above can be dried in 8 hours. Following this drying period the specimen is removed to a vacuum chamber for embedding in paraffin. The quality of the results, in terms of histological detail, appears equivalent to that of results obtained by the conventional vacuum freeze-drying approach. The efficiency of this method compares favorably with the best reported vacuum freeze-drying and is substantially superior to that of devices which do not have the cold trap in line of sight from the specimen.

The mechanical simplicity achieved through elimination of vacuum pumps and use of a vacuum-tight system is considerable. An additional advantage of this approach, however, appears to be in situations where a number of specimens are to be dried simultaneously, where the many specimens and their supports constitute physical obstacles which diminish the effectiveness of the vacuum system by preventing straightline passage of vapor to the trap. Above all, these experiments (5) confirm the supposition that passage of vapor through the dried specimen shell is primarily a matter of vapor pressure gradient rather than of total pressure in the system.

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- 5. I wish to acknowledge the skilled assistance of Raymond Long in the design and construction of the device described in this report.
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Photodynamic Inactivation of Monkey Kidney Cell Monolayers

Abstract. Monkey kidney cell monolayers exposed to white light in the presence of neutral red (1:40,000) undergo degeneration within 24 hours after exposure.

This communication concerns the effect of white light on monkey kidney cell monolayer cultures in the presence of the vital stain neutral red. In the course of attempts to photoreactivate ultraviolet-light-inactivated monkey kidney cell monolayers grown in 100-mm petri dishes, it was observed that unirradiated monolayers, when exposed to white light in the presence of the vital stain neutral red (1:40,000), degenerate within 24 hours after exposure. Similar plates exposed to the same light for the same time period, but in the absence of

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Table 1. Tissue degeneration with various dyes applied to the culture.

		*		
Time of ex- posure (min)	No dye	Neutral red (1:40,000)	Trypan blue (1:10,000)	Methylene blue (1:10,000)
0	0	0	0	0
2.5	0	0	†	†
5.0	0	0	ŧ	ŧ
10	0	1	Ó	0
20	0	3	†	÷
3 0	0	4	Ó	Ò
60	0	4	0	0
90	0	4	+	+
	Light	filtered throu	ıgh trypan b	lue
60	-	4		
1	Light fil	tered throug	h methylene	blue
6 0		4	•	
	Light	filtered through	ugh neutral	red
6 0	3	0	5	

* Degeneration: 100 percent, 4; 50 to 75 percent, 3; 0 to 25 percent, 1; none, 0. † Not determined.

the neutral red, showed no ill effects from the exposure. Neutral red applied to the cells after exposure to the white light did not produce cell degeneration and was picked up by the cytoplasm as in unexposed cells. If the light was first passed through a solution of neutral red (1:40,000), the particular wavelengths of light responsible for the damage were filtered out, and damage to the cells was prevented. However, when methylene blue (1:10,000) or trypan blue (1:10,-000) were used as filters, damage occurred as it did with unfiltered white light. The results are summarized in Table 1.

The source of the white light was a bank of three 20-watt fluorescent bulbs at a distance of 15 cm. The light was filtered through a 3-cm solution of 1-percent copper chloride, and the temperature of the air above the solution did not rise above 38° to 39° C.

It is suggested that perhaps one cause for the occasional degeneration of monkey kidney monolayers or for the loss in plaque count (1) seen after the addition of the nutrient-agar-neutral-red mixture used in the Dulbecco Plaque Technique is inadvertent exposure to white light. It is evident from these findings that in the presence of neutral red, the exposure of the tissue to white light should be kept to a minimm.

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Effects of Differential Infantile Handling upon Weight Gain and Mortality in the Rat and Mouse

Abstract. Animals were handled during ages 1 to 10, 11 to 20, or 1 to 20 days, or were nonhandled controls. Animals handled for 20 days weighed the most in adulthood, while the controls weighed the least. Animals handled on days 1 to 10 survived food and water deprivation the longest of any group. Mice handled for 20 days died earlier than controls, while the reverse was true for the rat.

The general procedure followed in studying the effects of infantile experience upon adult behavior has been to stimulate the organism from birth until weaning and then test for effects of the stimulation later in life. Since the rat and mouse undergo tremendous developmental changes during the pre-weaning period, it is likely that the same stimulation affects the organisms differently at different developmental stages. This is suggested by Scott's critical period hypothesis (1) and by the work of Levine and Lewis (2). Therefore, if animals are stimulated only during certain parts of the pre-weaning period, the stimulation may interact with maturational processes which are at different stages of development to differentially modify adult behavior.

Complete litters of rats descended from the Harvard Wistar strain, and C57BL/10Sc mice were randomly assigned to one of the following infantile experience groups: handled on days 1 to 10, handled on days 11 to 20, handled on days 1 to 20, and nonhandled controls (3). At least two litters were used per experimental treatment. Handling consisted of removing the pups from the home cage, placing them in a container (a 1-gal can filled with sawdust for the rat, and a wooden mouse box for the mouse) where they remained for 3 minutes, and then returning them to the home cage. This procedure was followed once daily on the appropriate days. All animals were weaned at 21 days and reared thereafter with like-sexed members of their own litter in small groups. Food and water were always available. At 69 days the rats were weighed and placed on total food and water deprivation in individual cages. The same was done with the mice at 54 days. All animals had received 10 days of testing of avoidance learning just prior to this. Hours until death occurred were recorded.

The group means for weight and mortality for both species are presented in Table 1. The data were analyzed in a 2×2 factorial design: presence or absence of handling on days 1 to 10 was one factor, while presence or absence of handling on days 11 to 20 was the