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Pregnancy alters glucose-6-phosphate dehydrogenase trafficking, cell metabolism, and oxidant release of maternal neutrophils

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Pregnancy is associated with changes in host susceptibility to infections and inflammatory disease. We hypothesize that metabolic enzyme trafficking affects maternal neutrophil activation. Specifically, immunofluorescence microscopy has shown that glucose-6-phosphate dehydrogenase (G-6-PDase), the rate-controlling step of the hexose monophosphate shunt (HMS), is located near the cell periphery in control neutrophils but is found near the microtubule-organizing centers in cells from pregnant women. Cytocchemical studies confirmed that the distribution of the G-6-PDase antigen is coincident with functional G-6-PDase activity. Metabolic oscillations within activated pregnancy neutrophils are higher in amplitude, though lower in frequency, than activated control neutrophils, suggesting limited HMS activity. Analysis of radioisotope-labeled carbon flux from glucose to CO2 indicates that the HMS is intact in leukocytes from pregnant women, but its level is not enhanced by cell stimulation. Using extracellular fluorescent markers, activated pregnancy neutrophils were found to release reactive oxygen metabolites (ROMs) at a lower rate than activated control neutrophils. However, basal levels of ROM production in polarized pregnancy neutrophils were greater than in control neutrophils. Microtubule-disrupting agents reversed the observed changes in G-6-PDase trafficking, metabolic oscillations, and ROM production by maternal neutrophils. G-6-PDase trafficking appears to be one mechanism regulating ROM production by maternal neutrophils.


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Conflict of interest: The authors have declared that no conflict of interest exists.

Nonstandard abbreviations used: reactive oxygen metabolite (ROM); hexose monophosphate shunt (HMS); glucose-6-phosphate (G-6-P); glucose-6-phosphate dehydrogenase (G-6-PDase); 6-aminonicotinamide (6-AN); phosphofructokinase (PFK); pyruvate kinase (PK); lactate dehydrogenase (LDH); N-formyl-met-leu-phe (FMLP); resonance energy transfer (RET); diketoenoxyethyleneamide (H2TMRos); tetraethylroamine (TMRos); microtubule-organizing center (MTOC).
coupling of glucose transport with hexokinase activity (21). During cell activation, hexokinase is translocated to the plasma membrane (23), where it accelerates glucose transport to provide glucose-6-phosphate (G-6-P) for glycolysis and the HMS (Figure 1). The HMS’s first and rate-controlling step is mediated by glucose-6-phosphate dehydrogenase (G-6-PDase), which converts G-6-P into 6-phosphogluconolactone with NADPH release. 6-Aminonicotinamide (6-AN), an inhibitor of G-6-PDase, inhibits ROM production but not metabolism (24). The HMS’s second step is catalyzed by 6-phosphogluconate dehydrogenase, which yields NADPH and ribose 5-phosphate. Phosphofructokinase (PFK) is a key allosteric enzyme of glycolysis. Pyruvate kinase (PK) converts phosphoenolpyruvate to pyruvate. Lactate dehydrogenase (LDH) converts pyruvate to lactate. Only about 1–2% of the carbon flux enters the citric acid cycle in neutrophils (25). These enzymatic steps are generally unidirectional in living cells and, by definition, not at equilibrium. Moreover, these steps do not occur in isolation but rather form a complex network of chemical reactions regulated by cooperative interactions. These conditions promote the emergence of novel patterns of enzyme activity, such as temporal oscillations and spatial waves (26–29).

This study shows that G-6-PDase undergoes anterograde transport in neutrophils from nonpregnant women, but retrograde transport in cells from pregnant women. Microtubule-dependent retrograde transport of G-6-PDase apparently limits HMS activity and the extracellular release of ROMs; this mechanism may account for changes in disease susceptibility during pregnancy.

Methods

Patients. Peripheral blood samples were obtained from nonpregnant women and pregnant women with written informed consent, and the study was approved by the institutional review board. The nonpregnant group consisted of women in the secretory phase of the menstrual cycle who were not taking oral contraceptives and who had no history of acute or chronic inflammatory conditions. Women with normal pregnancies (from 20 to 30 weeks to term) had no medical or obstetric complications. Eligible patients were approached at the Detroit Medical Center (Detroit, Michigan, USA).

Cell preparation. Neutrophils were isolated using Ficoll-Hypaque (Sigma Chemical Co., St. Louis, Missouri, USA) density gradient centrifugation (30). Neutrophil viability was greater than 95% as assessed by trypan blue exclusion.

Reagents. Colchicine, vincristine, vinblastine, taxol, cytochalasin D, LPS (serotype 026:B6), N-formyl-met-leu-phe (FMLP), 6-AN, PMA, and Brij-58 were obtained from Sigma Chemical Co. FITC and TRITC were obtained from Molecular Probes Inc. (Eugene, Oregon, USA). Rabbit anti-hexokinase, anti-G-6-PDase, and anti-PK polyclonal antibodies and a goat anti-LDH polyclonal antibody were obtained from Chemicon International Inc. (Temecula, California, USA). The anti-PFK antibody was the generous gift of R. Kemp (The Chicago Medical School, North Chicago, Illinois, USA). Anti-γ-tubulin and IFN-γ were obtained from R&D Systems Inc. (Minneapolis, Minnesota, USA). FITC- or TRITC-conjugated antibodies were prepared as described previously (30).

HMS activity. HMS activation was measured as described previously (31, 32). Cells (∼2 × 10⁶) were incubated in a total volume of 0.8 ml in media consisting of either [D-1-14C]-labeled or [D-6-14C]-labeled glucose (American Radiolabeled Chemicals Inc., St. Louis, Missouri, USA) at 0.5 µCi/ml with 1 mM glucose in PBS. The 14CO₂ produced was captured in a center well containing 0.5 ml hyamine hydroxide (Research Products International Corp., Mount Prospect, Illinois, USA) and a strip of filter paper. Incubations were performed in sealed containers at 37°C in a shaking water bath for 4 hours. CO₂ was released from the solution by addition of 1 ml of 7N trichloroacetic acid followed by incubation for 1 hour. Lastly, 3 ml of scintillation fluid was added to the center well, followed by liquid scintillation counting.

G-6-PDase cytochemistry. G-6-PDase activity was localized within cells using cytochemical methods (33). Briefly, the G-6-PDase incubation medium consisted of phosphate buffer (pH 7.4), 32% (wt/vol) polyvinyl alcohol, 5 mM G-6-P (VWR International, Batavia, Illinois, USA), 0.4 mM NADP⁺ (Calbiochem-Novabiochem Corp., San Diego, California, USA), 2.5 mM MgCl₂, 2.5 mM NaN₃, 0.16 mM 1-methoxy-5-methyl-phenazinium methyl sulfate (Sigma Chemical Co.), and 2.5 mM Nitro Blue Tetrazolium (VWR International). Cells were washed, fixed with 2% paraformaldehyde, incubated for 30 minutes, and then washed again. Samples were trans-illuminated using a 590 long-pass optical filter (Omega Optical Inc., Brattleboro, Vermont, USA) to enhance the reaction product’s contrast.

Immunofluorescence staining. Neutrophils were placed on glass coverslips, incubated with reagents as described below, then fixed as described previously (23). Briefly, cells were fixed with 2% paraformaldehyde, permeabilized with 1% Brij-58, and fixed with 2% paraformaldehyde at room temperature for 20 minutes. Cells were washed with HBSS, labeled with antibodies at 4°C for 30 minutes, then washed with HBSS at room temperature.

Fluorescence microscopy. Cells were observed with an Axiolab fluorescence microscope (Carl Zeiss Inc., Thornwood, New York, USA) with mercury illumination interfaced to a computer using Scion image-processing software (NIH, Bethesda, Maryland, USA) (27). A narrow bandpass filter set (Omega Optical Inc.) was used with excitation at 485/22 nm and emission at 530/30 nm for FITC, and excitation at 540/20 nm and emission at 590/30 nm for TRITC. Long-pass dichroic mirrors of 510 and 560 nm were used for FITC and TRITC, respectively. For resonance energy transfer (RET) imaging, 485/22-nm and 590/30-nm optical filters were used for excitation and emission, respectively, in conjunction with a 510-nm dichroic mirror. The fluorescence images were collected with an IMax ICCD (Princeton Instruments Inc., Princeton, New Jersey, USA). RET was also examined using microspectrophotometry (28).
NAD(P)H oscillations. NAD(P)H autofluorescence oscillations were detected as described previously (30, 34). An iris diaphragm was adjusted to exclude light from neighboring cells. A cooled photomultiplier tube held in a model D104 detection system (Photon Technology International Inc., Lawrenceville, New Jersey, USA) attached to a Carl Zeiss Inc. microscope was used (30, 34).

ROM production. Pericellular release of ROMs from single cells was monitored as described previously (35). Briefly, adherent neutrophils were surrounded in 2% gelatin containing 100 ng/ml dihydrotetramethylrosamine (H2TMRos) (Molecular Probes Inc.). ROMs, especially H2O2, released by the cells entered the gelatin matrix, where they oxidized H2TMRos to tetramethylrosamine (TMRos), which was detected by fluorescence microscopy. The internal accumulation of ROMs was detected as described previously (36).

Results

Altered trafficking of G-6-PDase in neutrophils from pregnant women. Since G-6-PDase is the HMS’s rate-controlling step, ROM release is decreased in activated cells from pregnant women (2, 3), and metabolic enzymes may undergo translocation during cell activation (23), we hypothesized that G-6-PDase trafficking is changed during pregnancy. To test this hypothesis, we examined the intracellular distribution of five metabolic enzymes in neutrophils from nonpregnant and pregnant women under unstimulated and LPS-activated conditions (Figure 2, second row). We found that G-6-PDase is normally detected in the plasma membrane’s vicinity in cells from nonpregnant women (Figure 2, e and f). In contrast, G-6-PDase was not found near the plasma membrane in neutrophils from pregnant women (Figure 2, g and h). Quantitatively, cells from pregnant women demonstrated 3% near the plasma membrane, 39% in the cytoplasm, and 58% as centrosome-associated fluorescence. G-6-PDase translocation was observed in all of the samples from normal pregnant shape, show a uniform cytoplasmic distribution of hexokinase. When neutrophils were exposed to LPS (50 ng/ml) for 20 minutes, hexokinase was translocated from the cytoplasm to the cell periphery (Figure 2, a and b). This confirms previous work showing that hexokinase is translocated to plasma membranes during rat peritoneal macrophage activation with PMA (23). To determine whether hexokinase translocation was deficient in maternal neutrophils, we assessed the intracellular distribution of hexokinase in LPS-activated cells. Pregnancy neutrophils were competent to translocate hexokinase to the cell periphery (Figure 2, c and d). Hence, this cannot account for the reported changes in neutrophil function.

We next evaluated the distribution of G-6-PDase in neutrophils from nonpregnant and pregnant women under unstimulated and LPS-activated conditions (Figure 2, second row). We found that G-6-PDase is normally detected in the plasma membrane’s vicinity in cells from nonpregnant women (Figure 2, e and f). In contrast, G-6-PDase was not found near the plasma membrane in neutrophils from pregnant women (Figure 2, g and h). Quantitatively, cells from pregnant women demonstrated 3% near the plasma membrane, 39% in the cytoplasm, and 58% as centrosome-associated fluorescence. G-6-PDase translocation was observed in all of the samples from normal pregnant women.
women. Thus, neutrophils from pregnant women are characterized by a largely central distribution of G-6-PDase, in contrast to the peripheral distribution seen in 100% of the cells from nonpregnant individuals.

To ascertain the specificity of these changes in G-6-PDase distribution, we performed experiments using antibodies directed against other enzymes. We examined the intracellular distribution of PFK, PK, and LDH in nonpregnancy and pregnancy neutrophils with and without LPS stimulation (Figure 2, third through fifth rows). PFK was chosen because it is a primary allosteric regulator and is specific for the glycolytic pathway (Figure 1). Carbon flux through glycolysis and the HMS reaches PK, whose activity is a rate-limiting step of glycolysis. Under the conditions tested, no major differences between the distribution of these enzymes could be found in control and pregnancy cells. PK accumulated at the lamellipodium (Figure 2, fourth row). Interestingly, the three enzymes that can act as rate-controlling steps in glycolysis and/or the HMS accumulate at the cell periphery and lamellipodium. Similar results concerning the intracellular distributions of hexokinase, G-6-PDase, PFK, PK, and LDH were obtained using 50 nM FMLP (data not shown), which indicates that these changes in enzyme distribution are not specific for LPS. Furthermore, the activating stimulus PMA (30 nM for 30 minutes at 37°C) had no effect on the distribution of G-6-PDase (data not shown), although some changes in metabolic oscillations and ROM production were noted (see below). Our experiments suggest that the trafficking of G-6-PDase is uniquely important in the regulation of neutrophil metabolism during pregnancy.

G-6-PDase accumulate near the centrosome in neutrophils from pregnant women. Cells from nonpregnant (NP) (a–h) and pregnant (P) (i–p) women were studied using direct immunofluorescence. Experiments were performed without (a–d, i–l, and r) and with (e–h, m–p, and s) prior incubation with LPS (50 ng/ml, 20 minutes). Cells were fixed, then stained with FITC-conjugated anti–γ-tubulin and TRITC-conjugated anti–G-6-PDase. Although γ-tubulin and G-6-PDase did not colocalize (b, c, f, and g) or exhibit RET (d and h) in control neutrophils, both colocalization (j, k, n, and o) and RET (l and p) were found in maternal neutrophils. RET was confirmed using fluorescence spectroscopy in r and s. FITC and TRITC controls are shown in q. Thus, G-6-PDase accumulates near the centrosome of neutrophils from pregnant, but not nonpregnant, women. n = 9. x550.

Figure 3

G-6-PDase traffics to the centrosome in cells from pregnant women. Cells from nonpregnant (NP) (a–h) and pregnant (P) (i–p) women were studied using direct immunofluorescence. Experiments were performed without (a–d, i–l, and r) and with (e–h, m–p, and s) prior incubation with LPS (50 ng/ml, 20 minutes). Cells were fixed, then stained with FITC-conjugated anti–γ-tubulin and TRITC-conjugated anti–G-6-PDase. Although γ-tubulin and G-6-PDase did not colocalize (b, c, f, and g) or exhibit RET (d and h) in control neutrophils, both colocalization (j, k, n, and o) and RET (l and p) were found in maternal neutrophils. RET was confirmed using fluorescence spectroscopy in r and s. FITC and TRITC controls are shown in q. Thus, G-6-PDase accumulates near the centrosome of neutrophils from pregnant, but not nonpregnant, women. n = 9. x550.

Figure 4

Colchicine reduces the pericentriolar distribution of G-6-PDase. LPS-stimulated cells from nonpregnant (a–l) and pregnant (j–r) women were studied using direct immunofluorescence in conjunction with microscopic imaging and spectrophotometry. Cells were fixed, then stained with FITC-conjugated anti–γ-tubulin and TRITC-conjugated anti–G-6-PDase. The properties of γ-tubulin (a, d, and g) and G-6-PDase (b, e, and h) and RET (c, f, and i) between these labels were not significantly affected by colchicine or cytochalasin D in control neutrophils. However, colchicine (m–o) caused the G-6-PDase to become more uniformly distributed within pregnancy cells and reduced the amount of RET (o). n = 5. x560.
labeled the centrosome. As indicated above, neutrophils from nonpregnant women exhibited peripheral G-6-PDase labeling (Figure 3, c and g). RET enables the detection of molecular proximity (≤ 7 nm) between two suitable fluorescent labels. Since anti-G-6-PDase is more abundant than γ-tubulin, we chose it as the acceptor label (TRITC), thereby optimizing RET. As expected, RET was not detected between these labels on neutrophils from nonpregnant women, using optical imaging (Figure 3d) or spectrophotometry (Figure 3r). Furthermore, LPS (50 ng/ml for 20 minutes) had no effect on these parameters (Figure 3, h and s). When cells from pregnant women were stained with anti-γ-tubulin and anti-G-6-PDase, the FITC/anti-γ-tubulin and TRITC/anti-G-6-PDase patterns overlapped substantially (Figure 3, j with k and n with o). These results indicate that G-6-PDase traffics to the MTOC in maternal cells. However, since immunofluorescence does not demonstrate molecular proximity, we performed RET experiments. RET between these two labels was observed in the absence and presence of LPS, using microscopic imaging (Figure 3, i and p) and spectrophotometry (Figure 3, r and s). Thus, G-6-PDase is translocated to the MTOC and is within approximately 7 nm of γ-tubulin in pregnancy, but not control, neutrophils.

G-6-PDase trafficking to the MTOC suggests that microtubules contribute to its intracellular localization. To examine this possibility, cells were treated with colchicine, which disrupts microtubules. Cells were treated with 50 μg/ml colchicine for 30 minutes at 37°C followed by LPS stimulation (50 ng/ml for 20 minutes). Although colchicine treatment had little effect on cells from nonpregnant women (Figure 4, d–f), colchicine changed the G-6-PDase distribution in cells from pregnant women (Figure 4, n vs. k). Furthermore, colchicine dramatically reduced RET between FITC/anti-γ-tubulin and TRITC/anti-G-6-PDase (Figure 4, o vs. l). These changes were not specific for colchicine, inasmuch as similar changes in G-6-PDase were noted after treatment of pregnancy cells with the microtubule-disrupting drugs vincristine (5 μM for 40 minutes at 37°C; n = 4) and vinblastine (80 nM for 40 minutes at 37°C; n = 3) (data not shown). However, the drug taxol (30 μM for 40 minutes at 37°C), which stabilizes microtubules, had no effect on pregnancy neutrophils. To determine whether these changes were specific for microtubules, we tested the ability of the microfilament-disrupting drug cytochalasin D to influence the intracellular trafficking of G-6-PDase. Cytochalasin D was chosen over cytochalasin B as the latter can interact with the glucose transporter, thereby potentially altering metabolism via an alternate route. Neutrophils from nonpregnant and pregnant women were incubated with 50 μg/ml cytochalasin D for 30 minutes at 37°C, conditions known to cause microfilament disassembly. In contrast to colchicine, cytochalasin D had no effect on the intracellular distribution of G-6-PDase in maternal neutrophils (Figure 4, p–r). Therefore, microtubules appear to play an important role in maintaining the intracellular distribution of G-6-PDase in cells from pregnant women.

Altered cytochemical localization of G-6-PDase activity in neutrophils of pregnant women. Our immunofluorescence microscopy experiments demonstrate a dramatic translocation of the G-6-PDase antigen in pregnancy neutrophils. To confirm the implicit assumption that distribution of G-6-PDase activity parallels its antigenic localization, we have employed cytochemical techniques to map G-6-PDase activity in cells (33). Figure 5 shows optical micrographs of neutrophils from pregnant and nonpregnant individuals, stained for G-6-PDase activity. In cells from nonpregnant women, G-6-PDase reaction product was predominantly found about a cell’s periphery (Figure 5,

**Figure 5**
Localization of G-6-PDase activity in neutrophils. Representative cells from nonpregnant women (a and b) and pregnant women (c and d) are shown. Cells were incubated in the absence (a and c) and presence (b and d) of LPS (50 ng/ml, 20 minutes). Cells were fixed and processed as described previously (22, 41). G-6-PDase activity is peripherally located in control cells (a and b) but was found to be enhanced near the centrosome in cells from pregnant women (c and d). n = 4. ×640.

**Figure 6**
HMS activity in cells from nonpregnant and pregnant women with and without treatment with 50 ng/ml LPS. The metabolism of 1-14C-glucose and 6-14C-glucose to 14CO2 was measured as described previously (39, 40). The cpm of 14CO2 formed from 6-14C-glucose was subtracted from the cpm of that formed from 1-14C-glucose. The cpm (HMS activity) is listed on the graph (mean ± SEM). LPS substantially increases HMS activity in cells from nonpregnant individuals (P < 0.001, paired two-tailed t test) but not from pregnant individuals (P > 0.01). n = 7.
a and b), as anticipated by the distribution of G-6-PDase antigen (Figure 2, e and f; Figure 3, c and g; and Figure 4b). G-6-PDase activity remained in a peripheral distribution in cells incubated with 50 ng/ml LPS for 40 minutes at 37°C (Figure 5b). However, G-6-PDase activity was centrally located in cells from pregnant women (Figure 5, c and d) with or without treatment with LPS. These findings are consistent with the localization of G-6-PDase antigen (Figure 2, g and h; Figure 3, k and o; and Figure 4k). The staining patterns were not due to a nonspecific effect. Staining could not be observed during control experiments that omitted substrate (G-6-P and NADP+). Thus, both structural and functional assays indicate a dramatic reorganization of G-6-PDase in pregnancy neutrophils. These findings also illustrate that G-6-PDase remains active after translocation to the centrosomal region. In situ kinetic quantitative microphotometry studies indicate that the rates of product deposition in these experiments do not differ in maternal and control cells (data not shown). Thus, a

**Figure 7**

Representative NAD(P)H oscillations in adherent neutrophils from nonpregnant (a–e) and pregnant (f–j) women. Cells were untreated (a and f) or treated with LPS (b and g), LPS plus 6-AN (c and h), LPS plus colchicine (d and i), or LPS plus colchicine and 6-AN (e and j). The amplitudes of the NAD(P)H oscillations are higher in pregnancy cells (a and f). LPS causes higher-frequency oscillations in cells from nonpregnant women (b), but not in cells from pregnant women (g), which appears to be due to an inability to properly engage the HMS, as suggested by the inhibitor 6-AN (c and h). However, colchicine is able to restore the normal high-frequency oscillations to pregnancy cells (i) in a 6-AN–sensitive manner, suggesting that the ability to engage normal NAD(P)H flux has been restored. See Table 1 for quantitative results. Bar = 10 seconds.

### Table 1

Quantitative analysis of metabolic oscillation properties of neutrophil populations from nonpregnant and pregnant women

<table>
<thead>
<tr>
<th>Neutrophils from pregnant women</th>
<th>Treatment</th>
<th>NAD(P)H properties</th>
<th>ROM release properties</th>
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</thead>
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<tr>
<td></td>
<td></td>
<td>HA/HF</td>
<td>HA/LF</td>
<td>LA/HF</td>
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<tr>
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<td>0</td>
<td>74 ± 3</td>
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<td>12 ± 4</td>
<td>61 ± 5</td>
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<td>0</td>
<td>60 ± 6</td>
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<td>6-AN</td>
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<td>0</td>
<td>75 ± 5</td>
<td>0</td>
</tr>
<tr>
<td>IFN</td>
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<td>89 ± 2</td>
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<td>62 ± 5</td>
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<th>Treatment</th>
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<td>PMA</td>
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<td>6-AN</td>
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<td>99 ± 1</td>
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<td>99 ± 1</td>
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The percentage of cells in each of the categories is shown. One hundred cells were measured for each treatment condition for each patient sample. The categories characterize the NAD(P)H oscillations or the time-dependence of ROM production as high amplitude/high frequency (HA/HF), high amplitude/low frequency (HA/LF), low amplitude/high frequency (LA/HF), and low amplitude/low frequency (LA/LF).
G-6-PDase remains functional during pregnancy, but its activity is localized to the centrosome.

Effect of pregnancy on HMS activity. Although the cytochemical findings clearly demonstrate the location of G-6-PDase activity in neutrophils, this does not necessarily reflect the HMS's activity in living cells. To ascertain HMS activity, we measured the formation of 14CO2 from 1-14C-glucose and 6-14C-glucose in the presence and absence of 50 ng/ml LPS in control and maternal cells. Subtracting the amount of 14CO2 formed during incubation with 6-14C-glucose from that formed in parallel experiments from 1-14C-glucose provides an indication of HMS activity (31). Figure 6 shows the HMS activity in leukocytes from pregnant and nonpregnant women in the presence and absence of LPS (50 ng/ml). A substantial level of HMS activity is present in unstimulated cells from pregnant and nonpregnant women. As previously reported by others (31), cells from nonpregnant individuals undergo a dramatic increase in HMS activity in response to stimuli activating the respiratory burst (Figure 6). However, no significant change in HMS activity was noted in samples from pregnant women incubated with LPS (Figure 6). Thus, unstimulated neutrophils from pregnant women possess HMS activity, but this activity is not upregulated by LPS.

Effect of pregnancy on oscillatory metabolic flux in neutrophils. Previous studies have shown that neutrophil metabolism oscillates in time and that these oscillations correlate with cell functions, including ROM production (26). Therefore, we sought to provide direct evidence that the dynamics of cell metabolism is affected during pregnancy. To do this, we measured the time dependence of NAD(P)H production, as judged by NAD(P)H autofluorescence, in adherent neutrophils. As previously reported (26, 29), polarized control neutrophils exhibit NAD(P)H oscillations with a period of about 20 seconds (Figure 7a). LPS stimulation decreases the oscillatory period to about 10 seconds (Figure 7b). 6-AN, an inhibitor of G-6-PDase (24), blocks the LPS-mediated enhancement of NAD(P)H oscillations (Figure 7c). This suggests that the 10-second metabolic period is associated with HMS activation. However, colchicine had no effect on the metabolic flux in control neutrophils (Figure 7d). In contrast to the effects of LPS, PMA treatment increased the amplitude of metabolic oscillations (Table 1). Thus, priming and/or activating stimuli can affect metabolism via multiple pathways.

When cells from normal pregnancy were examined, they demonstrated oscillations with a 20-second period, as did those from nonpregnant women. However, in contrast to control cells, about 70% of the pregnancy neutrophils demonstrated high-amplitude oscillations (Table 1), as if they had been primed by IFN-γ (34). Moreover, LPS and 6-AN had no effect on maternal neutrophils (Figure 7, f–h). However, PMA treatment did increase the proportion of pregnancy cells that exhibited high-amplitude oscillations from 74% to 98% (Table 1). When neutrophils of pregnant women were treated with colchicine (50 µg/ml for 30 minutes at 37°C), 100% of the cells acquired the approximately 10-second oscillations in response to LPS (Figure 7i) concomitant
The potential role of metabolic enzyme trafficking in ROM release is illustrated. Conditions associated with resting (a), activated (b), and pregnancy (c) neutrophils are shown. ROMs are produced during all of these conditions, but ROM production is maximal for activated cells from nonpregnant women. Hexokinase and G-6-PDase have peripheral distributions in activated cells from nonpregnant women, thus allowing maximal cooperation in ROM release. m.t., microtubule; m.f., microfilament.

Figure 10

The neutrophil’s respiratory burst is enhanced by IFN-γ (39). We have previously shown that IFN-γ promotes high-amplitude NAD(P)H oscillations in adherent cells, resulting in significant levels of NO (34) and superoxide release (our unpublished observations). Figure 8a, trace 3, shows that incubation with IFN-γ (10 µg/ml) for 2 hours at 37°C causes increased ROM release from adherent neutrophils. A similar response was found after PMA treatment (30 nM for 30 minutes) (Table 1). It should be noted that, in comparing traces 2 and 3, trace 2 rises at a higher frequency, whereas trace 3 increases with a greater step size or amplitude; both differences reflect the NAD(P)H oscillations (Figure 7). When combined, IFN-γ and LPS yield maximal ROM production (Figure 8a, trace 5). Further quantitative data are given in Table 1.

Figure 8b demonstrates that extracellular ROM release by adherent/polarized maternal neutrophils is not increased by LPS (trace 2) in comparison with untreated maternal cells (trace 1). Nonetheless, the basal ROM release rate is higher in adherent pregnancy neutrophils than in neutrophils from nonpregnant women (compare Figure 8a, trace 1, with Figure 8b, trace 1). A key difference between Figure 8a and Figure 8b is that the “step” frequency doubles when LPS is added to control neutrophils, but not when it is added to pregnancy neutrophils, which is consistent with the lack of 10-second-period NAD(P)H oscillations in pregnancy neutrophils (Figure 7g). The percentage of pregnancy neutrophils producing high-amplitude ROM release patterns increased from 74% to 98% after PMA treatment. In contrast to control neutrophils (Figure 8a, traces 3 and 6), ROM production by pregnancy neutrophils is not sensitive to 6-AN (Figure 8b, trace 4). Thus, the basal levels of unstimulated extracellular ROM release by adherent/polarized cells are enhanced in pregnancy to a level comparable to those in LPS-stimulated cells from nonpregnant individuals. However, the fully activated levels brought about by incubation with LPS plus IFN-γ in cells from nonpregnant individuals (Figure 8a, trace 5) cannot be achieved in pregnancy cells (Figure 8b, trace 5). LPS, IFN-γ, LPS plus IFN-γ, 6-AN, and 6-AN plus LPS did not alter ROM release from pregnancy cells in comparison with untreated cells. Hence, important changes in ROM production accompany pregnancy.

Microtubule-disrupting drugs rescue maximal ROM release from pregnancy neutrophils. Since colchicine normalizes with cytoplasmic randomization of G-6-PDase (Figure 4n). The metabolic oscillations of maternal neutrophils were similarly affected by vincristine and vinblastine (data not shown; n = 3). When control cells were incubated with LPS, colchicine, and 6-AN, they matched the 20-second oscillation periods of pregnancy neutrophils (Figure 7, e and j). Hence, neutrophils of pregnant women are simultaneously primed in the sense that the metabolic oscillations are higher in amplitude, but inhibited in that they cannot acquire the approximately 10-second metabolic period. Moreover, microtubules and the associated retrograde trafficking of G-6-PDase appear to be important in controlling metabolic flux.

Extracellular ROM release from neutrophils of pregnant and nonpregnant women. Given these alterations in G-6-PDase location and metabolism, we tested examined ROM production by neutrophils from pregnant and nonpregnant women. ROM production was monitored using H2TMRos, which becomes fluorescent upon exposure to ROMs. To distinguish between the extracellular release of ROMs and the intracellular accumulation of ROMs, probes were incorporated into either the extracellular environment or the cellular cytoplasm using previously established methods (35, 36). Figure 8 shows representative ROM release experiments on neutrophils migrating in an ECM containing H2TMRos. This single-cell ROM release assay using adherent cells in an ECM-like setting approximates the interstitial environment more closely than do cells in suspension. Furthermore, the phenotype of adherent neutrophils is distinct from that of suspended cells. For example, adherent cells produce enhanced quantities of ROMs (37). Adherent or polarized cells release low levels of ROMs (Figure 8a, trace 1) (35, 38). LPS, which activates the HMS and ROM release, enhanced ROM release from control neutrophils (Figure 8a, trace 2). However, LPS-mediated enhancement of ROM production is blocked by 6-AN (Figure 8a, trace 6), although 6-AN alone has no effect on basal levels of ROM release (Figure 8a, trace 4). Thus, LPS-mediated activation of ROM release from control neutrophils depends upon G-6-PDase.
the intracellular distribution of G-6-PDase and the LPS-stimulated metabolic dynamics of pregnancy cells, we tested the ability of colchicine to normalize ROM release from maternal neutrophils. We first tested the effects of colchicine on control neutrophils. When cells from nonpregnant women were treated with colchicine (50 μg/ml for 30 minutes at 37°C), no changes in ROM production were observed relative to that in buffer-treated cells (Figure 8c, trace 7 vs. trace 1). LPS-stimulated ROM release was not significantly affected by colchicine (Figure 8c, trace 8, vs. Figure 8a, trace 2), although 6-AN dramatically decreased ROM release from cells treated with LPS and colchicine (Figure 8c, trace 9). The higher basal rate of ROM release from pregnancy cells was not affected by colchicine (Figure 8d, traces 1 and 7). In contrast to control neutrophils, colchicine dramatically increased ROM release from LPS-stimulated pregnancy neutrophils (Figure 8d, trace 8). Enhanced ROM production is inhibited by 6-AN (Figure 8d, trace 9). Parallel results were obtained using vincristine (n = 3) and vinblastine (n = 4), but not taxol (n = 3), as described above (data not shown). Thus, microtubules restrain adherent/polarized maternal neutrophils from mounting a complete oxidative program in response to activating stimuli. We suggest that G-6-PDase trafficking influences H2O2 activation and ROM release.

Intracellular accumulation of ROMs within neutrophils of pregnant and nonpregnant women. ROM release into the extracellular environment is not necessarily the same as intracellular ROM accumulation. To develop a more comprehensive analysis of ROM responses during pregnancy, we next evaluated the intracellular oxidation of the dye H2TMRos in neutrophils from nonpregnant and pregnant women. H2TMRos-labeled neutrophils were incubated with buffer alone, LPS (50 ng/ml for 20 minutes), or IFN-γ (10 μg/ml for 2 hours at 37°C) followed by addition of LPS (50 ng/ml for 20 minutes). Figure 9a and b, shows the fluorescence emission spectra of TMRos-labeled neutrophils from nonpregnant and pregnant women, respectively. The peak height is the total amount of TMRos formed within cells. The amount of intracellular TMRos was very small for unstimulated neutrophils from nonpregnant women but was markedly increased by the addition of LPS or of LPS plus IFN-γ (Figure 9a). Neutrophils from pregnant women, however, did not respond with a significant change in internal ROM production. Nonetheless, the internal ROM accumulation in unstimulated cells from pregnant women was much higher than that in unstimulated neutrophils from nonpregnant women. The lack of response in maternal neutrophils was not due to saturation of the assay, since the addition of exogenous H2O2 caused the fluorescence level to rise dramatically (data not shown). Thus, the levels of external release and intracellular accumulation of ROM products from adherent/polarized neutrophils from pregnant women are comparable to the levels found for LPS-stimulated neutrophils from nonpregnant individuals. However, these levels cannot be enhanced by IFN-γ as they can be in cells from nonpregnant women.

Discussion
Neutrophil activation is associated with translocation of NADPH oxidase components, granules, protein kinase C, and hexokinase to membranes (23, 40, 41). We now propose a previously unknown regulatory step in neutrophil activation, retrograde translocation of G-6-PDase, which is only manifest during pregnancy. We suggest that G-6-PDase translocation is mediated by microtubules and that this redistribution accounts for diminished ROM release from maternal cells.

Several metabolic enzymes, including hexokinase, aldolase, PFK, GAPDH, and PK, bind to microfilaments and/or microtubules (23, 42–47). These interactions can alter both metabolic activity and cytoskeletal properties. Hexokinase translocates to the cell periphery during macrophage activation, where it increases the glucose influx rate (22, 23) and provides the H2O2 with G-6-P. The trafficking of metabolic enzymes to various cellular sites led to the concept of microcompartmentation of cell metabolism (42–47). One example of metabolic microcompartmentation is the preferential use of glycolysis as an energy source for plasma membrane functions (48–52). Although metabolic compartmentation may enhance certain cell functions, there are no unambiguous examples where different physiological states correspond to distinct compartmentation strategies. We suggest that G-6-PDase translocation in neutrophils from pregnant women constitutes a compartmentation switch regulating H2O2 performance.

We have identified what we believe to be a novel and fundamental difference in metabolic enzyme trafficking in neutrophils from pregnant women. Our studies have shown that G-6-PDase and G-6-PDase activity are peripherally located in control neutrophils but are located primarily in the centrosomal region of pregnancy neutrophils. This pregnancy-associated change in G-6-PDase distribution appears to be specific, as no significant differences were found for hexokinase, PFK, PK, and LDH. G-6-PDase often appeared to decorate the centrioles of stimulated cells. The apparent retrograde transport of G-6-PDase was confirmed using anti-γ-tubulin antibodies, which are specific for MTOCs (53). Colocalization and RET microscopy studies showed that G-6-PDase translocated to MTOCs in pregnancy, but not control, neutrophils.

We propose a simple model: ROM release is maximal when G-6-PDase traffics to the cell periphery, where its substrate G-6-P is readily available from hexokinase, which is also peripherally located in activated cells (Figure 10). The presence of G-6-PDase allows the H2O2 to be fully engaged with the local production of NADPH, the NADPH oxidase’s preferred substrate. Consequently, the NADPH oxidase is provided with maximal substrate levels to yield high levels of ROM release. On the other hand, during pregnancy, G-6-PDase traffics away from the cell periphery, thereby limiting H2O2 efficiency by allowing glycolytic conversion of G-6-P by phosphoglucose isomerase and PFK. This is reasonable since PFK’s conversion of F-6-P is irreversible and PFK maintains a
peripheral distribution in pregnancy cells (Figure 2, k and l). Perimembrane levels of NADPH would therefore be lower, making the NADPH oxidase less efficient and reducing ROM release. This hypothetical model accounts for the reduction in ROM release by maternal neutrophils.

Although signal-mediated regulation of NADPH oxidase function has been extensively studied, other mechanisms may also be involved. We suggest that NADPH availability is an important factor in ROM production. We have previously proposed substrate-level control of NADPH oxidase activity (26–29). For example, the intracellular production of NAD(P)H and the extracellular release of superoxide anions oscillate in rhythm with one another with the same frequencies and phases and respond in parallel to various perturbations. The communication between these and other oscillators in activated neutrophils has been thoroughly analyzed using appropriate mathematics and computer simulations (29). The correspondence between local NAD(P)H concentration and local superoxide release from polarized neutrophils has been shown using ultrafast microscopy and microspectrophotometry (28). The present study extends our work on substrate-level control of ROM release to include G-6-PDase translocation as a means of regulating local substrate levels by metabolic microcompartmentation.

If retrograde transport of G-6-PDase affects ROM release during pregnancy, it should be possible to alter the intracellular distribution of G-6-PDase with microtubule-disrupting drugs, thereby increasing ROM production. Indeed, we have found that colchicine, vincristine, and vinblastine normalize the properties of pregnancy neutrophils. Treatment of pregnancy neutrophils with these drugs causes G-6-PDase to become more uniformly distributed, brings about the return of rapid NAD(P)H oscillations in response to LPS, and restores robust ROM release. However, the drug taxol, which has the opposite effect on microtubules, did not influence the properties of pregnancy neutrophils. This supports the idea that the intracellular distribution of G-6-PDase is an important determinant of cell metabolism and ROM production.

Our studies have shown that the HMS is intact in leukocytes from pregnant women, but that it cannot be upregulated in response to activating stimuli. In contrast, under identical conditions, cells from nonpregnant women show a roughly threefold increase in HMS activity. Parallel changes in metabolic oscillations, which represent the dynamic flux of the NAD(P)H-NADP⁺ redox couple, were observed. Pregnancy cells demonstrated higher-amplitude oscillations than did cells of nonpregnant individuals. However, pregnancy neutrophils were unable to upregulate NAD(P)H oscillation frequency, which is closely tied to ROM production. These oscillations are likely important because (a) they encode information provided by the signal transduction apparatus, which is eventually decoded by the NADPH oxidase and NO synthase; (b) the delivery of electrons to the oxidase is compressed in time, thereby allowing short and intense bursts of activity; (c) they increase thermodynamic efficiency (54); and (d) they contribute to protecting the neutrophil from self-inflicted damage (55). Thus, pregnancy cells have HMS activity and metabolic oscillations, but neither can be upregulated in response to stimuli.

Although there are reports that superoxide release is decreased in pregnancy neutrophils (2–4), there are also reports that ROM production is increased in these cells (56). Studies reporting a decrease in ROM production measure the extracellular release of oxidants, while those reporting an increase in ROM production use an intracellular dye to measure ROM production. Our findings are consistent with several features of these studies, although our studies were conducted on adherent cells, not cells suspended in buffer. (The cytoskeletal reorganization associated with cell adherence may sufficiently redistribute metabolic enzymes to enhance ROM kinetics.) Our findings, like those of others (2–4), show that maternal neutrophils cannot be activated for extracellular ROM release as vigorously as neutrophils from nonpregnant individuals. We have also shown that cells from pregnant women have greater unstimulated levels of intracellular ROM accumulation than do controls. The ROM production rate of LPS-stimulated controls is similar to that of unstimulated pregnancy cells, as in other studies (56). Thus, whether pregnancy neutrophils are activated depends more on the definition of “activated” and the comparison being made than it does on the biology of the system.

The mechanism underlying retrograde transport of G-6-PDase in neutrophils during pregnancy is unknown. For example, G-6-PDase may be associated with a granule or supramolecular enzyme complex that undergoes differential translocation during nonpregnant and pregnant states. Retrograde transport of G-6-PDase apparently accounts for blunted ROM release in cells from pregnant women. This, in turn, may provide a partial explanation for the reduction in host resistance to infectious agents during pregnancy. Although retrograde G-6-PDase transport may diminish extracellular ROM release, it may enhance intracellular killing of certain microbes, since endosomes undergo retrograde transport as well. G-6-PDase translocation may affect other aspects of host defense and autoimmunity in pregnancy. For example, G-6-PDase is the rate-controlling step in ribose 5-phosphate synthesis, which is required for cell proliferation (57). Thus, the mechanism proposed might be more broadly involved in the control of immune responses. We are presently studying the mechanisms regulating G-6-PDase trafficking, with the long-range goal of improving disease management.

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