene   | Forward primer (5’-3’) | Reverse primer (5’-3’) |
|--------|------------------------|------------------------|
| β-actin| GGACTTCGAGAAGAGATGG   | AGCACCTGTGTGGCTTACAG  |
| COX-2  | TGAAAACCCACCTCAACACA   | GAGAAGCTTCCCAGCTTTT   |
| CYP1a1 | TAGACACTGATCGTGGCAG    | GGGAAAGCTTCAAGCATC    |
| IL-6   | GAACCTCCCTTCCACAAGCG   | TTTTCTGGCAGTGGCTTCTT  |
| IL-8   | CTGGCCGCAACAGAAATTA    | ATTTGACATCGGCCAAGCTTC |
| TNFa   | CAGAGGGGAAGGTGGCCAG    | CCTTGGCCTGTTAAGGACGC  |
| SAA1   | GGGCATGTAATGCGTCTCCT   | AGCGGAAGCTTCTTTGCAGT  |

Gel-mobility-shift assay (GMSA)
Nuclear extracts were isolated from U937 cells, as described previously [16]. In brief, 5 × 10⁶ cells were treated with PM or LPS for 90 min unless noted otherwise in the figure legends, and harvested in Dulbecco’s PBS containing 1 mM PMSF and 0.05 μg/μl of apro tinin. After centrifugation, the cell pellets were gently resuspended in 1 ml of hypotonic buffer (20 mM HEPES, 20 mM NaF, 1 mM Na₃VO₄, 1 mM Na₃P₂O₇, 1 mM EDTA, 1 mM EGTA, 0.5 mM PMSF, 0.13 μM ok adaic acid, 1 mM dithiothreitol, pH 7.9, and 1 μg/ml...
bands were quantified using a ChemiImager. Gels to X-ray films. For quantitative analysis, respective
lamellae gel and visualized by exposure of the dehydrated
plexes were resolved on a 4% nondenaturating polyacry-
excess of unlabeled DNA fragments. Protein-DNA com-
were performed in the presence of a 100-fold molar
room temperature for 20 min. Competition experiments
and 1
μ
NaCl, 1 mM EDTA, 0.5 mM dithiothreitol, 5% glycerol,
and 1 μg poly (dl-dC). The samples were incubated at
room temperature for 20 min. Competition experiments
were performed in the presence of a 100-fold molar
excess of unlabeled DNA fragments. Protein-DNA com-
plexes were resolved on a 4% nondenaturating polyacry-
lamide gel and visualized by exposure of the dehydrated
gels to X-ray films. For quantitative analysis, respective
bands were quantified using a Chemilumigter
(Alpha Innotech Corporation, San Leandro, CA).

Statistical analysis
All experiments were repeated a minimum of three
times, and data are expressed as mean ± SD. Differences
were considered significant for P < 0.05. Comparison of
two groups was made with an unpaired, two-tailed stu-
dent’s t-test. Comparison of multiple groups was made
with ANOVA followed by Dunnett or Tukey test.

Results
Dose-dependent effect of PM from dairy farms on
inflammatory factors
To address the dose-dependent effect of PM, the mRNA
expression of pro-inflammatory genes 6 hr after treat-
ment with various concentrations of PM was studied.
As shown in Table 2, treatment of U937 macrophages
for 6 h with the fine particles of PM fraction C with a
cutoff of 2.1 μm from dairy #57 in the range of 1, 5, or
10 μg/ml cell culture medium led to a dose-dependent
mRNA induction of cyclooxygenase 2 (COX-2), Tumor
neerosis factor-alpha (TNF-α), IL-6, IL-8, and serum
amyloid A (SAA1). In addition, COX-2, TNF-α, IL-6,
and IL-8 mRNA expression was significantly increased
by PM C compared to control at the low concentration
of 1 μg/ml. In contrast, SAA1 was significantly induced
only at 5 or 10 μg/ml fine PM. The most conspicuous
effect of fine PM from dairy #57 was found in the cases
of COX-2, TNF-α and IL-8 (over 35-fold increase),
followed by IL-6 and SAA1 expression.

To estimate the toxic potency, the effects of fine PM
were compared with LPS, which has been shown to be
an efficient inducer of inflammatory factors in U937
macrophages. As shown in Table 2, a concentration-
dependent increase of COX-2, TNFα, IL-6, IL-8, and
SAA1 mRNA expression showing a 14.5-, 9.4-, 6.2-,
12.8-, and 2.0-fold increase, respectively, at the lowest
concentration (0.1 μg/ml) of LPS tested was observed.
Time-dependent analysis of the mRNA increase of the
inflammatory markers after PM exposure showed a
maximum increase as early as 6 h after initial treatment,
which sustained over a period of 24 h (data not shown).
Therefore, U937 macrophages were treated for 6 h to
analyze mRNA expression of the target genes.

Effect of different size fractions of PM collected from
various dairy farms on IL-8 expression
The potency of PM collected from different dairies to
induce an inflammatory response determined by the
expression of IL-8 was tested. The pro-inflammatory
marker IL-8 was selected for the mechanistic studies
since IL-8 has been identified as a sensitive marker for
inflammation and PM exposure [18,19]. PM with a size
cutoff of 10.2 (fraction A), 4.2 (fraction B), 2.1 (fraction
C), and 1.3 μm (fraction D) were collected from five
different dairies in California. Both coarse (2.5-10 μm

| Gene  | PM (μg/ml) | LPS (μg/ml) |
|-------|------------|-------------|
|       | 1 | 5 | 10 | 0.1 | 0.5 | 1.0 |
| COX-2 | 10.6 ± 0.2a | 15.9 ± 1.4a | 43.5 ± 2.4a | 14.5 ± 1.3a | 48.2 ± 2.4a | 82.1 ± 4.1a |
| TNF-α | 8.1 ± 0.3a | 19.8 ± 0.6a | 425 ± 12a | 9.4 ± 1.4a | 168 ± 1.4a | 422 ± 2.6a |
| IL-6  | 3.2 ± 0.4a | 7.5 ± 0.4a | 221 ± 0.7a | 6.2 ± 0.4a | 14.2 ± 1.0a | 252 ± 1.1a |
| IL-8  | 9.3 ± 0.4a | 17.4 ± 2.1a | 55.6 ± 1.7a | 12.8 ± 1.4a | 225 ± 1.3a | 516 ± 1.3a |
| SAA1  | 1.6 ± 0.4 | 2.7 ± 0.4a | 4.6 ± 0.3a | 2.0 ± 0.2a | 3.2 ± 0.3a | 7.8 ± 1.2a |

U937 macrophages were treated with 1.0, 5.0, or 10.0 μg/ml PM C (size cutoff: 2.1 μm) from dairy #57 for 6 h. As a positive control, cells were treated with 0.1, 0.5, or 1.0 μg/ml LPS for 6 hours and mRNA was analyzed by real time RT-PCR. Control cells received 1% PBS. Results are normalized to b-actin and given as
fold increase of the mRNA levels in treated cells versus controls ( = 1). Values are given as mean ± SD of triplicates of three independent experiments.

*significantly increased compared to control cells (p < 0.05)
diameter) and fine (< 2.5 μm diameter) PM collected from the dairies had a significantly induced IL-8 mRNA in the human U937 macrophages (Figure 1A). PM from four out of the five dairies included in this study with a size cutoff of 4.2 μm tended to more actively induce IL-8 mRNA expression on an equal-dose basis than PM of 10.2, and 2.1, and 1.3 μm.

Treatment with 10 μg/ml coarse PM and fine PM collected from the California dairies caused a significantly stronger inflammatory response regarding the induction of IL-8 mRNA in the human U937 macrophages compared to 10 μg/ml PM collected from an urban area (Fresno, CA) in the same region as the dairies (Figure 1B). PM from dairy #57 were selected for mechanistic studies since dairy #57 represents a prototypical dairy of all five dairies investigated and is located in proximity to the Fresno area, where urban PM were collected for comparison. Urban PM with a size cut off of 4.2 and 2.1 μm (PM B and C) significantly induced IL-8 by 7- and 5-fold, respectively. Urban PM with a size cut off of 10.2 and 1.3 μm (PM A and D) induced IL-8 approximately 3-fold compared to control (Figure 1B).

**Effect of TLR4 and NF-κB inhibitors on PM-mediated induction of IL-8**

SC514 is a cell-permeable and selective IKK-2 inhibitor that blocks NF-κB-dependent gene expression. Pre-incubation of the U937 macrophages for 15 min with 5 μM SC514 blocked the induction of IL-8 mediated by fine PM (PM C) by approximately 70% (Figure 2). In order to neutralize the TLR4 pathway, U937 macrophages were incubated with an anti-hTLR4 antibody (5 μg/ml). After 1 h, the cells were treated with 10 μg/ml PM C from dairy #57 for 6 h. Pretreatment with anti-hTLR4 antibody blocked 65% of the PM-mediated induction of IL-8 (Figure 2). Furthermore, to neutralize the soluble fraction of endotoxin associated with the PM samples, we treated added Polymyxin B (PMB), an antibiotic that specifically binds LPS and prevents its binding to TLR4 receptors on the macrophages. PMB was added to the PM suspension at a concentration of 10 μg/mL and was pre-incubated for 30 min prior to addition of PM to the cells. PMB-treated PM C from dairy #57 effectively inhibited the induction of IL-8 by more than 50% compared to the untreated PM. To test whether PM collected from dairies contained components that activate
the Aryl hydrocarbon receptor (AhR), as shown for diesel PM from heavy-duty vehicles [19], the AhR antagonist MNF, which blocks ligand-dependent activation of the AhR, was used. The AhR is a cytosolic ligand-dependent transcription factor that translocates into the nucleus and binds the xenobiotic response element (XRE) located on the promoter of cytochrome P450 (CYP)1A1 gene. As shown in Figure 2, MNF had no significant effect on PM-mediated IL-8 induction in contrast to SC-514, PMB, or the anti-hTLR4 antibody.

PM from dairy farms induce NF-κB DNA-binding and NF-κB reporter activity

Activation of TLR4 is known to activate the NF-κB signaling pathway. Possible NF-κB complex binding to a NF-κB consensus element after treatment of U937 macrophages with dairy PM was investigated to determine if induction of IL-8 by PM is associated with NF-κB activation. U937 macrophages were treated for 1 h with 5 or 10 μg/ml PM C (2.1 μm) collected on dairy farm #57. Results show that both concentrations of PM C enhanced NF-κB binding compared to unexposed control cells (Figure 3A). Ten μg/ml PM C (lane 3) led to a higher increase of NF-κB binding than 5 μg/ml PM C (lane 2). As expected, LPS (positive control), significantly and dose-dependently enhanced DNA-binding activity to the NF-κB consensus element (lane 5 and 6). A 100-fold excess of cold NF-κB oligonucleotide completely abolished formation of PM C- or LPS-stimulated NF-κB complexes (Figure 3A, lanes 4 and 7).

In addition to GMSA, NF-κB reporter activity was investigated after PM exposure. As shown in Figure 3B, exposure of U937 macrophages to 10 μg/ml PM C from dairy #57 for 4 h significantly increased NF-κB luciferase reporter activity approximately 4-fold compared to control cells. Suppression of TLR4 with a neutralizing TLR4 antibody and the LPS-neutralizing antibiotic PMB decreased the PM-mediated reporter activity by about 35%. The effect of PM on XRE reporter activity was studied in order to test if PM from dairies would activate the AhR signaling pathway. Results in Figure 3B show that only diesel engine exhaust particles (DEP, 10 μg/ml) but not PM from dairy farm #57 were able to induce the XRE reporter activity. DEP are well known to contain polycyclic aromatic hydrocarbons (PAH), which can bind to and activate the AhR and induce XRE reporter activity and CYP1A1 gene expression [19].

Endotoxin units in PM

Previous studies, including our own, have found that PM collected from dairies are loaded with endotoxin [11]. In order to test the concentration of endotoxin, PM from five different dairies were collected, and the amount of endotoxin was compared to PM collected in an urban area of Fresno, CA. Due to the limited amount of PM available, only certain size fractions could be analyzed. The RFC analysis revealed an average of about 510 EU/mg including all PM fractions collected on five different dairy farms. With the exception of dairy #57, the endotoxin concentration tended to be higher in coarse
PM fractions with a size cutoff of 10.2 (PM A) and 4.2 μm (PM B) than in fine PM with a cutoff of 2.1 (PM C) and 1.3 μm (PM D) as shown in Figure 4. The PM collected in an urban area in Fresno, CA, contained an approximate mean of 40 EU/mg PM with the highest concentration of approximately 59 EU/mg in the coarse PM fraction. Fine particles within 1.3 μm contained significantly less endotoxin.

**Discussion**

Recent studies report that PM collected from urban pollution or diesel and gasoline engines contribute to cardiovascular morbidity and mortality and may induce inflammatory responses in vitro as well as in vivo, including in humans [20]. However, data from experiments investigating the effects of PM from agricultural facilities and dairies are scarce. The present study shows that treatment of human macrophages with PM collected from California dairies leads to increased expression of pro-inflammatory marker genes, such as IL-8, TNF-α, and COX-2, and upregulates acute phase proteins IL-6 and SAA1, which increase in concentration following infection, inflammation, or trauma.

Cells of the innate immune system, such as macrophages, recognize pathogens via pattern recognition receptors (PRR), such as TLR [21]. In LPS-sensitive U937 cells, LPS from E. coli is an agonist of TLR4. In this study, this led to a dose-dependent induction of the abovementioned inflammatory marker genes in the U937 macrophages. In addition, induction of the pro-inflammatory genes after exposure to LPS was associated with activation of NF-κB. The organic dust from a dairy farm contains microbial constituents that eminate from Gram positive as well as Gram negative bacteria, and gene activation may, therefore, be caused by the combined engagement of different TLRs.

To study the mechanism of the PM-induced inflammatory response, we selected IL-8 as a hallmark of inflammation. IL-8 has been demonstrated in chronic diseases including chronic obstructive pulmonary disease (COPD) [22,23] and ulcerative choliitis [24]. We have found that the receptor of IL-8 (CXC12) is involved in atherogenesis induced by environmental pollutants, such as dioxin [25], indicating the importance of IL-8 in the development of pathological endpoints.

The effect of ambient particles collected from an urban area (Fresno, CA) located in the San Joaquin Valley were significantly less potent in inducing IL-8 than PM from dairies, which contained 10-fold higher concentrations of endotoxin than the urban PM. Coarse PM fractions from the dairies with a cut off of 10.2 and 4.2 μm (PM A and PM B) contained more endotoxin than the fine fractions (PM C and PM D), which tended to correlate with a higher induction of IL-8 by PM A and B, although the effect was not statistically significant.

PM-activated gene expression of the inflammatory markers in this study was significantly suppressed by SC514, an inhibitor that blocks activation of NF-κB. This result indicates involvement of the NF-κB signaling pathway to mediate the effect of PM from dairies. Furthermore, exposure to dairy PM clearly activated NF-κB binding activity in the U937 macrophages. Addition of a TLR4 antibody to the culture medium before treatment with dairy PM neutralized approximately 60% of the PM-mediated effect on activation of NF-κB activity and the expression of IL-8. These data indicate that part of the PM components act through the TLR4 and that endotoxin is likely to be a critical component in the PM collected from dairies. The results also suggest that dairy PM-induced inflammation, such as IL-8 activation, is not entirely dependent on endotoxin and may include other components of dairy PM as perhaps double-stranded RNA or DNA of viruses, which may activate other TLR isoforms. The assumption, that other components in addition to endotoxin are triggering the inflammatory response, are also supported by the calculation of the endotoxin content in the dairy PM. Most dairy PM contained about 500 EU/mg which corresponds to 50 ng/mg. Accordingly, the exposure to 10 μg PM/ml would reflect an exposure to 0.5 ng endotoxin/ml however, the inflammatory response is still stronger compared to 100 ng LPS/ml. Interestingly, Poole et al. showed that agricultural PM may mediate their inflammatory response also by activating the TLR2 pathway [26], CD14 mediated responses [27] and effects of specific components of dairy PM such as muramic acid [28].

Treatment with LPS and activation of TLR4 is well known to activate NF-κB [29]. Thus, activation of NF-κB by PM treatment is likely to be mediated through...
the activation of TLR4, which is supported by the result that the addition of neutralizing TLR4 antibody inhibited approximately 35% of the PM-mediated activation of NF-κB in the luciferase reporter assay. In addition to the results from the GMSA and inhibitor studies, these data suggest that induction of IL-8, including other inflammatory genes, is mediated not only by TLR4, but also via NF-κB signaling. These results underline the involvement of NF-κB and indicate the supportive action of NF-κB on PM-mediated transcriptional activation of pro-inflammatory genes.

In contrast to PM derived from diesel engine exhaust (DEP), PM collected from dairies did not activate AhR regulated XRE activity, which is known to be activated by PM generated from traffic and combustion processes [19]. In contrast to NF-κB or TLR4, the classical AhR-XRE pathway is mostly responsible for rapid responses to xenobiotics and activation of genes encoding xenobiotic metabolizing enzymes, including CYP1A1, through XRE binding sites located on the promoter of the gene. More recently, studies have focused on investigating the toxicity of agricultural dust particles and human health effects. Animal excrement generates a large part of the pollution on dairies, and some facilities are used to xenobiotics and activation of genes encoding xenobiotic metabolizing enzymes, including CYP1A1, through XRE binding sites located on the promoter of the gene. More recently, studies have focused on investigating the toxicity of agricultural dust particles and human health effects.

In summary, the results from this study indicate that the most critical component of dairy farm dust is endotoxin, which may trigger local and systemic inflammatory reactions upon inhalation. In addition to endotoxin, allergens, microbial pathogens, bacterial toxins, fungal spores, and mycotoxins can attach to dust particles that, when inhaled, have the potential to cause local and systemic inflammatory reactions.

Conclusions

In conclusion, exposure to PM collected on dairy farms generates an inflammatory response in human macrophages partly mediated through activation of TLR4 and the NF-κB signaling cascade. The inflammatory response induced by the urban PM was significantly lower compared to PM from dairies, which has a higher concentration of endotoxin than the urban PM.

Acknowledgements

We like to thank Dr. Stephen Reynolds from Colorado State University for rIFC analysis of endotoxin. This work was supported by the NIOSH grant OH007550-06, the American Heart Association Western Affiliate grant 0765056Y, and NEHS grant R21ES15846-01A2. We would also like to thank Dr. Suzette Smiley-Jewell for her technical help with the manuscript.

Authors’ contributions

CV designed and performed experiments, analyzed the results and wrote the manuscript. JG helped with LPS analysis and writing the manuscript. DW, DA, AL, and PW performed experiments and assisted with data interpretation. YZ and NK managed PM collection and analysis. DB, MS, and FM designed study, providing overview of all the steps of sample collection, and helping write the manuscript. All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

Received: 11 October 2011 Accepted: 28 March 2012 Published: 28 March 2012

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