Beardmore, J. A. Genetical Institute, Haren, Netherlands. Medium dispensing machine.

Although several dispensers for Drosophila medium have been described, it may be of interest to some workers to know of the existence of a machine, which, without modification, can be used by relatively unskilled personnel. The reservoir holds 15 l of medium and this can be dispensed (lever operation, see accompanying photograph) very accurately and quickly in volumes varying from 1 - 80 ml; resetting of the volume control is very simple. The machine will take media varying widely in viscosity though it may be necessary to change the nose piece to avoid drips or clogging. Provided that it is scrupulously cleaned after use, the chance of appreciable wear or damage with normal use is very small as most moving parts are made of stainless steel.

"Perpetua" filling machine model H1, Engler Maschinenfabrik, Brunner und Co., Rissawegasse 12-14, Vienna X/75, Austria; cost in Europe c $230).


The following method has been found satisfactory for double-diffusion tests utilizing a single fly against appropriate antisera. Groundedge cover glasses (2" x 2") are flamed, coated with 2 ml of 0.5% Oxoid Ionagar No. 2 in distilled water, and dried overnight to a film. These precoated slides are then covered with 2 ml of a 1.0% solution of agar in buffered saline (0.005 M. phosphate, pH 7.4, 0.85% Na Cl, 0.1% thimerosal), and allowed to gel in a petri dish provided with a strip of wet filter paper. A central antiserum well and 6 outer antigen wells are cut with the aid of a template and cutter, and the gel is removed from the wells with a small spatula; no sealing gel is necessary.

Single etherized flies are crushed with a glass rod in individual Durham tubes. After about 100 strokes, 0.04 ml of buffered saline is added and homogenization continued to at least 200 strokes. This provides a 0.03 ml antigen dose for one well. The small beaker supporting the Durham tubes is kept in an ice bath until all of the antigen has been delivered to the plate. A total of 0.03 ml of antiserum is added to the central well and the plate is allowed to develop for 4 to 6 days.
If the distance between well centers is made 8 mm and the well diameter 4 mm, the reactants may be delivered in 3 doses of 0.01 ml at approximately 2-hour intervals. If the well geometry is increased to 10 or 13 mm to allow for a well diameter of 6 mm, the complete 0.03 ml of reactants may be delivered at one time.

When precipitation is complete the slide is washed in tap water to remove excess protein from the wells. Unprecipitated protein is eluted by soaking the slide in buffered saline for 24 hours, followed by 2 rinses of 1/2 hour each in distilled water. Slides are stained for 20-30 minutes in Crowle's triple stain (Immunodiffusion, 1961, Academic Press) or in dilute water-soluble nigrosin, destained in 1% acetic acid, and air dried to a film. The finished slide may be used directly in the photographic enlarger to prepare prints, and is in itself a convenient permanent record of the test.

Richardson, R. H. University of Texas. An improved technique for fecundity and hatchability tests.

A new technique of collecting eggs for fecundity or hatchability tests has been devised, which has the following advantages: homeogeneous egg laying surface resulting in uniform egg distribution, rapidly and easily dispensed medium, medium lacking extraneous food components (such as charcoal), transparent medium allowing scoring of burrowing larvae, and easily cleaned and reused equipment.

The medium consists of 1 g. Bacto-agar, 100 ml. water and 15 ml. white Karo syrup, which is dispensed with an automatic syringe while hot. This medium is then sprayed with a water suspension of bakers yeast immediately before use.

The equipment consists of two variations on the same theme. One variation supplies a black background to facilitate counting. The other presents a transparent background, which allows visual examination of eggs without the removal of the cap from the test bottle.

The test bottles are constructed from 40 dram Plastainer bottles (ca. 2" x 3 1/4") available from Owens-Illinois Glass Co., Toledo, Ohio, at a cost of about $5 per carton (6 dozen). Extra caps are available at about $20 per thousand. The screw caps are made of Teflon and the bottle of clear plastic. A hole is punched in the cap top with a die about 1 1/4" in diameter, and then a piece of plexiglass 1/16" thick is glued to the outer surface of the cap over the hole. The plexiglass may be either black or transparent, giving the two varieties of background. A critical factor in construction is the cement for glueing the cap and the plexiglass. The most satisfactory one tried was Eastman 910 adhesive, available from the Tennessee Eastman Company, Kingsport, Tennessee, at a cost of $8 per bottle. One bottle is sufficient to glue about 400 caps. Also the surface of the cap must be roughened with hardware cloth or a file before glueing. The glue is spread in a very thin band completely around the hole in order to get a water-tight seal. Leaks may be sealed with a band of Duco cement around the external cap-plexiglass junction.

Counting is easily accomplished by marking the agar surface into regions with a blunt needle under about 40X magnification or less. Eggs or larvae may be conveniently transferred to food bottles by transferring agar and eggs or larvae with a small spatula (eg., No. 19240, Curtin Cat. 40) bent at a convenient angle to work inside the cap. Larvae may crawl off the agar surface, but for caps changed every 24 hours or so, it is not a serious problem. Empty egg cases are easily distinguished from unhatched eggs.

An additional advantage of this technique is the practicality of a permanent photographic record of the egg production or hatchability, especially since the eggs are well spread over the surface. The quickest technique using the transparent plexiglass caps in a "contact print" of the cap on photographic paper (available in bulk rolls about 4 1/4" wide) where the shadow of the egg is recorded. Enlargement prints are possible by placing the cap in the film plane of a darkroom enlarger. More detailed records may be made by microfilming the black plexiglass caps with a 35 mm. camera. Examination of the negative either in a microfilm reader or under a dissecting scope allows easy egg counts, hatch scores, or even some egg development scores. It appears counts could even be made by visual scanners in use by automatic data processing systems.