ops for 4 hours, and then further cell division is blocked. The nuclei incorporate tritiated thymidine, but after further division is interrupted, the nuclei become vesicular and increase in size. Hydroxyurea can be removed from the solution up to 3 hours after fertilization and development proceeds. It thus may interfere with RNA production or protein synthesis. Urethane, which is somewhat similar to hydroxyurea in structure is far less active in the range of 20-30 mg/10cc and it acts to inhibit cleavage.

4. Whereas drugs which act later on development probably influence certain pathways for cell synthesis, others act in inhibiting the cleavage. Vincristine and vinblastine are very effective, in the range of $10-20 \frac{y}{10cc}$ of sea-water. Another compound, 8-mercapto-2-piperidino adenine was also an extremely active cleavage inhibitor, at doses in the range of $2\frac{y}{10cc}$ of sea-water. This is one of the most active cleavage inhibitors known; analogues of 8-mercapto-2-piperidino 2-piperidino adenine were also tested, but none approached it in activity.

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OXYGEN MOVEMENT ACROSS THE EEL GILL EPITHELIUM

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Measurements of the oxygen tension in fluid bathing the inside and outside surfaces of the isolated perfused eel gill epithelium were made by means of an oxygen electrode. It was found that under all conditions the oxygen tension was the same in both fluids. It would appear that oxygen is not actively secreted across the isolated perfused eel gill epithelium.

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SODIUM TRANSPORT BY THE EEL GILL in vivo

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Previous work in this laboratory with isolated perfused eel gill has demonstrated the importance of such factors as the rate of blood flow, the capillary pressure, and the colloid osmotic pressure of the fluid perfusing the blood vessels of the gill in determining the rate of salt and water movement across the gill epithelium. These earlier results also indicated that salt water adapted eel gills actively transport sodium and chloride ions outward but that fresh water adapted eel gills apparently do not actively transport salt inward. It seemed of interest to see if similar results would be obtained with intact eel. The U-tube technique of Krogh was used. Lightly anesthetized (MS-222) fresh water or salt water adapted eels were placed in a Tygon tube in such fashion that the anal papilla drained to one side of the tube while the mouth was in the other side of the U. Under proper conditions, the eel remained quiet and breathed regularly in this apparatus. Influx and outflux of sodium through the mouth end of the fish were estimated with the use of Na²². Control experiments in which the gut was blocked showed that the fluxes occurred primarily across the gill epithelium. It was found that fresh water adapted fish actively took up salt whereas salt water adapted eels actively excreted salt through the gills.