THE URINARY EXCRETION AND BIOLOGIC DECAY PERIODS OF RADIOMERCURY LABELING A MERCURIAL DIURETIC IN NORMAL AND DISEASED MAN

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In previous studies (1) some aspects of the mechanisms of renal excretion of radiomercury \(^1\) labeling an organic mercurial diuretic (Mercuhydrin \(^4\)) were observed in acute experiments in which renal venous blood and pelvic urine were obtained by means of catheters in a few subjects. The present observations are concerned with the rate of urinary excretion of the radiomercury of this diuretic in a larger number of subjects over a longer period of time. The percentage of injected radiomercury which was excreted in the urine, the time relationships of the diuretic effects and the urinary excretion of the mercury, the biologic decay rates of the isotopes, as well as other physiologic phenomena were determined from these experiments. Because of the extensive use of mercurial diuretics in clinical medicine and the problems of toxicity, the biologic decay rates are of importance.

MATERIALS AND METHODS

Eighty-three hospitalized subjects were studied. The diuretic was administered intravenously and intramuscularly to normal human subjects and to subjects with congestive heart failure. Table I shows the route of administration employed for the subjects of each clinical group. The control subjects were free from cardiovascular and renal diseases; most of them had primarily such illnesses as psychoneuroses or peptic ulcer or were convalescing from a respiratory infection. The subjects who were made to sweat had psychoneurosis or inactive peptic ulcer. They were placed in a hot and humid room (\( \approx 45^\circ \) C. and 95 per cent relative humidity) for periods varying from 20 to 90 minutes, the sweating procedure beginning from 0 to 90 minutes after injection of the diuretic. The patients with chronic congestive heart failure included groups in varying stages and degrees of failure with and without edema. The group of miscellaneous subjects had renal disease (subacute or chronic active glomerulonephritis) or hepatic cirrhosis.

One to 3 cc. of the mercurial diuretic, each containing 39 mg. of mercury, were administered by routes indicated in Table I. The quantity of radioactive material administered varied from 100 \( \mu \)c. in the earlier experiments to 10 \( \mu \)c. in the later ones.

Each voided specimen of urine was collected separately and its volume and radioactivity were determined (2). Samples of plasma or serum were obtained at varying intervals of time. Collections of urine and blood were made until there was no detectable radioactivity, \( \text{i.e.} \), until the counts reached background level.

Expression of data. The radiomercury in each fluid studied was expressed as counts per minute per cc. The time-course of the cumulative excretion of radiomercury was expressed as a percentage of the total counts excreted in the urine for all subjects in whom collections of

| TABLE I |
| Route of administration of radiomercury injected for various clinical groups |

<table>
<thead>
<tr>
<th>Subject groups</th>
<th>Route of administration</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>I. V.</td>
<td>I. M.</td>
</tr>
<tr>
<td>Control</td>
<td>35</td>
<td>8</td>
</tr>
<tr>
<td>Sweated</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>Congestive heart failure</td>
<td>20</td>
<td>7</td>
</tr>
<tr>
<td>Miscellaneous</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>68</td>
<td>15</td>
</tr>
</tbody>
</table>
all urinary output during the study were made. A mean curve was obtained for each group of subjects.

Since 88 per cent of the injected mercury was recovered for the control subjects, 100 per cent of the recovered mercury represents 8,800,000 CPM (counts per minute) or 68.64 mg. in a dose of 2 cc. This relationship is approximately the same for the subjects with congestive heart failure.

In the early studies errors existed due to loss by volatilization of mercury from the preparations of urine made for counting. Data obtained after methods of circumventing these difficulties were devised permitted determination of the percentage of the total administered dose excreted in the urine of 12 subjects. Because of the extreme variation in the chemical nature of urine, the error due to volatilization was inconstant from subject to subject but remained relatively constant for all of the samples of any one subject. The temporal and percentile relationships are therefore satisfactory. The problem of volatilization and other special features peculiar to tracer studies with mercury are presented in detail elsewhere (3).

It should be emphasized that only the radiomercury of the mercurial diuretic was traced regardless of its chemical state; the latter was unknown at all times.

RESULTS

To simplify the presentation of the data, results are divided into: (1) biologic decay periods, (2) time-course of urinary excretion, and (3) rate of clearance of radiomercury by the kidneys.

(1) Biologic decay periods (Results are summarized in Table II)

(a) \( \text{C}_4 \) values. The mean \( \text{C}_4 \) values, determined graphically, for the control subjects were smaller than those for the subjects with chronic congestive heart failure. \( \text{C}_4 \) for radiomercury was slightly smaller, when the diuretic was administered intravenously than when administered intramuscularly, although there was overlapping of values.

Although \( \text{C}_4 \) was previously described for radioiodine (4) and radiochlorine (5) as the time required for the concentration in the serum to reach half the concentration after equilibrium of distribution of the isotope had been reached, the radiomercury apparently reaches a steady state rather than equilibrium of distribution because of the rapid rate of excretion. However, in subjects with considerable impairment of renal function equilibrium of distribution was probably obtained.

(b) Time to reach background count. The time in hours required for the radiomercury in the plasma to reach a background count (\( \text{C}_6 \)) is

| TABLE II |
| Biologic decay periods of radiomercury |

<table>
<thead>
<tr>
<th>Group</th>
<th>No. of subjects</th>
<th>Mean ( \text{C}_4 ) hrs.</th>
<th>Mean ( \text{U}_4 ) hrs.</th>
<th>Mean ( \text{UE}_4 ) hrs.</th>
<th>Mean ( \text{U}_b ) hrs.</th>
<th>Mean ( \text{W} ) hrs.</th>
<th>Total recovery*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control, I. V.</td>
<td>25</td>
<td>2.2 (1.0–3.7)</td>
<td>2.3 (1.5–4.5)</td>
<td>2.2 (0.8–7.4)</td>
<td>27.2</td>
<td>4.8</td>
<td>88 (80.4–95.5)</td>
</tr>
<tr>
<td>Control, I. M.</td>
<td>5</td>
<td>2.5 (2.2–3.2)</td>
<td>4.1 (2.4–7.8)</td>
<td>37.2</td>
<td>7.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control Sweated, I. V.</td>
<td>7</td>
<td></td>
<td>2.1 (0.9–2.4)</td>
<td>36.7</td>
<td>14.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Congestive heart failure, I.V.</td>
<td>8</td>
<td>4.0 (2.2–7.0)</td>
<td>3.2 (1.6–6.0)</td>
<td>43.9</td>
<td>14.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Congestive heart failure, I. M.</td>
<td>6</td>
<td>5.0 (2.9–11.5)</td>
<td>4.6 (3.1–6.4)</td>
<td>45.7</td>
<td>12.8</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* For 12 control subjects in whom effects of volatilization did not offer difficulties (2).
indicated in Table III for the various groups of subjects. Since the samples of serum were collected at six to 24 hour intervals after the first six hours of study, the C₀ values are not sharply demarcated (Table III).

The serum counts reached background within the first 24 hours in 94 per cent of the control subjects; in only one was there radiomercury in the serum on the second day when the diuretic was administered intravenously. When administration was intramuscular, approximately the same amount of time was required for background to be reached. Thermal sweating apparently did not alter the concentration-time course of the radiomercury in the serum, whereas chronic congestive heart failure tended to prolong the latter (Table III).

(c) \( U₄ \). A mean \( U₄ \) value, the time required for half of the injected radiomercury to be excreted in the urine, of 2.3 hours (range 1.5 to 4.5 hours) was obtained for the 12 control subjects (Table II). This \( U₄ \) value is almost the same as the \( C₄ \) value of 2.2 hours for the same group, which is to be expected since almost all the radiomercury is excreted by way of the urine (6).

(d) \( UE₄ \). These values represent the time required for half the total radiomercury excreted in the urine to be eliminated by way of the urine and are summarized in Table II. As would be expected from the \( U₄ \) and \( C₄ \) values previously discussed, \( UE₄ \) was larger for subjects with chronic congestive heart failure than for the control subjects and larger for the intramuscular route of administration than for the intravenous (Table II).

(e) Time for urine to reach background, \( U₅ \). This parameter measures the time after injection when no more measurable radioactive mercury appears in the urine and is represented in Figure 1 by the time when the 100 per cent value was reached. The control subjects excreted "all" of the mercury more rapidly than did the subjects with congestive heart failure (Table II).

(f) \( WE₄ \). This parameter indicates the time required for half the total volume of urine containing measurable amounts of radiomercury to be excreted. These mean values as well as the \( UE₄ \) values (Table II) were obtained from the curves shown in Figure 1. The \( WE₄ \) values were larger for the subjects with chronic congestive heart fail-

FIG. 1. THE MEAN CUMULATIVE OUTPUT OF RADIOMERCURY AND WATER FOR FIVE GROUPS OF SUBJECTS

The 100 per cent value for radiomercury represents the total recovered mercury. The 100 per cent value for water output represents the total volume of urine excreted from the time of injection to the time when no more radioactivity was detectable in the urine. The curves may be expressed by the equation

\[
Rₜ = \sum_{s=1}^{t} X Rₛ
\]

where \( R \) = percentage excreted up to time \( t \), where \( s = \) any one unit of time, and where \( t = \) time in hours.
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PERCENTAGE EXCRETION OF RADIONERCUY AND WATER

FIG. 2. THE CUMULATIVE OUTPUT OF RADIONERCUY AND WATER FOR SUBJECT R. P. WITH CONGESTIVE HEART FAILURE AND RENAL INSUFFICIENCY

The almost linear curve of water excretion indicates an absence of diuresis.

Figure 1 are partially the result of differences in rate of excretion of radionercury, by virtue of the definition of $WE_i$.

Some differences in the slopes of the curves for water excretion are due to differences in diuretic response. There was less diuresis in the sweated than nonsweated subjects (Figure 1), and in subject R. P. (Figure 2) no diuretic response at all was noted, the curves of water excretion approaching a straight line. It should be emphasized that inasmuch as water intake was not controlled, the water curves have no volumetric quantitative value and are significant only in their temporal and percentile relationships from one group to the next.

(g) Particular subjects. Subjects with renal insufficiency excreted the radionercury slowly; the summarized data on one such subject (R. P.) are shown in Figure 2. The $C_4$ value was 45 hours, $UE_4$ 36 hours, and $U_b$ 191 hours or eight days. This subject excreted, in relatively large volumes of hyposthenuric urine, only approximately 19 per cent of the radionercury administered (Table IV). Another subject (J. E.) with renal insufficiency and oliguria who died 112 hours after the injection excreted only 2.4 per cent of the injected radionercury in that time. This subject had an estimated mass of edema fluid of 30.2 kg. with a concentration of 70 CPM/cc. of edema fluid or 2.1 million CPM, approximately 50 per cent of the total radionercury administered, or 69 per cent in the extracellular fluid when the serum is also included with the edema fluid. The edema can be a storage depot of a large quantity of mercury.

(2) Time-course of urinary excretion of radionercury

The time-course of urinary excretion of the radionercury was obtained by differentiating the curves shown in Figure 3. These mean curves

<table>
<thead>
<tr>
<th>Subject</th>
<th>$C_4$</th>
<th>$UE_4$</th>
<th>$WE_4$</th>
<th>$C_b$</th>
<th>$U_b$</th>
<th>Urine</th>
<th>BP</th>
<th>Edema</th>
</tr>
</thead>
<tbody>
<tr>
<td>R. P.</td>
<td>45</td>
<td>36</td>
<td>83</td>
<td>221</td>
<td>191</td>
<td>235/66</td>
<td>208/110</td>
<td>4+</td>
</tr>
<tr>
<td>J. E.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>85</td>
<td>258/158</td>
<td>4+</td>
</tr>
</tbody>
</table>
are essentially the same as the means of the actually observed curves for each subject. These studies, as indicated in a previous publication (6), show a concordant relationship between the concentration of radiomercury in the serum and in the urine.

The decline in rate of excretion of the radiomercury was exponential in nature, especially when the intravenous route of administration was employed, the general equation being

\[ R_{t2} = R_{t1}e^{-a(t_2 - t_1)}, \]

where \( R_{t2} \) = the rate of excretion at time \( t_2 \),
\( R_{t1} \) = the rate of excretion at an earlier time \( t_1 \),
\( a \) = the fraction of decrease in the rate of excretion per unit of time,
\( e \) = log base.

This equation holds only after a maximum rate of excretion of the isotope has been achieved. The maximum was reached more rapidly following intravenous than intramuscular administration (Figure 3). Figures 4 and 5 show the slow rate of urinary excretion of the radiomercury for the two subjects with renal insufficiency (Figure 3 and Table IV). The maximum rate of excretion for one of the subjects (J. E.) was essentially 0.005 that of the subjects with congestive heart failure, the mean rate of excretion for this subject being less than 0.001 of the mean rate of even the subjects with congestive heart failure.

(3) Rate of radiomercury clearance by the kidneys

The rate of excretion of the radiomercury may also be expressed by the rate of renal clearance of the serum of radiomercury. The values were ob-

### CHANGES IN SERUM CONCENTRATION AND RATES OF EXCRETION OF MERCURY AND WATER

![Graph](image)

**Fig. 3. The Mean Serum Concentration and Mean Rates of Excretion of Radiomercury and Water for the Five Groups Studied**

The rates of urinary excretion are expressed as percentage of the total recovered mercury excreted per minute. These curves of urinary excretion are differentials of the cumulative curves of Figure 1, or \( \% / \text{min} = d\% / dt \), where \( t \) is expressed in minutes. Each 0.1 per cent for the control group equals 8,800 CPM of the radiomercury or 0.06864 mg. of regular mercury of the diuretic. This relationship is only approximate for the subjects with congestive heart failure.
CHANGES IN SERUM CONCENTRATION AND RATES OF EXCRETION OF MERCURY AND WATER

Subject R.P.—Cardiorenal Failure

\[
\begin{align*}
\text{Hg}^{203, 205} & \quad \% / \text{Min.} \\
\text{H}_2\text{O} & \quad \% / \text{Min.} \\
\text{Serum Conc.} & \quad \text{CPM/CC.}
\end{align*}
\]

FIG. 4. THE PLASMA CONCENTRATION AND RATES OF EXCRETION OF RADIOMERCURY AND WATER FOR SUBJECT R. P.

Each 0.1 per cent represents approximately 1,900 CPM of the radiomercury or 0.015 mg. of the regular mercury in the diuretic.

CHANGES IN SERUM CONCENTRATION AND RATES OF EXCRETION OF MERCURY AND WATER

Subject J.E.—Uremia, Terminal

\[
\begin{align*}
\text{Hg}^{203, 205} & \quad \text{CPM/Min.} \\
\text{H}_2\text{O} & \quad \text{CC/Min.} \\
\text{Serum Conc.} & \quad \text{CPM/CC.}
\end{align*}
\]

FIG. 5. THE PLASMA CONCENTRATION AND RATES OF EXCRETION OF RADIOMERCURY AND OF WATER FOR SUBJECT J. E., A SUBJECT WITH EXTREME OLIGURIA WHO DIED IN UREMIA

The rates of excretion are expressed as CPM/min. and cc./min., since this subject died before the urine reached background for the radiomercury.
with impaired renal function retained considerable quantities of the administered mercury even though urinary volume was relatively high. Measurable quantities of radiomercury were present in the serum of one subject for eight days, even though urinary volume was not reduced nor was there any diuretic response to the drug. Another subject with renal insufficiency excreted only about 2.5 per cent of the injected mercury in about four and one-half days. These observations indicate the need for continual evaluation of renal function to ascertain whether mercury is being retained in excessive quantities.

It is evident from the data that normal subjects excrete the mercury of the mercurial diuretic rapidly, half of it being excreted in a little over two hours whether the drug is administered intravenously or intramuscularly. On the other hand the average control subject still had mercury in the urine after 24 hours, which indicates considerable accumulation when daily doses are administered. This accumulation is even greater in the presence of disease states which reduce the rate of excretion of the mercury, such as congestive heart failure (Table II). It must also be remembered that these individual variations may be considerable and this accumulation will be even greater in certain people even though their clinical state may appear to be similar.

If a constantly elevated level of mercury in the serum is beneficial, then such an occurrence should cause no concern. If, however, accumulation of mercury at a rate of approximately 4 per cent of the injected dose per day is detrimental, then caution should be exercised and repetition of the injections should proceed at greater intervals.

The time required for the radiomercury to be "completely" excreted, or for the radiomercury to be too low in concentration to be measured in the serum, would be expected to be about ten times that required for the serum concentration to reach half-concentration after a steady state is attained. Since the regression of mercury from the serum is exponential in nature, ten times the one-half value would be 1/1,024 the initial level (1000–2000 CPM/cc.) or 1 to 2 CPM/cc. after about 25 hours.

The importance of the C₁ and C₂ of the radiomercury or mercury values in therapy with mer-
curial diuretics is self-evident. Individual variations in excretion of the mercury are considerable, especially when renal disease exists. The mercury tends to be excreted more slowly when the diuretic is administered intramuscularly than when administered intravenously. This must be due in part to delay in absorption from the tissues. Other unknown factors may also be concerned with the difference.

Congestive heart failure results in delay in excretion of the mercury, as evidenced by greater $C_1$ and $C_2$ values for these subjects than for the control subjects. The mechanism of this delay is not known; it may be related to disturbances in the circulation itself as well as to reduction in renal function. Associated impairment of renal function or failure of diuresis to develop may result in further retention of mercury.

Acute thermal sweating for a relatively short period of time (approximately 20 to 90 minutes) resulted in a depression of the rate of water excretion but did not alter the excretion of mercury in the urine. The mechanism for this can only be conjectured.

Failure to collect 100 per cent of the injected mercury in the urine is probably due to losses of the element through other body fluids and volatilization during preparation of samples for counting (2) and to errors in measurement which occur when background levels of counts are approached.

**SUMMARY**

1. The mercury of a mercurial diuretic administered intramuscularly was excreted rapidly when the cardiovascular and renal functions were normal, one-half being excreted in approximately one to eight hours (mean, about three hours). The rate of excretion was slightly less rapid when the drug was administered intramuscularly than when administered intravenously.

2. Chronic congestive heart failure tended to diminish the rate of excretion, although individual variations were large. There was overlapping of the values for the control subjects and those with chronic congestive heart failure. The state and phase of the failure influenced the rate of excretion.

3. The rate of excretion of the radiomercury was considerably impaired by renal insufficiency; the degree of impairment may be great enough to result in accumulation of toxic quantities of mercury with frequent administration of the drug. The importance of this impairment in excretion during clinical therapy is evident.

4. Detailed biologic decay rates, rates of clearance, and other excretory factors are presented.

**BIBLIOGRAPHY**


