The Chemical Composition of Lambda in *Paramecium aurelia*, Stock 299

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(Received 25 February 1963)

SUMMARY

*Paramecium aurelia* containing λ particles was grown in an axenic medium; the λ particles were purified by passing homogenized organisms through a column of the anion exchange resin ECTEOLA. The λ particles were stained heavily with pyronin Y and weakly with the Feulgen procedure, *in vivo* and *in vitro*. Chemical analysis of the λ particles revealed that a single particle contained: 0.026 picogram (pg.) DNA, 0.230 pg. RNA, 4.0 pg. protein, 0.350 pg. carbohydrate, 5.6 pg. phospholipid. The RNA/DNA ratio in the λ particle was 8.8 as compared with a RNA/DNA ratio of 11.3 in the κ particle. Both particles contain about the same amount (0.0049 pg., and 0.0045 pg./μ³, respectively) of DNA, yet the λ particles stained very faintly with the Feulgen procedure, while the κ particles stained very heavily.

INTRODUCTION

Several stocks of *Paramecium aurelia* have been shown to possess certain cytoplasmic inclusions named κ, π, μ and λ. The longest known and most widely studied particle κ (Sonneborn, 1959) is a Feulgen-positive rod-shaped body, randomly scattered throughout the cytoplasm; its presence confers protection against, and results in the liberation of, a toxic agent. Particle π is a mutant form of κ; its presence neither confers protection against, nor results in, the liberation of a toxic substance. Beale & Jurand (1960) published a study of the μ particle, which occurs in syngen 1, stock 540. This stock is a mate-killer, having the same properties as the mate-killers earlier described by Siegel (1953). The presence of a cytoplasmic particle, λ, in stock 299 of syngen 8, was discovered by Schneller (1958); the λ-containing killer organisms cause sensitive organisms of stock 139 to lyse within 20 min. Ever since the discovery of κ particles there have been discussions about their nature. The features used for distinguishing endogenous genetic particles from intracellular symbionts have proved to be indecisive criteria. As part of an investigation to try to establish the phylogenetic relations of the various cytoplasmic particles, the determination of the chemical constitution of the λ particle was undertaken, in order to compare it with the previously determined composition of κ (Smith & van Wagendonk, 1962). The advent of an axenic medium for the culture of particle-bearing paramecia (Soldo, 1961) made it possible to determine the chemical constitution of these particles without the danger of contamination by bacterial constituents.
METHODS

An axenic culture of *Paramecium aurelia* stock 299, syngen 8, containing λ particles was obtained from Dr A. T. Soldo (Biological Research Division, Schering Corporation, Bloomfield, New Jersey). The organisms were maintained at 27° in the axenic medium described by Soldo (1961). This medium consists of a mixture of salts, water-soluble vitamins, stigmasterol, proteose peptone, Edamine S (Sheffield Chemicals, Norwich, New York) and a non-dialysable fraction of a heated yeast extract (Miller & van Wagendonk, 1956). Subcultures were made every 5 days by inoculating 3.0 ml. fresh medium with 0.1-0.5 ml. from a previous old culture. Large cultures of paramecia were grown in 500 ml. Soldo medium contained in 2.5 l. Erlenmeyer flasks. The population density of these cultures after 5 days ranged from 7000 to 10,000 organisms/ml. The paramecia were periodically checked for the presence of λ particles by staining with aceto-orceine. Fixed preparations were stained by the methyl green-pyronin Y technique of Kurnick (1953) or by the Feulgen procedure as modified by Rafalco (1946). The time of hydrolysis in the latter procedure is critical and should not exceed 5 min.; otherwise negative results are obtained.

Removal of ribonucleic acid from the λ particles was accomplished by treating the organisms for 30 min. at 37° with 1% (w/v) solution of crystalline ribonuclease in water (previously heated for 10 min. at 90° to inactivate any contaminating deoxyribonuclease).

Large quantities of pure particles were obtained by a method modified from the one used by Smith (1962). Cultures (3-4 l.) of λ-bearing paramecia were centrifuged at 3000 g for 10 min.; the supernatant fraction was then subjected to centrifugation at 15,000 g for 20 min. to sediment λ particles which might have been liberated from ruptured organisms. This sediment was added to the first sediment, and the combined fractions suspended in 15 ml. sterile saline solution (0.9%, w/v). The above centrifugation procedure was repeated twice. The final sediment was suspended in 10 ml. 0·1 m-phosphate buffer (pH 6·9). A homogenate was prepared by using the syringe technique (Sonneborn, 1950) and the homogenate was passed through an ECTEOLA (epichlorohydrin triethanolamine-cellulose) column, 2 cm. in diameter and 1 cm. high, prepared 20 min. before use by suspending in 1·5 g. ECTEOLA in 50 ml. phosphate buffer. The effluent was passed several times through a freshly prepared column until the particles appeared clean when viewed under brightly illuminated phase-contrast.

The clean particles were separated from the phosphate buffer by centrifugation (25,000 g.) and suspended in isotonic salt solution for the determination of the total number present (by counting a measured sample) and for the cytochemical and chemical analyses. The procedure outlined by Paul (1958) was used for the determination of the major constituents of the λ particles. The amount of lipid present was obtained by a modification of Bloor's colorimetric procedure (1947). Organic phosphorus was determined on a measured sample of the ethanol + ether soluble fraction by the method of Griswold, Humoller & McIntyre (1931); this value multiplied by 25 gives the amount of phospholipid present (Lees, 1957). Carbohydrate was measured according to the procedure of Trevelyan & Harrison (1952). Total nucleic acid was estimated by determining the absorption at 260 m\( \mu \) of measured
<table>
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<tr>
<th>Expt. no.</th>
<th>Total no. of particles</th>
<th>Lipid phosphorus</th>
<th>Phospholipid</th>
<th>Carbohydrate</th>
<th>Nucleic acids</th>
<th>DNA</th>
<th>RNA</th>
<th>Protein</th>
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<td></td>
<td>T*</td>
<td>P*</td>
<td>T</td>
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<td>0.35</td>
<td>0.23</td>
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* T = total lipid in µg; P = lipid per particle in picograms, etc.
samples of the perchloric acid fraction by a Beckman D.U. spectrophotometer. The method of Ceriotti (1955) was used to determine deoxyribonucleic acid and ribonucleic acid. The residue of the fractionation procedure was digested with the acid mixture of Brüel, Holter, Linderstrom-Lang & Rozits (1947) and the amount of ammonia formed measured with Nessler reagent; the obtained value was multiplied by 6.25 to give the amount of protein present.

RESULTS

Cytochemical observations. Large numbers of rod-shaped particles, randomly distributed throughout the cytoplasm, were visualized when organisms of the killer stock 299 were stained with aceto-orcein (Pl. 1, fig. 1). These particles also stained deeply with pyronin Y (Pl. 1, fig. 2), indicating the presence of large amounts of ribonucleic acid. After treatment with ribonuclease no pyronin Y positive-particles were observed (Pl. 1, fig. 3). Results obtained with the Feulgen procedure were much less pronounced. A careful study of Pl. 1, fig. 4, shows that there were small Feulgen-positive rods scattered throughout the cytoplasm. The random distribution and the shape of these bodies suggest that the Feulgen-positive material was most probably associated with the λ particles. Particles obtained by the purification procedure and placed directly on slides stained positive with the pyronin Y + methyl green mixture.

Chemical determinations. The results of nine chemical analyses of purified λ particles are shown in Table 1. The particles are characterized by a relatively large content of protein and lipid. Most of the lipid was present as phospholipid, whose specific type is not known. The concentration of protein and lipid tended to fluctuate, possibly with the rate of reproduction of the particles. The amount of carbohydrate was low. The average content of deoxyribonucleic acid and ribonucleic acid/single particle was 0.026 and 0.23 pg., respectively.

DISCUSSION

The most interesting results obtained in this work are those concerning the deoxyribonucleic acid (DNA) and ribonucleic acid (RNA) content of the λ particle. A comparison of the cytochemical and chemical data on the constitution of the two killer particles κ and μ, with the data obtained for the λ particle, reveals that all three rod-shaped particles contain DNA. In the particles κ and μ, the DNA was first cytochemically demonstrated; from these results two conclusions were reached: (a) DNA is distributed uniformly throughout the κ particle; (b) the κ particle contains ‘massive amounts of DNA’ (Sonneborn, 1959). From cytochemical tests the composition of the λ particle appears to be fundamentally different, since the Feulgen stain indicates that DNA is present in very small amounts. The chemical analysis of the κ and λ particles revealed however that κ contains less DNA and more RNA per unit volume than does the λ particle.

Various explanations for the discrepancy between the cytochemical observations and the chemical data can be suggested: (a) the λ DNA is more resistant to acid treatment, like the chloroplast DNA described by Chiba & Sugahara (1957); (b) the λ DNA may be dispersed through the particle very dilutely; (c) the λ DNA may be in a precursor state, not stainable by the Feulgen procedure (Vendrely, 1955). The
first explanation is not acceptable because it was found in the present work that the 
λ DNA was very labile under acid treatment. It does not seem very likely that 
DNA would be uniformly distributed in the λ particle and therefore more dilute 
than in the κ particle. On the contrary, a careful examination of Pl. 1, fig. 4, shows 
that the λ DNA appears to be concentrated at one end of the particle. Chayen (1959) 
remarked that the drastic conditions of the Feulgen test might disrupt the cell 
organization and cause a re-location of the DNA or even a diffusion of cytoplasmic 
DNA into the nucleus. The uniform distribution of DNA throughout the κ particle 
may be a result of similar events and this might lead to an erroneous interpretation 

Table 2. Comparison of DNA and RNA concentration in κ and λ particles

<table>
<thead>
<tr>
<th></th>
<th>κ particle</th>
<th>λ particle</th>
</tr>
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<tbody>
<tr>
<td>Size</td>
<td>2.25 x 10^{-μ}</td>
<td>6.75 x 10^{-μ}</td>
</tr>
<tr>
<td>DNA/unit volume (μg)</td>
<td>0.0045 pg.</td>
<td>0.0040 pg.</td>
</tr>
<tr>
<td>RNA/unit volume (μg)</td>
<td>0.058 pg.</td>
<td>0.043 pg.</td>
</tr>
<tr>
<td>RNA/DNA ratio</td>
<td>11.3</td>
<td>8.8</td>
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</table>


of the cytochemical observations. The lability of λ DNA under acid treatment 
might have prevented the re-location of the DNA throughout the particle. The 
available evidence does not rule out the possibility that the lambda DNA is present 
in a precursor form. It seems clear from the present work and that by Smith & van 
Wagtendonk (1962) that the determination of the chemical composition of these 
cytoplasmic particles can at best furnish an ambiguous criterion for the biological 
status of these particles. Not until it can be shown that these cytoplasmic particles 
have a metabolism of their own, either independent of, or co-ordinated with, that 
of the host will it be possible to classify these particles as viruses, parasites (Sonne-
born, 1961) or as endogenous components of the cell.

This work was supported by grants from the U.S. National Institutes of Health 
(NIH E3844-C1-C2) and the National Science Foundation (NSF G14569). One of 
us (R.B.T.) held a National Defense Act Fellowship. This report is based on a thesis 
submitted by R. B. Tanguay in partial fulfilment of the requirements for the 
degree of Master of Science in the Graduate School of the University of Miami, 
Coral Gables, Florida, U.S.A.

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**EXPLANATION OF PLATE**

Fig. 1. *Paramecium aurelia*, stock 299. The rod-shaped particles found throughout the cytoplasm and overlying the deeply stained macronucleus are the λ particles. Stained with Aceto-orcein. Magnification, ×1000 on film.

Fig. 2. *Paramecium aurelia*, stock 299. The rod-shaped particles in the cytoplasm deeply stained with Methyl green-pyronin Y mixture represent the λ particles. Magnification ×500 on film.

Fig. 3. *Paramecium aurelia*, stock 299 treated with 1% RNA-ase solution for 30 min. at 37° and subsequently stained with Methyl green-pyronin Y mixture. No λ particles can be seen. Magnification, ×500 on film.

Fig. 4. *Paramecium aurelia*, stock 299, stained with the Feulgen procedure. Note the small dots of Feulgen-positive material throughout the cytoplasm. Magnification, ×500 on film.