A Search for the Bacterial Mucopeptide Component, Muramic Acid, in Chlamydia

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INTRODUCTION

Chlamydiae are obligate intracellular parasites. They multiply by binary fission in a complex developmental cycle that involves a series of morphological forms. The mature infectious form, the elementary body, has a rigid wall similar in ultrastructure and amino acid content to the walls of Gram-negative bacteria (Manire & Tamura, 1967; Murray, 1968; Tamura, Matsumoto, Manire & Higashi, 1971). The intermediate form in multiplication, the reticulate body, lacks a rigid wall and is non-infectious.

Perkins & Allison (1963) presented chemical evidence for the presence of muramic acid in several strains of Chlamydia: the P-4 pigeon strain of psittacosis, the Baker strain of feline pneumonitis, the T'ang strain of trachoma and the Nigg strain of mouse pneumonitis. The last-named strain was grown in tissue cell cultures, but the others were propagated in the yolk sacs of embryonate eggs. Jenkin (1960) reported traces of muramic acid in intact particles and walls of the Cal-10 strain of meningopneumonitis grown in the allantoic cavity of embryonate eggs, but no technical details were given. However, Manire & Tamura (1967) were unable to detect muramic acid in purified intact elementary bodies or walls of the same strain grown in L cell cultures. Their method could not detect less than 20 μg muramic acid and they concluded that the meningopneumonitis wall contained less than 0.2% of this compound.

The presence or absence of muramic acid in the genus Chlamydia has important implications for taxonomy, for theories of the replication of these organisms and for the mode of action of antibiotics to which they are susceptible. We have now used a very sensitive test to look for muramic acid in trachoma and meningopneumonitis agents grown in L cells.

METHODS

The fast-killing variant (Taverne, Blyth & Reeve, 1964) of trachoma agent strain TRIC/2/GB/MRC-4/ON was grown in L cell cultures in suspension (Garrett & Harrison, 1973). The infected cells from six batches of 4 l of infected L cells were homogenized in a Teflon grinder and yielded a total of 3.4 × 10¹² inclusion-forming units (i.f.u.) (Furness, Graham & Reeve, 1960; Taverne & Blyth, 1971). The agent was purified by a slight modification of the method of Tamura & Higashi (1963); before and after treatment with nucleases and trypsin, the preparation was centrifuged at 7500g for 60 min at 4 °C through a 6 cm column of 25%
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(w/v) sucrose in 0.03 M-tris-hydroxymethylaminomethane-HCl, pH 7.4. A total of $1.3 \times 10^{12}$ elementary bodies was recovered, as measured by dark field microscopy (Reeve & Taverne, 1962). Electron microscopy showed that the purified preparations contained approximately as many reticulate bodies as elementary bodies. The dry weight of the total yield was estimated at 25 to 50 mg.

The Cal-10 strain of meningopneumonitis was grown in L cells, and elementary bodies and walls were isolated and purified by the method of Manire & Tamura (1967) by Miss Tanaka at the University of North Carolina, Chapel Hill, U.S.A. Ninety mg dry weight of elementary bodies and 10 mg wall material were examined for muramic acid.

A modification of the methods of Park & Hancock (1960) and Mandelstam (1962) was used in an attempt to concentrate a mucoprotein fraction from elementary bodies and walls. All products were collected by centrifugation at 10000 g for 15 min at 4 °C. The chlamydia preparations were suspended in 5 % (w/v) trichloroacetic acid (TCA) at 0 °C for 1 h. The insoluble material was washed with water and with 75 % (v/v) ethanol. It was then heated in 5 % (w/v) TCA for 6 min at 90 °C. The product was washed with water and resuspended in ethanol:ether (3:1, v/v) at 20 °C; after 18 h it was treated with 10 M-urea, 0.29 M-2-mercaptoethanol and 0.54 M-iodoacetic acid according to the method of Mandelstam (1962), and then incubated with crystalline pepsin (Sigma London Chemical Co. Ltd, Kingston-upon-Thames) in 0.05 M-HCl for 18 h at 37 °C. The product was then washed once with 2 M-NH₃, once with water, three times with 90 % formic acid and twice with ether.

The preparations were hydrolysed in sealed ampoules in 4 M-hydrochloric acid at 100 °C for 4 h. Hydrochloric acid was removed by evaporation under reduced pressure. The hydrolysates were streaked on Whatman no. 3MM chromatography paper, with markers of authentic muramic acid, and were run overnight in butan-1-ol:acetic acid:water (63:10:27, by vol.). Areas of the chromatogram with the same $R_f$ as the muramic acid markers were eluted with water. When Escherichia coli strain B organisms were fractionated in this way the final preparations contained 2 % muramic acid, estimated colorimetrically by the method of Rondle & Morgan (1955). It was assumed that chlamydia elementary bodies would contain insufficient muramic acid for colorimetric analysis: the eluates from the chromatograms of chlamydia hydrolysates were therefore treated with aqueous potassium [3H]borohydride (The Radiochemical Centre, Amersham, Buckinghamshire) and examined for [3H]muramicitol. After incubation at 20 °C overnight the reaction mixture was acidified, dried under reduced pressure and washed four times with aqueous methanol. It was then run on a column of Dowex 50 (H+) (×8, 200–400 mesh) equilibrated and eluted with 0.1 M-pyridine–acetic acid, pH 2.8 (Hughes, 1970). Authentic muramic acid yielded [3H]muramicitol. The radioactivity of each fraction was measured in an Intertechnique ABAC SL 40 scintillation counter. Under these conditions the activity of a peak of [3H]muramicitol from 0.1 μg authentic muramic acid was 200 counts/min. Eluates of the test material that showed activity in the same region as authentic [3H]muramicitol were streaked on either Whatman no. 1 or Whatman no. 4 paper with markers. They were run in butan-1-ol:pyridine:water (6:4:3, by vol.) overnight. The paper was cut into 0.5 cm strips to locate radioactivity with the same $R_f$ value as muramicitol. The efficiency of counting tritium on paper strips was 15 % of that in liquid.

RESULTS AND DISCUSSION

Hydrolysates of our trachoma agent preparations were fractionated and reduced as described. A very small amount of tritiated material that corresponded with [3H]muramicitol was detected by chromatography on a Dowex 50 (H+) column and on paper. Formation of
this product from a fraction co-chromatographing with muramic acid strongly suggested
the presence of muramic acid in trachoma agent but paucity of purified agent precluded all
the tests necessary for absolute proof. However, our results allow estimation of the maximum
amount of this sugar present in trachoma agent. The material eluted from the Dowex
column accounted for no more than 0.2 μg muramic acid. Therefore, if we assume that the
wall comprises 20% of the dry weight of trachoma elementary bodies, no more than 0.04% of
the wall is muramic acid. We found no muramic acid in whole elementary bodies or in
the purified walls of meningopneumonitis agent. Although mucopeptide certainly confers
rigidity on the walls of Gram-positive bacteria, it is doubtful whether this component, alone,
serves the same purpose in Gram-negative bacteria, in which only 5 to 10% of the wall is
mucopeptide (Ghuysen, 1968); in the chlamydiae we examined, mucopeptide accounted
for no more than 0.1% of the dry weight of the wall.

Although Perkins & Allison (1963) detected muramic acid in several strains of Chlamydia,
Manire & Tamura (1967) found none in meningopneumonitis elementary bodies or walls.
There are three possible explanations for this discrepancy:

(i) The assay method used by Manire and Tamura may have been insufficiently sensitive
to detect the amounts of muramic acid present; they estimated that there could be no more
than 0.2%, muramic acid in purified meningopneumonitis walls. Perkins & Allison gave no
estimation of the amount of muramic acid in their preparations but their methods detected
smaller amounts of the sugar.

(ii) Autolysis during preparation of the walls used by Manire and Tamura may have
dissolved the mucopeptide that was then lost during subsequent purification steps.

(iii) Reticulate bodies may contain mucopeptide, and elementary bodies little or none.
Envelopes of bacterial spores differ from those of the vegetative form and contain less muco-
peptide (Gledhill, 1967); the transition from reticulate to elementary bodies may resemble
sporulation in this respect at least. Our preparations of trachoma agent contained reticulate
bodies and elementary bodies; the meningopneumonitis agent preparation contained only
elementary bodies. The Chlamydia preparations used by Perkins & Allison (1963) probably
contained reticulate bodies; Lawn, Blyth & Taverne (1973) observed all the developmental
forms of trachoma agent in yolk sac preparations which had been partially purified by
trypsin treatment and differential centrifugation, and in trachoma-infected BHK-21 cells.

Mucopeptide may play a role in maintaining the integrity of the wall during development
of chlamydiae but it is extremely unlikely that it acts as a mechanical support. Tribby (1970)
proposed that reticulate bodies of meningopneumonitis agent have flexible walls containing
uncrosslinked mucopeptide. She also suggested that crosslinking of the mucopeptide is
necessary for the formation of the more rigid walls of elementary bodies, and that pencillin
interferes with this reaction. However, Tamura & Manire (1968) showed that pencillin
inhibits the replication of reticulate bodies and not only the formation of elementary bodies.
All pencillin-sensitive bacteria studied so far have the same general mucopeptide structure
of polysaccharide ‘backbones’ composed of alternate residues of N-acetylglucosamine and
N-acetylmuramic acid joined by a network of crosslinked peptides. Pencillin is now believed
to inhibit transpeptidation of the mucopeptide of these bacteria (Strominger, Izaki,
Matsuhashi & Tipper, 1967) and growing cells synthesize a defective wall and eventually
lyse. Pencillin could affect chlamydiae in a similar way, but it may inhibit the crosslinking
of peptides that are not attached to a polysaccharide containing muramic acid. Now that it
is agreed that chlamydiae are more closely related to bacteria than to viruses, some authors
too readily ascribe well-established properties of free-living bacteria to them. Only gross
chemical analyses of the envelope of one strain of meningopneumonitis have been done; the
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presence of mucoprotein is still in doubt, and it could well be that in these unique microorganisms penicillin acts in a unique way. Until more is known about the chemical structure and the biosynthesis of chlamydial cell walls, we should avoid drawing too close an analogy between Chlamydia and bacteria.

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