DROSOPHILA POLYRIBOSOMES

The Characterization of Two Populations by Cell Fractionation and Isotopic Labeling with Nucleic Acid and Protein Precursors

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ABSTRACT

Two populations of polyribosomes have been isolated from third instar larvae of D. melanogaster. One population appeared to be soluble while the second seemed membrane-bound. Short-term labeling of the two RNP fractions with radioactive nucleic acid and protein precursors was achieved by using a feeding stimulant. RNA was extracted from both polyribosomal fractions following 25, 40, and 60 min of in vivo uridine-3H incorporation. Soluble polyribosomes exhibited more rapid uptake of uridine into ribosomal and heterogeneous RNA fractions than did membrane-bound polyribosomes at comparable time periods. In vivo amino acid incorporation into the two polyribosomal populations was examined after 10, 20, 40, 60, and 80 min of incubation in leucine-3H. In this case, the membrane-bound polyribosomes reached a higher specific activity than did the soluble ones. These functional differences confirmed the observation, based on cellular fractionation studies, that the two classes of polyribosomes represented functionally distinct populations. These data have been compared with those from studies on other metazoan systems. In addition, dithiothreitol has been demonstrated to be a powerful ribonuclease inhibitor.

INTRODUCTION

Combining biochemical with genetic analysis to elucidate mechanisms of regulating protein synthesis has been highly successful in prokaryotes and certain eukaryotic systems. Drosophila melanogaster needs no introduction as an organism suitable for correlated biochemical and genetic studies (Ephrussi, 1942; Ritossa and Spiegelman, 1965; Ritossa and Atwood, 1966; Ritossa, Atwood, and Spiegelman, 1966a and b; Hennig, 1968). In order to utilize Drosophila for investigating the molecular basis of development and differentiation, it is essential to understand the regulation of protein synthesis and, concomitantly, the biochemistry of those RNA1 and RNP moieties involved in protein synthesis.

This report describes the isolation of soluble and...
membrane-bound polyribosomes from larvae of *Drosophila.* In addition, a method has been developed for rapidly labeling larval RNA and protein. Short-term incorporation of uridine into the RNA's of the two polyribosomal fractions has been investigated. Different rates of synthesis of the several classes of RNA's suggests an operational distinction between soluble and membrane-bound polyribosomes. Moreover, in vivo amino acid incorporation studies in both polyribosome preparations support this functional distinction between membrane-bound and soluble populations.

**MATERIALS AND METHODS**

**Fly Culturing**

Stocks of *D. melanogaster* Oregon R. were raised under standard laboratory conditions on a modified version of Carpenter's yeast-agar medium (1950).

**Sterile Culture:** All radioactive incorporation studies have been conducted with axenic cultures. With short-term radioactive labeling of RNA from nonsterile cultures, the bulk of uridine-3H appeared in size classes of r-RNA (23S and 16S) characteristic of prokaryotic organisms. Only after 24 hr did the radioactivity coemigrate with the optical density peaks. On the other hand, 16S and 23S peaks never appeared in rapidly labeled axenic larvae (Bush, 1969).

Laboratory cultures of *Drosophila* were made aseptically following the protocol of Horikawa and Fox (1964). The dechorionated, sterilized eggs, collected on Whatman GF/A glass filter pads (Arthur H. Thomas Co., Philadelphia, Pa.), were transferred aseptically to sterile food vials (see below) and maintained at 25°C in a sterile chamber. Following eclosion, the flies were transferred to other sterile vials every 48 hr. Eight separate lines were maintained in this fashion.

To assay for contamination, contents of used culture vials were periodically plated on three different growth media: Wort agar, pH 4.8 (Fisher Scientific Co., Chicago, Ill.), selects for molds and yeast; tryptone glucose extract agar, pH 7.0, is a general bacterial medium; and Sabouraud's dextrose agar, pH 5.6, selects for mold and aciduric bacteria. (The latter two media are from Difco Laboratories, Detroit, Mich.) Mold was suppressed by occasionally raising a generation of flies under these conditions, apparently owing to the absence of live yeast.

Axenic flies were cultured on sterilized modified Carpenter's medium. 1 ml of Tegosept solution (10% Tegosept powder [Goldschmidt Chemical Co., New York] dissolved in 95% ethanol) was added to 100 ml of medium to inhibit mold. To inhibit bacteria, penicillin-streptomycin (Microbiological Associates, Inc., Bethesda, Md.), final concentration 100 units/ml penicillin and 100 units/ml streptomycin, was included.

**Radioactive Labeling:** Taking into account that *Drosophila* larvae are particulate feeders, they were incubated in either 0.50 ml of uridine-5-3H (20.0 c/mmole, Schwarz BioResearch, Orangeburg, N.Y.) or 0.50 ml of L-leucine-4, 5-2H (30-50 c/mmole, New England Nuclear, Boston, Mass.) containing 0.02 g of autoclaved dried brewer's yeast (Standard Brands, New York) which stimulates feeding. In this yeast-isotope mixture, larvae could be maintained for as long as 96 hr, and they could pupate normally.

**Extraction Procedures—Ribonucleoprotein Particles**

**Isolation of Total Polyribosomal Population:** Larvae were collected by dispersing the masticated Carpenter's medium in water and pouring the slurry into a large beaker. The larvae sink while the small food particles, dead adults, and pupae remained in suspension. After repeatedly decanting the debris, the larvae were collected by filtration on a Buchner funnel and weighed. Typically, 0.4 g of third instar larvae was suspended in 8 ml of solution A containing 0.01 mM MgCl₂, 0.01 M Tris-HCl (pH 7.6), 0.05 M KCl, 0.004 mM DTT (Cal Biochemical Company, Pasadena, Calif.), to which were added 0.25 mM sucrose (ribonuclease-free from Schwarz BioResearch Company) and 1% Triton X-100 (Rohm and Hass, Philadelphia, Pa.). This suspension was homogenized at 4°C in a Dounce tissue grinder (Allfrey, 1959) with 20 strokes of the loose-fitting A ball. A ratio of 20 ml of homogenization buffer/g of filter-dried larvae is optimal. The ratio permits fairly rapid homogenization, and it yields adequate material for optical density and radioactive labeling studies. The homogenate was centrifuged at 10,000 g in the 50 rotor of the Spinco Model L ultracentrifuge (Beckman Instruments, Inc., Palo Alto, Calif.) for 30 min. The 8 ml of supernatant was then divided in half and layered on the following step gradients: 3 ml of 0.5 M sucrose in solution A layered over 2 ml of 1.5 M sucrose in solution A. The tubes were centrifuged in 10 ml polypropylene tubes at 105,000 g for 150 min in the 50 rotor. The supernatant was carefully poured off, and the walls were wiped dry with paper. The pellet which contained both populations of polyribosomes was then gently resuspended in 1 ml of solution A without DTT.

**Isolation of Nonsedimentable (Soluble) and Sedimentable (Membrane-Bound) Poly-
**Sucrose Density Gradients:** Soluble polyribosomes were recovered by homogenizing larvae in 0.25 M sucrose in solution A with no Triton X-100. To disrupt the cells in the absence of detergent, this preparation must be rehomogenized with five strokes of the tight-fitting B ball after treatment with the A ball. Following a 10,000 g centrifugation, Triton X-100 was added to bring the supernatant to a final concentration of 1%. The 10,000 g pellet was rehomogenized with 8 ml of 0.25 M sucrose in solution A containing 1% Triton X-100. Ten strokes of the A ball was followed by another 10,000 g centrifugation. Both supernatants were layered over sucrose step gradients, followed by another 10,000 g centrifugation. Both supernatants were layered over sucrose step gradients, centrifuged, and the RNP pellets were resuspended as previously described for total polyribosomes.

**Extraction Procedure—Ribonucleic Acid**

**Bulk RNA:** Total RNA was extracted and deproteinized according to a modification of the method described by Ritossa and Spiegelman (1965). 0.004 M DTT was added to all solutions in place of 0.001% sodium heparin as a ribonuclease inhibitor. Yields of RNA ranged from 70 to 115 OD260 units/g wet weight of larvae. *Drosophila* bulk RNA E24 equals 200 (Boshes, 1969), indicating a yield of 3.5-5.6 mg bulk RNA/g wet weight of tissue. Church and Robertson (1966) found in their biochemical study that wet weight of *D. melanogaster* is a valid parameter for standardizing data for comparison.

**RNA Extracted from Polyribosomal Fractions:** RNA was extracted from polyribosome-containing supernatants by utilizing a modification of Pennman's method for extracting RNA from HeLa nuclei (1966). Each supernatant was made 1% with SDS (Fisher Scientific Co.), homogenized in a Potter-Elvehjem grinder fitted with a Teflon pestle (Alfrey, 1959), and extracted with an equal volume of phenol saturated with a modified "high salt" buffer containing 0.5 M NaCl, 0.005 M MgCl2, and 0.004 M DTT, pH adjusted to 5.1 with 0.01 M acetate buffer. The RNA was further deproteinized and isolated as described by Ritossa and Spiegelman (1965).

**Analyses**

**Sucrose Density Gradients:** RNA and polyribosomes were analyzed on either linear or exponential sucrose density gradients. Linear gradients for polyribosomes consisted of 14.5 ml of 1.5 M sucrose in solution A (minus DTT) and 15 ml of 0.5 M sucrose in solution A (minus DTT). The gradients were precooled for a minimum of 2 hr before centrifugation. Polyribosomes were centrifuged for 2 hr at 25,000 rpm in the SW 25.1 rotor of the Spinco Model L at 4°C. The tubes were fractionated by means of the ISCO density gradient fractionator (Instrumentation Specialties Co., Lincoln, Neb.), and their optical density at 254 mÅ was recorded through an ISCO UV analyzer. Fractionation was performed at 2 ml/min, and 1 ml fractions were collected. To calibrate the UV analyzer chart, several tubes from the gradient, selected at points where the absorbance tracing is not inflecting, were read in the Zeiss PMQ II spectrophotometer at 254 mÅ against a blank of appropriate buffer.

Convex exponential gradients were made according to the procedure of Noll (1967), using 0.5 M sucrose in solution A and 1.25 M sucrose in solution A. The highest concentration of sucrose in the gradient was 1.0 M. Exponential gradients were spun for 2.5 hr at 25,000 rpm in the SW 25.1 rotor.

RNA was centrifuged on linear sucrose density gradients in buffer containing 0.1 M NaCl and 0.01 M acetate adjusted to pH 5.1. The sucrose concentrations ranged from 0.3 to 0.125 M. RNA was centrifuged at either 23,000 rpm for 16 hr or 25,000 rpm for 12 hr at 4°C in the SW 25.1 rotor. In all linear gradients, sedimentation values have been assigned according to the procedure of Martin and Ames (1961) for comparing an unknown peak to a known standard with similar sedimentation properties. Hastings and Kirby (1966) reported $S_{20,w}$ values of 29.4S and 18.8S for the major components of *Drosophila* ribosomal RNA. The majority of r-RNA from eukaryotic material has been assigned sedimentation constants of 28 and 18S. These latter values are used for ease of comparison. The nominal 28S r-RNA peak has been used as the referent peak in calculating S values of other peaks.

For purposes of comparing specific activities (dpm's/OD254) of RNA fractions between and within the several RNP populations, the preparation and collection of sucrose gradients were standardized. In nearly all cases, RNA greater than 28S includes tubes 32-23, 28S r-RNA includes tubes 22-16, and 18S r-RNA includes tubes 15-9, heterogeneous RNA includes tubes 8-5, and 4S RNA includes tubes 4-1.

**RNAse Digestion:** Polyribosomal preparations were treated with crystalline pancreatic RNase A (Worthington Biochemical Co., Freehold, N.J.). The RNA, dissolved in solution A without DTT, was treated for 20 min at 4°C with 37.4 units (10 μg) of RNase/ml.

**Radioactive Counting:** All samples of homogenized tissue, RNP, or RNA preparations were precipitated by adding 30% ice-cold TCA to a final concentration of 5%. The precipitate was collected on 2.4 cm GF/A glass filters and measured by liquid scintillation counting in a toluene-based fluor. All counts were converted to disintegrations by using the Channel's ratio method to estimate efficiency of counting.

**Kinetic Studies of Leucine-3H Incorporation**
RATION INTO POLYRIBOSOMAL FRACTIONS: In order to study the kinetics of amino acid incorporation into the two polyribosomal fractions, larvae were incubated in leucine-\(^{3}\)H; aliquots were removed after 10, 20, 40, and 80 min; and these samples were fractionated. Free and membrane-bound polyribosome-containing supernatants were layered on step gradients and centrifuged for 2.5 hr at 105,000 \(g\) as described above. The polyribosomal pellets, as well as the supernatants from the step gradient, were recovered. The pellets were redissolved in buffer, read at 254 \(\text{m}\mu\) in the Zeiss PMQ II spectrophotometer, and counted for radioactivity as described above. The supernatants through which these RNP aggregates had sedimented were also counted for radioactivity, providing an indicator of cumulative protein labeling. The relative activities of the supernatants were normalized to their respective RNP pellets, in order to compare the changes in supernatant activities at the several time points (see Fig. 7). Cytoplasmic proteins are a potential contamination of isolated RNP pellets and could confuse the data on incorporation into the RNP. If a constant percentage of contaminating cellular protein became associated with each RNP pellet isolated in the course of the kinetic study, newly synthesized cytoplasmic proteins would be contributing progressively more radioactivity with time. By monitoring relative activity in both supernatants which contain these discharged proteins, radioactivity specifically associated with the two RNP pellets could be estimated.

RESULTS

Analysis of Polyribosome Isolation

TOTAL POLYRIBOSOMAL EXTRACTION: Fig. 1 exhibits polyribosomes isolated from \textit{D. melanogaster} by homogenizing larvae in buffer containing Triton X-100. This procedure enhanced the recovery of polyribosomes relative to monosomes by eliminating a mechanical homogenization step—grinding with the tight-fitting B ball. (cf. Figs. 1 and 2). The yield of ribonucleoprotein from such preparations was similar to that of RNA.
recovering from the ribosomal components of a bulk RNA isolation. 54 OD_{260} units/g larvae were recovered from the gradients illustrated in Fig. 1, nearly 3 mg/g tissue. A yield of 3 mg r-RNA/g larvae is typical for cold-phenol-extracted bulk RNA (Boshes, 1969; Ritossa and Spiegelman, 1965).

**Extraction of Soluble and Membrane-Bound Polyribosomes:** Sequentially extracted soluble and membrane-bound polyribosomes are shown in Fig. 2. Both profiles exhibit the effects of more extensive mechanical manipulations. The yield of RNP from the two profiles in Fig. 2 was 3.2 mg RNP/g larvae, which is similar to total polyribosome extractions. Following homogenization in the absence of Triton X-100, the first 10,000 g supernatant contained one soluble, “nonsedimentable” polyribosome fraction. A gentle rehomogenization of the first 10,000 g pellet in the presence of detergent yielded a second polysome population. This second polysome class was considered to be membrane-bound on the basis of the following criteria: (a) it sedimented during the initial homogenization procedure; (b) it was solubilized by detergent, and (c) electron microscopic examination of the initial pellet showed the presence of ribosomes bound to membrane (Boshes, 1969).

**RNase Treatment of Polyribosomes:** Fig. 1 exhibits polyribosomes treated with RNase A in the presence and absence of 0.004 M DTT to illustrate the efficacy of the latter as an RNase inhibitor. The definition of the several RNP classes has been somewhat obscured by RNase treatment, but the over-all ratio of polysomes to monosomes is virtually unchanged. RNase exposure in the absence of DTT failed to degrade quantitatively polysomal aggregates to the monosome region, but did convert heavier polyribosomes to the di-, tri-, and tetrasome region of the gradient. The majority, however, did not appear under the monosome peak.

**Analysis of RNA Isolated from RNP Fractionation**

Ribosomal-RNA constituted 62% of the RNA recovered from the first 10,000 g supernatant (Fig. 3). However, the distribution of 28S:18S r-RNA was not typical of an RNA extraction from Drosophila. The expected relationship between the components should be 1.92:1, on the basis of measurements of Drosophila 28 and 18S r-RNA’s having molecular weights of $1.40 \times 10^6$ and $0.73 \times 10^6$, respectively (Loening, 1968), and the two classes being present in equimolar amounts.
having been synthesized coordinately (Brown and Littna, 1964; Darnell, 1968). The observed ratio of 28S:18S of 1.57 (se = ±0.05): 1 for the first supernatant was low in comparison with that from the membrane-bound population. The large OD 

FIGURE 3 Sedimentation profiles of RNA extracted from RNP recovered from first 10,000 g supernatant (soluble RNP) of fractionated D. melanogaster larvae incubated in uridine-3H. RNA was analyzed on 0.125-0.5 M sucrose linear density gradients made up in solution A without DTT and centrifuged at 4°C for 16 hr at 23,000 rpm in a Spinco SW 25.1 rotor. Optical density was continuously monitored at 254 nm. 1 ml fractions were collected for counting as described in Materials and Methods. A, 35 min incubation; B, 40 min incubation; C, 60 min incubation.

bound population at comparable time periods. This difference provides another operational criterion for distinguishing membrane-bound from soluble polyribosomes.

Uridine-3H incorporation into the 4S region was nearly identical in both populations (Figs. 5 A and B). This similarity in 4S specific activity between the two fractions is markedly contrasted by the other classes of RNA. These counts cannot be attributed to terminal cytidine labeling of transfer RNA since uridine cannot be converted to cytidine by Drosophila (Dr. Charles Laird, personal communication).

Amino Acid Incorporation into Polyribosomal Fractions

Fig. 6 demonstrates soluble and membrane-bound polyribosomes extracted from the same larvae following 60 min of incubation in leucine-3H. The OD 

The specific activities of amino acids incorporated into the polyribosomes were substantially higher in the membrane-bound than in the free population. The specific activities of the monosome regions were both low—less than 1000 dpm/OD 

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activity (9700 dpm/OD254) of comparable areas from the soluble fraction (5033 dpm/OD254). This region included aggregates up to approximately 10 ribosomes. The difference in specific activities in the very large polyribosome portion of the profiles was less than twofold.

Fig. 7 exhibits specific activity accretions of membrane-bound and soluble RNP pellets as well as net accumulation of TCA-precipitable counts into the respective supernatant fractions. These latter data reflect newly synthesized proteins in the cytoplasm.

Both polyribosomal fractions reached a steady state of amino acid incorporation by 40 min as evidenced by the plateaus in the curves of radioactive uptake. By this time, the membrane-bound fraction had approximately four times the specific activity of the soluble population. The supernatant
recovered from the membrane-bound RNP did not accumulate significant radioactivity until 20 min, indicating that label in membrane-bound RNP pellets was not artificial contamination of its associated supernatant.

After 40 min, the step-gradient through which the membrane-bound polysomes had been pelleted contained over 85,000 dpm/50 µg pelleted RNP, while the step-gradient of the soluble polysomes had 65,000 dpm/50 µg pelleted RNP. By 80 min, however, this supernatant had a higher specific activity than the comparable supernatant from the membrane-bound polyribosomes, in spite of the fact that these latter polyribosomes, which had pelleted through this less active supernatant, contained over three times the radioactivity of the soluble polysome pellets. These results suggest that the pelleted RNP's are not contaminating or being contaminated by supernatants of differing specific activities, and they provide further support for an operational distinction between the two populations of polyribosomes.

**DISCUSSION**

**Polyribosome Isolation**

**EVALUATION OF METHODOLOGY:** Criteria for evaluating a satisfactory polyribosome isolation procedure are not clearly defined. The primary criterion has been the proportion of aggregates of polyribosomes relative to monosomes, disomes, and trisomes, judged to be degradation products by many investigators (Warner et al., 1963; Wettstein et al., 1963; Rifkind et al., 1964). In this study the total yield of RNP relative to bulk RNA, as well as the quality of the optical density profile, have been monitored in evaluating the progress of the extraction procedure. In order to achieve the quantitative isolation of apparently undegraded polyribosomes, several modifications of previously published protocols were introduced. A Dounce tissue grinder was utilized to extract *Drosophila* polyribosomes without shearing damage. The concentration of Mg++ had been set at 5 mM according to many published protocols. 10 mM Mg++ was, however, essential for isolating large aggregates of *Drosophila* polyribosomes. Triton X-100 replaced DOC as the detergent since it appeared to be less injurious to *Drosophila* polyribosomes than ionic detergents.

While soluble and membrane-bound polyribosomes may be distinguished on the basis of isolation procedure and labeling characteristics, it is important to recognize the limitations of such distinctions. The soluble fraction is, in operational terms, a population of RNP that did not sediment after 30 min at 10,000 g. Within this class may be polyribosomes which are truly soluble, polyribosomes attached to intact membranes whose entire structure is too light to sediment, and polyribosomes attached to originally heavy vesicles which have become nonsedimentable owing to disruption in the course of isolation. 

![Figure 6](image-url) Sedimentation profiles of soluble and membrane-bound polyribosomes extracted from *D. melanogaster* larvae incubated for 60 min in leucine-3H. 0 A, non-sedimentable, soluble, polyribosomes; 0 B, sedimentable, membrane-bound, polyribosomes.
polyribosomes, on the other hand, may contain polyribosomes bound to endoplasmic reticulum or secretory vesicles, as well as RNP associated with the nuclear envelope.

The possibility of finding functional relationships between soluble and membrane-bound polyribosomes is tempting in a study of this kind. However, these two populations of Drosophila polyribosomes might have been isolated from differing cell types. Drosophila secretory tissue such as the salivary gland and ring gland may contain different ratios of membrane-bound to soluble polyribosomes than other rapidly growing larval tissues or rapidly dividing imaginal disc tissue (Palade, 1959; Siekevitz and Palade, 1959; Redman and Sabatini, 1966a and b; Goldberg and Green, 1967).

RNase Studies: DTT was selected as the RNase inhibitor of choice because it was highly effective in low concentration. Nuclease inhibitors such as sodium heparin or mercaptoethanol had been added in order to inhibit endogenous ac-

Figure 7 Specific activity accretion of soluble and membrane-bound RNP pellets and their supernatants following incubation in leucine-\(^{3}H\). At 10, 20, 40, and 80 min, aliquots of larvae were removed and homogenized according to the procedure for recovering two populations of polyribosomes. The RNP pellets, which had been purified by sedimentation through a step gradient, were resuspended in solution A, read at 254 nm, and counted for radioactivity. The step gradients, through which the RNP had been pelleted, were also counted for radioactivity.
tivity, but with no appreciable improvement. DTT, by virtue of the stability of the intramolecular disulfide bridge formed by oxidation, is a powerful sulfhydryl reducing compound, and hence a potent RNase inhibitor (Fig. 1). To the best of my knowledge, DTT has not previously been recognized as a ribonuclease inhibitor (see Calbiochem publication Cleland's Reagent—A Current Bibliography).

RNase treatment, in the absence of DTT, however, does not completely eliminate polyribosomal aggregates consisting of two to four ribosomes (Fig. 1). RNase-resistant polyribosomes from Ascaris lumbricoides have been described (Kaulenas and Fairbairn, 1966). After preincubation with trypsin, these workers reported a complete breakdown of their material to monosomes. RNase-insensitive, trypsin-sensitive polysomes have also been extracted from heart muscle of chick embryos (Rabinowitz et al., 1964). I have tried to disrupt Drosophilapolyribosomes by using trypsin with and without subsequent RNase treatment, with no particular effect (Boshes, 1969).

Whitney et al. (1968) reported in an abstract the extraction of polysomes from Drosophila. These authors claimed that 0.1 M K+ in their buffer provided an ionic environment which permitted thorough degradation of all RNP aggregates to monosomes; heterodisperse polysome profiles were converted to a monodisperse peak of approximately 80S. Martin et al. (1969) have demonstrated that ionic conditions similar to those in many polyribosome extraction buffers strongly influence dissociation and reaggregation of ribosomes and their subunits in eukaryotic material. Their observations might explain the persistence of di- and trisomes following RNase treatment of my preparations, since the buffer contains 10 mM Mg++. RNA Isolated from Soluble and Membrane-Bound Polyribosomes

The difference in ratio between ribosomal RNA's recoverable from the soluble and membrane-bound polyribosomes has already been noted; a consistently higher ratio of 28S:18S RNA was obtained from the membrane-bound population than from the soluble one. This observation cannot be attributed to degradation of the more extensively treated RNA since an opposite result, more 18S relative to 28S, would be anticipated by breakdown of ribosomal RNA. Sabatini et al. (1966) have reported the differential release of membrane-bound ribosomal subunits into the cell sap following EDTA treatment, the 40S RNP particle being more easily displaced than the 60S particle in guinea pig hepatic microsomes. Vesco and Pennman (1969) have also found an altered 28S:18S ratio in RNA extracted from a resuspended "mitochondrial pellet." They too attributed their unusual 28S:18S ratio, which was much higher than total cellular RNA, to be the result of selective loss of the 18S-containing ribosomal subunit from membranes in the presence of 0.001 M EDTA. Homogenization of Drosophila larvae with the Dounce grinder might have removed less firmly bound 18S-containing subunits from membranes. This partitioning artifact would account for the first 10,000 g supernatant's containing a low 28S:18S ratio owing to contribution from 18S-containing subunits from the membrane fraction, whereas the membrane-containing fraction has a higher 28S:18S ratio owing to the reciprocal absence of 18S-containing subunits. Alternatively, there may be more 18S-containing subunits normally circulating in the cytoplasm than 28S-containing subunits which are more tightly bound to the membrane. So far, these alternative hypotheses have not been critically tested.

Isotope Incorporation into Larvae

The procedure for inducing larvae to incorporate isotope in the presence of a feeding stimulant makes possible genuine short-term labeling studies in Drosophila. Unlike injections of individual animals this technique is limited by neither the size nor the number of treated individuals. In addition, one may anticipate that labeling the first or second instar larvae will have the same success that has been demonstrated with third instar larvae. After 25 min, a short labeling period for metazoan tissue (Girard et al., 1965; Henshaw et al., 1965; Latham and Darnell, 1965), over 120,000 dpm/g larvae were incorporated into total cellular RNA. The specific activities of several of the fractions, particularly the 16-5S heterogeneous RNA, indicate that even shorter incubations may be feasible.

Analysis of Radioactive Labeling of RNA Derived from Polyribosomes

The results of radioactive precursor incorporation into ribosomal RNA's of both soluble and
membrane-bound polyribosomal fractions are consistent with Greenberg's studies on total RNA labeling in *D. virilis* (1969): *Drosophila* behaves like HeLa cells with respect to r-RNA synthesis and processing (see Darnell, 1968, for review). Results similar to those described here have been demonstrated in comparable fractionation procedures utilizing a rat liver system (Hallinan and Munro, 1965) and rat spleen system (Talal and Kalreider, 1968). These groups reported similar label distribution between soluble and membrane-bound populations, as well as a predominance of uridine incorporation into the 16-5S region of RNA isolated from the soluble polyribosomal fraction. The noncoordinate appearance of 18S and 28S molecules in the cytoplasm suggests that ribosomes normally circulate in the form of free subunits (Girard et al., 1965; Joklik and Becker, 1965a and b). The accumulation of isotope in the ribosomal RNA peaks of both polyribosomal populations from *Drosophila* is consistent with this hypothesis. The 18S-containing subunit is rapidly exported, while the slowly labeling 28S-containing subunit does not reach the specific activity of the 18S r-RNA by 60 min (Figs. 3 and 4). Although the over-all specific activity of r-RNA from the soluble polyribosomes is higher than that of r-RNA from the membrane-bound fraction (Figs. 3 and 4), the rate at which the membrane-bound fraction equilibrates (equilibration is defined as specific activity of 28S r-RNA : specific activity of 18S r-RNA = 1) is faster than the rate for the soluble one. These data can best be explained by assuming that a small part of the total membrane-bound ribosomal subunit population turns over more rapidly than does the soluble subunit population (see Boshes, 1969, for discussion).

The 28S r-RNA-component cosedimented with the OD254 peak from the time of its first appearance, 40 min, indicating that it had been completely processed before exportation from the nucleolus (see Darnell, 1968, for review). The heterogeneous RNA fraction from the 16 to 5S region of the soluble polysomes exhibited the highest specific activity throughout all time periods (Fig. 5). This region contains the size class of molecules thought to correspond to m-RNA molecules (Latham and Darnell, 1965; Brown and Gurdon, 1966).

The specific activities of the 16-5S region as well as the 18S and 28S r-RNA's from the membrane-bound population of polyribosomes were all lower than those of their respective populations from the soluble polyribosome fraction. On the other hand, the specific activity of the 4S fraction from the membrane-bound population was practically identical to that of the 4S RNA recovered from the first 10,000 g supernatant, although the first supernatant contained most of the 4S RNA on the basis of the OD254 profile (Figs. 3–5). Since transfer RNA is a cytoplasmic component, one would anticipate that the bulk of it would appear in the first 10,000 g supernatant, the cell sap. The small amount of 4S RNA which appears in the second fraction probably represents the same population, on the basis of its parallel isotope incorporation. A small portion of this material may be t-RNA bound to membrane-bound polyribosomes. Trapping of soluble t-RNA by the sedimentable material of the homogenate may also account for some of the radioactivity in this region.

**Analysis of Amino Acid Incorporation into Polyribosomes**

The pattern of incorporation of leucine-3H into polyribosomes suggests the labeling of nascent proteins. Moreover, large aggregates appeared to have been isolated without substantial degradation (Fig. 6) since there was little radioactivity under the mono- and disome peaks where nascent peptides attached to degraded polysomal fragments would be expected to accumulate. These data are consistent with reports that the specific activity of amino acid incorporation increases throughout the dimer and trimer gradient regions, reaching a constant value in the area of larger polyribosomes (N = 4–7 [Goldberg and Green, 1967]). The heavier aggregates therefore appear to constitute the majority of the actively synthesizing populations in *Drosophila* as in other metazoan cell types (see Goldberg and Green, 1967, for review).

Labeled 18 and 28S r-RNA's indicate that newly completed ribosomal subunits are present in the cytoplasm by 60 min (Figs. 3 and 4). It is plausible that some ribosomal proteins may have also been synthesized within that time. The absence of radioactivity under the monosome region, therefore, may indicate that newly fabricated ribosomal subunits first appear in both fractions in the polyribosome, not the monosome, population. The monosomes may represent older, dissociated polysomal aggregates. Similar arguments
have been made for other eukaryotes (Girardet et al., 1965; Joklik and Becker, 1965a; Darnell, 1968).

One aspect of the kinetic study of amino acid incorporation (Fig. 7) deserves elaboration. Radioactive leucine sediments in the first 10,000 g pellet but is later released by Triton X-100 treatment of this pellet and appears in the supernatant of the membrane-bound polysomal fraction. Such behavior would be predicted from proteins bound in membranes. Redman and Sabatini (1966a and b) proposed that secretory proteins are sequestered in sedimentable endoplasmic reticulum and are exported via this membrane-transport system. While Redman and Sabatini’s microsomal fraction was derived from a 105,000 g pellet, Drosophila membrane-bound polyribosomes were isolated from a 10,000 g centrifugation. But this 10,000 g pellet from Drosophila did contain over 40% of the total ribosomal RNA; and electron microscopic examination revealed multiple polyribosomal aggregates bound to membranes (Boshes, 1969). Therefore, while the membrane-bound RNP fraction cannot be considered operationally identical to the microsomal preparation from guinea pig liver studied by Redman and Sabatini, the observed data seem to be best rationalized by an analogous argument.

CONCLUSION

Two populations of polyribosomes have been isolated in reproducible amounts from third instar larvae of Drosophila melanogaster. One is present in the cell sap, while the second appears to be membrane-bound. Both conform to physical criteria for polyribosomes: rapidly sedimenting RNP aggregates with peaks of absorbancy at 254 m\(\mu\) corresponding to multiple units of monosomes. Both are sensitive to mild ribonuclease digestion, which shifts the rapidly sedimenting polysomes to the region of the gradient where lighter RNP structures are found.

Short-term isotopic labeling of larvae with protein and nucleic acid precursors has been achieved by combining a feeding stimulant of dried yeast with mild dehydration of the larvae by filtration. A functional distinction between these soluble and membrane-bound polyribosomes has been demonstrated. The soluble population accumulates RNA precursors more rapidly than does the membrane-bound one while the latter exhibits a significantly higher rate of amino acid incorporation. The distribution of leucine-\(^3\)H confirms that both populations of RNP are functional polyribosomes and not aggregation artifacts. Whether there be any functional relationship between soluble and membrane-bound polyribosomes, whether there be a unique set of proteins synthesized by the respective classes, or whether these represent RNP from differing cell populations all await further investigation.

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