Human Phagocytes Employ the Myeloperoxidase-Hydrogen Peroxide System to Synthesize Dityrosine, Trityrosine, Pulcherosine, and Isodityrosine by a Tyrosyl Radical-dependent Pathway*

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Myeloperoxidase, a heme protein secreted by activated phagocytes, may be a catalyst for lipoprotein oxidation in vivo. Active myeloperoxidase is a component of human atherosclerotic lesions, and atherosclerotic tissue exhibits selective enrichment of protein dityrosine cross-links, a well characterized product of myeloperoxidase. Tyrosylation of lipoproteins with peroxidase-generated tyrosyl radical generates multiple protein-bound tyrosine oxidation products in addition to dityrosine. The structural characterization of these products would thus serve as an important step in determining the role of myeloperoxidase in lipoprotein oxidation in the artery wall. We now report the identification and characterization of four distinct tyrosyl radical addition products generated by human phagocytes. Activated neutrophils synthesized three major fluorescent products from L-tyrosine; on reverse phase HPLC, each compound coeluted with fluorescent oxidation products formed by myeloperoxidase. We purified the oxidation products to apparent homogeneity by cation and anion exchange chromatography and identified the compounds as dityrosine (3,3'-dityrosine), trityrosine (3,3',5,3'-trityrosine) and pulcherosine (5-[4-[2-carboxy-2-aminoethyl]phenoxy]3,3'-dityrosine) by high resolution NMR spectroscopy and mass spectrometry. Additionally, we have found that dityrosine is a precursor to trityrosine, but not to pulcherosine. In a search for a precursor to pulcherosine, we identified isodityrosine (3-[4'-(2-carboxy-2-aminoethyl)phenoxy]tyrosine), a nonfluorescent product of L-tyrosine oxidation by human phagocytes. Our results represent the first identification of this family of tyrosyl radical addition products in a mammalian system. Moreover, these compounds may serve as markers specific for tyrosyl radical-mediated oxidative damage in atherosclerosis and other inflammatory conditions.

Oxidized low density lipoprotein (LDL)1 may play a pivotal role in the pathogenesis of atherosclerosis (1–4), but the mechanisms that damage LDL in vivo have yet to be identified. A potential pathway involves myeloperoxidase, a heme protein secreted by neutrophils and monocytes (4). Phagocytes employ the myeloperoxidase-H2O2 system to generate potent cytotoxins that kill invading pathogens and tumor cells (5–8); under pathological conditions, normal tissue may also be a target for damage. Active myeloperoxidase is a component of human atherosclerotic lesions, where it co-localizes with monocytes and macrophages (9). The pattern of immunostaining for the enzyme in human atherosclerotic lesions (9) is remarkably similar to that for protein-bound lipid oxidation products in rabbit lesions (10), implicating myeloperoxidase as a potential pathway for LDL oxidation in vivo.

Hypochlorous acid is the best characterized product of myeloperoxidase (11), which is the only human enzyme known to produce this oxidant under physiological conditions (12). Hypochlorous acid is a potent cytotoxin that oxidatively bleaches heme proteins (13) and chlorinates proteins (14, 15) and unsaturated lipids (16, 17). LDL modified with reagent hypochlorous acid undergoes aggregation and causes macrophages to accumulate cholesterol ester (18); lipoproteins with similar properties have been isolated from atherosclerotic lesions (19–21). A monoclonal antibody specific for hypochlorous acid-modified proteins recognizes epitopes in human atherosclerotic lesions, as well as LDL-like particles isolated from aortic tissue, strongly suggesting that myeloperoxidase is promoting oxidation in the artery wall (22).

Another substrate for myeloperoxidase is L-tyrosine, which is converted to tyrosyl radical (Scheme I) (23). The major reaction of free tyrosyl radical is dimerization to form the intensely fluorescent amino acid 3,3'-dityrosine (23–26). Tyrosyl radical generated by peroxidase converts protein and lipoprotein tyrosyl residues to dityrosine (27, 28) and initiates LDL lipid peroxidation (29), which may make the lipoprotein atherogenic (1–4).

Tyrosine oxidation products, including dityrosine (30, 31), pulcherosine (32), isodityrosine (33), and trityrosine (30, 32), are common post-translational modification found in bacteria (34), yeast (35, 36), plants (33), and metazoans (30–33, 37). In

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1 The abbreviations used are: LDL, low density lipoprotein; dityrosine, 3,3'-dityrosine; trityrosine, 3,3',5,3'-trityrosine; pulcherosine, 5-[4'-(2-carboxy-2-aminoethyl)phenoxy]-3,3'-dityrosine; isodityrosine, 3-[4'-(2-carboxy-2-aminoethyl)phenoxy]tyrosine; DTPA, diethylenetriaminepentaacetic acid; HPLC, high pressure liquid chromatography; GC, gas chromatography; MS, mass spectrometry; PMA, phorbol 12-myristate 13-acetate.
addition of catalase (400 nM) and the reaction mixture was stored at 20°C until analysis. After a 2-h incubation at 37°C, the reaction was terminated by acidification to pH 3.0 with HBr (3.5 M) in methanol. Concentrated reaction mixtures were then prepared by incubation of dried propylated amino acids (100 μl of ethyl acetate/acetone) at 60°C for 30 min. Derivatized amino acids were stored in reagent grade methanol until analysis.

**RESULTS**

Gas Chromatography-Mass Spectrometry (GC-MS)—FPLC-purified and desalted oxidation products were evaporated under N2 and esterified by incubation with 200 μl of HBr (3.5 M) in n-propanol (Cambridge Isotope Laboratories) at 100°C for 30 min. Pentafluoropropionyl derivatives were then prepared by incubation of dried propylated amino acids with 50 μl of ethyl acetate/pentafluoropropionic acid anhydride (4:1, v/v) at 60°C for 30 min. Derivatized amino acids were stored in reagent grade methanol until analysis.

GC and direct probe MS analyses were obtained with a HP-5988A mass spectrometer equipped with a Texitronic Vector 1 Data System. Capillary GC was performed using a DB-1 column (8 m, 0.33 mm inner diameter, 1-μm film thickness; J & W Scientific) in the splitless injection mode. The initial column temperature of 70°C was increased to 150°C at 60°C/min, and then to 250°C at 10°C/min. The detector, transfer line, and ion source temperature were set at 220°C, 250°C, and 150°C, respectively. Mass spectra were obtained in the negative chemical ionization mode using methane as the reagent gas.

**RESULTS**

Myeloperoxidase Generates a Family of Tyrosyl Radical Addition Products—Three major fluorescent peaks, termed compounds I, II, and III, were identified by Mono S chromatography when L-tyrosine was oxidized by the myeloperoxidase-H2O2 system (Fig. 1). Production of all three oxidation products required active myeloperoxidase and the H2O2 generating system and was inhibited by catalase (a scavenger of H2O2), azide (a heme poison), and ascorbate (which reacts with both H2O2 and tyrosyl radical). Compounds I, II, and III eluted from a Mono S cation exchange column at 60 min, 115 min, and 105 min, respectively; compound I coeluted with authentic dityrosine. Each of the products exhibited dityrosine-like fluorescence and mass spectra; however, their mass spectrometric fragmentation patterns were different from authentic dityrosine. The identity of the three compounds was determined by NMR spectroscopy—One- and two-dimensional NMR spectra were acquired at 25°C using a Varian Unity 500 spectrometer (499.843 MHz for 1H) interfaced to a Sun SPARC-2 workstation and equipped with a Nalorac triple-resonance indirect-detection probe. Prior to analysis, reactions were filtered and the eluent was evaporated under N2 and stored in reagent grade methanol. GC and direct probe MS analyses were obtained with a HP-5988A mass spectrometer equipped with a Texitronic Vector 1 Data System. Capillary GC was performed using a DB-1 column (8 m, 0.33 mm inner diameter, 1-μm film thickness; J & W Scientific) in the splitless injection mode. The initial column temperature of 70°C was increased to 150°C at 60°C/min, and then to 250°C at 10°C/min. The detector, transfer line, and ion source temperature were set at 220°C, 250°C, and 150°C, respectively. Mass spectra were obtained in the negative chemical ionization mode using methane as the reagent gas.

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Tyrosyl Radical Addition Products of Phagocytes

**EXPERIMENTAL PROCEDURES**

**Materials**

Glucose oxidase and crystalline catalase (from beef liver; thymol-free) were purchased from Boehringer Mannheim. L-[14C]Tyrosine (>99% pure) was from Isotopic. All other reagents were from either Sigma or Fisher Scientific unless otherwise indicated.

**Methods**

Isolation of Myeloperoxidase—Nonidet P-40-treated leukocytes obtained by leukopheresis from a patient with chronic myelogenous leukemia (39) were extracted with cetlytrimethylammonium bromide (40). Myeloperoxidase was then isolated by sequential lectin-affinity chromatography, ammonium sulfate precipitation, and gel filtration chromatography as described previously (23, 40). Myeloperoxidase concentration was determined spectrophotometrically (ε 430 = 170,000 M−1 cm−1; Ref. 41) and the isolated enzyme had an A430/A280 ratio of 0.71.

L-Tyrosine Oxidation by Myeloperoxidase—Reactions were carried out in buffer A (50 mM phosphate, 100 μM DTPA, pH 7.4), which had been passed over a Chelex-100 column (Bio-Rad) to remove metal ions. After a 2-h incubation at 37°C, the reaction was terminated by addition of catalase (400 U) and the reaction mixture was stored at −20°C until analysis.

Isolation of L-Tyrosine Oxidation Products by Ion Exchange Chromatography—Reaction mixture (1 liter) was acidified to pH 3.0 with HCl and subjected to strong cation exchange chromatography (AG-50WX2; Bio-Rad) as a desalting step. The column was eluted with 2 M NH4OH. The resulting amino acid solution was lyophilized, dissolved in buffer B (10 mM acetic acid, pH 3.0), and then subjected to fast protein liquid chromatography (FPLC) on a Pharmacia Mono S HR 5/5 column equilibrated with buffer B. Amino acids were eluted from the cation exchange column with a 60-mL linear gradient of 0–0.25 M NaCl in buffer B at a flow rate of 1 ml/min. Collected peaks were desalted as above, dissolved in buffer C (10 mM Tris base, pH 9.0), subjected to FPLC on a Pharmacia Mono Q HR 5/5 anion exchange column equilibrated with buffer C, and eluted with a 60-mL linear gradient of 0–0.25 M NaCl in buffer C at 1 ml/min. Each amino acid was then desalted and again subjected to cation exchange chromatography as described above.
Tyrosyl Radical Addition Products of Phagocytes

Fig. 1. Cation exchange FPLC analysis of the L-tyrosine oxidation products generated by myeloperoxidase. L-Tyrosine (2 mM) in buffer A (1 liter) was incubated with myeloperoxidase (4 nM) and 100 ng/ml glucose oxidase and 0.1 mg/ml glucose oxidase for 2 h at 37 °C. The reaction was terminated by the addition of catalase (400 nM). The reaction mixture was desalted and subjected to FPLC on a Mono S HR 5/5 column as described under "Methods." The peaks of material labeled I, II, III, and IV were purified to apparent homogeneity by sequential chromatography on anion exchange (Mono Q HR5/5) and cation exchange (Mono S HR5/5) columns, and subsequently identified as dityrosine, pulcherosine, trityrosine, and iso-dityrosine, respectively, in the structural studies described under "Results."

Structural Characterization of the Fluorescent L-Tyrosine Oxidation Products Generated by Myeloperoxidase—H NMR resonances from the aromatic region (6.8–7.4 ppm) of compounds I, II, and III and their structural assignments are shown in Fig. 2. This region of compound I (Fig. 2A) was very similar to that previously reported for 3,3′-dityrosine isolated from yeast spore coats (32) and contained 6 protons with chemical shifts of: 6.99 (H3), 7.14 (H6), and 7.23 ppm (H7). Compound II exhibited 8 protons in the aromatic region (Fig. 2B): 6.98 (H3), 7.24 (H6), 7.33 (H7), and 7.36 ppm (H8) and is consistent with the structure of trityrosine. The spectrum of compound II (Fig. 2C) is consonant with the structure of pulcherosine and demonstrated nine aromatic protons: 6.98 (H3), 7.00 (H6), 7.02 (H7), 7.13 (H8), 7.23 (H9), 7.29 (H10), and 7.31 ppm (H11). To confirm the assignments of the protons in compound II, the cross-peak labels separated by commas represent through-bond correlations between protons resulting from scalar couplings. The spectrum was processed with pseudo-echo apodization with a line broadening of 6 Hz and a Gaussian time constant of 0.04 s. Number of transients, 64; number of time-domain points, 8192; spectral width, 6000 Hz.

Novel compounds were purified by reverse phase HPLC; each compound eluted as a single, symmetrical peak as monitored by both fluorescence and absorbance. Cation exchange FPLC analysis of the L-tyrosine oxidation products generated by myeloperoxidase—H NMR resonances from the aromatic region (6.8–7.4 ppm) of compounds I, II, and III and their structural assignments are shown in Fig. 2.

The negative-ion chemical ionization mode. Because it was relatively non-volatile, derivatized trityrosine was analyzed using direct probe MS. Dityrosine exhibited a low abundance ion peak at mass-to-charge (m/z) 1028, the calculated molecular ion (M+)) for the derivative. Prominent ions were observed at m/z 1008 (M−HF) and 880 (M−CF3 CF2CHO). The mass spectrum of derivatized pulcherosine demonstrated a low abundance ion at m/z 1395 (M+) and major fragment ions at m/z 1375 (M−HF) and 1247 (M−CF3 CF2CHO). The trityrosine derivative exhibited a low abundance ion at m/z...
Spectroscopic properties of dityrosine, trityrosine, and pulcherosine

UV spectra were obtained in 0.1 M HCl or 0.1 M NH₄OH. Fluorescence excitation and emission spectra were obtained in 0.1 M NaOH with excitation and emission slits of 5 and 10 nm, respectively.

|               | Absorbance | Fluorescence (λ<sub>em</sub>) |
|---------------|------------|-------------------------------|
|               | λ<sub>max</sub> | λ<sub>max</sub> |<br>nm | nm | nm |
| Dityrosine    |            |                               |       |     |     |
| Acid          | 5,400      | 284                           | 284   | 409 |
| Base          | 8,600      | 316                           | 317   | 407 |
| Trityrosine   |            |                               |       |     |     |
| Acid          | 11,000     | 286                           | 286   | 409 |
| Base          | 11,500     | 322                           | 319   | 416 |
| Pulcherosine  |            |                               |       |     |     |
| Acid          | 7,700      | 282                           | 283   | 416 |
| Base          | 9,500      | 315                           | 320   | 414 |

1541 (M⁻), as well as major ions expected from loss of HF and CF₃CF₂CHO at m/z 1521 and 1393, respectively. The mass spectra of compounds I, II, and III are thus in excellent agreement with the structures determined using multinuclear NMR spectroscopy.

Fluorescence excitation/emission spectra were obtained in 0.1 M HCl or 0.1 M NH₄OH. Fluorescence excitation/emission spectra of the compounds were then determined at acid and alkaline pH. These data, which are in good agreement with previously reported results for tyrosine oxidation products isolated from sea urchins, insects, and yeast (24, 25, 32, 36), are summarized in Table I.

Isolation and Identification of Isodityrosine—A Non-fluorescent L-Tyrosine Radical Addition Product Generated by Myeloperoxidase—Initial fractionation of the myeloperoxidase-dityrosine-H₂O₂ reaction mixture revealed a small shoulder of material, which eluted just after dityrosine on cation exchange chromatography (Fig. 1, peak IV). This material exhibited tyrosine-like absorbance but did not dityrosine-like fluorescence. To investigate the structure of this compound, a [13C₁₂]dityrosine in buffer A was exposed to the myeloperoxidase-H₂O₂ system, and the product was isolated by sequential cation, anion, and cation exchange FPLC as described above for compounds I–III. The isolated compound, which eluted as a single, symmetrical peak on reverse phase HPLC, was subjected to heteronuclear single quantum coherence NMR spectroscopy (Fig. 4).

Production of dityrosine, trityrosine, and pulcherosine was readily identified by GC-MS analysis, but was below the UV detection limit on HPLC.

Production of dityrosine, trityrosine, and pulcherosine required phagocytes, L-tyrosine, and activation of the cells by phorbol ester (Table II). L-Tyrosine oxidation by the cells was independent of iron or copper because the buffer contained DTPA, a potent inhibitor of metal-catalyzed reactions (44). As with the reaction catalyzed by purified myeloperoxidase, the neutrophil reaction was inhibited by catalase, by the heme poisons azide and cyanide, and by ascorbate (Table II). Collectively, these results suggest that the family of tyrosyl radical addition products generated by myeloperoxidase constitutes molecular markers of phagocyte activation.

Tyrosyl Radical Addition Products of Phagocytes

|                        | Absorbance | Fluorescence (λ<sub>em</sub>) |
|------------------------|------------|-------------------------------|
|                        | λ<sub>max</sub> | λ<sub>max</sub> |<br>nm | nm | nm |
| Dityrosine             |            |                               |       |     |     |
| Acid                   | 5,400      | 284                           | 284   | 409 |
| Base                   | 8,600      | 316                           | 317   | 407 |
| Trityrosine            |            |                               |       |     |     |
| Acid                   | 11,000     | 286                           | 286   | 409 |
| Base                   | 11,500     | 322                           | 319   | 416 |
| Pulcherosine           |            |                               |       |     |     |
| Acid                   | 7,700      | 282                           | 283   | 416 |
| Base                   | 9,500      | 315                           | 320   | 414 |

1521 (M⁻), as well as major ions expected from loss of HF and CF₃CF₂CHO at m/z 1521 and 1393, respectively. The mass spectra of compounds I, II, and III are thus in excellent agreement with the structures determined using multinuclear NMR spectroscopy.

Ultrasound Absorption and Fluorescence Spectra with Molar Extinction Coefficient Determinations of the Major Tyrosyl Radical Addition Products—Radio-labeled dityrosine, trityrosine, and pulcherosine were isolated by FPLC from an L-[14C]tyrosine solution of known specific activity oxidized with myeloperoxidase, and their concentrations determined by scintillation spectrometry. The extinction coefficients, ultraviolet absorption spectra, and fluorescence excitation/emission spectra of the compounds were then determined at acid and alkaline pH. These data, which are in good agreement with previously reported results for tyrosine oxidation products isolated from sea urchins, insects, and yeast (24, 25, 32, 36), are summarized in Table I.

Isolation and Identification of Isodityrosine, a Non-fluorescent L-Tyrosine Radical Addition Product Generated by Myeloperoxidase—Initial fractionation of the myeloperoxidase-dityrosine-H₂O₂ reaction mixture revealed a small shoulder of material, which eluted just after dityrosine on cation exchange chromatography (Fig. 1, peak IV). This material exhibited tyrosine-like absorbance but did not dityrosine-like fluorescence. To investigate the structure of this compound, a [13C₁₂]dityrosine in buffer A was exposed to the myeloperoxidase-H₂O₂ system, and the product was isolated by sequential cation, anion, and cation exchange FPLC as described above for compounds I–III. The isolated compound, which eluted as a single, symmetrical peak on reverse phase HPLC, was subjected to heteronuclear single quantum coherence NMR spectroscopy (Fig. 4).

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Dityrosine Promotes the Synthesis of Trityrosine by Neutrophils—Studies with isolated myeloperoxidase suggested that isodityrosine and dityrosine were precursors for pulcherosine and trityrosine, respectively. To test the hypothesis that dityrosine was similarly employed by the myeloperoxidase system of activated human neutrophils to generate trityrosine, we supplemented the medium of neutrophils with dityrosine (0.01 mM) and L-tyrosine (1 mM) and then monitored the kinetics of trityrosine and pulcherosine synthesis. The presence of dity-
Tyrosyl Radical Addition Products of Phagocytes

![Diagram](image)

**FIG. 5.** Reverse phase HPLC analysis of fluorescent tyrosine oxidation products generated by myeloperoxidase (A) and activated human neutrophils (B). L-Tyrosine was oxidized either with the myeloperoxidase-H$_2$O$_2$ system as described in the legend to Fig. 1 or with activated human neutrophils as described in the legend to Fig. 5. After a 2-h incubation at 37 °C, the reactions were terminated by addition of catalase (400 nM) and subjected to analysis by reverse phase HPLC as described under “Methods.” Tyrosine oxidation products were detected by monitoring fluorescence ($\lambda_{\text{em}}$ 296 nm, $\lambda_{\text{ex}}$ 414 nm).

**TABLE II**

| Requirement for dityrosine, trityrosine, and pulcherosine synthesis by neutrophils |
|---------------------------------|----------------|----------------|
| Oxidation of L-tyrosine by neutrophils (5 × 10$^5$ cells/ml) was carried out in medium A supplemented with 0.1 mM DTPA, 1 mg/ml D-glucose, and 1 mM L-tyrosine. Neutrophils were activated by the addition of 200 nM PMA. After a 2-h incubation at 37 °C, the reaction was stopped by the addition of 400 nM catalase. Samples were analyzed by reverse phase HPLC as described under “Methods.” The limit of detection for each compound was less than 1 pmol. Values are expressed as pmol/cell and represent the mean of duplicate determinations. Similar results were found in two independent experiments. |

| Dityrosine | Trityrosine | Pulcherosine |
|------------|-------------|--------------|
| pmol       | pmol        | pmol         |
| Complete system | 691 | 86 | 20 |
| Complete system plus: | | | |
| Superoxide dismutase (150 nM) | 1531 | 1024 | 256 |
| Catalase (400 nM) | 26 | <1 | <1 |
| CN$^-$ (5 mM) | 88 | <1 | <1 |
| N$_2$O (0.6 mM) | 112 | <1 | <1 |
| Ascorbate (0.5 mM) | 20 | <1 | <1 |
| Complete system minus: | | | |
| L-Tyrosine | 55 | <1 | <1 |
| Cells | 38 | <1 | <1 |
| Phorbol myristate acetate | 55 | <1 | <1 |

**FIG. 6.** Progress curve for trityrosine (A) and pulcherosine synthesis (B) by human neutrophils in the presence and absence of dityrosine. Neutrophils (5 × 10$^5$ cells/ml) were incubated at 37 °C in medium A supplemented with 1 mM L-tyrosine, 0.1 mM DTPA and 1 mg/ml D-glucose. Where indicated, dityrosine (DIY; 10 µM) was included. After stimulation of the cells with 200 nM PMA, aliquots of medium were removed at the indicated times and trityrosine and pulcherosine production quantified by reverse phase HPLC as described under “Methods.” Values are expressed as picomoles per 5 × 10$^5$ cells and represent the mean of duplicate determinations.

Dityrosine led to a decrease in the lag phase for trityrosine synthesis (Fig. 6, upper panel). Moreover, there was a dramatic 8-fold increase in the yield of trityrosine. These results strongly suggest that dityrosine is a precursor in the generation of trityrosine. In contrast to trityrosine, the kinetics and yield of pulcherosine (Fig. 6, lower panel) were unaffected by the presence of dityrosine, suggesting that pulcherosine generation involves another intermediate. These results suggest that tyrosyl radical reacts by two different pathways to generate trityrosine and pulcherosine.

Superoxide Dismutase Stimulates L-Tyrosine Oxidation by Activated Neutrophils—The initial product of the neutrophil oxidase is O$_2^-$, the one-electron reduced form of molecular oxygen (45). O$_2^-$ scavenges tyrosyl radical by acting as an electron donor (46, 47). This reaction is blocked by superoxide dismutase, which catalyzes the dismutation of O$_2^-$ to H$_2$O$_2$ (46, 47). To explore the role of O$_2^-$ in L-tyrosine oxidation, the kinetics of dityrosine, trityrosine, and pulcherosine generation by activated neutrophils in the presence and absence of superoxide dismutase were monitored. Neutrophil activation resulted in production after a lag phase of ~2 min (data not shown). The progress curve of dityrosine production was similarly nonlinear but with a longer lag phase of ~6 min (Fig. 7, top panel). In contrast, the lag phases for O$_2^-$ generation and dityrosine synthesis were virtually identical when superoxide dismutase was included. Catalase inhibited the production of dityrosine, trityrosine, and pulcherosine by neutrophils (Fig. 7), confirming that H$_2$O$_2$ is necessary for their synthesis.

The progress curves for both pulcherosine and trityrosine production by activated neutrophils exhibited lag phases that were greater than that observed for dityrosine (Fig. 7). As with dityrosine, superoxide dismutase significantly shortened the lag phases for trityrosine and pulcherosine production (Fig. 7, middle and bottom panels). Superoxide dismutase led to a 1.5-fold increase in dityrosine and 8- and 4-fold increases in trityrosine and pulcherosine, respectively. Superoxide dismutase may have stimulated the yield of L-tyrosine oxidation products by blocking the scavenging of tyrosyl radical by O$_2^-$ (46, 47), by preventing the formation of an inactive form of myeloperoxidase (48), or by increasing the yield of H$_2$O$_2$ from O$_2^-$ (45).
with tyrosyl radical is blocked by superoxide dismutase (46, 47). The results of this reaction were identified as dityrosine, isodityrosine, trityrosine, and pulcherosine by activated neutrophils. First, purified myeloperoxidase converted L-tyrosine to tyrosyl radical, which then undergoes radical addition reactions to form dityrosine, isodityrosine, trityrosine, and pulcherosine.

**DISCUSSION**

Several lines of evidence indicate that human neutrophils employ the myeloperoxidase-H$_2$O$_2$ system to convert L-tyrosine to tyrosyl radical, which then undergoes radical addition reactions to form dityrosine, isodityrosine, trityrosine, and pulcherosine. First, purified myeloperoxidase converted L-tyrosine to three fluorescent products and one non-fluorescent product by a reaction that required H$_2$O$_2$ and was sensitive to inhibition by heme poisons. The products of this reaction were identified as dityrosine, isodityrosine, trityrosine, and pulcherosine by NMR and mass spectroscopy. Second, activation of human neutrophils with phorbol ester led to the formation of the identical family of tyrosyl radical addition products. As with myeloperoxidase, synthesis of the compounds required H$_2$O$_2$ and was inhibited by heme poisons. Finally, the formation of dityrosine, trityrosine and pulcherosine by activated neutrophils was inhibited by ascorbic acid and stimulated by superoxide dismutase. Both ascorbate and O$_2^-$, the initial product of the neutrophil oxidase (45), are known to repair tyrosyl radical, and the reaction of O$_2^-$ with tyrosyl radical is blocked by superoxide dismutase (46, 47). These results represent the first identification of this family of tyrosyl radical addition products in a mammalian system and suggest their potential as specific markers for phagocyte-mediated oxidative damage.

The one-electron oxidation of L-tyrosine by myeloperoxidase bears striking parallels with the chemistry of phenoxyl radical (49, 50). In both systems, the spectrum of products is readily explained by radical-radical coupling at the predominant sites of electron density on the oxygen and aromatic carbons (Scheme I). Substituents in the para position of phenoxyl radical block both O$p$ and p,p$'$ dimerization (49, 50). L-Tyrosine oxidation by myeloperoxidase generates the products expected from para-substituted phenoxyl radical, including compounds with carbon-carbon cross-links (dityrosine, trityrosine, pulcherosine) and carbon-oxygen cross-links (pulcherosine and isodityrosine), strongly implicating tyrosyl radical in the reaction pathway.

Three observations suggest that dityrosine itself might be oxidized by activated neutrophils to dityrosyl radical, which then reacts with tyrosyl radical to form trityrosine. First, the lag phase for the conversion of L-tyrosine to trityrosine by the cells was significantly greater than that observed for dityrosine, suggesting that trityrosine production was dependent upon dityrosine formation. Second, the lag phases for the synthesis of both dityrosine and trityrosine were shortened by superoxide dismutase; this enzyme scavenges O$_2^-$, which is known to repair tyrosyl radical (46, 47). Third, when a low concentration of dityrosine was included together with L-tyrosine in the medium of the cells, the lag phase for trityrosine synthesis was shortened and the yield of the compound was increased 8-fold. In contrast, dityrosine had little effect on either the progress curve or the yield of pulcherosine. Scheme II is consistent with these results and the chemistry of phenoxyl radical (49, 50).

Oxidation of LDL may represent an important step in conferring atherogenic potential to the modified lipoprotein. However, the biochemical pathways that mediate LDL oxidation in vivo have yet to be identified (1–4). Previously, we have shown that active myeloperoxidase is expressed in human atherosclerotic lesions (9) and that myeloperoxidase-derived tyrosyl radical initiates LDL lipid peroxidation (29). LDL isolated from vascular tissue exhibits evidence of lipid peroxidation (19–21) and is aggregated (21), suggesting that tyrosyl radical may account in part for intermolecular cross-linking of artery wall...
lipoproteins. L-Tyrosine oxidation may also play a role in stimulating the cellular uptake of lipoproteins because cross-linking causes LDL to aggregate and stimulates its uptake by macrophages (18, 21, 22, 52).

Tyrosyl radical generated by activated phagocytes causes protein tyrosylation (28), and human atherosclerotic lesions contain elevated levels of protein-bound dityrosine, implicating peroxidative pathways as one mechanism for oxidative damage of the vascular wall. The L-tyrosine oxidation products we have described are stable to acid hydrolysis, intensely fluorescent, and readily detected by mass spectrometry, making them attractive candidates for monitoring protein oxidative damage. Moreover, dityrosine accounts for only a third of the L-tyrosine covalently incorporated into the apolipoproteins of lipoproteins. L-Tyrosine oxidation may also play a role in stimulating the cellular uptake of lipoproteins because cross-linking causes LDL to aggregate and stimulates its uptake by macrophages (18, 21, 22, 52).

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