



Fig. 1. Kinetics of the hydrolysis of cyanovinyl phosphate at 100°C.

cyanovinyl phosphate) by heating 0.7 ml of 0.75M sodium cyanovinyl phosphate and 1.6M uridine (containing 0.1 curie of C¹⁴-uridine) at 60°C for 18 days. The UMP was detected by the coincidence of the radioactivity with that of an authentic UMP sample on paper chromatography (10). The ultraviolet spectrum of a sample eluted from the paper chromatogram was identical with that of UMP (λ_{\max} 260 m μ).

To assess the prebiotic significance of cyanovinyl phosphate as a phosphorylating agent, the relative ease with which it transfers orthophosphate to uridine and water must be determined. If the selectivity for uridine is high then cyanovinyl phosphate may have prebiotic significance. If it is low then cyanovinyl phosphate will only undergo hydrolysis in dilute solution without effecting any phosphorylation. This selectivity factor, which was first defined by Lohrmann and Orgel (11), may be formulated as shown, with M_u the molarity of uridine, X the percent yield of UMP, and assuming the molarity of water to be 55:

$$\text{selectivity} \left(\frac{\text{uridine}}{\text{water}} \right) = \frac{55}{M_u} \left(\frac{X}{100 - X} \right)$$

This selectivity factor is only 1.3 for uridine, suggesting that very little UMP was formed prebiotically in homogeneous solution from uridine and cyanovinyl phosphate. This result is in agreement with the usual observation that water and alcohols are phosphorylated at about the same rate (11, 12).

However, orthophosphate is phosphorylated 4.5 to 9.2 times as efficiently as water, suggesting that cyanovinyl phosphate may have been the prebiotic source of the pyrophosphate bond. For example, 1 percent and 0.1 percent

conversions to pyrophosphate would be obtained starting with 0.1M and 0.01M phosphate, respectively, and assuming a selectivity factor of 5.5.

A potential unified synthesis of pyrimidines, amino acids, and a high-energy phosphate compound from one source makes cyanovinyl phosphate attractive as a potential prebiological phosphorylating agent.

J. P. FERRIS

Department of Chemistry,
Rensselaer Polytechnic Institute,
Troy, New York 12181

References and Notes

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- I thank Dr. A. Beck for performing these experiments. The analytical procedure of R. H. Kolloff [*Anal. Chem.* **33**, 373 (1961)] was used.
- Descending paper chromatograms on Whatman 3 MM paper, using 7 parts by volume of 95 percent ethanol and 3 parts of 1M ammonium acetate, pH 7.5.
- I thank Dr. R. Lohrmann for his assistance with the UMP analyses. The analytical procedures will be described in detail in a forthcoming publication (R. Lohrmann and L. E. Orgel, *J. Am. Chem. Soc.*, in press).
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Transplantation of Marrow to Extramedullary Sites

Abstract. Autologous fragments of transplanted marrow have survived in various extramedullary sites in the rat, rabbit, and dog. Survival of the fragments occurs with a complete reconstruction of the hemopoietic and adventitial structures. The process originates from a network of surviving reticular cells which proliferate and differentiate into osteoblasts and give rise to trabecular bone. Later, the reticular cells reconstruct the marrow's microcirculation. Hemopoietic repopulation of the marrow implant takes place only after its sinusoidal microcirculation has been established.

The failure of implants of autogenous bone marrow to survive in extramedullary sites has led to a conclusion that hemopoiesis cannot be sustained except in the marrow cavity. This concept is supported by indications that hemopoiesis requires the unique microcirculation of the marrow. The work of Knospe, Blom, and Crosby (1) demonstrates the relation between the two major elements of bone marrow, the blood forming elements and the adventitial elements which comprise the microcirculation of marrow. Hemopoiesis cannot be sustained except when sinusoidal blood vessels are present and functioning normally. In earlier attempts to transplant fragments of marrow or aspirates, with one exception (2), this specialized adventitia was evidently not retained (3, 4). Our success in transplanting the bone marrow of rat, rabbit, and dog into extramedullary tissues evidently requires the use of a relatively large piece of undisrupted marrow to permit a reconstruction of the sinusoids.

More than 200 Wistar albino rats (300 to 500 g) of both sexes were used. A window was cut by a dental drill in the anterior plate of the tibia, the sequestrum was removed, and the marrow was lifted out on the tip of a spatula; by this method we obtained a relatively large piece of marrow without disrupting the tissue. These fragments were implanted into splenic, renal, subcutaneous, hepatic, muscular, and omental tissues. They were removed periodically; after obtaining touch imprints on slides for cytological and cytochemical examinations, we fixed the tissue in 10 percent buffered formalin for at least 3 days and decalcified it in Prenny's solution. Sections were stained

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J. P. Ferris

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