The Association of the Urobilin “Early Peak” and Erythropoiesis in Man *

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In a normal individual the administration of isotopically labeled glycine (1, 2) results in two peaks of radioactivity in the excreted fecal bile pigment. The first or “early” labeled peak of fecal bile pigment excretion occurs within the first few days and cannot be explained on the basis of the breakdown of normal circulating red cells. The second peak occurs 100 to 130 days after administration of the isotopic precursor and coincides with the termination of the life-span of the group of red cells that contains the labeled precursor. When the conversion of intravenously administered glycine-2-14C to serum bilirubin-14C was studied, two peaks of bilirubin specific activity were found, the first at 24 hours and the second at the third to fourth day (3). It must be recognized that the single observed fecal urobilin early peak is a result of fusion of the two early labeled peaks in the gastrointestinal tract and that the changes measured in these patients could, in theory, be due to a quantitative change in either component.

Several theories have been proposed to explain the first or early fecal peak. The most widely accepted is that this pigment is derived from red cells that are destroyed before or shortly after their release from the bone marrow. For this reason the early peak has been thought of as representing “ineffective” erythropoiesis. With this hypothesis, London, West, Shemin, and Rittenberg (1) and Gray, Neuberger, and Sneath (2) calculated that in a normal individual 11 to 20% of erythropoiesis is represented in this peak. However, recent studies have been interpreted as showing that the early peak might arise by a direct synthetic pathway from a common precursor pool that does not require heme as an intermediate and, thus, is not necessarily related to erythropoiesis (4).

James and Abbott (5) reported the presence of an elevated early peak in two patients who eventually developed acute granulocytic leukemia. One of these patients had erythroid hyperplasia of the bone marrow, and the other had erythroid aplasia. From these observations it was concluded that the major portion of the early peak could not be associated with erythropoiesis. However, this conclusion is at variance with studies in which the amount of labeled urobilin in the early peak did increase after an experimental hemorrhage or was found to be increased in disease states associated with increased rates of erythropoiesis (6).

The present study was undertaken in an attempt to define the relationship between erythropoiesis and the early labeled peak of urobilin by studying a given individual at two different rates of red cell production. With the exception of two patients who were studied before and after an experimental hemorrhage (7, 8), this has not been done previously. Four patients with hematologic diseases that might respond favorably to therapy were selected for study. Before and after treatment these patients were given glycine-2-14C, and the specific activity of urobilin-14C was determined. In addition, erythrokinetic function was evaluated by using either 59Fe or glycine-2-14C as a precursor of heme. We feel that the results of these studies support the concept that in man most of the early labeled peak is associated with erythropoiesis.

Methods

Fecal urobilin isolation. For each study the patients were given intravenous injections of 100 µc glycine-2-14C (SA 2.6 to 4.0 µc per µmole) in a volume of 1 to 5 ml over a 10-second period. Stool samples were collected daily and stored at −20° C until analysis. Early in the study Watson’s original method for the isolation of fecal urobilin was used in a modified form (7, 9), but the most
recent method of Watson and his co-workers (10) was used in later isolations.

The term "urobilin" is used generically in this report to include the closely related pigments urobilin IXa, d-urobilin, and l-urobilin (sterobilin), which are identical from a physiologic point of view and which all may be isolated with the above method (10). However, in any given individual only one species of urobilin is usually present. Except for patient W.G., the specific optical rotation of the urobilin crystals of each sample was determined in chloroform, and the values obtained were consistently within the accepted limits of purity for l (inactive), d-, or l-urobilin (11). In addition, the specific activity of urobilin crystals obtained by these methods was found to be constant after crystallization from ethyl acetate. To determine the specific activity of the urobilin crystals, we plated 1 to 2 mg on a preweighed 5-cm planchet with the aid of a turntable and counted in a low background thin-window β-counter with an efficiency of 22% and a background of 2 cpm. All samples were counted for at least a half-hour or until a minimum of 1,000 counts had been observed.

**Hemin isolation.** The method of Fischer (12) was used for most of the samples. The crystalline material was plated and counted for 14C content as with urobilin. However, for patients S.S. and J.F., hemoglobin was extracted, and calculations were made to convert the data to the specific activity of hemin (13). The values given for each patient represent maximal specific activities at 10 to 20 days.

**Ferrokinetics.** Twenty μg of 55Fe in the form of ferrous citrate with a specific activity of 2 to 40 mc per mg was used in each study. The method of Huff and co-workers (14) was used to determine the plasma radioiron disappearance half-time and the red cell radioiron uptake.

**Blood volume.** The chromium method of Sterling and Gray (15), as modified by Read (16), was used.

**DFP red cell life-span.** Trinitiated diisopropylfluorophosphate (DFP-3H) was used as a red cell label as previously described (17).

**Isotopic measurements.** Samples containing 52Cr and 55Fe were counted in a well-type scintillation counter with a sodium iodide (thallium-activated) crystal and a single channel spectrometer. When both 52Cr and 55Fe were in the same sample, a correction was made for 55Fe activity in the 52Cr counting window.

Crystalline urobilin-14C or hemin-14C was counted by plating the material infinitely thin on a 5-cm diameter planchet and counting in a thin window low background counter. Planchets plated with hemin that contained both 55Fe and 14C were counted twice, with sufficient time between the two countings to allow at least half of the 55Fe to decay. Assuming that the radioactivity due to 14C remained constant during this period, one may then calculate the radioactivity due to 14C alone.

### TABLE I

Erythrokinetic data for four patients with varying rates of erythropoiesis and for two hematologically normal patients *

<table>
<thead>
<tr>
<th>Patient</th>
<th>Study</th>
<th>Plasma iron turnover</th>
<th>RBC iron turnover</th>
<th>Serum iron</th>
<th>Total iron-binding capacity</th>
<th>Plasma radioiron uptake</th>
<th>Red cell life-span</th>
<th>Fecal urobilinogen</th>
<th>Hemin-14C day 9-20</th>
<th>Urobilin-14C maximal value</th>
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<tbody>
<tr>
<td>R.P.</td>
<td>Normal</td>
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<td></td>
<td></td>
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<tr>
<td>W.G.</td>
<td>Normal</td>
<td>0.53</td>
<td>0.43</td>
<td>90</td>
<td>290</td>
<td>68</td>
<td>83</td>
<td>153</td>
<td>110</td>
<td>142</td>
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<td>S.S.</td>
<td>September 1961</td>
<td>0.40</td>
<td>0.37</td>
<td>62</td>
<td>284</td>
<td>52</td>
<td>91</td>
<td>154</td>
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<tr>
<td>A.H.</td>
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<td>0.31</td>
<td>0.005</td>
<td>271</td>
<td>271</td>
<td>390</td>
<td>1.6</td>
<td>400</td>
<td>400</td>
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<tr>
<td>March 1963</td>
<td>0.74</td>
<td>0.610</td>
<td></td>
<td>80</td>
<td>146</td>
<td>57</td>
<td>82</td>
<td>44</td>
<td>404</td>
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<tr>
<td>July 1964</td>
<td>0.63</td>
<td>0.384</td>
<td></td>
<td>213</td>
<td>213</td>
<td>131</td>
<td>61</td>
<td>86</td>
<td>293</td>
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<tr>
<td>J.F.</td>
<td>April 1962</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>800†</td>
<td>9</td>
<td>15</td>
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<tr>
<td>L.F.</td>
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<td>1.06</td>
<td>0.15</td>
<td>202</td>
<td>202</td>
<td>95</td>
<td>14</td>
<td>72</td>
<td>400‡</td>
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<tr>
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<td>0.72</td>
<td>0.43</td>
<td>60</td>
<td>140</td>
<td>93</td>
<td>60</td>
<td>131</td>
<td>220‡</td>
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<tr>
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<td>181</td>
<td>181</td>
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<td>40</td>
<td>33</td>
<td>81</td>
<td>110</td>
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<tr>
<td>January 1964</td>
<td>0.76</td>
<td>0.11</td>
<td>185</td>
<td>207</td>
<td>120</td>
<td>15</td>
<td>14</td>
<td>419</td>
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* Abbreviations: RBC = red blood cell, Hb = hemoglobin, and DFP-3H = trinitiated diisopropylfluorophosphate.
† Calculated from transfusion requirement, red cell life-span, or both.
‡ Two-day pool.
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Fig. 1. Erythokinetic studies of two hematologically normal patients, R.P. and W.G. The radioiron plasma disappearance and red cell (RBC) uptake, used here as an index of the rate of erythropoiesis, are shown as lines connecting the circles; the bar graphs of urobilin-\(^{14}\)C specific activity represent the "early peak."

Fig. 2. Erythokinetic studies and clinical course of patient A.H. The three studies illustrated above, from left to right, indicate the radioiron and urobilin-\(^{14}\)C data obtained before treatment of erythroid hypoplasia, after successful treatment, and 18 months later. The three small arrows at the top margin of the clinical data indicate the times when the glycine-\(^{2}\)\(^{14}\)C and radioiron studies were performed. The specific activity of the early peak of urobilin-\(^{14}\)C, represented by the bar graphs, varied with changes in the rates of plasma clearance and red cell incorporation of radioiron.
Results

Clinical course and erythrokinetic data for each patient may be found in Figures 1–6. Case histories are in the Appendix. The data have also been tabulated in Table I.

Three of the four patients were studied during two or more phases of their illnesses. For comparison, two hematologically normal individuals, R.P. and W.G., were also studied (Figure 1). R.P. had a localized carcinoma of the tongue, and W.G. had psoriasis. Their respective ferrokinetic studies were as follows: plasma radioiron disappearance half-time, 68 and 52 minutes; and maximal red cell $^{59}$Fe incorporation, 83 and 91%. Plasma iron turnover (PIT) values were 0.53 and 0.40 mg per kg per day, and red cell iron turnover (RCIT) was 0.43 and 0.37 mg per kg per day. Two weeks after injection of 100 $\mu$C of glycine-2-14C, the hemin-14C specific activities were, respectively, 110 and 153 dpm per mg, and in two other hematologically normal patients the values were 115 and 152 dpm per mg. The peak specific activities of the urobinin-14C were 142 and 223 dpm per mg. These maximal values for the urobinin-14C specific activity in hematologically normal individuals compare favorably with the maximal specific activity of 91 dpm per mg obtained by White (18) in a normal adult after the intravenous injection of 100 $\mu$C glycine-2-14C. Two additional hematologically normal subjects have been studied by Gray and Scott (7) and

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**Fig. 3. Erythrokinetic Studies and Clinical Course of Patient S.S.** At the time of the first study, when the patient had profound erythroid hypoplasia, there was very little incorporation of isotopic precursors into red cells or the early peak. However, after treatment, red cell production returned, and the amount of isotopic glycine incorporated into hemin and into the urobinin-14C early peak was greater than normal.
Israels and Zipursky (19); one was given 25 μc glycine-2-14C orally and the other, 50 μc intravenously. The maximal observed fecal urobilin specific activities can be extrapolated to a dose of 100 μc, assuming factors of 4 and 2, respectively, to yield calculated peak fecal urobilin specific activities of approximately 75 and 286 dpm per mg, respectively.

Patient A.H. (Figure 2), a 57-year-old man with chronic lymphocytic leukemia, was found to have almost complete erythroid aplasia of the bone marrow at the time of the initial study (January 1963). The plasma radioiron disappearance half-time was 390 minutes, and there was virtually no incorporation of radioiron into red cells and of glycine-14C into urobilin and hemin. The specific activity of the urobilin was 22 dpm per mg, and that of the hemin was 0 dpm per mg. PIT was 0.31 and RCIT 0.005, mg per kg per day. However, after treatment with prednisone and testosterone a reticulocytosis occurred, and a second study was performed (March 1963). Plasma radioiron disappearance half-time was 57 minutes, with a maximal radioiron red cell uptake of 82%. PIT was 0.74 and RCIT 0.61, mg per kg per day. The specific activity of hemin-14C after 2 weeks was 404 dpm per mg, and the maximal specific activity of urobilin-14C was 1,291 dpm per mg, which is greatly elevated. The patient was studied again in July 1964. Plasma radioiron disappearance half-time was 131 minutes, and maximal radioiron red cell uptake was 61%. PIT was 0.63 and RCIT 0.38, mg per kg per day. The specific activity of the hemin-14C was 196 dpm per mg, and hemin-14C red cell life-span was 86 days. The specific activity of urobilin-14C reached a maximum of 596 dpm per mg.

Patient S.S. was a 45-year-old woman with chronic lymphocytic leukemia. In the first study (Figure 3) her bone marrow showed almost complete erythroid aplasia. Ferrokinetic studies showed a prolonged plasma radioiron disappearance half-time of 293 minutes, with no radioiron incorporation into circulating red cells. The maximal specific activity of urobilin-14C was 48 dpm per mg. The second study was done shortly after the peak of reticulocytosis, when the bone marrow contained 10% erythroid precursors, many of which showed megaloblastoid changes. A radioiron study was not performed, but the maximal specific activity of hemin-14C was 461 dpm per mg, which is greater than normal (20). The specific activity of the urobilin-14C early peak was also greater than normal, with a maximal value of 459 dpm per mg.

J.F., a patient with aplastic anemia (Figure 4), was found to have hypoplastic bone marrow. Glycine-14C incorporation into hemin decreased markedly (9 dpm per mg). Maximal urobilin-14C specific activity was very low (15 dpm per mg).

Patient L.F., had refractory anemia with erythroid hyperplasia of the bone marrow, but at autopsy a diagnosis of multiple myeloma was made. Over 2½ years, four studies were performed on this patient. In the first (Figure 5, November

**Fig. 4.** Erythrokinetic studies and clinical course of patient J.F. This patient with aplastic anemia was studied on only one occasion. The maximal specific activities of the hemin-14C and urobilin-14C were both approximately 10% of the values found in normal individuals after a similar amount of glycine-2-14C.
1961), plasma radioiron disappearance half-time was normal (87 minutes), but red cell radioiron uptake was low (14%). PIT was 1.06 and RCIT 0.15, mg per kg per day. The maximal specific activity of urobin-14C was 1,650 dpm per mg, which is markedly elevated. A hemin-14C red cell life-span was 52 days, with a maximal specific activity of 30 dpm per mg. In November 1962 (Figure 5), after treatment with testosterone, the patient's plasma radioiron uptake was 60%. PIT at this time was 0.72 and RCIT 0.43, mg per kg per day. Red cell life-span (DFP-5H) was 131 days. The specific activity of urobin-14C was 125 dpm per mg, which is within the normal range, and the maximal specific activity of hemin-14C was slightly low (70 dpm per mg) (20). In May 1963 (Figure 6), when the patient's rate of erythropoiesis had decreased, plasma radioiron disappearance half-time was 154 minutes, and red cell uptake was 40%. PIT was 0.40, and RCIT 0.18, mg per kg per day. The 51chromium red cell survival half-time was 33 days. Maximal specific activity of urobin-14C was 343 dpm per mg, and that of hemin-14C was 110 dpm per mg. In January 1964 (Figure 6), when the patient had a large transfusion requirement, plasma radioiron disappearance half-time was 120 minutes, with a red cell radioiron uptake of 15%. PIT was 0.76 and
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![Graphs showing erythrogenetic studies and clinical course of patient L.F.]

**Fig. 6. Erythrogenetic studies and clinical course of patient L.F.** The third and fourth studies in L.F., who had previously responded to treatment, were performed when red cell production decreased and the patient became refractory to treatment. Concomitantly, the specific activity of the urobilin-\(^{14}\)C early peak increased to levels greater than normal.

RCIT 0.10, mg per kg per day. Maximal specific activity of urobilin-\(^{14}\)C was 502 dpm per mg, and that of hemin-\(^{14}\)C was 52 dpm per mg.

**Discussion**

The early labeled peak was first observed by Gray and co-workers (2) and London and co-workers (1) in 1950. Both groups felt that the early peak was probably related to erythropoiesis. Subsequent studies in disease states with increased erythropoiesis and after an experimental hemorrhage indicated that the maximal specific activity was increased when the rate of erythropoiesis was increased (6). However, James and Abbott (5) reported studies performed on a patient in which an increased early labeled peak was found at a time when the bone marrow showed erythroid aplasia.

The present study was designed to determine whether the early peak in man is in fact related to erythropoiesis by measuring the incorporation of glycine-2-\(^{14}\)C into the urobilin-\(^{14}\)C early peak at a time when erythropoiesis was markedly reduced or absent as judged by bone marrow examination, reticulocyte count, radioiron plasma clearance and red cell uptake, and glycine-2-\(^{14}\)C incorporation into red cell heme.
Before treatment three patients (A.H., S.S., and J.F.) were found to have a very small early labeled peak when erythropoiesis was markedly reduced and erythroid hypoplasia of the bone marrow was present. After treatment had restored erythropoiesis in two of these patients (A.H. and S.S.), the early labeled peak of urobiolin was found to be increased above normal and markedly increased in comparison with the studies performed when erythropoiesis was absent. The third patient (J.F.) did not respond to therapy.

Patient L.F., in contrast to the preceding three patients, initially had erythroid hyperplasia of the bone marrow. In his first study (Figure 5, November 1961), the presence of anemia and the decreased incorporation of radioiron into circulating red cells, together with a normal plasma radioiron disappearance half-time, a markedly elevated early peak, and erythroid hyperplasia, suggest that a large amount of ineffective erythropoiesis was present. Maximal specific activity of the early peak was 1,650 dpm per mg, which is much higher than the normal range of 75 to 286. A year later (November 1962), the second study was performed. The patient did not require transfusions, and the ferrokinetic studies were normal, suggesting that the synthesis of red cells had returned toward normal. The early peak had also fallen to normal levels. In the ensuing months the patient's effective red cell production slowly decreased, as evidenced by anemia and an increasing transfusion requirement. Concomitantly, the studies in May 1963 and January 1964 (Figure 6) showed an increasing urobiolin-14C specific activity, suggesting that the bone marrow was synthesizing red cells in adequate numbers but that there were defective cells that were catabolized either in the marrow or shortly after their release into the peripheral blood. The hemoglobin-14C contained in the defective red cells was presumably converted to urobiolin-14C, and thus an increased isotope content of the early peak was observed. This combination of findings can be designated as ineffective erythropoiesis.

When patients A.H., S.S., and J.F. had erythroid hypoplasia of the bone marrow, essentially no erythropoiesis, and a low early peak, one may speculate that the metabolic defect was in the synthesis of hemoglobin. However, in patient L.F. with bone marrow erythroid hyperplasia, an elevated early peak, and a normal radioiron plasma clearance, it is likely that the defect occurred at a later stage and could involve the mechanism of nuclear extrusion. Electron microscopic studies by Bessis, Breton-Gorius, and Thiery (21) suggested that 5 to 10% of the reticulocyte cytoplasmic hemoglobin is lost at the time of nuclear extrusion, and Hammel and Bessman (22) have indicated that the nucleus itself may be the site of a significant amount of hemoglobin. It is possible that if the nuclear extrusion mechanism went awry, large amounts of labeled hemoglobin would be found. It is also possible that in the normal individual the hemoglobin contained in the perinuclear cytoplasm and in the nucleus itself could represent a major source of the early peak of labeled urobiolin, a possibility that would be consistent with our studies in which changes in the amounts of labeled precursor in the early peak were associated with changes in the rate of effective erythropoiesis.

After the injection of glycine-2-14C, Yamamoto, Skanderbeg, Zipursky, and Israels (23) found two peaks of isotopic incorporation in plasma bilirubin, one occurring within the first day and the second during the fourth day. The occurrence of the second peak corresponds to the time of nuclear extrusion of the group of red cells that would contain the largest amount of an isotopic precursor (24). Evidence that labeled bilirubin appears in dog fistula bile before labeled heme can be detected in the bone marrow has also been presented (4), suggesting that a nonheme pathway for labeled bile pigment might exist. However, it is more likely that various heme-containing enzymes with rapid turnover times, such as those in the liver, may contribute to the early labeled peak, and it may be this source that accounts for the first of the two plasma bilirubin peaks. This concept is supported by recent studies in bile-fistula dogs (25, 26) in which the early peak of bile bilirubin was separated into erythropoietic and nonerythropoietic fractions by stimulating or suppressing erythropoiesis. It was found that in the normal state one-half to two-thirds of the early peak was of erythropoietic origin, and that the remainder could be accounted for by the radioactivity present as hepatic nonhemo- globin heme. The presence of a hepatic contribution to the early peak of labeled urobiolin in man is suggested by studies in which an early peak ap-
peared in a normal individual after the ingestion of delta-aminolevulinic acid-\(^{14}\)C, a poor \textit{in vivo} precursor of red cell heme (27), and also by the initial studies in patients A.H. and S.S., in whom urobilin specific activity was found to be very low but never absent, whereas virtually no radioiron was incorporated into the red cells (Figures 2 and 3). In man, however, pigment production associated with erythropoiesis certainly constitutes the bulk of the fecal urobilin early peak, as shown by the extremely low urobilin specific activity in the absence of erythropoiesis in patients A.H., S.S., and J.F.

Theoretically, once the specific activity of the urobilin is known, it should be possible to calculate the total amount of precursor in the early peak if the total urobilin excretion is also known. However, the quantitative fecal urobilinogen determinations is inexact, and meaningful calculations cannot be made (6, 8). Alternatively, the amount of labeled precursor in the early and late peaks could be compared, but this would require steady state conditions for more than 120 days and would be impossible in patients without a “late” peak, such as those with rapid rates of hemolysis or those in whom the amount of labeled precursor incorporated into circulating red cells is very small. These difficulties make it necessary for one to evaluate the early peak of labeled urobilin excretion in terms of the maximal specific activity. This is a valid concept if the precursor is given intravenously over a short period of time and if fecal samples are analyzed in 1-day blocks as was done in this study. This method of comparison requires that the area under the curve be proportional to the maximal value and that the urobilin production be constant, thus making the maximal value the principal variable reflecting the total amount of isotope in the peak. With these assumptions, the contribution to the early peak in man that is not due to erythropoiesis may be estimated as approximately 7 to 10% (J.F.), 10 to 15% (A.H.), and 22 to 34% (S.S.).

The possibility that the differences observed in the urobilin specific activities before and after treatment could be the result of changes in the total daily excretion of bile pigment was also examined. Although fecal urobilinogen determinations were not obtained in every case, it was possible to estimate the rate of pigment excretion from the transfusion requirement and the rate of change of venous hemoglobin concentration. When we made these calculations, we found that the maximal change in total bile pigment excretion was twofold, whereas the smallest change in the pre- and post-therapy urobilin maximal specific activities was ninefold. Thus—despite some variation in the total pigment excretion—because of the large change in the specific activity before and after treatment, it is clear that the maximal urobilin specific activities are a presently acceptable reflection of the rate of isotope excretion.

**Summary**

An “early peak” of labeled fecal urobilin excretion occurs during the first few days after the intravenous administration of glycine-\(^{2-14}\)C to man. To determine if this early peak is associated with erythropoiesis, we gave four patients with hematologic diseases and two hematologically normal patients glycine-\(^{2-14}\)C, a precursor of bile pigment, and thereafter made determinations of the amount of radioactivity in the fecal urobilin. Three patients were studied when erythropoiesis was virtually absent, and in all three instances the specific activity of the fecal urobilin-\(^{14}\)C was markedly reduced when compared to hematologically normal subjects. Two patients (S.S. and A.H.) were studied a second time during a period of reticulocytosis and rising hemoglobin concentration. The maximal specific activity in the urobilin-\(^{14}\)C increased approximately tenfold in patient S.S. and 60-fold in A.H. Thus, the maximal urobilin-\(^{14}\)C specific activity was shown to be low initially when compared with normal and to increase markedly when erythropoiesis was active. This suggests that a small early labeled peak is present in man in the absence of erythropoiesis but that the bulk of the peak is associated with red cell formation in the marrow in these patients. A fourth patient with anemia and normoblastic hyperplasia of the marrow was studied four times. In this patient the maximal urobilin-\(^{14}\)C specific activity was shown to decrease by approximately 13-fold concomitant with return of the bone marrow toward normal and return of effective erythropoiesis.
Acknowledgments

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Appendix

Case histories of patients

A.H. (NIH no. 02-63-51), a 57-year-old white man, was found in 1959 to have a leukocyte count of 290,000 per mm³, with 90% lymphocytes, and a bone marrow characteristic of chronic lymphocytic leukemia. He was treated in 1960 and 1961 with a total of 160 röentgens of whole body X-irradiation. In July 1962, the hemoglobin (Hb) was 12.0 g per 100 ml, but by November it had dropped to 5.0, and in January 1963, at the time of initial study, it was found to be 2.9 g per 100 ml. His bone marrow showed 90% lymphocytes and no erythroid cells. He was treated with prednisone, but because there was no apparent response after 3 weeks, intramuscular injections of testosterone were begun. After 2 weeks of this treatment, an increase in reticulocyte count was noted, and a second study was performed. The bone marrow was hypercellular with 35% lymphocytes and 45% erythroid precursors. One and one-half years later, at the time of the third study, the Hb was 12.6 g per 100 ml, reticulocyte count was 1.5%, and bone marrow aspiration revealed a hyperplastic marrow packed with lymphocytes and containing abundant reticulin and collagen fibrosis. Occasional normoblasts were seen.

S.S. (NIH no. 01-49-65), a 41-year-old white woman, first sought medical attention in 1956 because of fatigue. She was found to have cervical adenopathy, splenomegaly, and profound anemia. The leukocyte count was 1,000 per mm³, with 80% lymphocytes, and the reticulocyte count was 0%. Her bone marrow showed normal cellularity with infiltration by small lymphocytes; a diagnosis of chronic lymphocytic leukemia was made. Chemotherapy was given, and subsequently the cellular elements of the blood increased and transfusions were no longer necessary. In late 1959 pancytopenia again developed but was successfully treated by splenectomy. In mid-1960, however, transfusions again became necessary. The bone marrow showed a striking decrease in the number of erythroid precursors, with a myeloid:erythro ratio of 20:1. During the summer of 1961, the patient was treated with a testosterone preparation with no apparent effect. The first of the patient's two studies was done at this time. In October the patient began on triamcinolone and 1 week later reticulocytosis began. The second study was done during reticulocytosis. The patient developed chronic otitis media and died with a septicemia in February 1962.

J.F. (NIH no. 04-19-59), a 17-year-old Negro male student, was admitted in April 1962 with a diagnosis of idiopathic aplastic anemia. No abdominal organs were palpable, and an axillary lymph node biopsy was normal. The Hb was 8.7 g per 100 ml, the platelet count was 5,000 per mm³, and the leukocyte count was 800 per mm³, with 90% lymphocytes and 8% polymorphonuclear leukocytes (FMN). The reticulocyte count was 0, and the bone marrow was markedly hypocellular, with 20% normoblasts and 66% lymphocytes. Prednisone and testosterone were given without apparent response. The patient died several weeks after discharge.

L.F. (NIH no. 03-92-55), a 65-year-old white engineer, was first admitted in November 1961, with refractory anemia. There was no adenopathy or organomegaly. The transfusion requirement was 2 U per month, the reticulocyte count was 1%, the platelet count was 82,000 per mm³, and the leukocyte count was 3,000 per mm³, with 30% PMN and 70% lymphocytes. There were 75 nucleated red cells per 100 leukocytes. Moderate anisocytosis and poikilocytosis of the red cells were present, and the mean cell volume was 108 μ³. The bone marrow was moderately hypercellular, with occasional megaloblastoid cells. Seventy-three per cent of the marrow cells were in the erythroid series, with abundant sideroblasts. No periodic acid-Schiff positive cytoplasmic inclusions were seen in the erythroid cells. Injections of a long acting testosterone preparation were begun, and prednisone was given until May 1962.

The patient was readmitted in November 1962. No transfusions had been necessary during the preceding 5 months. The Hb was 15.2 g per 100 ml, the reticulocyte count was 0.5%, and the leukocyte count was 3,800 per mm³, with 60% PMN and 28% lymphocytes. The mean cell volume was 110 μ³. The bone marrow hypercellularity was less marked, few sideroblasts were seen, and the maturation of the granulocytic and erythrocytic series was normal. The testosterone therapy was stopped.

The patient was admitted again in May 1963. In the interim the Hb had fallen slowly to 12 g per 100 ml. The fecal urobilinogen was 81 mg per day, and serum protein electrophoresis revealed a total protein of 6.3, albumin of 3.6, and gamma globulin of 1.2 g per 100 ml. The bone marrow contained many sideroblasts, and there was a mild shift to the left with 27% myeloblasts. Testosterone injections were reinstituted.

Transfusions became necessary in October 1963, at the rate of 3 to 4 U per month. The patient was readmitted in January 1964. The fecal urobilinogen was 419 mg per day. Bone marrow was hypercellular, with 40% of the cells present as abnormal erythroid elements with megaloblastoid changes. Many sideroblasts were present. The total protein was 6.6 g per 100 ml, and the albumin was 3.5 g per 100 ml. Unsuccessful trials of a number of hematologic agents were given, and the patient was then begun on prednisone in addition to testosterone. After 4 months no improvement was noted on this regimen. Be-
cause of dysuria the patient was admitted to another hospital, where serum protein electrophoresis revealed a total protein of 7.8 g per 100 ml, with 32% albumin and 39% gamma globulin (broad band). The patient died suddenly after a prostatic biopsy. Autopsy examination revealed a hyperplastic marrow with many plasma cells, and a diagnosis of multiple myeloma was made.

References
18. White, P. Personal communication.