Separation of Sublethal and Lethal Effects of Polymorphonuclear Leukocytes on Escherichia coli

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Abstract

Escherichia coli ingested by PMN promptly stop growing and form no colonies in nutrient agar, but metabolize near normally for up to several hours. The bactericidal/permeability increasing protein (BPI) of PMN also inhibits E. coli growth without initial metabolic impairment. We recently showed that BPI-treated E. coli, although unable to grow in normal nutrient agar, can form colonies in this medium plus 0.1% BSA, as long as their metabolism is maintained, indicating that biochemical impairment is a better indicator of death than growth arrest (1990. J. Clin. Invest. 85:853–860). We have now reexamined the fate of ingested E. coli. Rabbit PMN ingest > 85% of several rough E. coli strains in 15 min, but > 80% of these bacteria, while unable to form colonies in conventional agar, grow normally on agar plus 0.1% BSA. Thus, the PMN under these conditions promptly stop growth of ingested E. coli without killing. Adding nonlethal concentrations of normal human serum (NHS) before, but not after ingestion, accelerates killing and, in parallel, loss of bacterial metabolism (t1/2 < 0.5 h vs. > 3 h, respectively, with and without NHS). The rapid killing of both rough and smooth E. coli pretreated with NHS is lost after C7 depletion (C7-D) and restored when C7 is replenished. Similar results are obtained with human PMN. In contrast, ingested Staphylococcus epidermidis, opsonized with either NHS or C7-D serum rapidly stops metabolizing and do not form colonies in nutrient agar with or without BSA. Respiratory burst activity is the same during ingestion of E. coli (with or without NHS) and S. epidermidis. Killing of E. coli JS (however, not of O111-B4) by BPI is also accelerated by pretreatment with NHS but not C7-D human serum. These findings indicate that late complement components are needed for efficient killing of both rough and smooth E. coli by PMN, and that BPI is the principal intracellular agent acting on ingested rough E. coli. (J. Clin. Invest. 1990. 86:631–641.) Key words: bactericidal/permeability increasing protein • complement • growth inhibition • serum albumin

Introduction

A paradox in the apparent killing of gram-negative bacteria by both intact PMN and the purified bactericidal/permeability increasing protein (BPI)1 (1, 2), located in the primary granules of both human and rabbit PMN (3), is that prompt growth arrest, as determined by direct counting in a bacterial counting chamber or by colony formation in nutrient agar plates, is not accompanied by appreciable structural and functional damage for up to several hours (1, 2, 4–6). We have recently reported that the level of metabolic activity by Escherichia coli treated with BPI correlates closely with the number of bacteria that are capable of resuming growth in the presence of 0.1% BSA, revealing that BPI initially produces only sublethal effects, and thus lack of colony formation, as determined by generally accepted assays of bacterial viability, need not reflect bacterial death (7).

Since we have shown before that the fates of E. coli exposed to purified BPI or ingested by intact PMN are nearly indistinguishable with respect to alterations in the outer envelope and lack of initial biochemical and structural damage (1, 2, 4–6), we explored in this study the possibility that E. coli, although incapable of growing under the conventional culture conditions after ingestion by PMN, may in fact also not be dead. The results show that indeed PMN are unexpectedly inefficient killers of both rough and smooth strains of E. coli, and that pretreatment of these gram-negative bacteria with nonlethal concentrations of normal human serum (NHS), but not C7-depleted (C7-D) serum, dramatically enhances the efficiency of killing of E. coli by the PMN.

Methods

Preparation of PMN. Rabbit PMN were obtained from sterile peritoneal exudates elicited in New Zealand white rabbits by injection of glycogen in physiological saline as previously described (8). Cells were collected 12–14 h later. Human PMN were collected as described (9) from heparinized venous blood obtained after informed consent from healthy volunteers. The cells were sedimented at 50g for 10 min, washed twice with HBSS without phenol red (Microbiological Associates, Inc., Walkersville, MD), and resuspended in the same medium at a concentration of 1 × 108 PMN/ml.

Radiolabeling of human C7. 200 μg purified human C7 (Calbiochem-Behring Corp., San Diego, CA) was dialyzed against a 1,000× vol of 40 mM Na2HPO4/NaH2PO4 buffer, pH 7.4, and added to a tube precoated with 75 μg iodogen (Pierce Chemical Co., Rockford, IL) (10). 1 μl sodium iodide (125I, 482 μCi, 17 Ci/mg; New England Nuclear, Boston, MA) was added, followed after 15 min on ice by 1:20 vol of 2 M potassium iodide, and the mixture was passed through a Sephadex G-25 column (Pharmacia Fine Chemicals, Piscataway, NJ) pre-equilibrated with 40 mM Na2HPO4/NaH2PO4 buffer, pH 7.4.

Preparation of human serum. Venous blood was collected from healthy volunteers after informed consent. The serum was collected after clot formation, centrifuged at 10,000 g for 20 min to remove any debris, and stored at −70°C. C7-D serum was prepared by immunoaf-

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1. Abbreviations used in this paper: BPI, bactericidal/permeability increasing protein; C7-D, complement component 7–depleted; NHS, normal human serum.
finity chromatography of freshly prepared human serum on Aff-Gel 10 (Bio-Rad Laboratories, Richmond, CA) containing covalently coupled goat anti-human C7 IgG (Calbiochem-Behring Corp.), prepared according to the instructions of the manufacturer. Recovery of unbound human serum protein (> 95%) was followed by measurement of A_{280} using a spectrophotometer (model DU-7; Beckman Instruments, Inc., Palo Alto, CA). Depletion of C7 was monitored by two independent methods: (a) by following the disappearance of 125I-labeled C7 during the chromatographic procedure, and (b) by following the loss of the ability of the serum to kill a rough strain of E. coli (15), and the restoration of the bactericidal activity toward the level of untreated serum after supplementing the depleted serum with purified C7 to a final concentration of 50 μg/ml.

**Purification of BPI.** Rabbit BPI was purified as previously described (2) and stored in 50 mM sodium acetate/acetate acid buffer, pH 4.0.

**Bacterial strains and growth conditions.** The rough E. coli strain 15, its parent smooth strain O111:B4, and the smooth strain ML-35 were grown in triethanolamine-buffered (pH 7.7–7.9) minimal salts medium (11). Other rough strains of E. coli, S15, 1602 (pldA, wild-type strain), 1303, lacking the principal envelope phospholipase, and 1303pPl232 (pldA+), a phospholipase-rich strain containing the wild-type pldA gene in a multicopy plasmid (12) were grown in physiologic saline supplemented with 0.8% (wt/vol) nutrient broth (Difco Laboratories Inc., Detroit, MI). *Staphylococcus epidermidis* (ATCC 14990; American Type Culture Collection, Rockville, MD) was grown in trypticase soy broth (BBL Microbiology Systems, Cockeysville, MD). Growth media were supplemented with 50 μg/ml chloramphenicol (Parke-Davis, Morris Plains, NJ) for growth of *E. coli* 1303pPl232 to select for bacteria containing the plasmid encoding the pldA gene product and chloramphenicol resistance. Stationary phase overnight cultures were transferred to fresh medium (diluted 1:20) and grown to mid-to-late logarithmic phase (about 3 h) at 37°C. Where indicated, the growth medium was supplemented with 1 μCi/ml [1-14C]oleic acid (57 mCi/mmol; New England Nuclear), or 5 μCi/ml [methyl-3H]thyminidine (20 Ci/mmol; New England Nuclear) plus 2 mM uridine (Sigma Chemical Co., St. Louis, MO) to label bacterial phospholipids or DNA, respectively, during growth of the bacteria in subculture. Bacterial concentrations were determined by measuring the OD_{660} in a spectrophotometer (Junior model, Coleman Instruments, Inc., Maywood, IL). The bacteria were sedimented in a clinical centrifuge for 12 min and resuspended to a final concentration of 1 × 10^7 cells/ml in sterile isotonic (0.9%) saline. In the case of radiolabeled bacteria, the cell pellet was resuspended in fresh growth medium without added radiolabeled precursors and further incubated for 30 min at 37°C before resuspension in isotonic saline. In the case of phospholipid-labeled bacteria, the bacteria were washed once with 1% (wt/vol) BSA at the end of the chase period to remove any unesterified radiolabeled precursors.

**Incubation of E. coli with BPI.** Purified rabbit BPI (2–4 μg) was added to bacteria (1 × 10^9) in 0.25 ml of physiologic saline that also contained 10% (vol/vol) HBSS, 40 mM Tris/ HCl, pH 7.5, and 0.1% (wt/vol) vitamin-free casein amino acids (Difco Laboratories) and incubated at 37°C. Where indicated, the bacteria were first incubated with the indicated preparation of fresh NHS (30 min at 37°C), centrifuged at 13,000 g for 1 min to remove unbound serum constituents, and resuspended in the same medium before adding BPI.

**Incubation of bacteria with PMN.** Typical incubation mixtures contained bacteria (5 × 10^5 in 50 μl of sterile isotonic saline) and PMN (5 × 10^5 in 50 μl of HBSS), added sequentially to 150 μl sterile isotonic saline containing 10% HBSS buffered with either 20 mM Hepes, pH 7.3, 20 mM NaHPO_4/NaH_2PO_4, pH 7.3, or 20 mM bicarbonate buffer, pH 7.3, as indicated, and were incubated at 37°C with shaking. Where indicated, human serum was also added at the given concentration after the bacteria but before the PMN. In some experiments, either the bacteria or the PMN were first exposed to serum (30 min at 37°C), followed by removal of the serum by centrifugation before incubation of bacteria plus PMN. At the indicated time points the incubation was stopped by adding an equal volume of ice-cold isotonic saline to each sample.

**Measurement of ingestion of bacteria by PMN.** *E. coli* prelabeled with either [3H]thyminidine or [14C]oleic acid were incubated at 37°C with PMN in the standard incubation mixtures. After interruption of phagocytosis at various times by adding ice-cold physiologic saline, the samples were spun at 100 g for 7 min to separate cell-associated and extracellular bacteria, and the cell pellet was washed 2 times with 1% BSA/HBSS to remove trapped and loosely adherent bacteria. In the case of [14C]oleic acid–labeled bacteria, the supernatant was then subjected to centrifugation at 13,000 g for 5 min to separate the uningested bacteria from the released products of phospholipid hydrolysis complexed by the added BSA (13). The distribution of the radiolabeled bacteria was determined by measuring aliquots of the recovered supernatant, the resuspended bacterial pellet, and the resuspended PMN cell pellet by liquid scintillation counting (5). There was no detectable conversion of radiolabeled material from TCA-precipitable to TCA-soluble form during the course of the incubation. Uptake (ingestion) is expressed as the percent of the total recovered radioactivity that is associated with the PMN cell pellet in the case of [3H]thyminidine-labeled bacteria and of the sum of the radioactivity in cell pellet plus supernatant (< 5% of total radioactivity was present in the supernatant during the first 30 min) fractions in the case of [14C]oleic acid–labeled bacteria. Recovery of total added radioactivity was always > 90%.

**Assay of bacterial viability.** After the indicated incubation at 37°C, samples were subjected to a brief (6-s pulse on ice) sonication using a micro-ultrasonic cell disruptor (Kontes Co., Vineland, NJ) at 75% of full strength. Aliquots of the sonicated suspension were serially diluted 40,000-fold in sterile isotonic saline containing 4 mM MgCl_2 and 400 μg/ml BSA (United States Biochemical Corp., Cleveland, OH). A 25-μl aliquot of the diluted sample was transferred to 5 ml of 1.3% (wt/vol) molten (47°C) Bacto-agar (Difco Laboratories) containing 0.8% (wt/vol) nutrient broth and 0.5% (wt/vol) NaCl and poured into a Petri dish. Where indicated, the molten agar was supplemented with 1 mg/ml BSA. The agar was allowed to solidify at room temperature, and bacterial viability was measured as the number of colonies formed after incubation at 37°C for 18–24 h.

**Measurement of bacterial protein synthesis.** Bacteria were incubated with PMN as described above except that the medium was supplemented with 0.5 mM cycloheximide to block protein synthesis by the PMN. At the indicated time points, [3H]-l-lysine (1 μCi/ml final concentration) were added, and the mixtures were incubated an additional 15 min. Reactions were terminated by the addition of ice-cold TCA to a final concentration of 10%, and the tubes were left on ice for 30 min. TCA-precipitable radioactive material was collected by filtration through a 0.45-μm HAWP membrane (Millipore/Continental Water Systems, Bedford, MA), washed three times with 1 ml 10% ice-cold TCA, and measured by liquid scintillation counting (5). Protein synthesis is expressed as a percentage of the total amino acid incorporation into TCA-precipitable material by untreated *E. coli* during the same 15-min period.

**Measurement of hexose monophosphatase shunt activity.** For measurement of hexose monophosphatase shunt activity of PMN during phagocytosis of bacteria, the same incubation procedure was carried out as above (1 × 10^9 PMN and 1 × 10^8 bacteria in a total volume of 0.5 ml) except that 15 ml polypropylene tubes sealed with a rubber stopper were used, and the suspension was supplemented with 5 mM KCN to block bacterial respiration and 0.03 μCi of [1-14C]-glucose (45 mCi/mmol; New England Nuclear). Incubations were started after addition of bacteria to the PMN, and hexose monophosphatase shunt activity was measured as conversion of [1-14C]glucose to [14C]CO_2 (14). Evolved [14C]CO_2 was collected in polystyrene cups suspended from rubber stoppers. At the end of the incubation, the reaction was stopped by injecting 0.2 ml of 10 N H_2SO_4 through the rubber stopper with a 21-gauge needle. After 15 min, hydroxide of hyamine (0.3 ml) (Packard Instrument Co., Inc. Downers Grove, IL) was added to the collection cups, and incubation at 37°C was continued for an additional 60 min. At this time the cups were removed, placed in counting vials, and
vigorously shaken with 9 ml of toluene-BBOT (2,5-bis-2-(5-tert-butylbenzoazolyl)-thiophene; (Packard Instrument Co., Inc.) scintillation mixture and immediately counted as described (5).

Preparation of samples for transmission electron microscopy. For electron microscopy, the incubation of PMN and bacteria was terminated by the addition of ice-cold 0.7% glutaraldehyde in Kellenberger's veronal acetate buffer (15), pH 6.1, and the samples were incubated an additional 30 min on ice. The PMN were then sedimented by centrifugation (100 g/1 min) and the pellets were resuspended in 2.5% glutaraldehyde in veronal acetate buffer and incubated for 2 h at 10°C. The cells were then washed three times with the veronal acetate buffer and postfixed with 1% osmium tetroxide in the same buffer for 16 h at 10°C. Cells were stained with uranyl magnesium acetate (0.5%) for 1 h at 20°C and embedded in 1% agar. The agar-embedded blocks were dehydrated in the presence of increasing concentrations of ethyl alcohol and replaced with propylene oxide as a last step. Specimens were infiltrated and embedded in a low viscosity epoxy resin (Polysciences, Inc., Warrington, PA) and thin sections were made with an ultratome (LKB Instruments, Inc., Gaithersburg, MD) using a diamond knife. Sections were mounted on Formvar-carbon-coated grids and double-stained with uranyl acetate and lead citrate by the method of Frasca and Parks (16). Stained samples were viewed in an Elmiskop 1A electron microscope (Siemens Corp., Iselin, NJ), and images were recorded on film 4489 (Eastman Kodak Co., Rochester, NY).

Results

Prolonged intracellular survival of E. coli J5 after ingestion by rabbit PMN. As we have shown before in studies of the fate, during phagocytosis, of other rough strains of E. coli and Salmonella typhimurium, incubation of E. coli J5 with rabbit PMN causes damage to the outer envelope (4–6) and rapid inhibition of colony formation on nutrient agar, yet the bacteria continue to incorporate 14C-amino acids into acid-precipitable material at nearly the same rate as untreated bacteria (Fig. 1, left). When the bacteria are plated in nutrient agar supplemented with 0.1% BSA instead of normal nutrient agar, nearly all the added E. coli now are able to form colonies, demonstrating that these bacteria actually have not been killed. Bacterial colony-forming ability on BSA-supplemented, but not unsupplemented, nutrient agar and incorporation of radiolabeled amino acids are preserved for several hours (Fig. 1, left), indicating that the bacteria, although jured (4, 5) and growth inhibited, are in fact metabolically active and viable. Identical results are obtained in Hepes, NaH2PO4/Na2HPO4, or bicarbonate-buffered cell suspensions (not shown).

Although demonstration of bacterial colony formation in BSA-supplemented nutrient agar required lysis of the PMN, the surprising survival of E. coli during prolonged incubation with PMN prompted us to seek further verification that these bacteria had indeed been ingested. Separation of PMN-associated from extracellular (noningested) bacteria by low speed centrifugation showed that > 85% of the viable bacteria were PMN associated (Fig. 1, right). Maximum bacterial uptake was achieved within 15 min of incubation, and the bacteria remained cell associated throughout 3-h incubations. Examination of thin sections of PMN pellets by transmission electron microscopy confirmed that PMN-associated E. coli were intracellular and also demonstrated the ultrastructural integrity of these bacteria, even after prolonged intraphagosome residence (Fig. 2).

Effect of bacterial load on intracellular survival of E. coli. To determine whether the ability of E. coli to survive after ingestion by PMN depends on the mass of bacteria ingested, the number of E. coli added to the PMN was varied over a 100-fold range, from 0.5 to 50 bacteria per PMN. Ingestion and nearly complete inhibition of growth in nutrient agar were equally efficient (> 85% within 15 min) at all ratios. After 30-min incubation at all ratios tested, nearly all the added bacteria had been ingested by the PMN and did not form colonies in nutrient agar. At this time, at least 75% of the bacteria could still form colonies in BSA-supplemented nutrient agar, independent of the bacterial load (Fig. 3). However, at later time points a slow decrease in colony formation in BSA-supplemented nutrient agar was evident at low bacterial loads (t1/2 ~ 70 and 120 min at 0.5 and 2 E. coli/PMN, respectively), but not at bacteria/PMN ratios ≥ 10 (Fig. 1, left, and Fig. 3).

Fate of other E. coli strains after ingestion by rabbit PMN. To determine whether the prolonged intracellular survival of ingested, sublethally injured E. coli J5 is representative of the fate of the E. coli species in general, we examined the effects of rabbit PMN on several other strains of E. coli (all at a bacte-
ria/PMN ratio of 10:1). The findings with five other rough strains tested, exemplified by E. coli S15, 1303, and 1602 (Fig. 4, a–c), were virtually identical to those shown for E. coli J5; i.e., phagocytosis caused almost immediate arrest of colony formation in nutrient agar, but the bacteria continued near-normal protein synthesis and colony formation in BSA-supplemented nutrient agar for at least 2 h. The one exception was E. coli 1303pP232, a derivative of E. coli 1303 that carries a high copy plasmid that bears the wild-type gene (pldA) for the outer membrane phospholipase A, resulting in a 20-fold increase in bacterial phospholipase A content (12) and a doubling of bacterial phospholipid degradation during phagocytosis (13). After ingestion, this strain showed a much more rapid (t1/2 ~ 30 min) loss of colony-forming ability in BSA-containing nutrient agar and a parallel inhibition of protein biosynthesis (Fig. 4 d). This confirms our recent study showing a role of bacterial and host phospholipases in the rate of destruction of ingested E. coli (13).

In contrast to the ability of most rough strains of E. coli to survive for extended periods of time within PMN, neither of the two smooth strains of E. coli examined survived intracellularly. Ingested E. coli ML-35 and O111:B4 promptly lost their ability to form colonies in nutrient agar both in the absence and presence of BSA, with a coincident inhibition of bacterial protein synthesis (Fig. 4 e and f). The more rapid killing of ML-35 is attributable to faster uptake of this strain.

Role of serum in intracellular killing of E. coli by rabbit PMN. The much more rapid killing, after ingestion, of smooth E. coli compared with rough E. coli was unexpected because smooth strains are generally more resistant to many of the elements of the host defense apparatus (17, 18). The more phagocytosis-resistant smooth E. coli, but not rough E. coli, require serum for ingestion by rabbit PMN (18, 19). Serum might, therefore, play a role in the more efficient killing of smooth E. coli. Indeed, addition of nonlethal amounts (Table I) of NHS to suspensions of PMN and either of two rough strains, E. coli 1303 (Fig. 5) or E. coli J5 (Table II) markedly accelerated killing (i.e., inhibition of colony formation in BSA-supplemented nutrient agar). This effect of serum was dose dependent and followed uptake (Fig. 5), indicating that the lethal damage occurred intracellularly. Pretreatment of the bacteria but not the PMN was needed for the serum effect; thus, washed E. coli that had been pretreated with serum were rapidly killed after ingestion by PMN, whereas untreated E. coli still exhibited prolonged intracellular survival after phagocytosis by serum-pretreated and washed PMN (Table II). Further, the ingested E. coli remained viable when serum was added to the PMN after phagocytosis was complete (Table II).

Role of the late components of complement in the serum-dependent killing of E. coli by rabbit PMN. Because the antibacterial properties of nonimmune serum toward E. coli are mainly attributable to the cytotoxic C5-9 complex (17), we examined the effect of interference with its formation by depleting NHS of C7. Fig. 6 (solid lines) shows that the enhancing effect of serum on PMN-mediated killing of E. coli J5 was almost completely eliminated by the removal of C7 from the serum (compare solid squares with solid triangles). Repletion of C7 to the C7-D serum fully restored the enhancing activity.
In all circumstances, the impairment of protein synthesis by the ingested bacteria closely paralleled the diminishing bacterial colony formation in BSA-supplemented nutrient agar (Fig. 6, dotted lines). A small reduction in colony formation in BSA-supplemented nutrient agar was observed after ingestion in the presence of C7-D serum (compare Figs. 1 and 6), suggesting that other serum factors may play a minor role in enhancing killing by the PMN.

Depletion of C7 also resulted in prolonged intracellular survival of smooth *E. coli*. Ingestion of *E. coli* ML-35 after opsonization with C7-D serum promptly triggered loss of colony formation in nutrient agar but not in BSA-supplemented growth medium (Fig. 7, left). Even more striking, ingested *E. coli* O111:B4 remained fully capable of growth in nutrient agar both in the presence and absence of BSA (Fig. 7, right) despite efficient uptake (Fig. 7, right, dotted line). Under these conditions, *E. coli* O111:B4, unlike ingested rough *E. coli* (4–6), also retain their outer membrane permeability barrier to antimycin D and do not undergo appreciable phospholipid degradation (data not shown), suggesting that *E. coli* O111:B4 also escape all detectable envelope alterations.

Role of late components of complement in killing of *E. coli* by human PMN. Similar experiments were carried out with human PMN to determine whether these phagocytes also require late components of complement for killing of ingested *E. coli*. As shown for rabbit PMN, after ingestion in the presence of nonlethal concentrations of NHS, the smooth *E. coli* strain O111:B4 promptly stopped forming colonies in nutrient agar either without or with BSA (Fig. 8, right, circles), but these bacteria ingested after opsonization with C7-D serum retained colony-forming ability in both growth media (Fig. 8, right, squares). Rough *E. coli* J5, opsonized with nonlethal doses of NHS, were also rapidly killed after phagocytosis (Fig. 8, left, circles). After opsonization with C7-D serum, however, a substantial fraction of these bacteria ingested by human PMN, although not growing in normal nutrient agar, remained viable in BSA-supplemented nutrient agar (Fig. 8, left, squares). As opposed to rabbit PMN (either from peritoneal exudates or peripheral blood), human PMN do not ingest rough *E. coli* in the absence of serum opsonins (9, 18). Thus, the fate of rough *E. coli* ingested by human PMN in the absence of serum could not be determined.

Killing of *S. epidermidis* by rabbit PMN. In contrast to the gram-negative bacteria tested, the gram-positive *S. epidermidis*, ingested after opsonization with either NHS or C7-D serum, rapidly lose their ability to form colonies (in both normal and BSA-supplemented nutrient agar) and, in parallel, cease protein synthesis (Fig. 9). Therefore, PMN effectively kill these gram-positive bacteria without a need for the late complement components.

Relation of intracellular bacterial killing to the respiratory burst of the PMN. Phagocytosis of rough *E. coli* by rabbit PMN in the absence of serum triggered a 10-fold increase in glucose oxidation via the hexose monophosphate shunt (Table III), reflecting activation of the respiratory burst, within the range reported in the literature (20–23). Addition of nonlethal amounts of serum to promote killing of *E. coli* did not increase the magnitude of this response, nor did ingestion and rapid killing of *S. epidermidis*. Hence, differences in bacterial killing by PMN do not reflect differences in oxidative response.

Effect of serum on killing of *E. coli* by purified BPI. BPI is a major component of the PMN’s arsenal against *E. coli*, and the early effects of BPI and intact PMN on *E. coli* are closely similar (1, 2, 6, 9, 19, 24). We therefore examined the effect of NHS on colony formation by BPI-treated *E. coli* J5 or O111:B4. Colony formation was almost immediately arrested when *E. coli* J5 were plated in nutrient agar (not shown; reference 7), but only slowly (t_{1/2} > 2 h) when plated in nutrient agar plus BSA (Fig. 10, left, open circles). Pretreatment of *E. coli* J5 with nonlethal concentrations of NHS before adding BPI accelerated loss of colony formation in BSA-supplemented nutrient agar (t_{1/2} 30–40 min) (Fig. 10, left, solid circles). This effect of serum on killing by BPI, as shown above for intact PMN, is abolished after removal of C7 (Fig. 10, left, solid squares) and is restored when the C7-D serum is repleted with purified C7 (Fig. 10, left, triangles). In contrast, pretreatment of *E. coli* O111:B4 with either C7-D or normal serum has no effect on the rate of killing by BPI (Fig. 10, right).

Discussion

The PMN is equipped with many O2-dependent as well as O2-independent antimicrobial agents, acting on a wide range of pathogens (6, 25). Despite this broad spectrum and seem-
Figure 4. Rescue of various bacteria by BSA after ingestion by rabbit PMN. *E. coli* S15, 1303, 1602, 1303pPl232, ML-35, or O111:B4 (5 × 10⁷ in 250 µl) were incubated with rabbit PMN (5 × 10⁶) at 37°C in the standard incubation medium (supplemented with 10% NHS for the latter two strains). At the indicated time points after adding PMN, samples were placed on ice. Bacterial protein synthesis (dotted lines) and viability with (solid circles) or without (open circles) BSA (1 mg/ml) in the growth medium were measured as described in Methods. The data shown represent the mean of at least two closely similar experiments. In the absence of PMN, bacterial viability was > 90%, and bacterial ¹⁴C-amino acid incorporation into acid-precipitable material ranged from 10,000 to 15,000 cpm in 15 min.
ingly redundant antimicrobial arsenal, *E. coli* that have been ingested by PMN, although unable to form colonies on nutrient agar, continue to incorporate radiolabeled precursors into macromolecules and respond appropriately to environmental changes with adjustments in biochemical activity (4–6). Because we have found that loss of colony formation, but continued biosynthesis, by *E. coli* after treatment with an antibacterial protein (BPI) of the PMN reflects bacteriostasis and not bacterial death (7), we have reexamined in this study the fate of *E. coli* during phagocytosis. We show that of the ingested *E. coli* that do not grow on nutrient agar, large portions (> 75%) are, in fact, not dead but can resume growth when the culture medium is supplemented with 0.1% BSA. Rescue is possible for several hours after ingestion is complete and parallels the extent of preservation of metabolic activity (Figs. 1 and 3).

The surprising finding that most of the apparently nonviable *E. coli* can be rescued several hours after their sequestration by PMN must raise questions about the adequacy of the performance of the PMN in vitro. However, we have found no evidence in these experiments that the PMN failed in any of its characteristic responses to added bacteria: (a) Ingestion: ingestion was prompt and essentially complete over a wide range of bacteria/PMN ratios. That the *E. coli* were truly intracellular was verified by electron microscopy (Fig. 2). Moreover, demonstration of rescue required lysis of the PMN by sonication. Further, the serum-mediated acceleration of irreversible growth inhibition (see also below) was only produced when the

![Graphs](image)

**Figure 5.** Effect of nonlethal concentrations of NHS on uptake and killing of *E. coli* 1303 by rabbit PMN. [14C]Olate-labeled *E. coli* 1303 (5 × 10^9 in 250 μl) were preincubated for 10 min at 37°C in the presence of the indicated concentration of NHS, at which time rabbit PMN (5 × 10^5) were added (t = 0 min) and the incubation was continued at 37°C. At the indicated time points after adding PMN, bacterial uptake (left) and viability with or without BSA (1 mg/ml) in the growth medium (right) were measured as described in Methods. In all cases, bacterial viability was < 1% in the absence of BSA in the growth medium, and bacterial viability in the absence of PMN was > 90%. The data shown represent the results of one of three closely similar experiments.

**Table I. Sensitivity of *E. coli* J5 to Serum Preparations**

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<thead>
<tr>
<th>Bacterial CFU</th>
<th>NHS</th>
<th>C7-D serum</th>
<th>C7-reconstituted serum</th>
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<tr>
<td>1%</td>
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<td>99.2±0.3</td>
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<td>2%</td>
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<td>4%</td>
<td>54.5±2.5</td>
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<tr>
<td>5%</td>
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<td>20%</td>
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<td>91.3±3</td>
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*E. coli* J5 (2 × 10^9/ml) were incubated at 37°C for 30 min in the presence of increasing amounts of NHS, C7-D serum, or C7-D serum to which had been added purified C7 to a final concentration of 50 μg/ml (C7-reconstituted serum). Bacterial viability was then determined as described in Methods. Data represent the mean±SEM of four experiments.

**Table II. Effect of NHS and/or PMN on Viability of *E. coli* J5**

<table>
<thead>
<tr>
<th>Bacterial CFU after 60-min incubation</th>
<th>E. coli J5 plus:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>−BSA</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>−</td>
<td>100</td>
</tr>
<tr>
<td>PMN</td>
<td>2</td>
</tr>
<tr>
<td>NHS</td>
<td>94</td>
</tr>
<tr>
<td>PMN + NHS</td>
<td>3</td>
</tr>
<tr>
<td>NHS pretreatment + PMN</td>
<td>1</td>
</tr>
<tr>
<td>NHS-pretreated PMN</td>
<td>4</td>
</tr>
<tr>
<td>PMN, followed by NHS after 30 min</td>
<td>1</td>
</tr>
</tbody>
</table>

*E. coli* J5 (2 × 10^9/ml final concentration) were included in the standard incubation mixture with PMN (2 × 10^7/ml final concentration) and/or 2% NHS. Where indicated, either the bacteria or the PMN were first treated with 2% NHS for 30 min at 37°C, followed by removal of the serum by centrifugation before beginning the *E. coli*–PMN incubation. Bacterial viability was determined after 60 min as described in Methods. Data represent the mean of at least two closely similar experiments.
serum was added to the incubation mixture before, not after, ingestion had taken place (Table II). (b) Respiratory burst: the stimulation of $^{14}$CO$_2$ production from $[1-^{14}$C]glucose is $\sim$ 10-fold over resting values (i.e., within the range found by other investigators studying antimicrobial activity of PMN in vitro [20–23]), and the same upon addition of E. coli with or without NHS, or of S. epidermidis (Table III). (c) Bactericidal activity: in contrast to the intracellular survival of both rough and smooth E. coli, S. epidermidis promptly lost viability (both in nutrient agar and in nutrient agar supplemented with BSA) and protein–biosynthetic activity (Fig. 9). We cannot attribute the prolonged intracellular survival of E. coli merely to limiting amounts of antimicrobial factors under our experimental conditions of relatively high bacteria/PMN ratios, because even at 0.5:1, half of the ingested E. coli can be rescued after 1 h (Fig. 3). Thus, the conclusion seems justified that the PMN, while effective at bacteriostasis of E. coli, is unexpectedly inefficient at killing these bacteria, raising new questions about host-defense against gram-negative bacterial infection.

Others have shown before that the fate of intracellular bacteria can be affected by serum (26, 27). This study confirms and extends these earlier observations showing that pretreatment of a wide range of E. coli strains with nonlethal concentrations of NHS contribute to the subsequent killing by rabbit and human PMN. This effect of serum is also apparent at low bacteria/PMN ratios (not shown), indicating that killing of the relatively small numbers of bacteria ingested under the usual clinical conditions is also rendered more efficient by serum. The close temporal association between loss of bacterial metabolic activity and the progressive loss of viability in albumin-supplemented nutrient agar lends credence to our contention that colony formation in this growth medium is a valid measure of bacterial death (Figs. 4, 6, and 9). Tedesco et al. have implicated the late components of complement in the killing of E. coli (O111:B4) by human PMN (27). The ineffectiveness of C7-D serum in enhancing the bactericidal activity of rabbit PMN towards several rough and smooth E. coli strains coupled with its restoration when C7 is re-added (Figs. 6 and 7) also points to the importance of the

Figure 6. Fate of E. coli 15 incubated with rabbit PMN in the presence of normal, C7-D, and C7-reconstituted human serum. E. coli 15 ($5 \times 10^8$ in 250 μl) were incubated with rabbit PMN ($5 \times 10^6$) at 37°C in the standard incubation medium with 2% NHS (circles), 4% C7-D serum (squares), or 4% C7-D serum to which purified C7 was added (final concentration = 50 μg/ml; triangles). At the indicated time points after adding PMN, bacterial viability with BSA (1 mg/ml) in the growth medium (solid lines) and protein synthesis (dotted lines) were measured as described in Methods. The data shown represent the mean±SEM of at least four experiments in the case of bacterial viability and the mean of two closely similar experiments in the case of protein synthesis. In all cases, bacterial viability was <1% in normal nutrient agar and >90% in the absence of PMN, and $^{14}$C-amino acid incorporation into acid-precipitable material in the absence of PMN was $\sim$ 12,000 cpm in 15 min.

Figure 7. Fate of E. coli ML 35 and O111:B4 incubated with rabbit PMN in the presence of C7-D serum. E. coli ML-35 (left) or E. coli O111:B4 prelabeled with $[^{14}$C]-oleic acid, $5 \times 10^8$ in 250 μl (right) were incubated with rabbit PMN ($5 \times 10^6$) at 37°C in the standard incubation medium in the presence of C7-D serum (10% for E. coli ML-35 and 20% for E. coli O111:B4). At the indicated time points after adding PMN, bacterial uptake (dotted lines) and viability with (solid squares) or without (open squares) BSA (1 mg/ml) in the growth medium were measured as described in Methods. The data shown represent the mean of at least two closely similar experiments. Bacterial viability in the absence of PMN was >90%.

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formation of a cytotoxic complement complex in the extracellular environment to the intracellular fate of ingested E. coli. This effect of the late components of complement must precede ingestion; addition of NHS after phagocytosis is complete does not result in more rapid killing (Fig. 5). These findings differ from observations on the enhancement of intracellular killing by human monocytes of Staphylococcus aureus (26), which has been attributed to the interaction of complement components with receptors on the phagocyte surface. However, in our experiments with PMN the serum effect need not depend solely on interaction with the bacteria (Table II), but may also include subsequent effects of serum components fixed to the bacteria on the PMN.

How nonlethal concentrations of NHS prepare the bacteria for more rapid intracellular destruction has not been determined. It appears likely that the cytotoxic complement complex can produce some outer membrane alterations that, while insufficient to interfere with bacterial viability, render the E. coli more vulnerable to the PMN’s antimicrobial agents (28, 29). Other envelope-damaging factors may influence the rate of intracellular destruction of gram-negative bacteria. An E. coli strain (1303pP2323), enriched in an outer membrane-associated phospholipase A (12) that hydrolyzes large portions of the bacterial phospholipids when activated during phagocytosis, also undergoes accelerated killing (Fig. 4 d; reference 13).

The remarkably similar separation of sublethal and lethal effects (and their reduction by added nonlethal amounts of NHS) when rough E. coli strains are exposed to intact PMN
and purified BPI (Fig. 10, left) is consistent with a major role for BPI in the action of PMN on these BPI-sensitive bacteria. A primary role of BPI is further supported by our recent observations showing the preferential binding to E. coli of BPI in crude PMN extracts that contain a multitude of antibacterial proteins and peptides (30), and by the finding that the separation of sublethal and lethal effects is not evident when BPI-insensitive S. epidermidis are ingested by PMN (Fig. 9). It is still unclear, however, why in the absence of serum (or late complement components) the lethal action of intact PMN on ingested E. coli, in particular smooth E. coli O1111:B4, is more limited than that of purified BPI (Figs. 4 f and 10, right) and why the late complement components dramatically enhance killing of these smooth bacteria by PMN, but not by isolated BPI.

Although the addition of serum albumin to the growth medium (nutrient agar) permits the resumption of growth by many ingested E. coli after their release from the PMN, the presence of albumin in the extracellular medium of the incubation mixture has no effect on the fate of the ingested bacteria. Thus, the length of intracellular survival of rough E. coli ingested by rabbit PMN in the absence of serum was the same whether or not albumin was added. Further, NHS accelerates the killing of E. coli by rabbit PMN despite the high albumin content of serum. Perhaps not enough albumin, in contrast to certain other serum constituents, can reach the phagolysosome. In addition, the effect of albumin may be overcome by other, more potent serum constituents.

In conclusion, these studies demonstrate that the traditional assay for bacterial viability in nutrient agar may fail to measure adequately the bactericidal activity of host-defense systems. We have shown that a simple modification of the growth medium can restore growth by phagocytosed bacteria unable to grow on nutrient agar. Therefore, PMN can promptly inhibit multiplication of ingested E. coli without actually causing death. The continued metabolic activity of ingested E. coli, and its termination when the bacteria are pretreated with nonlethal NHS, shows that loss of metabolic activity, which coincides with apparently irreversible growth inhibition, provides a valid measure of bacterial killing. The results also show that factors (late components of complement) in serum are needed to improve the relatively inefficient bactericidal activity of the PMN against gram-negative bacteria. Preliminary observations show the same enhancement by inflammatory exudate fluid, further demonstrating the integration of extracellular factors in intracellular microbicidal events. Thus, the importance of the complement system, in addition to its established role in mobilizing the cellular elements and promoting phagocytosis, includes the amplification of the destructive capabilities of the phagocyte toward ingested microorganisms. These studies set the stage for a further exam-
ination of cellular and extracellular elements in antibacterial host defense, not only in the killing, but also in the ultimate digestion and disassembly of microbial invaders.

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References


