Identification and Clearance Involved in the Formation of Glucuronides of RT-3003, a New Peripheral Blood Flow Enhancer, and Its Metabolite in Rats

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Glucuronides of RT-3003 and its metabolite (9-OH-RT-3003), which was hydroxylated at the 9 position on the benzene ring, were separated by HPLC and identified by liquid chromatography (LC)/MS/MS and NMR. The conjugation sites of these glucuronides were determined by nuclear Overhauser effects (NOE) irradiation; RT-3003 was conjugated at an alcoholic hydroxyl group of the hydroxymethyl moiety, and 9-OH-RT-3003 at a phenolic hydroxyl group on a benzene ring and at an alcoholic hydroxyl group of a hydroxymethyl moiety. On a reversed-phase HPLC of 9-OH-RT-3003, alcoholic glucuronide was eluted later than phenolic glucuronide, indicating the high hydrophilicity of alcoholic glucuronide. Clearance for the glucuronides (CLc) of RT-3003 was lower than the summation of CLc for two types of glucuronidation of 9-OH-RT-3003. CLc of 9-OH-RT-3003 was high in phenolic glucuronide. The activity of UDP-glucurononitransferase (UDPGT) for RT-3003 was 9.63 times that for 9-OH-RT-3003, and the activity ratio of the two types of glucuronidation of 9-OH-RT-3003 was similar to the ratio of the corresponding CLc. The difference between CLc and UDPGT activity is discussed in association with clearance for the hydroxylation and interaction of substrates with UDPGT.

Key words: glucuronidation site; nuclear Overhauser effect; liquid chromatography (LC)/MS/MS; clearance

RT-3003, (-)-1β-Ethyl-1α-hydroxymethyl-1,2,3,4,6,7,12b-octahydroindol[2,3-ajquinolizine which was developed by Gedeon Richter, Ltd. (Hungary) has selective peripheral blood flow enhancing properties in experimental animals. The chemical structure of RT-3003 is shown in Fig. 1. This compound has a symmetric center at positions 1 and 12b; RT-3003 is trans-form, referring to the relative position of the C1-C12 group and the C12b-H bond. Szombathelyi et al. found that the pharmacological effect of trans-form was higher than that of cis-form.

The structure of RT-3003 suggests various types of glucuronidations: glucuronidation at the hydroxymethyl group, secondary amine, tertiary amine and hydroxyl group on the benzene ring formed by monooxidation. The formation rates of these glucuronides are assumed to vary with the types of glucuronides produced by the catalysis of various isozymes of UDP-glucurononitransferase (UDPGT). Therefore, identification of the structures of glucuronides and pharmacokinetic analyses of glucuronidation are useful in clarifying the elimination of this drug from the body.

The purpose of this study is to identify the various types of glucuronide produced from RT-3003 following the oral administration of 14C-RT-3003 into rats, and to analyze the clearance for glucuronidation associated with the elimination of RT-3003 and its metabolites.

MATERIALS AND METHODS

Labeled Compound 14C-RT-3003 (specific activity; 14.7 mCi/mmol) was obtained from Gedeon Richter, Ltd. (Hungary). 14C-9-OH-RT-3003 (specific activity; 9.5 mCi/mmol) was obtained by the hydrolysis of biliary excretes: biliary excretes were incubated in excised cecum at 37°C for 1 h and extracted with ethylacetate, followed by purification using HPLC on 300×25 mm, μBondapak C18 125 Å (Waters) using solvent A, 10 mM ammonium acetate buffer (pH 8.0) and solvent B, acetonitrile with a linear gradient to 10—30% B in 30 min.

Animal Treatment Male Wistar rats weighing 240—260 g (8 weeks old) were purchased from Clea Japan, Inc (Tokyo). The bile ducts of all animals were cannulated with polyethylene tubes. To collect urinary samples, vinyl tubes (4.0 mm i.d.×4.8 mm o.d.) were adhered to abdominal skin in a fashion such that the urinary outlets were placed in the tubes.

Purification Procedure Bile was collected for 8 h following the oral administration of 14C-RT-3003 at a dose of 3 mg/kg. Bile samples were extracted from Sep-Pak C18 (Waters) with 30% and 40% methanol. HPLC of these fractions was performed on an LC-10A system (Shimadzu, Japan) equipped with an LC-10A pump and SPD-10A UV detector. Radioactivity collected with a fraction collector was assayed with a Model 903 liquid scintillation counter (Aloka). The 30% methanol extracts were first purified by HPLC on 300×25 mm, μBondapak C18 125 Å (Waters). The solvent system was as follows: A, 10 mM ammonium acetate buffer

* Position labeled with 14C.
(pH 5.0) and B, acetonitrile, with a linear gradient to 0—30% B in 15 min, followed by a 15 min hold at 30% B at a flow rate of 1 ml/min. Further purification was achieved by HPLC with μBondapak C18 10 μm (300×7.8 mm, Waters). The column was eluted with an isocratic phase of A and B (95:5). Purification of the 40% methanol extracts was first carried out by using the first HPLC condition for the 30% methanol extracts. Further purification was performed by HPLC with μPorasil 10 μm (300×7.8 mm, Waters). The solvent system was as follows: dichloromethane and methanol containing 0.1% triethylamine, with a linear gradient to 20—100% methanol containing 0.1% triethylamine in 15 min.

**Assay of UDPT Activity** The assay of the activity of hepatic microsomal UDPT was carried out as described previously. Briefly, microsomes (0.15 mg protein) were incubated with 4 μmol of UDP-glucuronic acid, 0.4 μmol of MgCl₂ and 0.5 nmol of 1⁴C-RT-3003 or 1⁴C-9-OH-RT-3003 in 0.1 ml of 20 mM Tris–HCl buffer (pH 7.8) for 20 min at 37°C.

**TLC Plates** (Merck Silica gel F₂₅₄) were developed in toluene–methanol–triethylamine (8:1:1). The areas of radioactivity were assayed.

**NMR Spectrometry** NMR spectra were recorded at 270 MHz using a JEOL EX270 spectrometer (JEOL). Samples were dissolved in 0.5 ml of dimethylsulfoxide-d₆.

**LC/MS/MS Analysis** A Micromass tandem quadrupole mass spectrometer (Quattro II, Micromass UK Ltd.) equipped with an electrospray ionization source system, a Hewlett Packard HPLC system (Series 1100, CA, U.S.A.), was employed. HPLC conditions are as follows: TSK-gel ODS-80T's column (150×2 mm) was eluted with solvent A, 10 mm ammonium formate buffer (pH 8.0) and B, acetonitrile, with a linear gradient to 0—34% B in 30 min.

**Calculation of Clearance** For the determination of Cl₀ of RT-3003, 1⁴C-RT-3003 was administered orally with 3 mg (10.6 μmol)/kg, and urine and bile were collected for 24 h. The blood samples were taken by cutting carotid arteries at various times until 24 h. To determine the Cl₀ of 9-OH-RT-3003, 1⁴C-9-OH-RT-3003 was injected intravenously with 3.7 μmol/kg. Biological samples were collected in the same way as 1⁴C-RT-3003. Clearance for glucuronidation (Cl₀) was determined as follows:

\[
\text{Cl₀} = \text{amount of glucuronide in the bile and urine}/\text{AUC}
\]

where \text{AUC} is the area under the curve of the plasma concentration of corresponding aglycon; \text{AUC} was obtained by adding all the trapezoidal areas from the measured values.

**Measurement of Biological Sample** The plasma obtained by centrifugation of the blood, urine and bile was extracted from Sep-Pak C18 (Waters) with 50% methanol. An aliquot of the extracts was injected on HPLC with a column of μBondapak C18 10 μm (300×7.8 mm, Waters) and eluted with 10 mM ammonium acetate buffer (pH 5.0) and acetonitrile (95:5). The eluate was collected with a fraction collector, followed by the radioactivity assay with a liquid scintillation counter.

**RESULTS AND DISCUSSION**

Following the oral administration of ¹⁴C-RT-3003 to rats, the excreted radioactivity in bile appeared to be 3.8 times that in urine. A thin layer chromatogram of these excretes showed one radioactive spot on origin, indicating the presence of polar metabolites; thus, these polar metabolites in the bile were purified chromatographically. The metabolites were separated into two fractions by solid phase extraction: fractions extracted with 30% methanol and with 40% methanol. On HPLC for the further purification of 30% methanol extracts, two radioactive peaks appeared at 20 min (M2) and 35 min (M1). The fractions including these peaks were subjected to analysis by LC/MS/MS and NMR.

M1 and M2 gave a simplified mass spectrum in which an ion of m/z 477 was observed as a base peak corresponding to the protonated molecular ion (M+H)⁺ of the glucuronide of 9-OH-RT-3003. The product ion spectrum from (M+H)⁺ of m/z 477 for both metabolites showed a fragment ion of m/z 301 which corresponded to the loss of a 176-unit moiety from 477, a process that is characteristic of glucuronide conjugation (Fig. 2). The ion at m/z 301, 16 mass units higher than RT-3003, further suggested monooxidation to RT-3003.

**Figure 3** shows ¹H–¹H correlation spectroscopy (COSY) of M1. The aromatic region of M1 shows the presence of three protons and an upfield shift of the signals of two protons of H-8 (6.66 ppm) and H-10 (6.50 ppm) compared with RT-3003 (H-8, 7.00 ppm; H-10, 6.96 ppm), suggesting hydroxylation of the aromatic ring. The characteristics of the amoneric proton (H-1') of the glucuronic acid suggested a doublet (J₁'-₂=8.0 Hz) at 4.45 ppm. The COSY spectrum showed cross-peaks corresponding to the coupling of the amoneric proton of H-1' to H-2' (3.30 ppm), as well as cou-
pling among other protons of the glucuronidic acid moiety (3.2–3.4 ppm). Signals of other regions were similar to those of RT-3003, indicating that no metabolic reaction occurred in other regions. The COSY spectrum of M2 resembled that of M1 except that the signals of H-8 (7.00 ppm) and H-10 (6.78 ppm) in the aromatic region exhibited a downfield shift.

From the spectrum of LC/MS/MS and NMR, M1 and M2 were identified as glucuronides of 9-OH-RT-3003. The results that both glucuronides showed differences in their retention time of HPLC and in the chemical shift of protons in the aromatic region suggest that these glucuronides were conjugated at different sites. Thus, the assignment of the site was achieved by nuclear Overhauser effects (NOE) difference spectroscopy (Fig. 4). When a proton of the hydroxymethyl moiety of M1 was irradiated, NOE enhancement was observed at an anomic proton of the glucuronidic acid moiety, indicating that M1 (9-OH-RT-3003-15G) is conjugated at an alcoholic hydroxyl group at the 15 position. Irradiation of the proton attached to C-8 of M2 resulted in NOE enhancement at an anomic proton of the glucuronidic acid moiety, indicating that the glucuronidation site of M2 (9-OH-RT-3003-9G) was the phenolic hydroxyl group at the 9 position on the benzene ring.

On the reversed-phase HPLC, 9-OH-RT-3003-15G was eluted later than 9-OH-RT-3003-9G, indicating that 9-OH-RT-3003-15G was more hydrophobic than 9-OH-RT-3003-9G. Leo et al.\(^5\) reported that a chain hydroxyl group contributes greatly to the hydrophilicity of a compound comparing with phenolic hydroxyl groups; thus, the late elution of 9-OH-RT-3003-15G would be due partly to the reaction of the alcoholic hydroxyl group with glucuronic acid. To explain the remarkable difference in retention time, however, it is necessary to take into account the possibility that the glucuronidic acid moiety of 9-OH-RT-3003-15G interacts with other regions of this molecule in a fashion such that the hydrophilicity of this compound decreases.

The MS spectrum of the metabolite (M3) from 40% methanol extracts also gave a simplified mass spectrum in which the ion of m/z 461 corresponding to (M+H)\(^+\) was observed as a base peak. The product ion from (M+H)\(^+\) of m/z 461 showed a signal at m/z 285 (M+H–176)\(^+\) which corresponded to protonated RT-3003 formed by the cleavage of the glucuronidic acid moiety. After the hydrolysis of this metabolite with β-glucuronidase/aryl sulfatase, the products were analyzed by TLC, and showed a radioactive spot corresponding to RT-3003. Irradiation of the anomic proton showed NOE enhancement at a proton at the 15 position. From these results, M3 was identified as a glucuronide of RT-3003 (RT-3003-15G), which was conjugated at a hydroxyl group of a hydroxymethyl moiety.

Table 1 shows the ratio of glucuronides in bile and urine after the oral administration of \(^{14}\)C-RT-3003. Three species of glucuronides were mainly excreted into the bile. To examine the effects of the hydroxylation and glucuronidation site on the distribution in both excretion pathways, the urine/bile excretion ratio of glucuronic was calculated; however, no significant difference in the excretion ratio among these glucuronides was observed. The total amount of glucuronic in the excretes was 30.2% of dose in RT-3003-15G, 20.5% in 9-OH-RT-3003-9G and 13.2% in 9-OH-RT-3003-15G, showing that glucuronidation played an important role in elimination.
Table 2. Clearance for Glucuronidation and UDPGT Activity of RT-3003 and Its Metabolites

<table>
<thead>
<tr>
<th>Formed glucuronide</th>
<th>Clearance (l/kg)</th>
<th>UDPGT activity (pmol/min mg protein)</th>
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<tbody>
<tr>
<td>RT-3003-15G</td>
<td>3.53±0.47&lt;sup&gt;a&lt;/sup&gt;</td>
<td>91.5±8.9&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>9-OH-3003-9G</td>
<td>3.36±0.55&lt;sup&gt;b&lt;/sup&gt;</td>
<td>5.72±0.61&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>9-OH-3003-15G</td>
<td>2.17±0.31</td>
<td>3.78±0.42</td>
</tr>
<tr>
<td></td>
<td>(5.53±0.86)&lt;sup&gt;c&lt;/sup&gt;</td>
<td>(9.50±0.70)&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

Data are expressed as mean±S.D. of 5 experiments. <sup>a</sup>p<0.05 compared with the value of the summation of the two types of glucuronide of 9-OH-RT-3003. <sup>b</sup>p<0.05 compared with the value of 9-OH-RT-3003-15G. <sup>c</sup>Summation of the two types of glucuronide of 9-OH-RT-3003.

from the body. Thus, $Cl_G$ for the formation of these glucuronides was determined. Considering that 9-OH-RT-3003 is produced in the body, the $Cl_G$ of 9-OH-RT-3003 was calculated using the data obtained following the intravenous injection of <sup>14</sup>C-9-OH-RT-3003. The dose was designed to be 35% of the oral dose of <sup>14</sup>C-RT-3003 from the result that 33.7% of <sup>14</sup>C-RT-3003 appeared to be hydroxylated at the 9 position until 24 h.

As shown in Table 2, $Cl_G$ of RT-3003 was 3.53 l/kg, 63.8% of the summation of $Cl_G$ of 9-OH-RT-3003. $Cl_G$ for 9-OH-RT-3003-9G was 1.55 times that for 9-OH-RT-3003-15G. The values of $Cl_G$ are assumed to depend on UDPGT activities. Thus, UDPGT activities toward RT-3003 and 9-OH-RT-3003 were measured using hepatic microsomes; substrate concentrations were designed to cover the plasma concentrations. The activity toward RT-3003 was 9.63 times the total activity toward 9-OH-RT-3003, showing a high ratio compared with the ratio of the corresponding $Cl_G$ (Table 2). On the other hand, the UDPGT activity ratio in both types of glucuronidation of 9-OH-RT-3003 was 1.51, similar to the ratio of the corresponding value of $Cl_G$. This similarity suggests that the in vivo activity of UDPGT is higher in RT-3003 than in 9-OH-RT-3003, and the activity toward RT-3003 is competitively inhibited by the formation of 9-OH-RT-3003 through hydroxylation, resulting in the low value of $Cl_G$. To clarify the effect of this hydroxylation, the clearance of the hydroxylation was calculated using the AUC of RT-3003 and the excreted amounts of the glucuronides of 9-OH-RT-3003 instead of its aglycon, since 9-OH-RT-3003 was excreted as glucuronide in the excretes. The clearance obtained was 3.17 l/kg, which was similar to $Cl_G$ of RT-3003; therefore, the formation of 9-OH-RT-3003 is assumed to not greatly decrease the glucuronidation. This study could not explain the difference between $Cl_G$ and UDPGT activity. To clarify this difference, it would be necessary to examine the interaction between substrates and UDPGT, in association with the physicochemical properties of microsomal membranes and substrates.

Three species of glucuronide in the bile are assumed to be hydrolyzed in the digestive tract, followed by reabsorption into the blood. Reabsorbed RT-3003 would again be glucuronidated with the values of $Cl_G$ in this study. $Cl_G$ of reabsorbed 9-OH-RT-3003 would be different from the values which were obtained from the intravenous injection, since these values did not evaluate the first pass effect.

REFERENCES