The Unsuitability of the Uridine Incorporation Assay for the Measurement of Phagocytosis of Escherichia coli

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The uridine incorporation technique for assaying phagocytosis is based on the fact that polymorphonuclear leucocytes are impermeable to labelled uridine, and therefore ingested bacteria inside phagocytic vacuoles will be unable to take it up. Extracellular bacteria, including those adherent to the phagocytic cell surface, can do so however. Differences in uptake between bacteria alone and in the presence of phagocytic cells can be used to measure ingestion. The present paper describes the application of this technique to Escherichia coli O-86 as the test organism. It appears that with this test species, the method is unsuccessful, because exposure of the non-ingested bacteria to some soluble product of the triggered polymorphonuclear leucocytes causes a large increase in their uridine uptake rates, over that of the control bacteria. The nature of the product responsible is unknown. It is unconnected with change in the pH of the medium, is heat stable, and is only produced by polymorphonuclear leucocytes which are actively phagocytosing. It may be that a release of phagolysosome contents is responsible.

INTRODUCTION

An alternative technique to the classical ingestion test for assaying phagocytosis, which is able to discriminate between intracellular and surface adherent extracellular test bacteria, was developed by Foroozanfar et al. (1976) and Yamamura et al. (1977). This assay is based on the incorporation of a labelled tracer such as thymidine or uridine, into uningested bacteria during the post-phagocytic period. A successful assay required the system to have the following properties. (a) Tracer uptake should be linearly correlated with number of test bacteria used. (b) Phagocytic cells should be impermeable to the tracer, so that surviving intracellular bacteria are unable to take it up. Comparison of incorporation in the presence with that in the absence of phagocytic cells would then allow an estimate of phagocytic ingestion.

The technique has been used successfully with Staphylococcus aureus and rabbit leucocytes (Lam & Mathison, 1979) and the yeasts Candida albicans (Yamamura et al., 1977) and Torulopsis glabrata (Thong & Ferrante, 1977). Braconier & Odeberg (1979) modified the technique to enable both phagocytosis and intracellular killing of S. aureus and Streptococcus pneumoniae to be assayed simultaneously. They combined incorporation measurements with total bacterial viable counts following recovery of viable intracellular bacteria from phagocytic cells by osmotic lysis. Although an increase in thymidine incorporation by bacteria exposed to either intact granulocytes, or extracts of such cells was observed, it was claimed that the procedure could be made very reliable by decreasing the number of phagocytic cells used.

Similarly Bridges et al. (1970) also modified the procedure to allow simultaneous estimation of ingestion and killing of C. albicans by measurements in the presence and absence of sodium deoxycholate, which disrupts the phagocytic cells permitting viable intracellular yeasts to take up the extracellular radiolabelled uridine provided.
In the present paper, the validity of the technique of Foroozanfar (1976) was investigated using Escherichia coli as the test organism to assay phagocytosis by human polymorphonuclear leucocytes (PMNL).

METHODS

Bacteria and polymorphonuclear leucocytes. Escherichia coli O-86 was obtained from St Stephen’s Hospital, Fulham Road, London and maintained on nutrient agar slopes, and grown for experiments on Tryptic Soy broth at 37 °C. Cells were harvested in early-stationary phase, and washed in Hank's balanced salts solution (HBSS) then opsonized by 30 min incubation at 37 °C in 1 ml human serum followed by washing and resuspension in HBSS.

PMNL suspensions in HBSS were prepared by a modification of the method of Boyum (1968). Preservative-free heparin was added to 20 ml blood to a concentration of 10 μl ml⁻¹. One part of 6% (w/v) dextran in saline was added to four parts of blood by volume, and after 30 min the leucocyte-rich plasma was separated by centrifuging at 300 g for 7 min. After resuspension in 4 ml HBSS, pellets were layered on to 3 ml Ficoll-Paque, and centrifuged for 30 min at 400 g. The PMNL containing pellet was resuspended in 0-83% Tris buffered ammonium chloride, pH 7.2 for hypotonic lysis of residual erythrocytes, and after 5 min the pellet washed twice in HBSS and adjusted to the required cell density. Further details of the procedure are given in Fazeli (1982).

Phagocytosis assay by uptake of tritiated uridine. Unless otherwise stated, assays were set up in sterile siliconized screw-capped glass bottles. Incubation mixtures containing human PMNL from healthy volunteers and preopsonized E. coli in HBSS were incubated on a shaking incubator at 37 °C. Similar control tubes lacking either bacteria or PMNL were also incubated. After allowing phagocytosis to proceed for 20 min, further incubation for 60 min with 1 μCi [5,6-3H]uridine ml⁻¹ (37 kBq ml⁻¹; 27 pmol ml⁻¹) was carried out to allow incorporation of the radiolabel. Samples (1 ml) were removed at timed intervals during this incubation, and PMNL and bacteria were recovered on cellulose nitrate membrane filters (0.45 μm pore-size, 2.5 cm diameter) in a vacuum filtration manifold. Both the filters and the filtrates were saved, and the filters washed with 20 ml HBSS, and dissolved in 5 ml Bray's scintillation fluid (Bray, 1960). Samples (500 μl) of the filtrates were dissolved in 10 ml Bray's fluid and radioactivity was quantitated using either a Beckman LS233 liquid scintillation counter (at 8 °C), or a Packard 460 CD liquid scintillation counter at room temperature. Samples were counted twice for 10 min each, and observed count rates corrected for sample quenching by the external standard method.

Uridine permeability of E. coli exposed to resting or phagocytosing PMNL. Mixtures of PMNL and bacteria, or PMNL alone were incubated for 20 min at 37 °C, then centrifuged at 10000 g and room temperature for 10 min in a Micro-angle centrifuge (Baird Tatlock, London). Supernatants were mixed with pellets of opsonized bacteria, and after 20 min at 37 °C the mixtures were further incubated with [3H]uridine and treated as described previously.

RESULTS

Uridine incorporation by E. coli O-86

Varying concentrations of opsonized bacteria in a total volume of 2 ml HBSS were mixed with [3H]uridine at 37 °C for 60 min, and at intervals samples were taken and processed as described in Methods.

Uridine incorporation after 30 min incubation, as a function of bacterial numbers is shown in Fig. 1. The uridine incorporation per cell by opsonized E. coli after 30 min incubation varied between 0.106 and 0.128 amol. The average concentration factor (intracellular concentration divided by extracellular concentration) inside the bacterial cells was 25-6 × 10³ on the assumption of an internal bacterial cell volume of 0.78 μm³ all assumed to be available to the intracellular uridine, and assuming insignificant consumption of the uridine by the metabolism or for macromolecular synthesis during the incorporation period.

Uptake by the heat-killed bacteria (56 °C for 90 min) was only about 1% of that by the living bacteria in the same time (Fig. 2).

Uridine uptake by E. coli in a phagocytosis mixture

Phagocytic mixtures containing 5 × 10⁶ PMNL and 5 × 10⁷ c.f.u. E. coli O-86 in total volume of 2.5 ml HBSS were incubated in duplicate. After 20 min at 37 °C one tube had [3H]-uridine added as described above, and the other was centrifuged at 300 g for 5 min, and the pellet washed twice with HBSS to remove unattached extracellular bacteria, resuspended in the original volume of HBSS, then reincubated with [3H]uridine as before. Control mixtures of
Phagocytosis assayed by uridine incorporation

Fig. 1. Incorporation of $^3H$uridine by varying numbers of E. coli O-86 after 30 min incubation at 37 °C. Stationary phase opsonized E. coli were resuspended in HBSS with labelled uridine at 20 nM. Samples were withdrawn at timed intervals, filtered and washed. Filter and filtrate incorporated radioactivity was measured as described in the text.

![Graph](image)

**Fig. 1.** Incorporation of $^3H$uridine by varying numbers of E. coli O-86 after 30 min incubation at 37 °C. Stationary phase opsonized E. coli were resuspended in HBSS with labelled uridine at 20 nM. Samples were withdrawn at timed intervals, filtered and washed. Filter and filtrate incorporated radioactivity was measured as described in the text.

![Graphs](image)

**Fig. 2.** Incorporation of $^3H$uridine into a phagocytic mixture. Time course of incorporation of uridine at 20 nM into a mixture containing $5 \times 10^6$ PMNL ml$^{-1}$ and $5 \times 10^7$ E. coli O-86 ml$^{-1}$ (○), opsonized E. coli alone (●), PMNL alone (■), E. coli attached to the PMNL surface but not ingested (□), and an equal number of heat killed E. coli (△).

**Fig. 3.** $^3H$uridine incorporation by phagocytic mixture. $^3H$Uridine incorporation by $5 \times 10^7$ S. aureus ml$^{-1}$ (●), by $5 \times 10^6$ PMNL ml$^{-1}$ (■), and by a phagocytic mixture containing both (○). Experimental conditions identical to those in which E. coli was used as test organism.

either bacteria alone or PMNL alone were treated similarly (Fig. 2). Incorporation into bacteria in phagocytic mixtures was always higher than that into the bacteria alone, although the PMNL were impermeable to the labelled uridine. There was a variation in uridine uptake by PMNL observed with PMNL from different donor individuals, but such uptake was always insignificant compared with that by the bacteria. Uptake by PMNL plus associated extra-
Fig. 4. Effect of supernatant from phagocytosis mixture on [3H]uridine uptake by E. coli. Incorporation by E. coli O-86 (5 x 10^7) suspended in fresh HBSS (○), or the supernatant from a phagocytosis mixture (●). Each point is the mean of five experiments, the error bars show the S.E.

Fig. 5. Influence of the suspending medium on [3H]uridine uptake by E. coli O-86. Uptake by 5 x 10^7 stationary phase opsonized E. coli in HBSS containing 1 μCi [3H]uridine (20 nM) in a total volume of 2 ml. Assay mixtures were PMNL alone (■), bacteria plus PMNL which had been incubated 20 min before addition of uridine (○), and bacteria alone suspended in fresh HBSS (□), supernatant from resting PMNL (▲), supernatant from phagocytosing PMNL (●), and heat-treated supernatant from phagocytosing PMNL (△).

cellular bacteria was also higher than that by PMNL alone, and this is attributed to uptake by the adherent bacteria on the PMNL surface.

_Uridine incorporation by Staphylococcus aureus C-18 in the phagocytic system_

As a control for the assay procedure, an identical set of experiments was carried out using _S. aureus_ C-18 as the test organism. This species has previously been used successfully in this type of assay (Lam & Mathison, 1979).

The result is shown in Fig. 3, from which it can be seen that the uptake by phagocytic mixtures was substantially less than that by bacteria alone, as would be expected. The amount of incorporation by a given number of _S. aureus_ was also considerably less than that by a corresponding number of _E. coli_ in an identical experiment.

_Uridine incorporation by uningested _E. coli_ exposed to PMNL_

To investigate the influence of exposure to granulocytes on the permeability of _E. coli_ to uridine, 5 x 10^7 stationary phase opsonized bacteria were suspended in cell-free supernatant from a phagocytosis mixture as described above. The incorporation of labelled uridine by such a suspension was greatly increased compared with the incorporation by a corresponding suspension in unused HBSS (Fig. 4). Heating of the HBSS supernatant from phagocytosis mixtures to 56 °C for 60 min produced no loss of the ability to increase the bacterial permeability to uridine (Fig. 5), and the same effect on _E. coli_ permeability was also produced by supernatants from phagocytosis mixtures in which PMNL were ingesting _S. aureus_ C-18.

The supernatants from resting PMNL suspensions after 30 min incubation at 37 °C caused no
such enhancement of uridine uptake (Fig. 5), thus the effect may be attributed to some soluble extracellular, heat-stable product of the triggered phagocyte.

**DISCUSSION**

The uridine (or thymidine) incorporation assay has been used with a variety of test organisms. The present study was undertaken to assess the validity of the technique for phagocytosis of *E. coli*. However, the degree of incorporation of radiolabelled uridine into bacteria which had been exposed to triggered PMNL far exceeded that by unexposed *E. coli* due presumably to the action of some product of the active PMNL metabolism, and was independent of the bacteria : PMNL ratios (data not shown). In contrast, using *S. aureus* as the test organism in an identical series of experiments, the technique worked well, and allowed an estimate of ingestion from the reduction in uptake in the presence of phagocytic cells. It thus appears that the technique cannot be used as it stands with *E. coli* as the test organism, and further, that it is necessary to test, when using the technique for the first time with a new species, that a similar effect does not take place. It is possible that the assay could be modified by comparing the uptake in phagocytosis mixtures not with that by bacteria alone, but rather with the uptake by a suspension of bacteria incubated in the supernatant from an identical phagocytosis mixture. However, since the uningested bacteria in the mixture are only affected by a product of triggered PMNL, and not of resting cells, presumably the agent responsible for the effect is only produced from the start of the incubation, and is changing in concentration during the assay. It is thus difficult to see how a matched control could be created to compare with the uptake by the mixture.

The effect on uptake rates cannot be attributed to growth of the bacteria, since the *E. coli* used were at stationary phase, and exhibited no growth at least for 3 h in this medium (Fazeli, 1982). An unequivocal increase in the rate of uptake of [3H]uridine by *E. coli* was produced by exposure to phagocytosing PMNL, but not by resting PMNL. Since it was possible to induce such an increase by exposure of the bacteria to the supernatant of phagocytosing PMNL alone, it is clear that a soluble product of the metabolism of triggered PMNL is responsible. The responsible product appears to be produced (and to be active against *E. coli*) irrespective of the nature of the test organism used to trigger the PMNL, that is, it appears to be a general property of the phagocytic response. The agent responsible is stable to heat at 56 °C for 1 h, which suggests that it is not a heat-labile protein. Further the effect is not one of simple alteration of the medium pH, since there was no change in the pH of the medium for at least 2 h of phagocytosis (Fazeli, 1982).

A phagocyte-induced change in membrane permeability of Gram-negative bacteria has previously been reported using intact PMNL (Beckerdite et al., 1974), disrupted PMNL, or fractions from PMNL cells (Elsbach et al., 1974; Weiss et al., 1976) and by a combination of serum and antibiotics (Dutcher et al., 1978). The killing of *E. coli* by PMNL is accompanied by an increase in permeability of the bacterial envelope. This may be due to enzyme attack or degradation of structural components of the membrane by reactive species, and may take place intraphagosomally or (as must be the case here) extracellularly by release of the responsible factors into the suspending medium. Factors known to be released into the medium during phagocytosis include plasminogen activator (Granelli-Piperno et al., 1977), lysozyme and β-glucuronidase (Wright & Malawista, 1972; Okamura et al., 1979), N-acetyl-β-glucosaminidase (Johnston & Lehmeyer, 1976) lactoferrin and myeloperoxidase (Leffell & Spitznagel, 1975), hydrogen peroxide and superoxide anion (Root et al., 1975; Hafeman & Lucas, 1979). Weiss et al. (1976) have reported a highly purified rabbit PMNL fraction able to increase *E. coli* envelope permeability, and demonstrated that it consisted of cationic proteins and phospholipase A2 (Elsbach et al., 1979).

Phagocytic vacuole contents may be released simply as a result of cell death, or by regurgitation during phagocytosis, reverse endocytosis (Goldstein, 1976), or may be induced by complement and immunoglobulin (Henson, 1971). It is probably unlikely that the *E. coli* envelope damage results from antibody/complement action, since the effect is not observed with
supernatants from resting PMNL and opsonized bacteria. It could be a result of the myeloperoxidase/hydrogen peroxide system however (Sips & Hamers, 1981).

Elsbach et al. (1974) found that exposure of E. coli to granulocyte extracts increased the rate of incorporation of labelled precursors into RNA by more than 15-fold, and into protein and DNA about twofold. This effect took place even 30 min after the bacteria had lost the ability to multiply.

Thus the present results, which mean that the assay technique is not suitable for measuring phagocytosis of E. coli are supported by these previous observations, and it appears that an increased rate of uridine incorporation in bacteria exposed to phagocytosing PMNL results from an effect on the envelope of cells of the non-ingested population brought about by one or more reactive species released from the PMNL.

REFERENCES


Phagocytosis assayed by uridine incorporation

