Macrophages that have ingested apoptotic cells in vitro inhibit proinflammatory cytokine production through autocrine/paracrine mechanisms involving TGF-beta, PGE2, and PAF.

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Abstract

Apoptosis in vivo is followed almost inevitably by rapid uptake into adjacent phagocytic cells, a critical process in tissue remodeling, regulation of the immune response, or resolution of inflammation. Phagocytosis of apoptotic cells by macrophages has been suggested to be a quiet process that does not lead to production of inflammatory mediators. Here we show that phagocytosis of apoptotic neutrophils (in contrast to immunoglobulin G–opsonized apoptotic cells) actively inhibited the production of interleukin (IL)-1β, IL-8, IL-10, granulocyte macrophage colony-stimulating factor, and tumor necrosis factor–α, as well as leukotriene C₄ and thromboxane B₂, by human monocyte-derived macrophages. In contrast, production of transforming growth factor (TGF)–β₁, prostaglandin E₂, and platelet-activating factor (PAF) was increased. The latter appeared to be involved in the inhibition of proinflammatory cytokine production because addition of exogenous TGF–β₁, prostaglandin E₂, or PAF resulted in inhibition of lipopolysaccharide-stimulated cytokine production. Furthermore, anti–TGF–β₁ antibody, indomethacin, or PAF receptor antagonists restored cytokine production in lipopolysaccharide-stimulated macrophages that had phagocytosed apoptotic cells. These results suggest that binding and/or phagocytosis of apoptotic cells induces active antiinflammatory or suppressive properties in human macrophages. Therefore, it is likely that resolution of inflammation depends not only on the removal of apoptotic cells but on active suppression of inflammatory mediator production. Disorders in either could result in chronic inflammatory diseases. (J. Clin. Invest. 1998. 101:890–898.)

Key words: apoptosis • inflammation • eicosanoids • mediators • phagocytosis

Introduction

Phagocytosis by macrophages or other cells is the final event in the lives of many cells undergoing apoptosis (1, 2). Apoptotic cells express cell surface changes which allow recognition and removal by macrophages. Removal occurs before lysis, which prevents the release of potentially toxic and immunogenic intracellular contents from the apoptotic cells into the surrounding tissue. Thus, in tissues such as the thymus in which apoptosis is ongoing, normal structure and function are maintained and inflammation is avoided. The removal of apoptotic cells also appears to be critical in the resolution of inflammation. Phagocytosis of apoptotic cells in inflammatory sites has been documented in vivo in experimental as well as clinical disease states, and disorders of apoptosis have been suggested to contribute to the persistence of chronic inflammatory conditions in the lung, kidney, and other organs (3–5).

The mechanisms by which apoptotic cells can be recognized and removed have received extensive study in the last few years. A number of receptors have been identified in vitro. These include an uncharacterized lectin inhibited by N-acetylglucosamine (6), the vitronectin receptor (αvβ3 integrin) which is thought to cooperate with CD36 in binding to thrombospondin on the surface of the apoptotic cell (7, 8), a phosphatidylerine (PS)–specific receptor (9–12), scavenger receptors (13–15), and the macrophage antigen identified by the mAb 61D3 (16), which may be identical to, or related to, CD14 (Devitt, A., C. Raykundalia, O.D. Moffatt, J.D. Capra, D.L. Simmons, and C.D. Gregory, manuscript submitted for publication). The ABC1 transporter has also been suggested to be involved in phagocytosis (17). The mechanisms by which any of these receptors facilitate apoptotic cell uptake are not known. The changes on the apoptotic cell which induce recognition by macrophages have also not been well characterized. Exposure of PS on the surface of apoptotic cells is associated with phagocytosis by macrophages of some phenotypes (10–12). Loss of membrane phospholipid asymmetry and early external exposure of PS have been documented on many different cell types undergoing apoptosis and the mechanisms mediating this membrane change are an area of active study (10, 18–23). Other changes reported to occur include expression of immature sugars on apoptotic thymocytes and hepatocytes (24, 25), the expression of ICAM-3 on apoptotic B cells which facilitates recognition by macrophages via a lymphocyte function-associated antigen-1–independent pathway (26), specific loss of surface expression of some (but not all) GPI-linked protein antigens, such as CD16 on apoptotic neutrophils (22, 27), and exposure of novel uncharacterized antigens recognized by mAbs on apoptotic cells in the embryonic chicken limb bud (28).

The effects of phagocytosis of apoptotic cells on the macrophage itself have not been characterized completely. Meagher et al. initially studied the effects of apoptotic neutrophil up-
take on the human monocyte-derived macrophage and found no induction of N-acetylgalactosaminidase or thromboxane B2 (TxB2) (29). If, however, the same apoptotic cells were opsonized with antibody and then fed to macrophages, a brisk induction of these substances was observed. After the phagocytosis of apoptotic eosinophils by human macrophages, there was no increase in TxB2 or GM-CSF (30). In contrast, secretion of TxB2 and GM-CSF increased after engulfment of opsonized apoptotic eosinophils or necrotic eosinophils. These investigators hypothesized that phagocytosis of apoptotic cells by the VnR/CD36 mechanism did not lead to a macrophage response, whereas uptake mediated by the FcR was stimulatory. Thus, the receptor involved in apoptotic cell uptake determined the response by the macrophage.

Several questions remain to be answered however. Is this truly a passive lack of response or could the macrophage be actively suppressing proinflammatory mediator production? Is the suppression peculiar to GM-CSF or are other proinflammatory cytokines involved? What about the effects on eosinosanoids other than TxB2? Based on our previous work, we knew that macrophages which had ingested digestible particulate stimuli secreted TGF-β (11, 31, 32). Since apoptotic cells are digestible particles (albeit large ones), it is reasonable to hypothesize that phagocytosis of apoptotic cells might induce the secretion of TGF-β, which would then act in an autocrine or paracrine fashion to inhibit the production of proinflammatory cytokines. TGF-β has been shown previously to downregulate production of proinflammatory mediators in macrophages (for reviews see references 33 and 34). Therefore, we studied the effects of phagocytosis of apoptotic human neutrophils on the production of TGF-β1, as well as GM-CSF, IL-1β, IL-8, IL-10, and TNF-α. Because the secretion of lipid inflammatory mediators could also be affected, the effects of apoptotic cell uptake on secretion of four eicosanoids, TxB2, LTC4, PGD2, and PGE2, were determined.

Methods

Reagents. DME and HBSS were purchased from Gibco BRL (Gaithersburg, MD); X-Vivo 10 medium was purchased from Bio-Whittaker (Walkersville, MD). Mouse anti-human CD45 clone 2D1, IgG1 was purchased from Becton-Dickson (San Jose, CA), or clone H130, IgG1, κ (PharMingen, San Diego, CA); rabbit anti–mouse IgG was purchased from Jackson ImmunoResearch Laboratories (West Grove, PA). LPS (Escherichia coli, serotype 0111:B14) was obtained from List Biological Laboratories (Campbell, CA); A23187 and zymosan were purchased from Cayman Chemical Co. (Ann Arbor, MI). Human TGF-β1, anti–human TGF-β antibody, human PDGF, anti–human PDGF, human IL-10, and anti–human IL-10 were purchased from R & D Systems (Minneapolis, MN). Platelet-activating factor (PAF, alkyl chain a mix of C18 and C16) was purchased from Avanti Polar Lipids (Alabaster, AL). Eicosanoids and indomethacin were purchased from Cayman Chemical Co. (Ann Arbor, MI).

Cells. Human macrophages were prepared from peripheral blood mononuclear cells, as previously published (35). The cells were plated at 4 × 10⁶/ml in 24-well plates for 60 min in DME, after which nonadherent cells were washed out with HBSS. The macrophages were cultured for 3 d in X-Vivo 10 medium containing 10% pooled human serum (five or more donors) prepared as described (36). The medium was changed at 3 d. For all experiments, macrophages had been cultured for 6–8 d before use. This yielded 1–2 million macrophages per well at the time of the assay.

Human neutrophils were used as a source of apoptotic cells. Apoptosis was induced by 10 min of UV irradiation at 254 nm followed by culture in DME + 10% heat-inactivated FBS (Gemini Bioproducts, Calabasas, CA) at 37°C, 5% CO₂ for 3–4 h by tumbling end-over-end in polypropylene tubes. At that time, apoptosis (assessed morphologically by light microscopy of stained cytocentrifuged cells) equaled or exceeded 60%, whereas necrosis (assessed by trypan blue uptake) was ≤ 1%. As a control particle, apoptotic human neutrophils were opsonized to induce uptake by Fc receptors for IgG. Apoptotic neutrophils were rotated end-over-end at 4°C with anti-CD45 for 30 min. After three washes, the secondary antibody rabbit anti–mouse IgG was added for an additional 30 min. The cells were washed again and resuspended in X-Vivo 10 medium without serum for use in the phagocytosis assay.

Jurkat cells. Jurkat T cells were cultured in RPMI medium containing 20% FBS, glutamine (2 mM), penicillin (100 U/ml), and streptomycin (100 μg/ml) at 37°C and 5% CO₂. When cell density reached 10⁶/ml, the culture was UV-irradiated for 10 min as described above, then cultured for 2–2.5 h. This routinely yielded populations containing 60–70% apoptotic cells and 5 × 10⁶ were added to each well (24-well plate) of macrophages. Uptake was confirmed by microscopy, as described below, except that Jurkat cells were not stained for myeloperoxidase (MPO).

Phagocytosis of apoptotic cells. Uptake was determined as described previously (7–11, 36). Apoptotic neutrophils were added to day 7–10 human monocyte-derived macrophages at 5 × 10⁶ cell/well of a 24-well plate in X-Vivo 10 medium and incubated for 1 h at 37°C 5% CO₂. The monolayer was then washed vigorously with ice-cold PBS to remove bound but uningested neutrophils, fixed with 1% formalin overnight, and treated with one part diansidine (1.25 mg/ml) to one part 0.05% H₂O₂ to stain for MPO as a marker of the ingested neutrophils (7, 8, 36). The macrophages themselves were routinely negative for peroxidase staining. Using ×40 phase-contrast microscopy, uptake of apoptotic neutrophils was assessed, counting two replicate wells for each condition of 200 macrophages per well. Phagocytic index was calculated as the percentage of phagocytosing macrophages multiplied by the average number of neutrophils ingested per macrophage. Macrophages that showed discrete, round, MPO-positive inclusion(s) were scored as having ingested one (or more) apoptotic neutrophils. Neutrophils with margins extended > 50% beyond the edge of the macrophage cell membrane were scored as noningested. (This convention may have underestimated the number of phagocytosed neutrophils, because ultrastructural analyses have shown macrophages to engulf such neutrophils with a thin wall of cytoplasm [data not shown].) Phagocytosis as assessed by this method was confirmed in preliminary experiments by electron microscopy. For uptake of opsonized apoptotic cells, the same technique was used; however, varying numbers (1–5 million) were added per well to determine what number was needed to achieve equivalent phagocytosis to that for unopsonized apoptotic cells.

Collection of supernatants. Before use, macrophage monolayers were washed with HBSS, then X-Vivo 10 medium without human serum was added. Apoptotic cells or control particles were added and supernatants were collected 18 h later. This time point was chosen because preliminary experiments showed that cytokine secretion stimulated by LPS or zymosan was maximal. Phagocytosis was assessed by microscopy (see above) to ensure that uptake was equivalent for the two different particles. For some experiments, the apoptotic cells were washed out after 1 h and fresh X-Vivo 10 medium was added for another 18 h. Supernatants were centrifuged at 2,000 rpm to remove particulate debris, then were stored in aliquots at −70°C. In some experiments, LPS at 1 ng/ml or zymosan at 50 μg/ml was added to stimulate cytokine production. Apoptotic cells or opsonized apoptotic cells were added at the same time of the stimulus.

Analysis of cytokines. Cytokine concentrations in the culture supernatants were determined by ELISA, using the Quantikine immunoassays manufactured by R & D Systems. The cytokines analyzed were GM-CSF, IL-1β, IL-8, IL-10, TGF-β1, and TNF-α. Assays were performed according to the instructions provided with each kit. For TGF-β1, the supernatants were acid-activated according to the manu-
facturer’s instructions before the assay. Color development was as-
sewed using the microplate autoreader (EL309) by Bio-Tech In-
struments (Winooski, VT). Data were analyzed using a log/log curve fit op-
tion from Delta Soft 3 ELISA analysis software for the Macintosh

Analysis of eicosanoids. PGD2, PGE2, LTC4, and TxB2 were
quantified by enzyme immunoassays using acetylcholinesterase-con-
jugated tracers as described previously (37, 38). The rabbit polyclonal
antibody for PGD2 was a gift from Dr. R.W. Kelly (Edinburgh, UK),
while the rabbit polyclonal antibody for TxB2 was a gift from Dr.
Frank Fitzpatrick (Pharmacia-Upjohn, Piscataway, NJ). The mouse
mAb for the peptidoleukotrienes was cross reactive with LTC4,
LTD4, and LTE4 and was purchased from Perceptive Bioreresearch
Products (Cambridge, MA). LTC4 was used as the reference stan-
dard in assays and the peptidoleukotriene products referred to as
LTC4. The mouse mAb used for the quantification of PGE2 was pur-
chased from Cayman Chemical Co. Supernatant samples for LTC4,
PGE2, and TxB2 were used without purification at dilutions of 1:5 to
1:10. Samples for PGD2 were derivatized with a methoxyamine re-
agent (50 μl of sample and 50 μl of reagent) by heating at 60°C for 1 h
(39), and diluted to a final concentration of 1:5. Standard curves and
samples were analyzed using the four-parameter curve fit option on
Delta Soft 3 software. The sensitivity of the assays ranged from 4 to
16 pg/ml.

Analysis of IL-8 and TNF-α mRNA levels by quantitative RT-
PCR. Total RNA was isolated from each macrophage population
using the Micro-Scale Total RNA Separator kit produced by CLON-
TECH (Palo Alto, CA). Then, 2 μg of product was checked for in-
tegrity by electrophoresis through a 1% formaldehyde agarose gel
containing ethidium bromide. If intact, 1.2 μg of the total RNA was
used for first-strand cDNA synthesis, using CLONTECH’s Advan-
tage RT-for-PCR kit. PCR products were quantitated using competi-
tive PCR with CLONTECH’s PCR Mimics kits for IL-8 and TNF-α;
minor modifications included the use of 4 μl of cDNA and MIMICS,
rather than 2 μl, and electrophoresis of 12 μl of sample in 2% ethid-
ium bromide/agarose gel for detection and quantification. β-actin was
used as a control and was not found to vary from experiment to ex-
periment.

Statistical analysis. The means of the concentrations for each cy-
tokine were analyzed using ANOVA for multiple comparisons; when
ANOVA indicated significance, the Tukey-Kramer HSD test for all
pairs was used to compare groups. For experiments in which exoge-
nous inhibitors were used (see Figs. 5–7), Dunnnett’s method was
used, with LPS-treated macrophages as the control. All data were an-
alyzed using JMP (version 3) Statistical Software for the Macintosh
(SAS Institute, Inc., Cary, NC).

Results

Phagocytosis of apoptotic cells does not stimulate the produc-
tion of proinflammatory cytokines by human macrophages.
Human monocyte-derived macrophages were given apoptotic
human neutrophils or opsonized apoptotic neutrophils in a se-
rum-free medium (X-Vivo 10). Preliminary experiments were
performed to determine the ratio of opsonized cells to mac-
rophages that should be added to achieve equivalent phagocy-
tosis for both types of cells. In five experiments we found that
5 × 10⁶ apoptotic cells added to each well of macrophages gave
an average phagocytic index of 43.1 ± 4.9 SEM, where the
phagocytic index is defined as the product of the percentage of
macrophages positive for phagocytosis (39.2) and the number
of cells per macrophage (1.1). A similar phagocytic index for
opsonized apoptotic cells (39.1 ± 5.7) was achieved using 10⁶
cells, with the percentage of positive macrophages equaling
35.5 and the number of neutrophils per macrophage equaling
1.1. Therefore, these cell numbers were used routinely for the
cytokine experiments. Fig. 1 shows a photomicrograph of apo-
totic neutrophils stained for MPO that have been ingested by
human macrophages, using the same culture conditions as
were used for assessment of cytokine production.

To determine what effects uptake of apoptotic cells would have,
unstimulated macrophages were exposed to apoptotic
cells for 18 h and supernatants were analyzed for cytokines.
Preliminary data had indicated that IL-1β, IL-8, IL-10, GM-
CSF, and TNF-α levels stimulated by opsonized zymosan or
opsonized apoptotic cells were maximal around this time (data
not shown). After 18 h, macrophages which had phagocytosed
apoptotic cells failed to increase production of GM-CSF, IL-
1β, IL-8, IL-10, or TNF-α, but stimulated release of TGF-β1 to
levels approximately three times that of control (Fig. 2).
A slight increase (40%) in TGF-β1 levels induced by apoptotic
cells was apparent as early as 1 h after addition of apoptotic
cells (data not shown). In contrast, opsonized apoptotic cells
stimulated increased production of all the cytokines except for
TGF-β1. As an additional control, we incubated apoptotic
neutrophils alone for 18 h and measured cytokine levels in the
harvested supernatant. Levels were undetectable for all of the
cytokines (data not shown).

One of the potential problems with allowing the apoptotic
neutrophils to remain in contact with the macrophages for 18 h
was the possibility that they would undergo secondary necrosis
and influence the results. This possibility was examined in two
ways. First, apoptotic neutrophils that had become necrotic ei-
ther spontaneously after time in culture or by heating to 56°C
for 5–10 min were fed to macrophages. These experiments
were repeated six times and the results were inconsistent. In
three out of six experiments, the necrotic cells stimulated cy-
tokine production, as was seen for necrotic eosinophils (31); however, in three experiments, there was no stimulation of cy-
tokine secretion. This disparity may be due to variations in the
necrotic cell preparation, since the neutrophils tended to clump together when they became necrotic. Therefore, we
took a second approach. Macrophages were allowed to phago-
cyte apoptotic or freshly isolated neutrophils for 1 h, at
which time residual cells were removed by washing, and fresh
medium (X-Vivo 10) was added. Supernatants were then col-
Apoptotic Cells Actively Suppress Inflammatory Cytokine Production

lected 18 h later. Although the total cytokine concentrations were lower, TGF-β1 was stimulated by apoptotic neutrophils while the other cytokines were inhibited (data not shown). In contrast, freshly isolated neutrophils either showed no inhibition or stimulated proinflammatory cytokine production.

Inhibition of cytokine production after uptake of apoptotic cells. Phagocytosis of apoptotic cells appeared to decrease the production of all cytokines by macrophages except TGF-β1 (Fig. 2); however, the decreases were not statistically significant since the levels of cytokine produced by unstimulated macrophages were low. Therefore, we wondered whether the binding and phagocytosis of apoptotic cells could inhibit cytokine production by stimulated macrophages. The possibility of active suppression was investigated by studying the effect of apoptotic cell uptake by macrophages stimulated to produce cytokines with LPS or zymosan. Macrophages were treated with either LPS (1 ng/ml) or zymosan (50 μg/ml) and at the same time exposed to apoptotic or opsonized apoptotic cells. As shown in Fig. 3, the phagocytosis of apoptotic cells by LPS-stimulated macrophages was associated with decreased cytokine production except for TGF-β1, which was enhanced significantly \((P = 0.003)\) when compared with LPS-treated macrophages or those fed opsonized cells. GM-CSF was decreased significantly from macrophages treated with LPS alone or with opsonized apoptotic cells \((P = 0.0001)\), IL-10 was decreased significantly compared with LPS treatment alone \((P = 0.002)\), and both IL-1β and TNF-α were decreased significantly compared with macrophages treated with both LPS and opsonized apoptotic cells \((P = 0.002)\). Similar results were observed with macrophages stimulated by zymosan (Fig. 3). One potential explanation for these results is that the phagocytosis of apoptotic cells resulted in macrophage loss. The macrophages were counted 18 h after each treatment, by lysing the cells with Zapoglobin and counting the nuclei. In five experiments, the average number of macrophages per well after stimulation with LPS was 1.4 (±0.2 SEM) million and after treatment with LPS and apoptotic cells 1.3 (±0.2 SEM) million.

An additional question was whether the effect of apoptotic cells was restricted to neutrophils or whether other apoptotic cells would have the same effect. Therefore, we fed the LPS-stimulated macrophages UV-induced apoptotic Jurkat cells and measured the effects on TNF-α, IL-8, IL-10, and TGF-β.
In five experiments, there was no statistical difference between the effects of the neutrophils compared with the Jurkat cells. Both populations had equivalent numbers of apoptotic cells and both caused decreases in TNF-α, IL-8, and IL-10 and an increase in TGF-β (data not shown). To determine whether the inhibition occurred at the level of transcription, quantitative RT-PCR was used to determine mRNA levels for IL-8 and TNF-α in macrophages treated with LPS, or LPS and apoptotic neutrophils. As expected, LPS increased steady-state mRNA levels for both cytokines compared with control macrophages (Fig. 5). If, however, the macrophages were exposed to apoptotic cells at the same time they were stimulated with LPS, the mRNA levels for both cytokines were decreased 1, 4, and 18 h later (Fig. 5). There was no change in β-actin levels (not shown), which provided additional evidence that macrophages had not died or been lost. These data are most compatible with a decrease in transcription of cytokine message or possibly a decrease in stability of the message.

Ingestion of apoptotic cells by macrophages induces soluble inhibitors of cytokine production. Since macrophages fed apoptotic cells were able to suppress cytokine production at both the mRNA and protein levels, it was important to determine whether this was due to a soluble factor or factors. Macrophages were fed apoptotic cells and supernatants were collected 18 h later. This conditioned medium was then added to macrophages at the same time as either LPS or zymosan. The inhibition of proinflammatory cytokines was even more profound than when apoptotic cells were added at the same time as the stimulus. Levels of IL-1β, GM-CSF, TNF-α, and IL-10 were undetectable; IL-8 was decreased to ≤ 10% of the levels seen with LPS or zymosan alone (data not shown). TGF-β1 was increased to levels similar to those seen from macrophages fed apoptotic cells (data not shown). These results suggested that one or more soluble inhibitors were produced by the macrophages after binding and/or phagocytosing apoptotic cells. Given that TGF-β1 was the only cytokine which was increased after addition of apoptotic cells or conditioned media from macrophages fed apoptotic cells, it seemed a likely candidate.

Direct addition of TGF-β1 to LPS-stimulated macrophages partially inhibited production of proinflammatory cytokines, which achieved statistical significance for IL-1β, IL-8, GM-CSF, and TNF-α (Fig. 6). Although the concentration used in the experiments shown was 10 ng/ml, we found suppression of GM-CSF, IL-1β, IL-10, and TNF-α at concentrations as low as 100 pg/ml (data not shown), which is similar to what we observed produced after the phagocytosis of apoptotic cells.

Figure 4. Phagocytosis of apoptotic cells inhibits zymosan-induced proinflammatory cytokine production but stimulates TGF-β1. Experiments were performed as described for Fig. 2, except that zymosan at 50 μg/ml was used to induce cytokine secretion. The mean cytokine production ± SEM is shown for seven experiments; *significantly different from zymosan-treated macrophages fed opsonized apoptotic cells (P < 0.05). Control, Unstimulated cytokine levels.

Figure 5. Phagocytosis of apoptotic cells decreases mRNA levels for IL-8 and TNF-α in LPS-stimulated human macrophages. The macrophages were treated exactly as described in Fig. 2, then RNA was isolated at the time points shown. Levels of mRNA for each cytokine were determined by RT-PCR as described in Methods. Data are expressed as mRNA levels (±SEM) relative to LPS-stimulated cells which are indicated as 100% (n = 3). White bars, Control; gray bars, LPS; black bars, LPS plus apoptotic cells.
In addition, we found that anti–TGF-β antibody added to LPS-stimulated macrophages fed apoptotic cells restored the inflammatory cytokine production, suggesting that endogenously produced TGF-β played a role in inhibiting the generation of inflammatory cytokines (Fig. 6). Finally, anti–TGF-β antibody was also found to partially block the inhibitory action of supernatants derived from macrophages which had ingested apoptotic cells (data not shown). As an additional control, PDGF (which is also secreted when mouse bone marrow macrophages ingest digestible particles, see reference 31) was added to LPS-stimulated macrophages. PDGF had little to no effect on cytokine secretion, and the addition of anti-PDGF antibody to LPS-stimulated macrophages fed apoptotic cells failed to restore cytokine production (data not shown).

Because IL-10 secretion was decreased by phagocytosis of apoptotic cells, we assumed it played no role in the inhibition of cytokine production. To confirm this, we studied the effects of adding exogenous IL-10 to LPS-stimulated macrophages, and anti–IL-10 antibody to LPS-stimulated macrophages which had phagocytosed apoptotic cells. Exogenously added IL-10 was able to inhibit production of TNF-α by 68% and IL-8 by 42% when added at 10 ng/ml. When IL-10 was added at 1 ng/ml, it inhibited TNF-α by 60% but had no effect on IL-8. However, anti–IL-10 antibody failed to restore cytokine production inhibited by the phagocytosis of apoptotic cells.

Because TGF-β did not appear to mediate all the inhibition and IL-10 was not involved, other candidate inhibitors were examined. We have shown previously that uptake of digestible particles by mouse macrophages leads to production of PAF (32). Although PAF is generally considered a proinflammatory mediator, its role in this system was examined. Addition of PAF to macrophages along with LPS led to a decrease in all cytokines except for TGF-β, which was stimulated threefold ($P = 0.006$) compared with macrophages treated with LPS alone (Fig. 7). Inhibition by PAF achieved statistical significance for IL-8, IL-10, and TNF-α ($P < 0.05$). In support of a role for PAF, the PAF receptor antagonist WEB 2086 led to restoration of cytokine production and a decrease in TGF-β secretion in LPS-stimulated macrophages ingesting apoptotic cells (Fig. 7). Two other PAF receptor antagonists (CV-6209 and PCA-4248) also restored cytokine production (data not shown). Taken together, these data implicated PAF in the inhibition of cytokine production after phagocytosis of apoptotic cells.

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has been shown to decrease macrophage production of cytokines such as TNF-α (e.g., references 40 and 41). As shown in Fig. 8, the exogenous addition of PGE2 to LPS-stimulated macrophages decreased their production of all the cytokines except TGF-β1, which was stimulated (Fig. 8). By adding indomethacin to LPS-stimulated macrophages that had ingested apoptotic cells, proinflammatory cytokine production was restored and TGF-β1 production was inhibited slightly.

Phagocytosis of apoptotic cells increased PGE2 production while decreasing TxB2 and LTC4. Since PGE2 affected cytokine production in this system, the ability of these macrophages to produce PGE2 and other eicosanoids was examined. Supernatants were collected from unstimulated human macrophages 1, 4, and 18 h after addition of apoptotic cells or opsonized apoptotic cells, and analyzed for PGE2, PGD2, LTC4, and TxB2. PGD2 levels were undetectable or very low in all cases (< 170 pg/ml). Opsonized apoptotic cells induced a 5-fold increase in PGE2 levels, but unopsonized apoptotic cells induced as much as a 15-fold increase detectable 1 h after the cells were added (Fig. 8). Compared with those from control macrophages, TxB2 levels were decreased at all time points by the addition of apoptotic cells, and LTC4 levels were decreased at 4 and 18 h; opsonized cells were stimulatory for both eicosanoids (Fig. 9).

Discussion

Phagocytosis of apoptotic cells was shown to inhibit macrophage production of GM-CSF, IL-1β, IL-8, IL-10, TNF-α, TxB2, and LTC4, but to increase production of TGF-β1, PGE2, and PAF. At this point, we have not determined whether these effects result from binding to macrophage receptors for apoptotic cells or from the actual uptake process itself. These effects were apparent in unstimulated macrophages as well as those stimulated with LPS or zymosan. Furthermore, TGF-β1, PAF, and PGE2 appeared to mediate the increase in inflammatory cytokine production, because addition of each of these to LPS-stimulated cells decreased cytokine production, and because addition of anti-TGF-β antibody, PAF receptor antagonists, or indomethacin to LPS-treated macrophages which had ingested apoptotic cells restored cytokine production.

We propose that the lack of proinflammatory mediator production results not from a passive lack of stimulation, but from an active suppressive mechanism involving the autocrine/paracrine action of TGF-β, PGE2, and PAF. The precise nature of the interaction between these three mediators is under further study. The data presented here suggest the possibility that PAF and/or PGE2 can stimulate the production of TGF-β1. Therefore, TGF-β may be the most important antiinflam-
flammatory mediator ultimately generated or PAF and PGE2 can have both direct and indirect effects on proinflammatory cytokine production. We have done preliminary experiments to assess these possibilities by adding anti-TGF-β antibody to LPS-stimulated macrophages that had been treated with either PAF or PGE2 to look for effects on TNF-α and IL-8 production; however, results were not consistent. The regulation appears to be quite complex and needs to be studied further before conclusions are drawn.

Initially, IL-10 seemed a likely candidate for potential anti-inflammatory activity in this system, based on observations that it is a potent suppressor of inflammatory cytokines including IL-1, IL-6, IL-8, and TNF-α, and it can also suppress production of superoxide anions and nitrous oxide (42–46). However, production of IL-10 was inhibited by the phagocytosis of apoptotic cells, and anti–IL-10 antibody failed to restore the production of cytokines inhibited by apoptotic cells. This does not rule out a role for IL-10 in vivo, but suggests that macrophages would be an unlikely source.

The mechanisms by which TGF-β, PAF, and PGE2 are produced by macrophages after phagocytosis of apoptotic cells and how they downregulate inflammatory cytokine and eicosanoid production are under active investigation. The decreased mRNA levels for IL-8 and TNF-α observed after ingestion of apoptotic cells are compatible with decreased transcription of cytokine messages; however, we cannot rule out alterations in message stability. The regulation is likely to be complex, given that we have at least three candidates that appear to affect proinflammatory cytokine production. TGF-β itself has been shown to inhibit cytokine production on a number of levels. It is known to increase transcription of its own message, but its effects on other cytokines are more complicated. TGF-β has been reported to have no effect on TNF-α mRNA levels, release of preformed TNF-α, or its degradation; rather it inhibited translation of TNF-α mRNA (47). TGF-β has also been shown to inhibit the release of IL-1, to downregulate the number of IL-1 receptors, and to increase production of IL-1ra (33, 34). Therefore, it may act as an antiinflammatory agent on a variety of levels. The importance of TGF-β in the regulation of inflammation is illustrated by the observations that knockout mice have severe and generalized inflammatory disorders (48, 49).

In this system, we studied macrophages which primarily used the VnR and CD36 mechanism for recognition of apoptotic cells. It is still not known whether the other receptors mediating uptake of apoptotic cells would have the same suppressive effect; however, preliminary data suggest that uptake mediated by the PS recognition mechanism also inhibits proinflammatory cytokine production. As we found for mouse bone marrow macrophages, treating human macrophages with particulate β-glucan for 48 h led to recognition of apoptotic cells through a PS-inhibitable mechanism (Warner, M.L., V.A. Fadok, D.L. Bratton, and P.M. Henson, manuscript submitted for publication). After phagocytosis of apoptotic cells, glucan-treated macrophages also produced increased levels of TGF-β1 and decreased levels of GM-CSF, IL-1β, IL-8, IL-10, and TNF-α. These results support the hypothesis that removal of apoptotic cells by “apoptotic” receptors rather than opsonic receptors such as FcR or CR3 leads to suppression of inflammatory cytokine production.

The selective effect of apoptotic cell uptake on macrophage eicosanoid generation suggests either that these macrophages use different PGH synthase enzymes for TxA2 and PGE2 production and/or undergo a switch in metabolism of PGH2 by PGE2 synthase versus thromboxane synthase. In support of the latter possibility, we have observed that TNF-α induced PGE2 synthase activity in mouse bone marrow–derived macrophages fed zymosan compared with macrophages which had not been pretreated with TNF-α (Fournier, T., V.A. Fadok, and P.M. Henson, manuscript submitted for publication). The somewhat delayed effect of apoptotic cell uptake on LTC4 may indicate that an effect on lipoxygenase pathways (5 lipoygenase or LTC4 synthase) takes longer for implementation. It is also possible (though less likely) that differential effects are mediated by alterations in local substrate (arachidonate and/or PGH2) distribution.

It seems likely that the nature and duration of the inflammatory response is determined by competition between proinflammatory and antiinflammatory uptake mechanisms. Our observations suggest that phagocytosis of apoptotic cells not only prevents the release of toxic and immunogenic intracellular contents, but also stimulates the macrophages to express an antiinflammatory or suppressive phenotype. Both mechanisms are likely to be critical in the resolution of inflammation. If this is true, then disorders in either uptake or response to apoptotic cells by macrophages could contribute to chronic inflammation.

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