Metabolism of Reverse Triiodothyronine by Isolated Rat Hepatocytes

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Abstract
Reverse triiodothyronine (rT₃) is metabolized predominantly by outer ring deiodination to 3,3'-diiodothyronine (3,3'-T₂) in the liver. Metabolism of rT₃ and 3,3'-T₂ by isolated rat hepatocytes was analyzed by Sephadex LH-20 chromatography, high performance liquid chromatography, and radioimmunoassay, with closely agreeing results. Deiodinase activity was inhibited with propylthiouracil (PTU) and sulftoltrerase activity by sulfate depletion or addition of salicylamide or dichloronitrophenol. Normally, little 3,3'-T₂ production from rT₃ was observed, and ¹²⁵I-rT₃ was the main product of both 3,3',5'-I₂T₃ and 3,3',5'-I₂T₄. PTU inhibited rT₃ metabolism but did not affect 3,3'-T₂ clearance as explained by accumulation of 3,3'-T₂ sulfate. Inhibition of sulfation did not affect rT₃ clearance but 3,3'-T₂ metabolism was greatly diminished. The decrease in ¹²⁵I formation from rT₃ was compensated by an increased recovery of 3,3'-T₂ up to 70% of rT₃ metabolized. In conclusion, significant production of 3,3'-T₂ from rT₃ by rat hepatocytes is only observed if further sulfation is inhibited.

Introduction
In euthyroid subjects the main secretory product of the thyroid is thyroxine (T₄).¹ Some 3,3',5-triiodothyronine (T₃) is secreted as well, but thyroidal production of 3,3',5'-triiodothyronine (rT₃) is negligible. More than 97.5% of plasma rT₃ and ~80% of plasma T₃ originate from peripheral deiodination of T₄ (1). As thyroid hormone bioactivity is exerted largely through T₃, it is important to understand the regulatory mechanisms of iodothyronine metabolism.

In rats, three types of iodothyronine-deiodinating enzymes have been identified (2). Most likely, the type I deiodinase of liver catalyzes both inner ring deiodination (IRD) and outer ring deiodination (ORD) of iodothyronines (2). A similar enzyme is present in kidney and thyroid (2). Type II deiodinase has been localized in brain, pituitary, brown adipose tissue, and placenta.

It deiodinates only the outer ring of substrates such as T₄ and rT₃ (2-4). Type III enzyme is found in brain, placenta, and skin, and it is a specific inner ring deiodinase (2, 5). Type I deiodinase is inhibited by 6-propylthiouracil (PTU), while types II and III are PTU insensitive (2). Thus, there may be two sources of plasma rT₃, namely type I or type III IRD of T₄. There are also two pathways of rT₃ deiodination, i.e., type I and type II ORD to 3,3'-diiodothyronine (3,3'-T₂).

Considering the high rate of rT₃ ORD by the type I deiodinase (2), it is likely that little rT₃ produced from T₄ in the liver is released into the circulation. It would seem, therefore, that most plasma rT₃ is derived from type III deiodination of T₄, while it is cleared mainly by the liver. This hypothesis is substantiated by measurements of arterio-venous gradients of rT₃ across the liver in patients with mild liver failure.²

In vivo studies in normal rats have demonstrated that the type I deiodinase is the predominant site for the peripheral production of T₃ (6). Opposite variation in plasma T₃ and rT₃ concentrations has been observed in a number of clinical situations, in which changes are due to a decrease in both the production of plasma T₃ and the clearance of plasma rT₃ (7). To investigate the potential importance of changes in type I deiodinase activity for the regulation of thyroid hormone metabolism, we initiated studies of the deiodination of rT₃ by isolated rat hepatocytes.

Initial results, using outer ring ¹²⁵I-labeled rT₃, showed that radioliodoide was the main product, but little production of 3,3'-T₂ from unlabeled rT₃ could be detected by radioimmunoassay (RIA) (8). Further investigations have demonstrated rapid metabolism of added 3,3'-T₂ in rat hepatocytes by sulfation and subsequent ORD of the 3,3'-T₂ sulfate (3,3'-T₂S) formed (9). 3,3'-T₂S is a far better substrate for the type I deiodinase than 3,3'-T₂ itself (9). This may be the reason for our failure to detect significant production of 3,3'-T₂ from rT₃ by liver cells. If so, the yield of 3,3'-T₂ produced by this pathway should increase if its further sulfation is inhibited. This hypothesis was tested in the present study using rat hepatocytes with diminished phenol sulfotransferase activity.

Methods
The materials used are essentially the same as described previously (10). Carrier-free 3,3',5'-I₂T₃ and 3,3',5'-I₂T₄ were prepared in our laboratory by radiiodination of 3-iodothyronine or of 3,3'-T₂ (Henning GmbH, Berlin, Federal Republic of Germany) using the chloramine T method (11, 12) and purified by Sephadex LH-20 chromatography. Salicylamide (SAM) and 2,6-dichloro-4-nitrophenol (DCNP) were purchased from Riedel-de Haën AG, Hannover, Federal Republic of Germany. All other chemicals were of the highest quality commercially available.

Hepatocytes. Rat hepatocytes were prepared by collagenase perfusion (10). Monolayers of hepatocytes were obtained by seeding 10⁶ cells in 2 ml culture medium (10) into uncoated 3.5-cm wells of plastic 6-well dishes (Nunc, Roskilde, Denmark). The plates were kept for 4 h at 37°C

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in a culture stove under atmospheric conditions. Before each experiment, cell viability was tested by trypan blue exclusion and exceeded 85%. Nonviable cells were removed by aspiration of the medium.

General incubation procedures. Incubations were done for 60 min in 2 ml of Dulbecco’s balanced salt solution which contained 1 mM MgSO4, 0.1% bovine serum albumin (BSA), 2 mM glutamine, and 1 mM vitamin C. Experiments were carried out in triplicate under atmospheric conditions at 37°C. The dishes were placed on a slightly angled, slowly rotating plate. Substrate levels were 10 nM rT3, 3,3'-T2 with or without 0.1 μg [125I]rT3 or [123I]3,3'-T2, respectively. Sulfation was inhibited with 100 μM SAM or 100 μM DCNP and deiodination was inhibited with 10 μM PTU.

Sulfate depletion. Hepatocytes were preincubated for 30 min at 37°C in Dulbecco’s solution which contained 1 mM MgCl2, 2 mM glutamine, and 1 mM vitamin C with or without 100 μM SAM. Incubations were done as described above in medium without SO42-. Controls were preincubated and incubated in medium containing MgSO4.

Analysis of incubation medium. Incubation media with unlabeled rT3 and 3,3'-T2 were analyzed before and after hydrolysis with specific RIAs (11, 12). Hydrolysis of eventual S conjugates was achieved by addition of 250 μl 1 N HCl to 100-μl samples and treatment for 1 h at 80°C (13). Thereafter, 300 μl 1 N NaOH was added. The nonhydrolyzed samples were treated the same way but without heating. RIA was done in duplicate on 50-μl aliquots of the mixtures.

Incubation media with labeled rT3 and 3,3'-T2 were assayed by column chromatography. An equal volume of 1 N HCl was added to 500-μl samples and the mixtures were applied to small (bed volume 0.75 ml) Sephadex LH-20 columns equilibrated in 0.1 N HCl. Stepwise elution was done by successive application of 2 x 1 ml 0.1 N HCl, 6 x 1 ml H2O, 6 x 1 ml 0.1 N NaOH-ethanol (9:1, vol/vol), and 3 x 1 ml 0.1 N NaOH-ethanol (1:1, vol/vol).

High performance liquid chromatography (HPLC). Analysis of rT3 and 3,3'-T2 conjugates and native iodothyronines was accomplished by reverse-phase HPLC. For this we used a 10 x 0.3-cm CP Spher C18 column (Chrompack, Middelburg, The Netherlands), a model 6000 A solvent delivery system and a model 440 fixed wavelength detector (Waters Assoc., Millipore Corp., Milford, MA). Flow was 0.6 ml/min and absorbance was measured at 254 nm. For separation of conjugates a 20:80 vol/vol mixture of acetonitrile and 0.02 M ammonium acetate (pH 4) was used, and for separation of rT3 and 3,3'-T2 a 55:45 vol/vol mixture of methanol and 0.02 M ammonium acetate (pH 4) was employed.

To obtain the conjugates for HPLC analysis, the water fractions of the LH-20 chromatography were pooled, acidified, and rechromatographed. The columns were washed with 2 ml 0.1 N HCl, 0.5 ml H2O, and 0.5 ml 0.1 M ammonia in ethanol. Conjugates were then collected in a subsequent fraction of 1 ml 0.1 M ammonia in ethanol. The solvent was evaporated at 50°C under a stream of N2. To obtain the iodothyronine fraction, medium was processed as above and LH-20 chromatography was modified as follows. After the H2O fractions, columns were washed with 0.5 ml 0.1 M ammonia-ethanol (1:1, vol/vol), and iodothyronines were collected in 1 ml 0.1 M ammonia in ethanol. The solvent was evaporated as above.

Enzymatic analysis of rT3 and 3,3'-T2 conjugates. Another method by which the conjugates were identified was by enzymatic hydrolysis with glucurononidase and sulfatase in the presence or absence of saccharic acid lactone (10). Hydrolysis was quantified by LH-20 chromatography and the liberated iodothyronines were identified by HPLC.

Analysis of cell content. Cell-associated radioactivity was extracted after removing the medium by immediate addition of 1 ml 0.1 N NaOH. After centrifugation, 0.5 ml of supernatant was chromatographed on LH-20 as described above for medium.

Miscellaneous. The influence of SAM (10-1000 μM) and DCNP (1-100 μM) on the type 1 deiodinase was tested by measurement of their effect on the release of 125I from 10 nM [125I]rT3 in incubations with rat liver microsomes (14) in 0.1 M phosphate buffer (pH 7.2), 2 mM EDTA, and 5 mM dithiothreitol (DTT). The free fractions of rT3 and 3,3'-T2 in incubation medium were determined by equilibrium dialysis.

Cellular ATP content was measured according to the method described by Jaworek et al. (15).

Data analysis. In each experiment, 1- production was corrected for the amount of 1- recovered from control incubations, while the 3,3'-T2 production from rT3 was corrected for the slight contamination of the 3,3'-T2 fraction with rT3. Production of unlabeled 3,3'-T2 was corrected for rT3 crossreactivity (0.03%) in the 3,3'-T2 RIA. Statistical analysis was done by Student’s t test for unpaired data.

Results

Sephadex LH-20 chromatography. The Sephadex LH-20 chromatographic pattern of rT3, 3,3'-T2, 3,3'-T2, and rT3 is depicted in Fig. 1. More than 97% of 125I activity was found in fractions 1-4. Recovery of 3,3'-T2, eluting in the H2O fractions, was > 97.5%. All other 3,3'-T2 and rT3 conjugates eluted also in fractions 5-9. Approximately 97% of 3,3'-T2 was found in fractions 10-15, while fractions 16-18 contained at least 95% of rT3. Thus, there was little overlap of fractions containing the different metabolites, although some 4% of rT3 eluted in the 3,3'-T2 fractions. This was not due to contamination of rT3 tracer with 3,3'-T2 as checked by HPLC.

Time dependence of 3,3'-T2 and rT3 metabolism. Fig. 2 shows the LH-20 analysis of the main products in the medium generated from 3,3'-T2 and rT3 as a function of time of incubation in the absence or presence of SAM. In 3,3'-T2 incubations 1-
was the main metabolite, the amount of which was inversely correlated with that of remaining 3,3'-T₂. The semi-logarithmic plot of the latter against time demonstrated that the disappearance of 3,3'-T₂ followed first order kinetics with a rate constant of 0.031 min⁻¹. Addition of SAM strongly inhibited I⁻ formation. In the presence of this inhibitor 3,3'-T₂ clearance remained a first order process with a rate constant of 0.007 min⁻¹.

In rT₃ incubations I⁻ was also the main metabolite found, while 3,3'-T₂ formation was undetectable under control conditions. Addition of SAM resulted in the inhibition of I⁻ release with a reciprocal accumulation of 3,3'-T₂. Clearance of rT₃ was also a first order process, characterized by a rate constant of 0.014 min⁻¹, which was not influenced by SAM. The identity of the 3,3'-T₂ produced was further tested by HPLC analysis (Fig. 3). Little or no 3,3'-T₂ formation could be detected in control incubations by HPLC, while in the presence of SAM 3,3'-T₂ accumulation was substantial. Relative to the quantity of rT₃ remaining after 30, 60, and 120 min, 3,3'-T₂ accumulation amounted to 11, 33, and 80%, respectively.

Dose-dependent effects of SAM (Fig. 4). As previously shown (10), addition of 10 μM PTU greatly inhibited ¹²⁵I⁻ production in incubations with 3,3'-T₂, but did not affect the clearance of this compound. Most radioactivity recovered from the medium now eluted in the water fractions of the LH-20 chromatography, which was shown to consist mainly of 3,3'-T₂S (see below). Addition of PTU also led to an increase in cell-associated radioactivity, from < 5% in the absence to 22% in the presence of the inhibitor. This radioactivity was predominantly in the form of conjugates and presumably represented the accumulation of 3,3'-T₂S within the cells. Addition of 10–1,000 μM SAM resulted in a progressive decrease in the clearance of 3,3'-T₂ independent of PTU. This was indicated by the parallel decrease of I⁻ formation without PTU and of 3,3'-T₂S accumulation with PTU. With the SAM-induced inhibition of 3,3'-T₂ clearance there was an increase in cell-associated 3,3'-T₂, irrespective of PTU.

In contrast to 3,3'-T₂, rT₃ metabolism was almost completely blocked by PTU but was not affected by even 1,000 μM SAM. In the absence of SAM, little 3,3'-T₂ production was observed in the medium once again, and no 3,3'-T₂ was detectable in the cells. With increasing SAM concentrations, 3,3'-T₂ accumulation in the medium was accompanied by a rise in cellular radioactivity from 5 to 12%, part of which was in the form of 3,3'-T₂. Based on the dose-effect relationship for SAM on the metabolism of both 3,3'-T₂ and rT₃, 100 μM was chosen as the near-maximal sulfotransferase inhibitory concentration.

3,3'-T₂ and rT₃ metabolism; correlation between LH-20 and RIA. Table 1 shows the effects of 100 μM SAM or DCNP or 10 μM PTU on 3,3'-T₂ and rT₃ metabolism as measured by LH-20 chromatography or by RIA. To compare the results obtained by these two methods it has to be taken into account that (a) LH-20 data are not corrected for cell-associated radioactivity.
(Fig. 4), and (b) the specific radioactivity of the 3,3'-T₂ produced is half that of added rT₃. With this in mind there exists a good correlation between the two methods.

After incubation for 60 min with [¹²³I]3,3'-T₂, 80% of radioactivity in the medium was recovered as [¹²³I]. SAM, DCNP, and PTU reduced I⁻ formation by 64, 57, and 95%, and 3,3'-T₂ clearance by 59, 54, and 10%, respectively. If remaining 3,3'-T₂ in medium was measured by RIA, inhibition of the clearance of 3,3'-T₂ by these treatments was 52, 62, and 1%, respectively. The 3,3'-T₂S accumulating in the presence of PTU as observed in the incubations with [¹²³I]3,3'-T₂ was largely recovered as immunoreactive 3,3'-T₂ after hydrolysis of medium from parallel incubations with unlabelled 3,3'-T₂. No 3,3'-T₂ was liberated by hydrolysis in all other experimental conditions.

### Table 1. Comparison of RIA and Sephadex LH-20 Analyses of the Effects of SAM, DCNP, and PTU on the Metabolism of 3,3'-T₂ and rT₃ by Rat Hepatocytes

<table>
<thead>
<tr>
<th>Products in medium</th>
<th>Condition</th>
<th>LH-20</th>
<th>RIA</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>n</td>
<td>3,3'-T₂</td>
</tr>
<tr>
<td>3,3'-T₂ 30 min</td>
<td>Control</td>
<td>9</td>
<td>48.2±4.0</td>
</tr>
<tr>
<td></td>
<td>SAM</td>
<td>8</td>
<td>11.5±1.9*</td>
</tr>
<tr>
<td></td>
<td>DCNP</td>
<td>6</td>
<td>17.4±2.1*</td>
</tr>
<tr>
<td></td>
<td>PTU</td>
<td>4</td>
<td>4.7±1.3*</td>
</tr>
<tr>
<td>3,3'-T₂ 60 min</td>
<td>Control</td>
<td>10</td>
<td>79.0±2.4</td>
</tr>
<tr>
<td></td>
<td>SAM</td>
<td>9</td>
<td>28.3±3.3*</td>
</tr>
<tr>
<td></td>
<td>DCNP</td>
<td>6</td>
<td>33.7±4.9*</td>
</tr>
<tr>
<td></td>
<td>PTU</td>
<td>5</td>
<td>4.4±1.3*</td>
</tr>
<tr>
<td>rT₃ 60 min</td>
<td>Control</td>
<td>11</td>
<td>47.7±3.9</td>
</tr>
<tr>
<td></td>
<td>SAM</td>
<td>10</td>
<td>34.5±3.2*</td>
</tr>
<tr>
<td></td>
<td>DCNP</td>
<td>6</td>
<td>44.5±4.3</td>
</tr>
<tr>
<td></td>
<td>PTU</td>
<td>5</td>
<td>1.8±0.6*</td>
</tr>
</tbody>
</table>

10 nM labeled or unlabeled 3,3'-T₂ or rT₃ were incubated for 30 or 60 min at 37°C with 10⁶ rat hepatocytes in 2 ml S-supplemented Dulbecco's medium with or without 100 µM SAM or DCNP or 10 µM PTU. Radioactive products were spotted on Sephadex LH-20 and unlabeled 3,3'-T₂, and rT₃ were measured by RIA before (−) and after (+) hydrolysis as described in Methods. Since acid treatment did not change rT₃ levels, mean values of these two measurements are RIA values expressed as percentage of iodothyronines measured after incubations without cells. The results are given as the mean±SE of the number of observations indicated. Statistical analysis of differences between experimental and control conditions was done by Student's unpaired t test: * P < 0.001. † P < 0.005. ‡ P < 0.01. § P < 0.025. ¶ P < 0.05.
After incubation for 60 min with $^{125}$I$rT_3$, 48% of radioactivity in the medium was accounted for by $^{125}$I$rT_3$, while the sum of the conjugate and 3,3'-T_2 fractions was only 2%. Again, rT_3 disappearance was not changed by SAM, but rT_3 formation was decreased by 13.2% (~2.6 nM rT_3) while 3,3'-T_2 accumulation was increased by 13.9% (~2.8 nM 3,3'-T_2). These results correspond with an increase of immunoassayable 3,3'-T_2 by 23.0% (~2.3 nM 3,3'-T_2). On the basis of the results obtained by LH-20 chromatography and by RIA, it was calculated that with SAM 51 and 54%, respectively, of the amount of rT_3 cleared was recovered as medium 3,3'-T_2. Similar results were obtained with DCNP, except that the acceleration of rT_3 metabolism seen with this inhibitor was different. PTU inhibited rT_3 disappearance from the medium by 77 and 67% as estimated by LH-20 and RIA, respectively.

S dependence of 3,3'-T_2 and rT_3 metabolism. As discussed above, 3,3'-T_2 was metabolized predominantly by successive sulfation and ORD. The small amount of conjugates found after incubation of 3,3'-T_2 in S-containing medium (Fig. 5) was shown by HPLC to consist of similar proportions of 3,3'-T_2 glucuronide (3,3'-T_2G) and 3,3'-T_2S (Fig. 6). In the presence of PTU, rT_3 formation was again strongly suppressed (Fig. 5), while radioactivity in the conjugate fraction almost entirely eluted in the position of 3,3'-T_2S with no change in the absolute amount of 3,3'-T_2G recovered (Fig. 6). If cells were preincubated with S-free medium and incubated with 3,3'-T_2 in this medium plus PTU, conjugation and clearance were diminished by 55%. A further reduction in 3,3'-T_2 conjugation to 30% was observed if SAM was added to the preincubation, while the experiment was otherwise conducted under the same S-free conditions (Fig. 5). HPLC analysis of the conjugate fraction demonstrated that 3,3'-T_2S formation was virtually blocked, and that now most radioactivity eluted in the position of 3,3'-T_2G (Fig. 6). In the absence of PTU, rT_3 release was decreased from 74 to 38% by omission of S from the medium, and further to 17% after addition of SAM to the preincubation. However, conjugates represented an increasing proportion of the radioactivity in the medium, i.e., 7, 10, and 15%, respectively (Fig. 5). HPLC revealed that this increase was totally accounted for by a rise in 3,3'-T_2G (Fig. 6).

Clearance of rT_3 was not S dependent. Iodide production, however, decreased from 48% in the presence of S to 38% in the absence of S, and further to 33% after preincubation with SAM. The loss of I was completely compensated for by the accumulation of 3,3'-T_2, which amounted to 1, 9, and 14% under these conditions, respectively (Fig. 5). In S-supplemented incubations the conjugate fraction contained small amounts of rT_3G, 3,3'-T_2G, and 3,3'-T_2S. S depletion induced an increase in conjugates mostly in the form of 3,3'-T_2G, although even after preincubation with SAM the conjugate fraction comprised only 5% of medium radioactivity. Inhibition of deiodinase by PTU augmented the appearance of conjugates in S-replete incubations (Table I, Fig. 6). In S-deplete cultures, addition of PTU led to a decrease in T_2G and an increase in rT_3G without a change in the total amount of conjugate formed (Fig. 6).

Effect of substrate concentration on rT_3 metabolism (Fig. 7). In S-replete incubations, progressive increases in unlabeled rT_3 concentration from 1 to 1,000 nM led to minimal increments in the recovery of immunoassayable 3,3'-T_2 from the medium. Medium 3,3'-T_2 ranged from undetectable after incubation with 1 nM rT_3 until maximally 1.9 pmol per dish (0.1%) at 1,000 nM rT_3. After addition of SAM, accumulation of 3,3'-T_2 rose from 0.8 pmol/dish (38%) at 1 nm rT_3 to 29.7 pmol/dish (1.5%) at 1,000 nM rT_3. The results indicated that 3,3'-T_2 accumulation under these conditions was a saturable process; it was half-maximal between 10 and 100 nM rT_3. Cell-associated 3,3'-T_2 was similarly dependent on substrate concentration.

Effects of SAM and DCNP on free substrate levels, cellular integrity, and deiodinase activity (Table II). In the absence of inhibitors, the free fraction of 3,3'-T_2 and rT_3 in Dulbecco's medium containing 0.1% BSA was 8.5 and 4.9%, respectively. Up to 1,000 µM, SAM did not influence the free 3,3'-T_2 and rT_3 fractions. DCNP had no effect on free 3,3'-T_2, while there was a dose-dependent increase in free rT_3 up to 7.8% at 100 µM DCNP.

Up to 100 µM, SAM did not inhibit rT_3 deiodination by rat liver microsomes, but a 25% reduction of T_3 release was observed at 1,000 µM SAM. In contrast, 100 µM DCNP resulted in a 65% lowering of deiodinase activity, while only slight inhibition was noted at 10 µM.

**Figure 5.** S dependence of 3,3'-T_2 and rT_3 metabolism. Monolayers of 10⁶ hepatocytes were preincubated with 30 min in S-supplemented medium or in S-free medium plus or minus 100 µM SAM. Subsequently, incubations were performed for 60 min with 10 nM [¹²⁵]I]3,3'-T_2 or [¹²⁵]I]rT_3 in S-replete or -deplete medium. PTU was added at 10 µM. Medium was analyzed by Sephadex LH-20 chromatography and data are expressed as mean±SE of the number of observations indicated. Completion of medium radioactivity to 100% represents the proportion of unaltered substrate. No correction is made for difference in specific radioactivity between substrate and products in rT_3 incubations. (c) 1; (m) conjugates; (a) 3,3'-T_2. * Significant difference with preincubation +SO₄²⁻, P < 0.001. ** Significant difference with preincubation −SO₄²⁻, P < 0.05.
Cell viability as assessed by ATP measurements was not affected by SAM. However, a 35% reduction of cellular ATP was found after incubation with 100 μM DCNP.

**Discussion**

The occurrence of rT3 and 3,3'-T2 in rat thyroid as well as in rat plasma has already been reported by Roche et al. in 1955 (16–18). These workers also provided evidence for the conversion of both T3 and rT3 to 3,3'-T2 in vivo by demonstration of the presence of radioactive 3,3'-T2 in the kidney after administration of labeled T3 or rT3 to thyroidectomized rats (19). In humans, most plasma 3,3'-T2 originates from IRD of T3 (20, 21). Plasma 3,3'-T2 (22) and rT3 (25) are cleared predominantly by the liver, which indicates that they are derived from extrahepatic IRD of T3 and T4, respectively. It is likely that these processes are catalyzed by the type III deiodinase, which specifically acts on the inner ring of iodothyronines and which has been localized in tissues such as brain and skin but not in liver (2, 5). In our studies we focused on the role of the type I deiodinase in the hepatic clearance of rT3.

Numerous studies have appeared on the characteristics of the deiodination of different iodothyronines by homogenates and subcellular fractions of rat liver (for a review, see reference 2). These studies have established the existence of a so-called type I deiodinase in the endoplasmic reticulum, which is an integral membrane protein and requires thiols for deiodination activity. The enzyme is a nonspecific deiodinase that is capable of removing iodines from either ring of iodothyronine substrates (2). However, the enzyme is most effective in the ORD of rT3 (2, 23). To test the possible regulatory function of this enzyme in peripheral thyroid hormone metabolism, we have therefore initiated studies of the breakdown of rT3 by isolated rat hepato-

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**Figure 6.** HPLC analysis of conjugates generated in incubations of (A) 3,3'-T2 and (B) rT3 with hepatocytes. Incubations were done with S-replete or -deplete cells after preincubation in medium with S or in medium without S plus 100 μM SAM as described in the legend to Fig. 5. Conjugates were isolated on Sephadex LH-20, concentrated on a second LH-20 column, and collected in ammonium acetate (pH 4) at a flow of 0.6 ml/min. Fractions of 0.3 ml were collected and counted for radioactivity. Elution position of S conjugates was determined using synthetic standards and conjugates were further identified by enzymatic liberation of 3,3'-T2 or rT3 after treatment with sulfatase or glucuronidase. Radioactivity eluting in the position of rT3S was not hydrolyzed by sulfatase, in agreement with previous observations using synthetic rT3S (13).

**Figure 7.** Conversion of rT3 to 3,3'-T2 by rat hepatocytes as a function of substrate concentration. Monolayers of 10^6 hepatocytes were incubated for 60 min with 1–1,000 nM unlabeled rT3 in Dulbecco's medium containing ∼S with or without 100 μM SAM. 3,3'-T2 content was measured by RIA in duplicate 50-μl aliquots of medium or 0.1 N NaOH extracts of cells. Data are taken from a representative experiment repeated on three different occasions with closely agreeing results.
Table II. Effects of SAM and DCNP on Medium1
Free rT3 and 3,3'-T2 Levels, on Microsomal Deiodinase Activity, and on Cellular ATP Content

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Free fraction (ng/ml)</th>
<th>Deiodinase activity</th>
<th>ATP (nM)</th>
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<tr>
<td></td>
<td>rT3</td>
<td>3,3'-T2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Control (%)</td>
<td>Control (%)</td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td>4.9</td>
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<td>100</td>
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<td>SAM</td>
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<tr>
<td></td>
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<td>4.6</td>
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<tr>
<td>DCNP</td>
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<tr>
<td></td>
<td>100 μM</td>
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</table>

* Free iodothyronine levels in medium were determined in triplicate by equilibrium dialysis.

1 Deiodinase activity was measured in triplicate in mixtures of 10 nM [125I]rT3, 2 or 5 μg microsomal protein/mL, and 5 mM DTT in 0.1 M phosphate (pH 7.2) and 2 mM EDTA (7). After incubation for 15 min at 37°C, reactions were halted by addition of serum, and 125I was released was measured by TCA precipitation. Data are expressed as the means of the six observations relative to iodide production in the absence of inhibitors.

Cells were incubated for 1 h at 37°C with Dulbecco’s medium plus serum and the indicated concentrations of SAM and DCNP. ATP was measured in 0.2-M perchloric acid extracts (0.5 ml/10⁶ cells) after centrifugation and neutralization with 2 M KOH according to the method of Jaworek et al. (15). Data are taken from a representative experiment repeated under different conditions with similar results.

tocytes. In spite of the high rate of rT3 to 3,3'-T2 conversion with isolated microsomes, and the stability of the product in such incubations (23), surprisingly little production of 3,3'-T2 from rT3 was observed in incubations with intact liver cells (8). That nevertheless rapid ORD took place in this system was demonstrated by the large amounts of radioiodide released from outer ring labeled rT3 (8). This implies that any 3,3'-T2 formed would have undergone further deiodination, as substantiated by analysis of the metabolism of added 3,3'-T2. Similar rapid degradation of both rT3 and 3,3'-T2 with extensive 11° formation was observed by Flock et al. (24) in isolated rat liver perfusions. The unexpected rapid deiodination of 3,3'-T2 in rat liver cells in contrast to microsomes was subsequently shown to be due to the efficient sulfation of 3,3'-T2 in hepatocytes yielding 3,3'-T2S, a highly effective substrate for the type I deiodinase (10). A similar faciatory effect of S conjugation has been described for the IRD of T3 and T4 (25, 26).

In the present study, we tested the hypothesis that it is possible to demonstrate production of 3,3'-T2 from rT3 in this system if further metabolism, especially sulfation, of 3,3'-T2 is prevented. In experiments with isolated rat hepatocytes three methods are available in principle to manipulate the sulfotransferase activity of the cells. Firstly, it has been demonstrated that there is a rapid equilibrium between medium and cellular inorganic S levels (27-29). Rat liver cells also have the capacity to generate S by oxidation of the sulfur-containing amino acids methionine and cysteine (28). Therefore, considerable S depletion is obtained by keeping cells in medium without S and possible precursors. Secondly, sulfation of phenolic substances may be prevented by competitive substrates (30). SAM has been shown to undergo extensive sulfation and at higher concentrations also glucuronidation in rat liver cells. For instance, Koike et al. (29) have reported that in isolated rat hepatocytes, sulfation of SAM is characterized by an apparent Km value of 6 μM with a Vmax value of 2 nmol/min per 10⁶ cells at 1.2 mM inorganic S in the medium. This level of S is similar to that used in our system (1 mM), providing near-maximal rates of S conjugation (9, 10). Glucuronidation of SAM became more pronounced if the competing pathway of sulfation was saturated at high SAM levels or prevented by S depletion (29). G conjugation occurred with a Km value of 0.2 mM and Vmax of 1.3 nmol/min per 10⁶ cells as determined under S-depleted conditions. Thirdly, specific inhibition of phenol sulfotransferase activity may be obtained using DCNP, a compound which in itself is not sulfated (31, 32). All three abovementioned ways of intervention with the further sulfation of 3,3'-T2 have been applied to our study of the deiodination of rT3 by rat hepatocytes.

Previous studies from our laboratory have demonstrated the feasibility of using small Sephadex LH-20 columns for separation of 3,3'-T2 and its main metabolites, 3,3'-T2S and I (10). We have adapted this method in the present investigation for the analysis of rT3 metabolites. With this modification it was possible to separate fractions containing I, conjugates, 3,3'-T2, and rT3, respectively. Since the fraction eluted with water may contain various conjugates, and also because there is a slight contamination of the 3,3'-T2 fraction with rT3, these analyses were extended by HPLC and RIA determinations. Despite the limitations of the Sephadex LH-20 method, a good correlation between 3,3'-T2 production rates as measured by the three different techniques was observed.

Time-course studies of the metabolism of 3,3'-T2 and rT3 demonstrated that under the conditions used, i.e., 10-nM substrate levels and S-supplemented cells, clearance of both substances follows first-order rate kinetics (Fig. 2). Such kinetics were also observed in the presence of 100 μM SAM, which did not affect the rate constant for rT3 but lowered that for 3,3'-T2 by 77%. These results show that conjugation is an important, rate-limiting step in the metabolism of 3,3'-T2, but not of rT3. SAM does not only inhibit degradation of added 3,3'-T2, but apparently also that of 3,3'-T2 produced from rT3. No evidence was obtained by HPLC for the generation of iodothyronines other than 3,3'-T2 such as 3,3'-T2 or 3'-iodothyronine (Fig. 3).

The 100-μM SAM concentration used mostly in our experiments was shown to have near-maximal effects on the clearance of 3,3'-T2. The generation of conjugates observed in the presence of PTU is inhibited to only a slightly greater extent at 1,000 μM SAM, while 10 μM of this drug already provided 60% inhibition. This value is in excellent agreement with the Michaelis constant (Km) value for SAM sulfation of 6 μM, which was reported by Koike et al. (29). Note that after incubation of 3,3'-T2 in the presence of PTU, substantial radioactivity in the form of 3,3'-T2S remains associated with the cells, which suggests that exit of cellular 3,3'-T2S is a relatively slow process. If the amount of cell-bound 3,3'-T2S is taken into account is is obvious that PTU does not affect clearance of 3,3'-T2, whereas metabolism of rT3 is strongly impaired (see below). In the absence of PTU, SAM induces a dose-dependent decrease in I formation from 3,3'-T2 that parallels the decrease of 3,3'-T2S in incubations with PTU. However, I production is not completely suppressed with even 1,000 μM SAM, suggesting that direct ORD of 3,3'-T2 may occur to some extent if sulfation is inhibited. Direct ORD of 3,3'-T2 has also been observed with isolated rat liver microsomes, albeit with only 2% of the efficiency of 3,3'-T2S ORD (9). Therefore,
with unimpeded sulfation, direct deiodination is a negligible pathway.

Also concerning the accumulation of 3,3'-T₂ produced from rT₃, 1,000 μM SAM shows only slightly greater effects than 100 μM of this drug, while roughly a half-maximal effect is observed with 10 μM SAM. At the highest SAM concentration, 3,3'-T₂ represents approximately one-third and 1' approximately two-thirds of the rT₁ metabolites, which corresponds to a 70% protection against further degradation of the 3,3'-T₂ produced. It should be stressed that clearance of rT₁ is not affected by even 1,000 μM SAM. If added to deiodinase assay mixtures of rT₁ with microsomes, inhibition was observed only with 1,000 μM SAM (Table II). The differential effects of 1,000 μM SAM on rT₁ deiodination by intact cells, and that by isolated microsomes, indicate that intracellular SAM levels are substantially lower than those added to the medium. The extremely rapid metabolism of SAM by both sulfation and glucuronidation (29) may be an important factor contributing to this difference. The lack of effect of SAM on medium free iodothyronine levels and cellular ATP (Table II) further underscores the usefulness of this compound in studies of the conjugation of thyroid hormone.

Table I illustrates that similar effects on the metabolism of 3,3'-T₂ and rT₁ are obtained with 100 μM DCNP as with 100 μM SAM. However, DCNP also exhibits some unwanted side effects. DCNP has been shown by Mulder and co-workers to be a specific inhibitor of sulfation in vitro as well as in vitro without being sulfated itself (31, 32). This compound does not affect glucuronidation and was reported to have no major toxic effects, which is in contrast with our findings that show a substantial decrease in cellular ATP by 100 μM DCNP. This effect on ATP may be related to the toxic effects on oxidative phosphorylation generally observed with phenols (33, 34) which may be promoted by the relative deficiency of nutrients and oxygen in our system. A further disadvantage of DCNP is that it may inhibit deiodinase activity directly, as demonstrated by the influence of DCNP in incubations with microsomes (Table II). Similar to SAM, however, intracellular DCNP concentrations may be substantially lower than the total concentrations added to the medium. One possible reason for this is that DCNP binds extensively to BSA (31). The decreased availability of DCNP to the cells, therefore, prevents direct inhibition of the deiodinase, and metabolism of rT₁ is even stimulated by 100 μM DCNP. This latter phenomenon is probably due to displacement of rT₁ from BSA (Table II), which causes a shift of rT₁ into the intracellular compartment. Nevertheless, results obtained with DCNP are in confirmation of the experiments with SAM which underscore the importance of sulfation for the metabolism of 3,3'-T₂ but not of rT₁.

The Sephadex LH-20 method used in this study does not permit the distinction of the different conjugates of 3,3'-T₂ and rT₁. Previous findings have indicated that sulfate conjugates do not crossreact in RIA's of non-conjugated iodothyronines (13). The RIA data presented in Table I are, therefore, consistent with the results obtained by LH-20 insofar as they indicate that clearance of 3,3'-T₂ is not inhibited by PTU. If the product that is formed from 3,3'-T₂ in the presence of PTU is subjected to acid hydrolysis, similar amounts of 3,3'-T₂ are recovered as those measured after incubation with SAM or DCNP. Since iodothyronine sulfates are prone to acid hydrolysis (13), while glucuronides are much more resistant (Oosterlaken, T., S. J. Eelkman Roorda, and T. J. Visser, unpublished observations), this indicates accumulation of 3,3'-T₂S in the presence of PTU. However, most direct evidence for the identity of the conjugates is provided by HPLC in combination with enzymatic analysis using sulfatase or glucuronidase. These experiments were made on samples from incubations with cells in different degrees of S depletion.

S-deplete hepatocytes were prepared by preincubation with medium lacking inorganic S and by using the same medium for incubation with substrate. The results obtained with 3,3'-T₂ in the presence of PTU showed that under these conditions S depletion was not complete, and led to some residual 3,3'-T₂S formation. Further exhaustion of cellular S stores was obtained by preincubation with SAM. A similar approach has been followed by other investigators to reveal the significance of the conjugation reactions in the elimination of drugs (35, 36). Thus, substantial lowering of hepatic UDP-glucuronic acid and adenosine 3'-phosphate 5'-phosphosulfate levels has been achieved by in vivo treatment with SAM due to the consumption of these co-factors in the glucuronidation and sulfation of the drug (35, 36). Application of a SAM load induces a prolonged lowering of body S stores with pronounced decrements in plasma S levels (35), but depletion of tissue UDP-glucuronic acid is only transient, and co-factor levels are rapidly restored (36). We also obtained evidence for the reversible nature of these effects. After preincubation with SAM, 3,3'-T₂ sulfation rates after readdition of SO₄²⁻ and T₂ glucuronidation rates (Eelkman Rooda, S. J., and T. J. Visser, unpublished observations) were completely normal.

In S-deplete cells, sulfation is by far the predominant pathway of 3,3'-T₂ metabolism, although little 3,3'-T₂S is recovered in the absence of PTU due to further deiodination. However, in S-deplete cells the contribution of the glucuronidation pathway increases, and results in augmented accumulation of 3,3'-T₂G from added 3,3'-T₂, as well as from added rT₁. The latter findings indicate that glucuronidation does not facilitate deiodination of 3,3'-T₂, which is in contrast to the stimulatory effects of sulfation. Although ORD is the foremost step in the hepatic metabolism of rT₁, some glucuronidation of this metabolite is observed, especially in the presence of PTU. The low rate of rT₁ sulfation in comparison with the highly efficient sulfation of 3,3'-T₂ is in agreement with data of the structure-activity relationship for the sulfation of iodothyronines by isolated phenol sulfotransferases (37).

The effects of substrate concentration on the conversion of rT₁ to 3,3'-T₂ by rat liver cells indicate that half-maximal 3,3'-T₂ production is obtained between 10 and 100 nM rT₁. This is in the same concentration range as the Kₘ value determined for rT₁ ORD in reactions with microsomes and DTT (23). However, the rT₁ levels in the culture medium are largely protein bound and certainly do not reflect intracellularly available substrate concentrations. Together with the uncertainty about the saturation kinetics of the type I deiodinase in its natural environment it is, therefore, impossible to interpret these findings in terms of a possible uphill gradient of rT₁ across the cell membrane (38).

In conclusion, our results demonstrate that in isolated rat hepatocytes, rT₁ is metabolized almost exclusively by ORD. However, the immediate product of this reaction, 3,3'-T₂, is not observed unless its further metabolism by successive sulfation and deiodination is inhibited. Evidence for formation of 3',5'-T₂ in these studies was not obtained. In different clinical situations, plasma T₃ and rT₁ levels change in opposite directions due to parallel alterations in the rate of T₃ production and rT₁ degradation. This is understandable, as both processes concern ORD reactions mediated by common enzymes among which the type I deiodinase of the liver plays an important function. It is expected, therefore, that a detailed knowledge of the ORD
of rT3 by intact liver cells will deepen our insight into the regulation of the bioavailability of thyroid hormone.

References


