Tetrahydrobiopterin, a Critical Factor in the Production and Role of Nitric Oxide in Mast Cells

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Mast cells (MC) are biologically potent, ubiquitously distributed immune cells with fundamental roles in host integrity and disease. MC diversity and function is regulated by exogenous nitric oxide; however, the production and function of endogenously produced NO in MC is enigmatic. We used rat peritoneal MC (PMC) as an in vivo model to examine intracellular NO production. Live cell confocal analysis of PMC using the NO-sensitive probe diaminofluorescein showed distinct patterns of intracellular NO formation with either antigen (Ag)/IgE (short term) or interferon-γ (IFN-γ) (long term). Ag/IgE-induced NO production is preceded by increased intracellular Ca\(^{2+}\), implying constitutive nitric-oxide synthase (NOS) activity. NO formation inhibits MC degranulation. NOS has obligate requirements for tetrahydrobiopterin (BH\(_4\)), a product of GTP-cyclohydrolase I (CHI), IFN-γ-stimulated PMC increased CHI mRNA, protein, and enzymatic activity, while decreasing CHI feedback regulatory protein mRNA, causing sustained NO production. Treatment with the CHI inhibitor, 2,4-diamino-6-hydroxypyrimidine, inhibited NO in both IFN-γ and Ag/IgE systems, increasing MC degranulation. Reconstitution with the exogenous BH\(_4\) substrate, sepiapterin, restored NO formation and inhibited exocytosis. Thus, Ag/IgE and IFN-γ induced intracellular NO plays a key role in MC mediator release, and alterations in NOS activity via BH\(_4\) availability may be critical to the heterogeneous responsiveness of MC.

Mast cells are immune effector cells located at strategic tissue sites, juxtaposed to vessels, epithelium, and nerves. These phenotypic and functionally heterogeneous cells release granule-associated mediators (e.g. histamine) and various newly formed molecules (e.g. phospholipid metabolites and cytokines) that play roles in innate and adaptive immune responses (1). Classically, MC\(^1\) can be activated via cross-linking of Fce-bound IgE molecules by specific antigen (Ag), and inhibition of MC activation is a key component in controlling allergic diseases such as asthma (2). The Th1 cytokine interferon-γ (IFN-γ) inhibits MC function, and dysregulation of IFN-γ contributes to atopy and asthma (2). IFN-γ modulation of cell phenotype is complex and involves regulation of gene expression (3) including the enzyme GTP-cyclohydrolase I (CHI), which produces tetrahydrobiopterin (BH\(_4\)) (4).

BH\(_4\) is a critical factor in the production of the neurotransmitters serotonin, catecholamines, and other cellular activities including proliferation, cell cycle regulation, and differentiation (4). The majority of cellular BH\(_4\) is synthesized de novo from GTP with CHI being rate-limiting (4). CHI activity is regulated at multiple levels, including transcription and phosphorylation, and can be inhibited by the CHI feedback regulatory protein (GFRP) (5, 6). BH\(_4\) is also an essential co-factor in NO formation from nitric-oxide synthase (NOS), and increases in NOS transcription and activity need to be accompanied by a concurrent increase in BH\(_4\) to sustain NO production (7).

NO is a reactive radical with pleotropic effects with both physiological and pathological functions (8). NO is formed from \(\text{L-arginine} \sim \text{BH}_4\) in the NO synthesis pathway (9), regulated by exogenous NO (9). We have recently shown that rat peritoneal MC (PMC) also constitutively express eNOS and can up-regulate iNOS and produce NO (10). More recent in vivo studies have implicated MC as a significant source of NO in vascular tissues, likely through nNOS (11). MC produce low amounts of NO, implying a paracrine, intracellular function. However, the roles for MC-derived NO remain poorly defined. Furthermore, our previous study showed that diverse stimuli induced differing levels of NO release from MC, despite similar induction of NOS (10). Clearly, other regulatory events are responsible for differences in the time course, quantities, and outcomes of MC-derived NO (12). Because no data are available on the role that BH\(_4\) plays in NO production by MC, we hypothesized that post-transcriptional regulation of NOS activity by BH\(_4\) is a critical determinant in NO production by MC.

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\(\text{BH}_4\), tetrahydrobiopterin; GFRP, CHI feedback regulatory protein; NOS, nitric-oxide synthase; eNOS, endothelial NOS; nNOS, neuronal NOS; iNOS, inducible NOS; PMC, peritoneal MC; eNOS, constitutive NOS; DAF, diaminofluorescein; \(\text{L-NAME}\), \(\text{N}^\text{N}\)-nitro-L-arginine methyl ester; DAHP, 2,4-diamino-6-hydroxypyrimidine; DAF-FM, 4,5-diaminofluorescein diacetate; WE, worm equivalents; RT, reverse transcription; HPLC, high pressure liquid chromatography; \(\beta\)-hex, \(\beta\)-hexosaminidase.
The purpose of this study was 2-fold. First, to characterize both short term (<5 min) Ca\(^{2+}\)-dependent constitutive NOS (cNOS) activity and sustained NO production (>18 h) in MC using the fluorescent NO-sensitive probe diaminofluorescein (DAF). Second, to investigate the effects of CHI activity and BH\(_2\)s levels in both short term (<5 min) and sustained (>18 h) NO production on MC reactivity. We demonstrate for the first time the contribution of rat MC to spontaneously activated intracellular Ca\(^{2+}\)-dependent NOS activity (<5 min) after Ag stimulation, which inhibits degranulation. Furthermore, IFN-γ treatment enhances long term (>18 h) intracellular NO production, which also inhibits MC degranulation. Constitutively expressing CHI mRNA and protein, and upon stimulation with IFN-γ, PMC up-regulate CHI mRNA and protein with a concordant increase in CHI activity. IFN-γ also decreases GFRP mRNA levels. The final outcome of these events is an increased level of NO production after IFN-γ treatment. Thus, modulation of MC BH\(_2\)s levels controls NO formation and influences MC degranulation.

**EXPERIMENTAL PROCEDURES**

**Animals**—Adult male Sprague-Dawley (Crl:CD (SD) BR) rats were obtained from Charles River Canada Inc. (Quebec, Canada). For experiments requiring antigen stimulation, the rats were sensitized with L\(_{31}\) larvae of *Nippostrongylus brasiliensis* 5–6 weeks before MC isolation (10). The experimental procedures were approved by the University Animal Care Committee and were in accordance with the guidelines of the Canadian Council on Animal Care.

**Chemicals**—The NOS inhibitor N\(^5\)-nitro-l-arginine methyl ester (l-NAME), NO donor S-nitrosoglutathione, and CHI inhibitor 2,4-diamino-6-hydroxypyrimidine (DAHP) were obtained from Calbiochem (San Diego, CA). The sepiapterin reductase inhibitor N-acetyl serotonin was obtained from Sigma (St. Louis, MO), and the NO probe 4,5-diaminofluorescein diacetate (DAF-FM) and Ca\(^{2+}\)-sensitive dye FURA-2 were obtained from Molecular Probes (Eugene, OR).

**Antigen**—Ag used to activate in vivo sensitized PMC was a collection of soluble excratory and secretory products of the nematode *N. brasilensis*, prepared as previously described (13). The antigen concentration was described as worm equivalents (WE/ml).

**MC Isolation and Stimulation**—PMC were obtained by peritoneal lavage followed by centrifugation through gradient Percoll as previously described (10). PMC purity was >98% as determined by staining with toluidine blue. Cell viability was >99%. For most experiments PMC from unsensitized rats were treated with medium containing IFN-γ (200 units/ml) (Invitrogen). For studies of MC secretion, PMC from *N. brasiliensis* sensitized rats were treated with Ag (5 WE/ml). *Live Cell Fluorescence Determination of Intracellular NO and Ca\(^{2+}\) Production*—NO production by PMC was assayed using DAF-FM, a cell-permeable NO-sensitive fluorescent dye (14). Coverslip-bottomed Petri dishes (Falcon) were coated with rat plasma fibronectin (10 \(\mu\)g/ml) (Sigma) in phosphate-buffered saline, pH 7.2, for 1 h at 37 °C. PMC were loaded with 10 \(\mu\)M of DAF-FM. For Ca\(^{2+}\) experiments, PMC were loaded with FURA-2 (5 \(\mu\)M) and then incubated for 1 h at 37 °C in the dark. The cells were resuspended in RPMI 1640 without phenol red and placed in dishes and then incubated for 1 h at 37 °C before use.

The cell images were obtained using a Zeiss confocal laser scanning microscope (LSM510) using a 488-nm (excitation) and 530-nm (emission) filter set for DAF and a 340-nm (excitation) and 505-nm (emission) filter set for FURA-2. The optimized annealing temperature was 48 °C for GFRP. The products were run on a 1.2% agarose gel and stained with ethidium bromide (Sigma).

**Cloning and Sequencing of cDNA Bands**—The amplified PCR products were subcloned into pCR1\(^{TM}\) plasmid vector using the TA cloning kit (Invitrogen). Plasmid DNA was isolated with the GenElute\textsuperscript{TM} Plasmid isolation kit (Sigma). Double-stranded DNA sequencing was performed using M13 forward and reverse primers. Sequencing was conducted using an ABI 373A automated sequencer (Applied Biosystems, Foster City, CA) by a dyeoxy chain termination method.

**Western Blot Analysis**—PMC were incubated in 24-well plates at 1 \(\times\) 10\(^6\) cells/well for 0–18 h in various experimental conditions. The cells were dissociated in RIPA buffer (phosphate-buffered saline, 1% Nonidet P-40). The total protein content was determined by the Bradford technique (Bio-Rad). Fifteen \(\mu\)g of protein from each sample was mixed with 1/500 (1 \(\mu\)l) and 1/5000 in horseradish peroxidase-conjugated goat anti-rat IgG (Sorotec, Raleigh, NC). Labeling was detected by chemiluminescence with SuperSignal substrate solution (Pierce). The resulting bands were scanned and quantified in a gel scanner (ImageMaster DTS; Amersham Biosciences).

**Immunofluorescence Detection of CHI**—Localization of CHI in PMC was performed on PMC fixed in 4% paraformaldehyde for 20 min and then permeabilized with 0.1% Triton X-100 in phosphate-buffered saline. The cells were blocked with 1% goat serum and 3% bovine serum albumin in phosphate-buffered saline for 30 min with 10 \(\mu\)M of YC2 and then incubated overnight at 4 °C with 1 µg/ml of monoclonal anti-CHI (6H11) (L. Ziegler, Munich, Germany) (5) diluted 1/5000 (1 \(\mu\)g/ml) and 1/5000 in horseradish peroxidase-conjugated goat anti-rat IgG (Sorotec, Raleigh, NC). The samples were separated on a 12% SDS-PAGE gel and transferred onto a polyvinylidene difluoride membrane (Bio-Rad). CHI was identified with rat monoclonal anti-CHI (6H11) (L. Ziegler, Munich, Germany) (5) diluted 1/5000 (1 \(\mu\)g/ml) and 1/5000 in horseradish peroxidase-conjugated goat anti-rat IgG (Sorotec, Raleigh, NC). Negative controls labeled with isotype (rat IgG1) antibody were run concurrently.

**Enzyme Assay for CHI Activity**—The activity of CHI was determined on homogenized cell extracts as previously described (5). Briefly, the reaction product was oxidized to neopterin triphosphate by acidic iodine solution. After reduction of excess iodine by ascorbic acid, the sample was immediately separated by ion pair reverse phase HPLC. Neopterin was detected by monitoring the excretion of the product, neopterin, which elutes at 6 min.

**Measurement of NO\(_2\)/NO\(_3\) Production**—NO\(_2\)/NO\(_3\) in culture (phenol red-free) supernatants was measured by the Griess reaction (10).

**β-Hexosaminidase Assay**—β-Hexosaminidase (β-hex) assay was performed using a bioluminescence assay. For the enzyme assay, 100 µl of each sample was incubated at 37 °C for 1 h and then incubated for 30 min with 10 µM of β-N-acetylgalactosamine (βGalNAc) to determine the baseline activity. The resulting activity was expressed as units per milligram protein (U/mg) and was determined by a standard titration curve.

**Enzyme Assay for β-Hexosaminidase**—For enzyme assay, the sample was incubated for 30 min at 37 °C with 3 units of β-hexosaminidase (Corning). The reaction was terminated by adding 3 ml of ice-cold trichloroacetic acid (TCA) and the resulting enzyme activity was determined fluorometrically at 450 nm.

**Statistical Analysis**—All of the experiments were performed at least three times. The data were analyzed using analysis of variance followed by the Bonferroni test for comparisons. p values < 0.01 were considered significant.

**On-line Supplemental Material**—Real-time images of degranulation and NO production in Ag/IgE-stimulated MC are shown in the on-line supplemental video. The video was obtained as described above, with 1-s exposures every 5 s. The video was compressed 10 times using Adobe Premiere (Adobe Systems Canada, Ottawa, Canada). The images were captured using a 40× objective on a Leica confocal laser scanning microscope. The video shows a flux of green DAF fluorescence in a
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RESULTS

Sustained (>18 h) Intracellular Production of NO in IFN-γ-treated PMC—To confirm that IFN-γ treated PMC produced NO, we utilized the NO-specific fluorescent molecule DAF. Confocal microscopic analysis showed that IFN-γ caused an increase in intracellular DAF fluorescence after 18 h of treatment compared with controls. The DAF fluorescence localized to cytoplasmic and occasionally to perinuclear sites (Fig. 1A).

Real Time Confocal Analysis of IgE-mediated Degranulation and NO Production—Next we wanted to assess constitutive NOS activity because there is evidence that MC possess cNOS activity (19). To study this short term (<5 min) NOS activity, its modulation, and association with degranulation, we developed a real time assay to detect DAF fluorescence and analyze dynamic changes in NO production by individual cells. PMC were isolated and loaded with DAF (10 μM). A large proportion of unstimulated PMC showed weak DAF fluorescence, with an occasional cell (~2%) having stronger cytoplasmic fluorescence. PMC showed no visible signs of degranulation, because their membranes were intact and showed no signs of granule release as noted in other MC live cell studies (16).

Stimulation with Ag (5 WE/ml) caused an increase in the number of degranulating cells. Degranulating PMC did not shown any increases in DAF fluorescence (Fig 1B and the supplemental video). Interestingly a proportion (33.1 ± 2.2%, mean ± S.E. of 116 total cells in eight different experiments) of PMC showed an immediate (~<2 min) increase in NO production (DAF fluorescence) and no visible signs of degranulation. DAF fluorescence accumulated in the cytoplasm of these cells, and a population of strong NO producing cells showed intense nuclear positivity (Fig. 1B and the supplemental video). Although the strong DAF fluorescence cells showed no sign of degranulation after 60 min of observation, some of the less positive cells eventually did show degranulation. Regression analysis comparing these weaker DAF fluorescent PMC to time of degranulation showed a highly significant correlation (R² = 0.7056, p < 0.001) (Fig. 1C). Equal loading of PMC with DAF was confirmed by adding the NO donor S-nitrosothioglutathione (100 μM); after 20 min almost all PMC showed strong cytoplasmic staining with DAF (data not shown). Interestingly, no nuclear localization of DAF fluorescence was seen after adding this exogenous NO source. Specificity of the effect was determined by pretreating PMC with the NOS inhibitor L-NAME (100 μM); these cells showed no flux of DAF fluorescence (data not shown). These results support the hypothesis that PMC possess cNOS activity that can be stimulated in the short term (~<5 min) in a proportion of cells by IgE cross-linking. Our data also suggest that immediate NO production by PMC causes inhibition of mediator release.

Ca²⁺ Flux and NO Production in PMC—The results above may involve constitutive NOS (eNOS or nNOS) activity in PMC (10). Because both eNOS and nNOS are Ca²⁺ dependent, we wished to determine whether increases in intracellular Ca²⁺ precede the flux of NO seen in Ag/IgE-treated PMC. PMC were isolated and loaded with both DAF (10 μM) and the Ca²⁺-sensitive dye FURA-2 (5 μM). As has been previously shown, upon stimulation with Ag (5 WE/ml), all PMC showed an increase in FURA-2 fluorescence beginning within 30 s (Fig. 1B) (20). DAF fluorescence in strong NO producing cells was delayed by almost 20 s on average compared with FURA-2 (Fig. 1D). However, DAF fluorescence in degranulating MC showed little change (Fig. 1E). Interestingly, although PMC were heterogeneous in their DAF positivity, all cells were homogenous in their Ca²⁺ flux (Fig. 1, D and E). This indicates that Ca²⁺ flux precedes short term NO production in nondegranulating, Ag-treated PMC.

GTP-CHI/GFRP mRNA Expression in PMC—We recently showed that rat PMC treated with IFN-γ could be stimulated to express iNOS mRNA (10); however, no data exist concerning the expression of CHI or GFRP in in vivo derived MC. mRNA production was thus determined using RT-PCR with gene-specific primers. PMC constitutively express significant CHI and GFRP mRNA (Fig. 2). All of the PCR’s were negative when the RT step was eliminated, indicating that there was no contamination from genomic DNA (data not shown). Cloned PCR amplicons were sequenced and showed >98% identity with published sequences by BLAST (NCBI) analysis.

Differential Regulation of GTP-CHI and GFRP mRNA—Because stimuli that increase NOS expression often induce a concordant increase in de novo BH₄ production (7), we looked at CHI and GFRP mRNA regulation by IFN-γ. Total RNA was extracted from unstimulated PMC (>98% pure) and from PMC treated in vitro with IFN-γ (200 units/ml). CHI and GFRP mRNA production was assessed by semiquantitative RT-PCR versus a standard curve constructed with known copy numbers of cloned inserts amplified under identical PCR conditions. Within 2 h following treatment with IFN-γ, the CHI signal in PMC increased ~3-fold. Levels of CHI mRNA reached maximal expression at 6 h and continued to increase to 18 h (Fig. 2A). No change in CHI mRNA was noted in unstimulated PMC at similar time points (data not shown). GFRP mRNA expression from IFN-γ-treated PMC was significantly down-regulated beginning at 2 h and continued until at least 18 h of treatment (Fig. 2B). The results are from PMC RNA obtained from three independent batches of cells.

CHI Protein Expression—To further evaluate whether the actions of IFN-γ on GTP-CHI mRNA levels were extended to protein expression, Western blot analysis was employed. Unstimulated PMC constitutively produced a band of 30 kDa detected by the antibody 6H11 (Fig. 3A). Exposure of PMC to IFN-γ produced a significant increase in CHI protein expression, with maximal stimulation between 6 and 18 h. Again unstimulated PMC cultured during the same time showed no increase in CHI protein expression. Furthermore, PMC stimulated with Ag/IgE also showed no detectable change in protein expression (data not shown).

Immunofluorescence Localization of CHI in Rat MC—Because the intracellular localization of CHI is poorly understood, we used confocal laser scanning microscopy to assess the cellular expression pattern of CHI protein in PMC. Unstimulated PMC from naïve rats showed a clear plasma membrane pattern with some nuclear immunofluorescence in occasional cells. Membrane staining was confirmed by imaging en face 0.5-μm sections through each cell. This stacked series of multiple images was deconvoluted and three-dimensionally reconstructed using Zeiss LSM 510 software. PMC stimulated with IFN-γ, on the other hand, showed a pronounced increase in cytosolic immunofluorescence while maintaining membrane and some nuclear labeling (Fig. 3B).

CHI Enzymatic Activity—To determine the functional consequences of IFN-γ treatment, we determined CHI enzymatic activity, employing a HPLC-based assay as previously described (5). Untreated PMC were compared with cells treated with IFN-γ or Ag/IgE. In unstimulated cells, CHI activity was low (~<5.0 mmol/mg/min) as previously described (5) but was markedly elevated upon treatment with IFN-γ with a significant increase observed at 6 h that continued to at least 18 h (Fig. 4). As expected from Western blot data, no increase in CHI activity was seen after Ag/IgE treatments (data not shown).

Control of PMC NO (>18 h) Production by BH₄—Previously,
FIG. 1. A, fluorescence detection of intracellular NO production in PMC. The cells were incubated with IFN-γ (200 units/ml) for 18 h and then stained with DAF for 20 min. DAF fluorescence (green) was visualized by confocal analysis. Untreated PMC are used as the control. Differential interference contrast (DIC) shows cellular morphology. Original magnification, ×400; bar, 10 μm. The values are representative of four independent experiments. B, real time confocal analysis of Ca²⁺ accumulation and NO production in Ag-stimulated PMC. PMC from sensitized rats were loaded with both DAF (10 μM) and FURA-2 (5 μM) for 1 h. The cells were stimulated with Ag, and images were simultaneously obtained every 5 s from both DAF and FURA-2 channels. The panels show representative DAF (green), FURA-2 (red), and combined images (yellow) obtained before the addition of Ag (t = 0) and at a time point when no further changes in PMC morphology were observed (t = 4 min). White arrowheads indicate degranulating cells, and white arrows indicate nuclear accumulation of DAF in PMC nucleus. Original magnification, ×600; bar, 10 μm. The results are representative of three independent experiments. The complete video is available as supplementary information. C, correlation between NO production (peak DAF fluorescence; arbitrary units) and time to degranulation (seconds) from Ag-stimulated MC that produce NO but later show signs of degranulation. The data shown are from 30 individual MC from five independent experiments. Linear regression analysis was
PMC have been shown to produce NO (10). Unstimulated PMC spontaneously released low levels of NO$_2$ (0.27 ± 0.03 μM). Activation with IFN-γ caused PMC to produce significantly greater (p < 0.01) amounts of NO$_2$ (3.6 ± 0.2 μM). (Fig. 5). To further define the role of CHI and BH$_4$ in MC NO production, we employed the pharmacological inhibitor of CHI (DAHP) (4). The addition of DAHP along with IFN-γ for 24 h inhibited NO production in a dose-dependent manner (Fig. 5). To further confirm that PMC cellular BH$_4$ levels were limiting in NO production, the cells were treated with an exogenous substrate for BH$_4$ production, sepiapterin. Sepiapterin treatment slightly (but not significantly) potentiated NO production in untreated cells and increased nitrite accumulation upon IFN-γ stimulation (4.6 ± 0.3 μM) (Fig. 5). Significant inhibition (p < 0.01) was attained with DAHP concentrations as low as 100 μM, with nitrite levels at or below sham values with concentrations of 500 μM. Because DAHP at high concentrations (500 μM) completely inhibit nitrite formation, there appears to be little contribution via the salvage pathway under these conditions. These results were confirmed using the sepiapterin reductase inhibitor N-acetyl serotonin (data not shown). However, co-administration of DAHP (500 μM) with 200 μM sepiapterin significantly (p < 0.01) reconstituted nitrite production (3.2 ± 0.1 μM) (Fig. 5).

Effects of BH$_4$ Modulation (>18 h) on MC Degranulation—

Given the above results showing that BH$_4$ levels are important in regulating MC NO production, we investigated the effects of IFN-γ and modulation of BH$_4$ on MC degranulation by measuring β-hex release. PMC were treated for 18 h with IFN-γ (200 units/ml), DAHP (500 μM), or sepiapterin (200 μM) and then
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stimulated with Ag (5 WE/ml) for 30 min. As Fig. 6A shows, Ag stimulation induced 21 ± 1.2% release, whereas IFN-γ treatment significantly (p < 0.05) inhibited Ag-induced release by over 40% (11 ± 1.5% release). Pretreatment of PMC with DAHP (500 μM) for 24 h resulted in a significant (p < 0.05) increase in Ag-induced β-hex release (from 21 ± 1.2% to 37 ± 2.3%) and abrogated the inhibitory effect of IFN-γ. Treatment with sepiapterin induced significant (p < 0.05) inhibition of β-hex release after Ag stimulation in IFN-γ and untreated PMC. Furthermore, modulation of BH₄ levels also influenced spontaneous β-hex release (Fig. 6B). PMC spontaneously release about 6% (6.3 ± 0.9%) β-hex. DAHP pretreatment significantly (p < 0.01) increased spontaneous β-hex release (16.1 ± 1.3%). Interestingly, sepiapterin also caused a modest but not significant inhibition of spontaneous release (Fig. 6B).

Effects of BH₄ Inhibition on Short Term (<5 min) NO Production/MC Degranulation—The results obtained above indicated that DAHP and sepiapterin modulate MC NO production long term (>18 h) and thus regulate degranulation. Because BH₄ is a critical co-factor for all NOS isoforms, we studied the effects of DAHP on Ca²⁺-dependent NOS activity and degranulation. PMC from sensitized rats were isolated and loaded with DAF (10 μM); some cells were first pretreated with DAHP (500 μM) for 6 h before loading. PMC were then treated with Ag and analyzed by real time confocal microscopy as outlined above. DAHP-treated MC showed increased membrane changes, with some degranulating MC clearly evident (Fig. 7). Furthermore, DAHP-treated PMC showed no NO flux compared with untreated controls, and most cells degranulated in <3 min.

**DISCUSSION**

NO is a known modulator of both pro- and anti-inflammatory MC functions, although regulation of its endogenous production is unknown (9). These diverse effects imply a complex regulatory mechanism to control the timing, amounts, and location of NO production (8). Because BH₄ is a critical NOS co-factor, we investigated the expression and regulation of BH₄ production in MC and its role in regulating long (inducible) and short term (constitutive) NOS activity and the resulting effect on MC secretion.

Using the fluorescent NO marker, DAF, we identified both long (iNOS) and short term (cNOS) intracellular NO production in MC. We have previously shown iNOS up-regulation in PMC following IFN-γ treatment (10). DAF staining in IFN-γ (long term) treated PMC showed a diffuse cytoplasmic pattern with possible Golgi staining in some cells. Earlier studies by ourselves and others have indicated that PMC may also have Ca²⁺-dependent NOS activity as well as eNOS or nNOS (10, 11). A proportion of PMC treated with Ag/IgE showed immediate (short term) NO formation preceded by a flux in Ca²⁺ levels. Because both cNOS isoforms are Ca²⁺-dependent for their activity, our results support a functional cNOS in MC.

IFN-γ is an important Th1 cytokine and is a potent inhibitor of Th2 type responses including those involving MC. Our data show a significant (8-fold) potentiation of CHI mRNA expres-
sion upon treatment with IFN-γ. Because CHI is a well known IFN-γ-inducible gene, these data are consistent with that shown for other cell types (21, 22). We also investigated the mRNA expression of a CHI regulatory protein, GFRP (23). IFN-γ significantly down-regulates GFRP mRNA production in PMC, which coincides with increased CHI production. By contrast, in C6 (glioblastoma) cells, IFN-γ or interleukin-1β caused an increase in CHI expression with no effect on GFRP, whereas in THP-1 (myelomonocytoma) cells, GFRP was unaltered by IFN-γ or interleukin-1β but instead was down-regulated by lipopolysaccharide (24, 25). IFN-γ thus has a unique regulatory effect on PMC that potentiates BH4 production through increased CHI and inhibiting GFRP expression.

The Western blot data confirm that IFN-γ up-regulates CHI protein expression in a similar manner. We also showed that IFN-γ treatment increased CHI enzymatic activity and BH4 accumulation in PMC as seen in other cell types (21). This increased expression is associated with altered localization of CHI in PMC. In untreated PMC there is a distinct membranous localization of CHI with occasional nuclear positivity but little cytoplasmic staining. Treatment with IFN-γ results in a pronounced increase in the cytoplasmic accumulation of CHI. Previous studies in neurons have shown both cytoplasmic and nuclear CHI staining (26). Our data are novel in that we show for the first time a membranous localization of CHI. A possible explanation for distinct localization patterns of CHI may be that BH4 is highly labile in vivo, and CHI may need to be in close proximity to NOS to maintain adequate BH4 levels and NO production (4). Such differences in localization may contribute to the observed effect of IFN-γ treatment.

BH4 is an important co-factor for all NOS isoforms. Using the CHI inhibitor DAHP and sepiapterin reductase inhibitor N-acetyl serotonin, we demonstrated that diminishing BH4 leads to decreased NO. Decreased NO production through direct inhibition of NOS has been previously shown to potentiate MC degranulation (28). We found similar results, because inhibition of BH4 production increased mediator release from PMC. Furthermore, the inhibitory effect of IFN-γ (18 h) treatment on PMC degranulation was removed by DAHP, implying that the IFN-γ effect was BH4-dependent. Our results using DAHP and N-acetyl serotonin are similar to those seen with mouse macrophages, although higher concentrations of inhibitor were necessary to inhibit NO in those studies (29–31). Because both cell types constitutively express CHI, this disparity may point to differences in compartmentalization of CHI and NOS (30).

We previously showed that NO production in naïve PMC likely relies on cNOS activity (10). PMC treated with DAHP, sepiapterin, or both but without a known secretagogue showed modulation of degranulation. Therefore regulation of BH4 levels is also important in cNOS activity in PMC. Our results using live cell analysis further support these findings because DAHP inhibited NO production and increased degranulation.

Levels of BH4 are a determining factor in the reactive radicals derived from NOS because the NOS dimer may produce superoxide (O2−) or peroxynitrite (ONOO−) rather than NO (32). Interestingly, O2− potentiates MC secretion, although the source of O2− is unknown (28). It is interesting to speculate that O2− arises in MC from NOS that is not saturated with BH4. Indeed, previous studies using the NOS inhibitor l-NAME showed decreased NO and a resulting increase in O2− production and MC degranulation (28, 33). Interestingly, l-NAME also inhibits the reductase region of the NOS enzyme, causing the preferential formation of O2− (34). Thus, NOS may retain a redox balance in MC by producing NO or ONOO to inhibit degranulation, and removal of BH4 leads to O2− production and degranulation. This hypothesis requires further study.

Previous studies have shown that Ag/IgE activation of MC causes a transient hyperphosphorylation of CHI and increase in cellular BH4 (5), and this was interpreted to be an activation step prior to degranulation, although the exact role for this increased BH4 production was unknown. Our live cell results add a caveat to these results, because this increased BH4 production appears to coincide with our noted NO production in MC after Ag/IgE activation. BH4 levels may thus play a key role in controlling MC degranulation via NO, and cellular localization of BH4 and its role in NO production require further investigation.

The nuclear localization of NO is to our knowledge the first identification of dynamic accumulation of NOS activity in the nucleus. Previous studies show eNOS protein in nuclear and perinuclear regions (35, 36). Furthermore, studies have shown both Ca2+ and calmodulin accumulation at nuclear sites in activated MC (20, 37). Because both Ca2+ and calmodulin are
critical in NOS activity, there may be a nuclear/NOS axis, although a role for NO in the nucleus is unknown (38).

Live cell imaging has allowed the investigation of MC functional heterogeneity and NO production at the single cell level. Furthermore, the differing localization of DAF positivity in IFN-γ- and Ag/IgE-treated MC implies involvement of distinct regulatory mechanisms. This pattern of DAF positivity was divergent from that seen when DAF-loaded MC were treated with an NO donor. Thus, interpretation of data in NO studies may have potentially divergent outcomes and molecular mechanisms.

In conclusion, we have characterized both short and long term intracellular NO production in MC and showed that increased intracellular NO production is associated with inhibition of Ag-induced degranulation. Furthermore, BH4 levels are critical in sustaining NO production in MC and can regulate MC degranulation. Furthermore, BH4 will help define novel molecular targets that modulate MC-related inflammatory and innate responses.

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