Control of the Oxidative Burst of Human Neutrophils by Staphylococcal Leukotoxins

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The ability of staphylococcal two-component leukotoxins to induce an oxidative burst and/or to prime human polymorphonuclear cells (PMNs) was studied by using spectrophotometry or flow cytometry. At sublytic concentrations, the HlgA–HlgB, HlgA–LukF–PV, LukS–PV–LukF–PV, and HlgC–LukF–PV combinations of leukotoxins, but not the LukS–PV–HlgB and HlgC–HlgB combinations, were able to induce H2O2 production similar to the H2O2 production induced by 1 μM N-formyl-Leu-Phe (fMLP). In addition, when added at sublytic concentrations, all of the leukotoxin combinations primed PMNs for H2O2 production induced by fMLP. Leukotoxin activation was dependent on the presence of Ca2+ and was inhibited by wortmannin, an inhibitor of phosphatidylinositol 3-kinase, but not by N-methyl-L-arginine, an inhibitor of NO generation, which eliminates the possibility that NO plays a role in the action of leukotoxins. At higher concentrations, all leukotoxins inhibited H2O2 production by PMNs activated by fMLP, phorbol 12-myristate 13-acetate (PMA), or the leukotoxins themselves. This inhibition was not related to the pore formation induced by leukotoxins. Intracellular release of H2O2 induced by fMLP and PMA was not primed by leukotoxins but was inhibited. It seems that leukotoxin inhibition of H2O2 release is independent of pore formation but secondary to an intracellular event, as yet unknown, triggered by leukotoxins.

Materials and Methods

Chemical reagents. Dichlorodihydrofluorescein diacetate (DCFH-DA), dihydrodorosamine 125 (DHR), and Fluor-3 AM were purchased from Molecular Probes (Eugene, Oreg.); wortmannin was purchased from Calbiochem (Meudon, France); fMLP, PMA, N-methyl-L-arginine (NMMA), and all salts were pur-
chased from Sigma-Aldrich (Saint Quentin Fallavier, France); and ethidium bromide was purchased from Interchim (Montluçon, France).

**Leukotoxin purification.** LukS-PV, HlgA, HlgC, LukF-PV, and HlgC were all produced from cultures of strain S. aureus V8 (≈ ATCC 49775) harvested at the stationary phase, as described previously (23). Briefly, the strain was grown for 17 h in YCP medium (3% [wt/vol] yeast extract, 2% [wt/vol] Bacto Casamino Acids [Difco], 2% [wt/vol] sodium pyruvate, 0.25% [wt/vol] Na2HPO4, 0.042% [wt/vol] KH2PO4; pH 7.0) at 37°C with vigorous shaking (11). The secreted proteins were concentrated by precipitation with 80% (wt/vol) ammonium sulfate and dialyzed against 30 mM sodium phosphate (pH 6.5). Positively charged proteins were enriched on a Sepharose SP Fast Flow column (Pharmacia, Uppsala, Sweden) and elution with 0.5 M NaCl. The eluted proteins were then subjected to cation-exchange MonoS fast-performance liquid chromatography (Pharmacia) and then to Alkyl-Superose fast-performance liquid chromatography (Pharmacia) as described previously (11). Proteins were purified to homogeneity controlled by sodium dodecyl sulfate-polyacrylamide gel electrophoresis, adjusted to a concentration of 0.6 mg/ml in 30 mM sodium phosphate–200 mM NaCl and stored at −80°C until they were used.

**Leukocyte preparation.** PMNs were prepared from buffy coats of healthy donors of either sex (Etablissement Français du Sang, Strasbourg, France) as described previously (21). Briefly, 40 ml of a 1/3 (vol/vol) dilution of blood cells in 0.9% NaCl was layered on 12 ml of J Prep (Techgen International, Voisins le Bretonneux, France). After centrifugation at 800 × g for 20 min, the pellet was suspended in 30 ml of 0.9% NaCl, added to 10 ml of 6% (wt/vol) dextran, and sedimented for 30 min. Thirty milliliters of the supernatant was centrifuged for 10 min at 800 × g. The pellet was suspended in HEPES buffer (140 mM NaCl, 5 mM KCl, 10 mM glucose, 0.1 mM EGTA, 1.1 mM CaCl2, 10 mM HEPES, 3 mM Tris; pH 7.3), and the contaminating erythrocytes were removed by hypotonic lysis for 45 s and subsequent washing in HEPES buffer. The final suspension was adjusted to a concentration of 6 × 106 PMNs/ml.

**Oxidative burst assay.** Three different techniques were used to determine superoxide anion production by PMNs in the presence of leukotoxins. All of them were based on fluorescence measurements with DCFDH-DA or DHR.

(i) **Microplate fluorometry.** Every well of a 96-well black tissue culture plate with a clear flat bottom (Costar, New York City, N.Y.) was filled with 200 μl of HEPES buffer containing 1.1 mM Ca2+, 2 × 106 PMNs, 2 μM DHR or 10 μM DCFDH-DA, leukotoxins at the concentrations indicated below, and 1 μM FMLP or 10 nM PMA. Wrightmann and NMMA were incubated with PMNs before they were added to wells. PMNs were incubated for 10 min with the leukotoxins before addition of FMLP or PMA. All experiments were conducted at 37°C. Plates were read (excitation wavelength, 485 nm; emission wavelength, 535 nm) with a SPECTRAfluor microplate fluorometer (Tecan France SA, Trappes, France) operated with the Biolise 2.0 data management software. Measurements were obtained every 2 min for 30 min, and the maximal rate was calculated from six successive values; means ± standard errors of four experiments are given below.

(ii) **Spectrofluorometry.** Continuous variations in fluorescence intensity were recorded by using a dual-excitation and dual-emission Deltasonic TM 4000 spectrofluorometer (Bioritet, PTL, Chamardance, France) with slit widths set at 4 nm. The HEPES buffer, incubation, DHR, and agonist concentrations were the same as those described above for microplates; the 1-cm-light-path cuvettes contained 2 ml of HEPES buffer with 1.1 mM CaCl2 and 2 × 106 PMNs/ml.

(iii) **Flow cytometry.** Flow cytometry data were obtained by using a FACSort cytometer (Becton Dickinson, Le Pont de Claux, France) equipped with a 15-mW argon laser tuned to 488 nm. PMNs were classically discriminated by forward and side light scattering, and DHR fluorescence was recorded in the FL1 channel (emission wavelength, 530 nm). The HEPES buffer, incubation, probe, and agonist concentrations were the same as those described above for microplates. The assay vials contained 2 ml (5 × 106 cells/ml).

**Pore formation determination.** The increase in the fluorescence of the ethidium cation added in the bromide form at a concentration of 100 μM was used as an indicator of the formation of transmembrane pores, as previously described (6), and was measured by using a Deltasonic spectrofluorometer (excitation wavelength, 540 nm; emission wavelength, 600 nm). The fluorescence intensity obtained for PMNs lysed by 0.2% (wt/vol) Triton X-100 added to the cuvette was defined as the maximal activity.

**RESULTS**

When incubated alone with PMNs, none of the five leukotoxin components was able to either induce or modify generation of superoxide anions in the absence or presence of an agonist with the DHR probe in a 96-well plate (data not shown). In contrast, when LukS-PV (10), HlgA, and HlgC (unpublished results) were applied at saturating concentrations and combined with different LukF-PV concentrations, they were able to induce release of H2O2 (Fig. 1). However, higher HlgB concentrations resulted in increased fluorescence intensity only when the HlgB was associated with HlgA. It should be noted that depending on the leukotoxin association considered, the amount of H2O2 released appeared to be above the baseline value when the class F component concentration reached a certain value and was suppressed when the class F component concentration was increased further. When the same leukotoxin concentrations were incubated with PMNs 10 min before addition of 1 μM FMLP, the production of H2O2 by FMLP was first enhanced and then totally inhibited by distinct concentrations of LukF-PV and HlgB (Fig. 2). Hence, the data suggest that within a fairly narrow concentration range, leukotoxins prime PMNs for FMLP activation. When 10 nM PMA was added instead of FMLP, the production of H2O2 by PMA was inhibited by the six combinations of leukotoxins in a concentration-dependent manner (Fig. 3).

Depending on the class F component concentration and the maximal priming obtained, the order of potency of the leukotoxin associations was as follows: HlgA–HlgB > HlgC–HlgB > LukS-PV–LukF-PV > HlgA–LukF-PV = HlgC–LukF-PV = LukS-PV–HlgB. Thus, it appears that the homologous associations of leukotoxins resulting from individual genetic loci were the most potent.
Figure 4A shows that in the absence of Ca\(^{2+}\), 1 nM LukS-PV–0.6 nM LukF-PV was not able to increase significantly the intracellular Ca\(^{2+}\) concentration and that the presence of PMA and fMLP did not change this lack of a response. In addition, no H\(_2\)O\(_2\) was produced (Fig. 4B, lines f and g). In the presence of 1 mM Ca\(^{2+}\), the same leukotoxin concentration induced an increase in the intracellular Ca\(^{2+}\) concentration (Fig. 4) and H\(_2\)O\(_2\) production (Fig. 4B, line a) greater than the H\(_2\)O\(_2\) production induced by 1 \(\mu\)M fMLP (Fig. 4B, line d). Furthermore, the H\(_2\)O\(_2\) generated by LukS-PV–LukF-PV combined with fMLP (Fig. 4B, line b) did not result from a simple additive effect of H\(_2\)O\(_2\) generated by leukotoxin and fMLP. Taken together, these results indicate that there was priming of PMNs by the LukS-PV–LukF-PV leukotoxin. At this LukS-PV–LukF-PV concentration no modification of the amplitude of the PMA response was observed (Fig. 4B, lines c and e), but the time lag was significantly reduced. These results confirm that extracellular Ca\(^{2+}\) is required for superoxide generation (2, 17) and show that modulation of the fMLP and PMA responses is not dependent on modification of the intracellular Ca\(^{2+}\) concentration; after the initial rise induced by leukotoxin, there is no further increase due to addition of fMLP or PMA (Fig. 4A).

As shown in Fig. 5A, the Fura-2 fluorescence increased as the leukotoxin concentration increased, while the DHR fluorescence increased at low concentrations but decreased at the highest leukotoxin concentrations (Fig. 5B). An increase in the intracellular Ca\(^{2+}\) concentration was necessary for induction of the oxygen burst, but there was no correlation between the level of intracellular Ca\(^{2+}\) and the level of H\(_2\)O\(_2\) generated. In conclusion, two effects were observed: (i) when an active concentration of leukotoxin is reached, there is superoxide anion...
production and priming of PMNs which is dependent on the presence of Ca\(^{2+}\); and (ii) a further increase in the leukotoxin concentration strongly inhibits superoxide ion production irrespective of the agonist added to the PMN suspension.

To test for a possible role of NO in the leukotoxin oxidative burst (19), we examined the effects of different concentrations of L-NMMA, an inhibitor of NO formation (14), on modulation of the fMLP and PMA responses. In both experiments no influence of the inhibitor was observed (data not shown).

The role of phosphatidylinositol 3-kinase (PI3-kinase) in activation of fMLP superoxide production (31) by LukS-PV–LukF-PV was determined by using wortmannin (Fig. 6), an inhibitor of PI3-kinase. Figure 6 shows that activation of the fMLP response by LukS-PV–LukF-PV was inhibited by high wortmannin concentrations.

It has been proposed that the oxygen burst inhibition induced by pore-forming toxins results from pore formation (3, 27). To test this hypothesis, we examined whether there is a correlation between pore formation and the level of the oxidative burst (i.e., between the variations in the fluorescence intensities of ethidium and DHR). Time courses for both ethidium entry and H\(_2\)O\(_2\) production were determined after addition of different concentrations of HlgA-HlgB and LukS-PV–LukF-PV to PMNs obtained from the same donor. Figure 7A shows curves obtained with concentrations of the two different leukotoxins that gave nearly identical pore formation but not comparable H\(_2\)O\(_2\) release values. Such determinations were extended to other leukotoxin concentrations and the average slope of the recorded curves of two responses calculated during the time delimited by dotted lines in Fig. 7A. The values for the average slopes obtained for different leukotoxin concentrations are plotted in Fig. 7B. The plots show that although the two leukotoxins induced equivalent pore formation, LukS-PV–LukF-PV induced a higher rate of H\(_2\)O\(_2\) production than HlgA-HlgB induced, which does not support the hypothesis that there is a link between the two events. Furthermore, although the lack of superoxide release has been attributed to a loss of cytoplasmic ATP after pore formation (4), addition of
10 mM ATP to PMN suspensions did not prevent inhibition of the oxygen burst (data not shown).

The fluorescence intensity recorded with the spectrofluorometer or the microplate fluorometer reflected both the intra- and extracellular H$_2$O$_2$ production. Conversely, the fluorescence recorded with the flow cytometer was emitted solely by DHR trapped inside PMNs and reflected the intracellular production of H$_2$O$_2$. The influence of LukS-PV–LukF-PV on intracellular superoxide anion formation in the absence of fMLP and PMA was tested under these conditions. Figure 8A shows that the intracellular H$_2$O$_2$ production induced by LukS-PV–LukF-PV was weak and inhibited at LukF-PV concentrations higher than 0.5 nM. LukS-PV–LukF-PV did not prime PMNs, but it inhibited the production of intracellular H$_2$O$_2$ by fMLP (Fig. 8B) and also inhibited the intracellular activity of PMA at LukF-PV concentrations higher than 0.1 nM (Fig. 8C).

**DISCUSSION**

DCFH-DA (7) and DHR are fluorescent probes for H$_2$O$_2$ and, to a much lesser extent, for other reactive oxygen species (26), and they have been shown to be suitable probes for determination by spectrofluorometry and flow cytometry of the production of the free radical superoxide which dismutates rapidly to H$_2$O$_2$ (12, 29), although DHR has a higher fluorescence intensity.

The present study demonstrated that leukotoxins from *S. aureus* are able to induce three kinds of effects on PMNs: (i) they generate concentration-dependent production of superoxide anion, although the production is limited, (ii) they prime PMNs at low concentrations for fMLP activation of superoxide anion production, like pseudomonal leukocidin (22) and *Escherichia coli* hemolysin (3), and (iii) at higher concentrations they inhibit the oxidative burst produced by fMLP or PMA. It seems likely that priming of fMLP activation is secondary to the increase in the intracellular Ca$^{2+}$ concentration induced by leukotoxins in the presence of extracellular Ca$^{2+}$, which is required for a PMN respiratory burst (2, 17).
It is generally thought that pore formation is the cause of inhibition of the respiratory burst by pore-forming toxins (3, 27) which induce leakage of ATP. However, the present study showed (i) that intracellular production and extracellular production linked to two separate pools of NADPH oxidase and localized to granule and plasma membranes, respectively (16), are modulated differently depending on the leukotoxin concentration which controls pore formation, (ii) that the amount of H$_2$O$_2$ released is not related to the number of pores formed, and (iii) that addition of ATP to a PMN suspension does not eliminate the inhibition by leukotoxins. Consequently, it appears that for staphyloccocal leukotoxins, inhibition of H$_2$O$_2$ production by PMNs and pore formation are two independent mechanisms resulting from the binding of these leukotoxins. It seems likely that the mediation of the respiratory burst by leukotoxins is a downstream signaling event secondary to the binding to cellular receptors of the toxins.

In conclusion, staphyloccocal leukotoxins at sublytic concentrations are able to induce a moderate oxygen burst and to prime PMNs for further activation. However, at higher concentrations they cause complete inhibition of superoxide anion production which is independent of pore formation but result in an intracellular signaling event that is currently under investigation. We hypothesize that under infectious conditions inhibition of the oxidative burst by leukotoxins produced by S. aureus could be a determinant for survival.

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