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Obesity-induced increase of CYP2E1 activity and its effect on disposition kinetics of chlorzoxazone in Zucker rats

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ABSTRACT

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This study was designed to investigate the induction of CYP2E1 in obese Zucker rats and its

effect on the disposition kinetics of chlorzoxazone (CZX). CZX 20 mg/kg was administered to

three groups of rats: normal Zucker rats fed a normal diet (ND), normal Zucker rats fed a high-fat diet (HF), and genetically obese Zucker rats fed a normal diet (OB). The values of the area under the plasma concentration–time curve from 0 to ∞ (AUC $_{\infty}$) of CZX were in the order of ND > HF > OB rats. The AUC $_{\infty}$ values of total 6-hydroxychlorzoxazone (60HCZX-T),

which is considered to be a CYP2E1 metabolic marker, were in the opposite order. The values

of the AUC $_{\infty}$ ratio (60HCZX–T/CZX) in ND, HF and OB rats were approximately 0.2, 0.3 and

0.4, respectively. The CZX concentration in fat was much higher than the concentrations in

plasma, liver and kidney in all groups. Induction of CYP2E1 protein was greater in both liver

and fat of OB rats than in those of HF rats. Microsomal activity of CYP2E1 in liver and fat was also in the order of OB > HF > NM rats. These results suggest that CYP2E1 may be induced in

liver and fat of obese patients, thereby potentially altering the disposition kinetics of not

only CZX, but also other lipophilic drugs metabolized by CYP2E1.

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Abbreviations:

6OHCZX, 6-hydroxychlorzoxazone 6OHCZX-G, glucuronidated 6-hydroxychlorzoxazone 6OHCZX-T, total 6-hydroxychlorzoxazone AST, aspartate aminotransferase CYP, cytochrome P450 CZX, chlorzoxazone CZX-G, glucuronidated chlorzoxazone PNP, *p*-nitrophenol UGT, UDP-glucuronosyltransferase

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1. Introduction

An increasingly sedentary lifestyle, together with a preference for refined foods containing high levels of meat and saturated fat, has resulted in a drastic increase in the incidence of metabolic abnormalities, obesity, and hypercholesterolemia, which are associated with premature mortality [1,2]. Therefore, animal models, such as genetically obese Zucker (fa/fa) rats, have been developed to study the mechanisms of physiological changes related to obesity. The Zucker (fa/fa) rat does not develop leptin receptors, resulting in impaired regulation of food intake and impaired energy homeostasis

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[3,4]. The overfed normal rat can also mimic physiological aspects of severe obesity in humans, such as hypercholesterolemia, hyperinsulinemia, etc. [5,6].

26 Changes in physiological status, such as obesity and 27 hypercholesterolemia, can modulate the metabolic activity of CYP2E1 [7-9]. Interestingly, CYP2E1 is a ubiquitous enzyme that 28 29 is distributed in various organs of animals, but obesity-related 30 increase of CYP2E1 activity is well documented only in liver. 31 There are several reports that the hydroxylation of chlorzox-32 azone (CZX) can be used as an indicator of CYP2E1 activity both 33 in vivo and in vitro [10,11]. CZX is a muscle relaxant that 34 primarily undergoes hydroxylation, catalyzed mainly by 35 CYP2E1, to 6-hydroxychlorzoxazone (6OHCZX), which is rapidly 36 glucuronidated and excreted in urine [12,13]. CZX is lipophilic 37 [14], and so may be a suitable probe to examine the activity of CYP2E1 in various tissues of obese animals. 38

This study was therefore designed to examine the induction of CYP2E1 in genetically obese Zucker rats fed a normal
diet (OB) and its effect on the disposition kinetics of CZX and
its metabolite 6OHCZX in liver, kidney and fat, compared with
those in normal Zucker rats fed a high-fat diet (HF) and normal
Zucker rats fed a normal diet (ND).

2. Materials and methods

2.1. Materials

47 CZX and 6OHCZX were purchased from Sigma–Aldrich Inc. (St.
48 Louis, MO). High-fat diet (LABO H Standard[®]) and normal diet
49 (LABO MR Stock[®]) were purchased from Nosan Corp.
50 (Yokohama, Japan). The high-fat diet contained 8.6% (w/w)
51 fat (total energy 346.7 kcal/100 g), while the normal diet
52 contained 4.1% (w/w) fat (total energy 259.2 kcal/100 g).

53 2.2. Animal treatment

54 Male 8-week-old Zucker (+/+) rats and genetically obese 55 Zucker (fa/fa) rats were purchased from Japan SLC Inc. 56 (Toyama, Japan). The rats were divided into three groups: normal Zucker (+/+) rats fed with normal diet (ND rats), normal 57 58 Zucker (+/+) rats fed with high-fat diet (HF rats) and genetically 59 obese Zucker (fa/fa) rats fed with normal diet (OB rats). The 60 animals were housed for 3 months in a climate- and lightcontrolled environment with free access to water and the 61 62 designated food. All animal procedures were in accordance 63 with the standards set forth in the guidelines for the care and use of laboratory animals at the Takara-machi Campus of 64 Kanazawa University. 65

66 2.3. Disposition kinetic of CZX

CZX (20 mg) was dissolved in 400 µl of 0.5N NaOH and diluted 67 68 with 600 μ l of normal saline solution (20 mg/ml). This solution 69 was administered to rats at a dose of 20 mg/kg by i.v. administration over 2 min via a lateral tail vein. Then, 70 71 approximately 0.2-0.5 ml of blood was collected from the tail 72 vein on the other side at 0.033, 0.083, 0.25, 0.5, 1, 2, 4, and 8 h after completion of the administration of CZX. The blood was 73 74 centrifuged at $3000 \times q$ for 10 min, and the plasma was collected. Some rats were killed by decapitation at 4 h after CZX administration, and the liver, kidney and abdominal fat were quickly excised, rinsed well with ice-cold saline, blotted dry, and weighed. Each tissue sample was homogenized with normal saline and stored at -80 °C until analysis.

2.4. Sample preparation

The concentrations of CZX and 6OHCZX in biological samples were analyzed as unchanged and glucuronidated fractions. The assay of glucuronidated CZX and 6OHCZX was performed according to Frye and Stiff [15] with slight modifications. Briefly, a 100 μ l sample of plasma or tissue homogenate was added to 300 μ l of 0.2 M phosphate buffer (pH 6.5) containing 500 units of β -glucuronidase. The mixture was incubated at 37 °C with shaking for 2 h, and then the reaction was stopped by adding 100 μ l of acetonitrile containing phenacetin (1 μ g) as an internal standard for HPLC analysis.

2.5. HPLC assay of CZX and 6OHCZX

To a 100 μ l of sample solution was added 5 ml of diethyl ether, then the mixture was shaken vigorously for 10 min, and centrifuged at 3000 \times *g* for 10 min. The ether layer was transferred to another tube for evaporation in a vacuum centrifugal concentrator. In the case of adipose tissue, reverse phase extraction from the case of homogenated samples was done by adding 0.5N NaOH to the samples and mixing. The aqueous was collected and titrated with an equal amount of 0.5N HCl. Next, diethyl ether (5 ml) was added to extract CZX and 60HCZX from aqueous phase, and the organic solution was further processed as described above.

The residue from evaporation was dissolved in 200 μ l of the mobile phase, and a 50 μ l aliquot was injected into an HPLC system (LC-9A, Shimadzu Co. Ltd., Kyoto, Japan) equipped with a CAPCELL PAK C18 column, 1.5 mm i.d. × 150 mm (Shiseido Co. Ltd., Tokyo, Japan). The mobile phase consisted of 25% (v/v) acetonitrile in 50 mM KH₂PO₄ (pH 4.0), pumped at a rate of 0.1 ml/min. The absorbance was detected at wavelengths of 295 and 287 nm for 6OHCZX and CZX, respectively [14,16]. The retention times of 6OHCZX, phenacetin and CZX were approximately 5, 12 and 20 min, respectively. Linear calibration curves (r > 0.999) were obtained for both compounds in plasma over the concentration range from 1 to 150 µg/ml. The limits of detection were estimated to be 0.25 µg/ml for both CZX and 6OHCZX.

2.6. Reverse transcriptase polymerase chain reaction (RT-PCR) assay

Total RNA was isolated from liver, kidney and fat with Isogen (Nippon Gene Co. Ltd., Toyama, Japan). Each RNA sample (1 μ g) was reversed-transcribed at 37 °C for 2 h, and the cDNA was amplified with a Peltier Thermal Cycler PTC-100 (Bio-Rad Laboratories Inc., Hercules, CA). The numbers of amplification cycles were 30 for fat cDNA and 25 for cDNAs from other tissues. PCR products were evaluated by electrophoresis on 2% (w/v) agarose gel stained with ethidium bromide, and photographed under UV trans-illumination. The product size was estimated by comparison with a 100 bp DNA ladder

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129 (Takara Bio Inc., Shiga, Japan). Control reactions to verify the absence of contaminants and genomic DNA were routinely 130 131 performed. Primers used for rat CYP2E1 were 5'-CTG ATT GGC TCCCCA CCC TGC-3' and 3'-GAA CACCTC GGC CAA AGT CAC-5', pop), and those for rat β -actin e 5'-TTC TAC AAT GAG 132 133 GTG CGT GTG GC-3' and 3'-CTC CTA GCT CTT CTC CAG GGA 134 135 GGA-5' (456 bp). PCR was run under the following conditions: 136 initial denaturation at 94 °C for 3 min, repeated denaturation 137 at 94 °C for 45 s, followed with annealing at 66 °C for 45 s for 138 CYP2E1 and 67 °C for 45 s for β -actin, primer extension at 72 °C for 45 s, and final extension at 72 °C for 3 min. The other 139 conditions for RT-PCR were as described previously [17]. 140

141 **2.7.** Preparation of microsomes

Tissue microsomes were prepared according to Yokogawa 142 et al. [17] with slight modifications. Liver, kidney or abdominal 143 fat was homogenized with phosphate buffer (50 mM K₂HPO₄ 144 containing 0.1 mM EDTA, pH 7.4). The homogenate was 145 146 centrifuged at 10,000 \times *q* for 30 min, and then the supernatant 147 was recentrifuged at $100,000 \times q$ for 60 min. The microsomal 148 pellet was collected, and resuspended in an appropriate volume of 50 mM Tris-acetate buffer (pH 7.4) containing 1 mM 149 EDTA and 20% glycerol. The protein concentration of the 150 151 microsomal solution was determined with a protein assay kit from Bio-Rad Laboratories Inc. (Hercules, CA). 152

153 2.8. Immunoblotting

Immunoblotting of the microsomes for CYP2E1 and β -actin 154 was carried out essentially as described by Yokogawa et al. 155 [18]. The microsomal protein was resolved by 10% SDS-PAGE, 156 and transferred to a polyvinylidene fluoride 157 Immobilon-P[®] (Millipore Co., Bedford, MA). The mbrane 158 159 was incubated with blocking buffer for 1 h (5%, w/v, non-fat 160 dried milk in PBS), followed with overnight incubation in a 161 2000-fold dilution of primary antibody (goat anti-rat CYP2E1 162 from Daiichi, Pure Chemicals Co., Ltd., Tokyo, Japan, and goat 163 anti-rat β -actin from Santa Cruz Biotechnology Inc., Santa

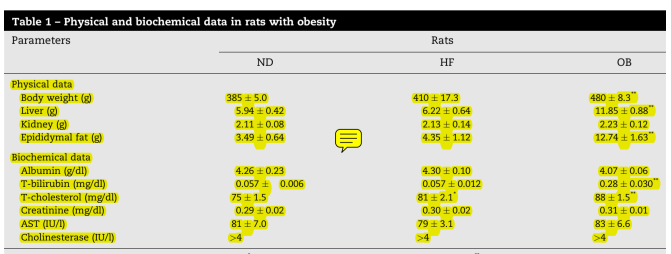
Cruz, CA). The membranes were then incubated with a 2000fold dilution of secondary antibody (rabbit anti-goat IgG from Santa Cruz Biotechnology Inc., Santa Cruz, CA). The immunopositive band was detected with an ECL chemiluminescence detection kit (Amersham Biosciences UK Ltd., Buckinghamshire, UK), and scanned with a Typhoon 9200 scanner (Amersham Biosciences Europe GmbH, Freiburg, Germany).

2.9. Measurement of hydroxylation activity of microsomal CYP2E1

The enzyme activity of microsomal CYP2E1 was determined by the measurement of 6OHCZX formation according to Chittur and Tracy [19], with minor modifications. A mixture of the microsomal solution (equivalent to 0.4 mg protein) and 50 mM phosphate buffer (pH 7.4) containing 50 μ M CZX and 1 mM NADPH (final, 200 μ l) was incubated at 37 °C for appropriate times. The reaction was stopped by adding 100 μ l of acetonitrile containing phenacetin (1 μ g) as an internal standard for HPLC assay. The 6OHCZX formed was extracted with diethyl ether and measured by HPLC as described above.

2.10. Measurement of glucuronidation activity of microsomal UGTs

The activity of UDP-glucuronosyltransferase (UGT) μ m microsomes was measured colorimetrically in 0.15 M trisphosphate buffer pH 7.4 containing 0.8 mM *p*-nitrophenol (PNP), 14 mM UDP glucuronic acid, 10 mM MgCl₂, and 1 mg microsomal protein [20]. The mixture (1.4 ml) was incubated at 37 °C for 30 min, then the reaction was stopped by adding 5 ml of 0.2 M glycine buffer (pH 10.4). The disappearance of PNP was quantified in terms of absorbance at 405 nm with a UV-vis spectrophotometer. The microsomal activities for glucuronidation of CZX and 6OHCZX were assayed similarly, except that the disappearance of CZX and 6OHCZX was determined by HPLC.



Data were presented as mean \pm S.D. of four rats. *Significant difference from ND rats at P < 0.05. *Significant difference from ND rats at P < 0.01.

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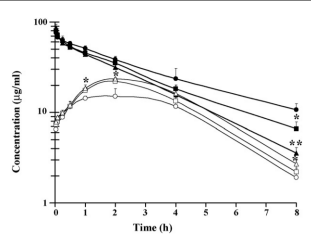


Fig. 1 – Plasma concentration-time courses of CZX (closed symbols) and 60HCZX-T (open symbols) after an i.v. administration of CZX (20 mg/kg) over 2 min in ND (\bigcirc), HF (\square) and OB (\triangle) rats. The concentration of CZX is shown as unchanged fraction, while 60HCZX is shown as total 60HCZX (unchanged fraction plus glucuronidated fraction). Each point and bar represents the mean + S.D. of four rats. 'Significant difference from ND rats at P < 0.05. "Significant difference from ND rats at P < 0.01.

199 2.11. Data analysis

The pharmacokinetic parameters were calculated according to 200 model-independent moment analysis as described by Yamaoka 201 202 et al. [21]. Electrophoregrams after RT-PCR and immunopositive 203 bands were evaluated in arbitrary units by using NIH Image 204 software. Comparisons of numerical data among groups were 205 made by one-way ANOVA, with P < 0.05 as the criterion of a 206 significant difference. For each significant effect, a multiple 207 comparison test was performed with Scheffe's test to verify the 208 difference between groups at P-values of 0.05 and 0.01, using 209 SPSS 10[®] from SPSS Inc. (Chicago, IL).

3. Results

3.1. Physical and biochemical data

Table 1 summarizes the physical data for ND, HF and OB rats. The body, liver and epididymal fat weights of OB rats were significantly higher than those of ND rats. The fat tissue weight of HF rats, although slightly higher, was not significantly different from that of ND rats. The value of total bilirubin of OB rats was significantly higher than that of ND rats, but liver function (AST) and kidney function (creatinine) were unaffected. Total cholesterol was significantly increased in HF rats and OB rats compared with normal rats.

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3.2. Disposition kinetics of CZX

Fig. 1 shows the plasma concentration-time courses of CZX and 6OHCZX after an i.v. administration of CZX 20 mg/kg in ND, HF and OB rats. The plasma concentrations of CZX linearly decreased in all cases, but the concentration at 8 h after administration was in the order of ND > HF > OB rats. In this figure, the concentration of 6OHCZX is presented as a total value of 6OHCZX (6OHCZX-T), i.e., the sum of free 6OHCZX and glucuronidated 60HCZX, after the administration of CZX. The amount of the glucuronide conjugate of CZX (CZX-G) was negligible (data not shown). It was found that the plasma concentration of 60HCZX-T increased gradually and reached a peak at about 2 h after administration. The peak concentration of 6OHCZX-T in OB rats was significantly higher than that in ND rats. Table 2 shows the pharmacokinetic parameters of CZX and 60HCZX-T after administration of CZX 20 mg/kg. In OB rats, the AUC_{\infty}, $T_{1/2}$ and MRT values of CZX were significantly smaller, while the CLtot and Vdss values of CZX were significantly larger than those of ND rats. The pharmacokinetic parameters in HF rats were intermediate between those in OB rats and ND rats. Moreover, the AUC_{∞} values of 6OHCZX-T in HF and OB rats were significantly higher than that of ND rats.

Table 2 – Pharmacokinetic parameters of CZX and 60HCZX after an i.v. administration of CZX 20 mg/kg over 2 min					
Parameters		Rats			
	ND	HF	OB		
CZX					
$AUC_{\infty} (\mu g/h/ml)^{a}$	204 ± 14	$\begin{array}{c} 181 \pm 12 \\ 1.38 \pm 0.19 \end{array}$	155 ± 21		
$T_{1/2} (h^{-1})^{b}$	1.67 ± 0.18		$1.19 \pm 0.23^*$		
$MRT (h^{-1})^{c}$	2.40 ± 0.19	2.30 ± 0.28	$2.00 \pm 0.17^*$		
CL _{tot} (1/h kg) ^d	0.104 ± 0.002	$0.122 \pm 0.010^*$	$0.145 \pm 0.013^{**}$		
Vd _{ss} (1/h) ^e	0.239 ± 0.014	0.277 ± 0.014"	0.285 ± 0.016**		
60HCZX-T					
AUC_{∞} (µg/h ml)	41.2 ± 4.7	60.3 ± 4.2	71.6 ± 5.9*		

Data were presented as mean \pm S.D. of four rats. *Significant difference from ND rats at P < 0.05. *Significant difference from ND rats at P < 0.01.

 $^{\rm a}\,$ AUC from 0 to $\infty.$

^b Half life.

^c Mean residence time.

^d Total clearance.

^e Volume of distribution at steady state.

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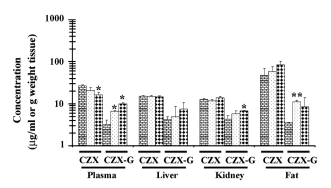


Fig. 2 – Tissue and plasma concentrations of CZX and its glucuronide (CZX-G) at 4 h after an i.v. administration of CZX (20 mg/kg) over 2 min in ND (), HF (⊡) and OB () rats. Each column and bar represents the mean + S.D. of four rats. [°]Significant difference from ND rats at P < 0.05. ^{°°}Significant difference from ND rats at P < 0.01.

244 Fig. 2 shows the tissue concentration of CZX compared with 245 the plasma concentration at 4 h after the i.v. administration of 246 CZX. The CZX concentration in fat tissue was considerably higher than those of plasma, kidney and liver. The concentra-247 248 tion of CZX-G was higher in HF rats and OB rats than ND rats. Fig. 3 shows the tissue concentrations of 60HCZX and its 249 glucuronide in fat, liver and kidney at 4 h after the i.v. 250 251 administration of CZX. The concentrations of both 6OHCZX and 60HCZX-G in plasma and all tissues tended to be higher in 252 253 the HF rats and OB rats than those in the ND rats. Interestingly, 254 the kidney concentration was much higher than those of other 255 tissues, and the concentration of 6OHCZX-G in kidney of OB rats was significantly higher than that of ND rats. 256

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3.3. Expression of CYP isoform mRNAs and proteins

The mRNA expression of CYP2E1 in liver of HF rats and OB rats was only slightly higher than that in ND rats, whereas, the relative expression of CYP2E1/ β -actin in fat tissue from HF rats and OB rats was apparently higher than that in ND rats (Fig. 4).

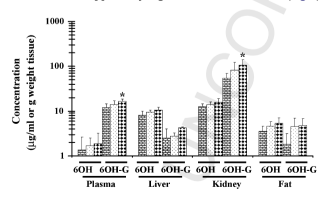


Fig. 3 – Tissue and plasma concentrations of 6OHCZX (6OH) and its glucuronide (6OH-G) at 4 h after an i.v. administration of CZX (20 mg/kg) over 2 min in ND (), HF (☉) and OB (↔) rats. Each column and bar represents the mean + S.D. of four rats. Significant difference from ND rats at P < 0.05. Significant difference from ND rats at P < 0.01.

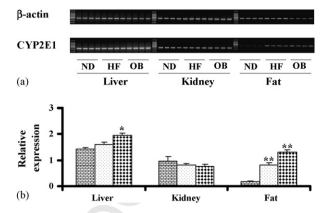


Fig. 4 – Effect of obesity on the expression of CYP2E1 mRNA compared with β -actin in liver, kidney and fat of ND (\equiv), HF (\equiv) and OB (\equiv) rats. Each column and bar represents the mean + S.D. of four rats. Significant difference from ND rats at P < 0.05. Significant difference from ND rats at P < 0.01.

The expression levels of CYP2E1 protein in the liver, kidney and fat tissue were examined by immunoblot analysis. The protein levels of CYP2E1 in the liver and fat of HF rats and OB rats were conspicuously increased compared with those of ND rats. Furthermore, the relative expression of CYP2E1/ β -actin in liver microsomes and fat microsomes of HF and OB rats was significant higher than that in ND rats (Fig. 5).

3.4. In vitro CYP2E1 and UGTs activity

Fig. 6 shows the hydroxylation activity of microsomal CYP2E1 in terms of 6OHCZX formation from CZX, in various tissues from the three groups of rats. The highest hydroxylation rate was found in the liver compared with kidney and fat tissue. The activities in liver and fat from HF rats and OB rats were significantly higher than those in ND rats, whereas, no difference was seen in the kidney.

Fig. 7 shows the glucuronidation activity of microsomal UGTs in liver and kidney towards PNP, CZX and 6OHCZX. The glucuronidation rates in the kidney of HF rats and OB rats were generally significantly higher than those in ND rats. The glucuronidation activity in fat was negligible in all groups. Interestingly, the glucuronidation of CZX was very much lower than that of 6OHCZX, although the enzyme activity in the liver was not affected by obesity.

4. Discussion

Zucker (fa/fa) rats at 20 weeks of age showed markedly increased body weight, accompanied with an enlarged liver and increased epididymal fat, compared with Zucker (+/+) rats. In contrast, Zucker (+/+) rats fed with high-fat diet (twice the level in normal diet) for 12 weeks did not show any significant physical or biochemical changes (Table 1).

CZX is well known to be a specific probe for CYP2E1, being hydroxylated to 6OHCZX [10,11], which in turn is rapidly glucuronidated to 6OHCZX-G [13]. Therefore, the appropriate

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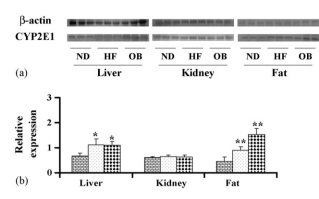


Fig. 5 – Effect of obesity on the content of CYP2E1 protein compared with β -actin in liver, kidney and fat of ND (\cong), HF (\cong) and OB (\cong) rats. Each column and bar represents the mean + S.D. of three rats. Significant difference from ND rats at P < 0.05. Significant difference from ND rats at P < 0.01.

295 indicator for determining CYP2E1 activity in vivo should be the 296 total amount of 60HCZX generated after the administration of 297 CZX, as reported in Fig. 1 and Table 2. The limited sampling time course of 8 h post-administration was contributed to the 298 299 detection limits of both CZX and 6OHCZX. Most of 6OHCZX in biological samples were lower than the detection limit, and 300 some of CZX levels were under the linearity of calibration 301 302 curves at 12 h after administration of CZX. The administration 303 of CZX 20 mg/kg to OB rats afforded lower values of AUC $_{\infty}$ and 304 $T_{1/2}$ in serum as compared with those in ND rats, while the 305 values of CL_{tot} and Vd_{ss} were significantly higher (Table 2). 306 Based on the Vd_{ss} and the tissue concentration of CZX (Fig. 2), 307 it appears that lipophilic substances, such as CZX, penetrate well into the fat reservoirs of OB rats. The fat to plasma ratio of 308 309 CZX was two to four folds both at 1 and 4 h of tissue sampling 310 time, and the OB rats showed the higher trend of accumulation 311 (data not shown for 1 h). Therefore, if CYP2E1 were not 312 induced in the fat of OB rats, it is likely that the elimination 313 rate of CZX would be delayed. However, the $T_{1/2}$ of CZX in OB 314 rats was shorter than in ND rats, and further, the value of

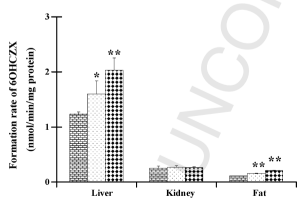


Fig. 6 – Effect of obesity on the hydroxylation activity of microsomal CYP2E1 in liver, kidney and fat of ND (\equiv), HF (\equiv) and OB (\cong) rats. Each column and bar represents the mean + S.D. of four rats. Significant difference from ND rats at P < 0.05. Significant difference from ND rats at P < 0.01.

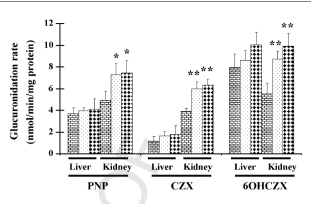


Fig. 7 – Effect of obesity on the glucuronidation activity of microsomal UGTs in liver and kidney of ND (\cong), HF (\cong) and OB (\cong) rats. Each column and bar represents the mean + S.D. of four rats. Significant difference from ND rats at P < 0.05. Significant difference from ND rats at P < 0.01.

60HCZX in fat higher than the plasma level (Fig. 3). It is unlikely that 60HCZX from blood would accumulate in fat, because 60HCZX is relatively hydrophilic and is rapidly glucuronidated by UGTs located in smooth endoplasmic reticulum, the same location as that of CYP2E1 [22]. Therefore, the induction of CYP2E1 in fat, in addition to liver, may play a pivotal role in determining the disposition kinetics of CZX in obese rats. We also found that the AUC_∞ ratio of 60HCZX–T/ CZX in HF rats and OB rats was 1.5–2 times higher than that in ND rats, reflecting the increased total activity of CYP2E1 in rats with obesity. Lucas et al. [7] reported that the 60HCZX–T/CZX ratio (0.4) in obese or hyperlipidemic patients was higher than that (0.3) in normal subjects, in agreement with our finding. These results suggest that obesity and feeding of a high-fat diet can induce CYP2E1 activity in both humans and rats.

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We found that both the protein content and activity of CYP2E1 were increased in microsomes of the liver from HF rats and OB rats, while there was no change in the kidney. Kobayashi et al. [23] reported that CZX was extensively metabolized in rat microsomes not only by CYP2E1, but also by CYP1A2 and CYP3A. Therefore, we also examined the mRNA expression and protein content of CYP1A2 and CYP3A, but found that they were unaffected by obese status (data not shown). Enriquez et al. and Irizar et al. [24,6] reported that CYP2E is poorly expressed in obese Zucker rats, which is consistent with our finding. However, Enriquez et al. [24] reported that CZX hydroxylase activity and CYP2E1 protein content were lower in Zucker (fa/fa) rats than in lean (+/?)littermates. There are various differences between their experimental conditions and ours, but one of the most important factors could be the influence of aging in obese Zucker (fa/fa) rats. Young obese Zucker (fa/fa) rats do not exhibit pathological conditions such as physical obesity, insulin resistance, etc., and usually have a lower CYP2E1 activity than their lean littermates, whereas, after the appearance of pathological symptoms at approximately 14-16 weeks of age, expression of CYP2E1 is increased. Therefore, we used Zucker (fa/fa) rats at 20 weeks of age in our study, when their pathological condition appeared to resemble that of severe obesity in humans. The report by Enriquez et al. [24]

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did not mention the physical condition of either the lean or 355 356 obese Zucker rats, so that it is difficult to compare their 357 findings and ours. Interestingly, the CYP2E1 mRNA isoform was expressed in fat tissue, and its expression level and 358 activity were significantly higher in HF rats and OB rats than in 359 ND rats (Figs. 4-6). Yoshinari et al. [25] and Wan et al. [26] 360 reported that the expression levels of both CYP2E1 mRNA and 361 362 protein were increased in adipose tissue of fasting rats. 363 However, the amount of adipose tissue in fasting rats was 364 small, and so CYP2E1 in fasting animals may contribute little to the pharmacokinetics of its substrates compared with the 365 situation in obese animals. 366

The hydroxylated metabolite of CZX (60HCZX) has been 367 reported to be excreted in bile to only a small extent; rather it 368 undergoes rapid glucuronidation with subsequent excretion 369 of the conjugate in urine [13]. We found that the glucuronida-370 tion activity in kidney microsomes was significantly higher in 371 HF rats and OB rats than ND rats (Fig. 7), and the concentration 372 373 of 6OHCZX glucuronide in the kidney was higher than that in 374 the liver (Fig. 3). Since there is no evidence that a specific UGT 375 subfamily in involved in glucuronide conjugation of CZX and 376 6OHCZX, we used PNP, a general marker for glucuronidation, 377 to confirm our results. The glucuronidation activity towards PNP in kidney microsomes of HF and OB rats showed the same 378 379 trend as did the activity towards CZX and 6OHCZX. However, the glucuronidation activity in liver microsomes was not 380 influenced by the high-fat diet or obese status, and was quite 381 different towards different substrates. This suggests the 382 presence of different UGTs isoforms in liver and kidney. 383 Further research will be required to identify the UGTs 384 isoform(s) responsible for 6OHCZX glucuronidation in tissues, 385 and the species that are affected by obese status. 386

In conclusion, CYP2E1 activity was induced in the liver 387 fat tissues of obese animals, and glucuronidation activity were 388 389 induced in the kidney. As a result, the disposition kinetics of 390 CZX was markedly changed in obesity, with an increase in 391 hydroxylation of CZX to form 60HCZX, and an acceleration of 392 6OHCZX glucuronidation, resulting in rapid excretion in urine. 393 Other drugs that are metabolized by CYP2E1 and UGTs may 394 show similar changes of disposition kinetics in obese patients, resulting in reduced potency and shorter duration of action. 395

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