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Effects of a Novel UGT2B Haplotype and UGT1A4*3 Allele Variants on Glucuronidation of Clozapine In vivo

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TITLE PAGE

Title: Effects of a novel *UGT2B* haplotype and *UGT1A4*3* allele variants on glucuronidation of clozapine *in vivo*

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Running title: Effects of various UGT polymorphism on clozapine glucuronides

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ABSTRACT

Background: Glucuronidation is an important metabolic pathway of clozapine (CLZ), but the impact of various uridine 5'diphospho-glucuronosyltransferases (*UGT*) polymorphisms on the exposure and metabolism of CLZ *in vivo* is unclear.

Objective: The objective of this study was to investigate the impact of UGT2B haplotype and UGT1A4*3 allele variants on the formation of CLZ glucuronide metabolites (5*N*- and *N*⁺-glucuronide) and CLZ exposure in patients' serum after a djusting for sex, age and smoking habits.

Methods: The study was based on serum samples from CLZ-treated patients (n=79) subjected to routine therapeutic drug monitoring (TDM) at Diakonhjemmet Hospital, Oslo, Norway. From the same patients the following *UGT* variants were genotyped using Real-Time PCR: *UGT2B:GA* haplotype (defined as *UGT2B:GA*; rs1513559A > G and rs416593T > A) and UGT1A4*3 (rs2011425T > G). Serum concentrations of CLZ 5*N*- and N^+ -glucuronide were measured by UPLC high-resolution mass spectrometry.

Results: None of the genotypes had significant impact on CLZ exposure (p>0.05). However, compared to UGT2B:AT/AT and UGT1A4*1/*1, the 5*N*-glucuronide exposure was reduced in UGT2B:GA/GA carriers (-75%, p=0.03) while the exposure was non-significantly increased in UGT1A4*3 carriers (+100%, p=0.14), respectively. The N^+ -glucuronide exposure was unchanged in UGT1A4*3 vs noncarriers (p=0.28), but significantly reduced in heterozygous (-50%, p=0.016) and homozygous carriers (-70%, p=0.021) of UGT2B:GA compared to UGT2B:AT/AT carriers, respectively.

Conclusion: The UGT2B:GA and UGT1A4*3 variants had no impact on CLZ exposure, but were associated with differences and preferences in CLZ glucuronidation. The latter might be of potential relevance for CLZ tolerability, since levels of the N^+ -glucuronide metabolite may reflect the generation and trapping of reactive metabolites involved in CLZ-induced toxicity.

Keywords: Clozapine, Uridine 5'-diphospho-glucuronosyltransferase, Therapeutic drug monitoring, Glucuronidation, Metabolism, High-resolution mass spectrometry.

1. INTRODUCTION

Cloza pine (CLZ) is the superior atypical antipsychotic drug with respect to symptom improvement in patients with schizophrenia [1]. The therapeutic use of CLZ is limited to patients with treatment resistant schizophrenia (TRS) due to the risk of serious CLZ-induced toxicity such as neutropenia, a granulocytosis and seizures [2, 3]. Despite this restriction in the use of CLZ, it is considered to be underutilized in the treatment of TRS patients [4]. A possible contribution to the observed underutilization may be the major interindividual variability in CLZ serum concentrations, which is challenging for obtaining an optimal dose within the narrow therapeutic range of 1071-1836 nM [5].

The metabolism of CLZ involves many enzymes, including several cytochrome P450 (CYP) enzymes and uridine 5'-diphospho-glucurosonyltransferases (UGTs) [6]. Of particular interest in CLZ metabolism are the CYP1A2, CYP3A4, CYP2D6, UGT1A and UGT2B enzymes [6, 7], in which the expressions of CYP1A2 and UGTs are induced by smoking [8, 9]. In two recent genome-wide association studies (GWASs), investigating associations between single nucleotide polymorphisms (SNPs) and *N*-desmethylclozapine (*N*-DMC)-to-CLZ ratios in patients' serum, significant loci were observed between the *UGT2B10/15* genes [10, 11]. The lead SNPs *rs1670747* and *rs2926038* were highly significant with p-values above 2.0 x 10^{-23} [10]. The minor allele frequencies (MAF) of *rs1670747* and *rs2926038* in the European population were 0.23 and 0.10 [12], respectively. The function of these SNPs beyond affecting CLZ metabolism is unknown. However, polymorphisms in the nearest genes, i.e. *UGT2B10* and *2B15* have been described to influence the glucuronidation rate of cotinine (nicotine metabolite) and postoperative anxiety after lorazepam premedication, respectively [13, 14]. Furthermore, much interests have been paid to the *UGT1A4*3* (142 T>G, L48V, MAF = 0.077) allele variant [15], which has been shown to alter the glucuronide formation of various compounds *in vitro* [15-18]. Particularly, the formation of CLZ 5*N*-glucuronide was increased in cell lines overexpressing *UGT1A4*3/*3*, suggesting an increased function phenotype of the *UGT1A4*3* allele variant in CLZ glucuronidation [19].

No studies have per date investigated the effect of the UGT1A4*3 and UGT2B allele variants (*rs1670747* and *rs2926038*) on the glucuronidation of CLZ *in vivo*. The aim of the present study was therefore to investigate the impact of UGT1A4*3 and UGT2B haplotype (based on *rs1670747G>A* and *rs2926038G>C*) variants on the formation of the two major glucuronides in human serum, i.e. CLZ 5*N*- and *N*⁺-glucuronide, and potential effect on CLZ exposure.

2. MATERIALS AND METHODS

2.2 Subjects

The study was based on serum samples of CLZ-treated patients from a routine therapeutic drug monitoring (TDM) service, who had also been subjected to genotyping, at the Center for Psychopharmacology, Diakonhjemmet Hospital (Oslo, Norway), during January 2018–January 2020.

Information about CLZ dose, concomitant medications and time interval between the last dose and sampling time was retrieved from the TDM requisition forms. Inclusion criteria were i) serum samples of CLZ drawn 10-24 hours after the last dose intake, ii) information about dosing of CLZ provided on the requisition forms, and iii) information about smoking. Samples were excluded if patient age was below 18 or above 75 years, or when review of the requisition forms disclosed concomitant use of CNS drugs interacting with the metabolism of CLZ, i.e. the CYP1A2 inhibitor fluvoxamine, antiepileptic CYP inducers (carbamazepine, phenytoin, and phenobarbital), valproic acid or the UGT1A4 substrate lamotrigine. In addition, samples were excluded if there was discrepancy between the smoking habits given on the requisition forms and cotinine levels (nicotine metabolite) in the accompanying serum samples. Since cotinine is present in small amounts in blood regardless of smoking, a conservative and pragmatic threshold value of cotinine areal-under-curve (AUC) was set to 1,00 x 10⁶ based on reviewing the cotinine levels in smokers vs. nonsmokers (supplementary table 1). Thus, patients who were smokers or nonsmokers based on the information on the requisition forms, would be excluded if the cotinine levels were below or above this threshold, respectively. The intention of using cotinine in the study was to confirm the information about smoking habits written on the requisition forms. The study included all samples (i.e. multiple samples per patients) obtaining the abovementioned criteria.

The study was approved by the Regional Committee for Medical and Health Research Ethics (2014/1185) and did not require informed patient consent as only historical data were applied without the potential to cause any harm.

2.2 Analyses of Clozapine, Clozapine N-oxide, N-desmethylclozapine, Clozapine 5N/N⁺-glucuronide and Cotinine serum concentrations

During the time course of retrospective data collection, a n ultra-high-performance LC (UHPLC)-high resolution mass spectrometry (HRMS) method was applied for identification and quantification of CLZ, CLZ *N*-oxide, *N*-DMC, CLZ $5N/N^+$ -glucuronide and cotinine. Briefly, the serum samples were prepared by protein precipitation in 96-deep well plates using a Microlab Star pipetting robot (Hamilton, Reno, NV, USA) in a semi-automated sample

preparation procedure. The LC system was a Vanquish-UHPLC (Thermo Fisher Scientific, Waltham, MA, USA), and chromatographic separation was performed by an XBridge BEH C18-column (2.6 μ m, 2.1x75 mm; Waters, Milford, MA, USA) using gradient elution at 35 °C with a mix of ammonium acetate buffer (pH = 4.8) and acetonitrile (20-52%). The retention times were 0.4, 0.8, 1.3, 1.5, 1.6, and 1.7 min for cotinine, CLZ *5N*-glucuronide, CLZ *N*⁺-glucuronide, *N*-DMC, CLZ *N*-oxide and CLZ, respectively. Detection used a QExactive Orbitrap mass spectrometer (Thermo Fisher Scientific, Waltham, MA, USA), operated in positive ionisation mode acquiring full scan data at a resolution of 70,000 within the 100-1,500 Da scan range. The compounds were quantified in full scan acquisition mode, while accurate data dependent MS2 analysis was simultaneously triggered to permit confirmation of their identification. The method for identification and quantification of CLZ, *N*-DMC and CLZ *N*-oxide are validated and used in routine analyses at Diakonhjemmet Hospital. CLZ and 13C-d3-CLZ (internal standard) were purchased from Sigma-Aldrich (St. Louis, MO, USA) and Alsachim (Illkirch-Graffenstaden, France), respectively.

For CLZ 5*N*- and *N*⁺-glucuronide and cotinine, their levels in the serum samples were assessed by retrospective reprocessing of the HRMS data file, as described elsewhere for other compounds [20], using the TraceFinder 4.1 software (Thermo Fisher Scientific, Waltham, MA, USA). The CLZ $5N/N^+$ -glucuronides and cotinine were tentatively identified by accurate mass (the protonated molecular ion was present within a mass tolerance of 5 ppm), isotope ratio and interpretation of the MS2 spectrum. Identification of CLZ 5N-glucuronide and CLZ N^+ -glucuronide were confirmed using retention time and matched MS2 spectrum by analysing reference standards purchased commercially (TLC Pharmaceutical Standards, Newmarket, ON, Canada). Thus, as the extracted-ion chromatogram for the exact mass for CLZ N^+ -glucuronide, respectively, by analyses of reference standards.

2.3 UGT1A4*3 and UGT2B (rs1670747G>A and rs2926038G>C) Genotyping

Genomic DNA was extracted from blood samples using a MagNA Pure 24 Instrument with MagNA Pure 24 Total NA Isolation kit (Roche Diagnostics, Germany).

Due to high levels of DNA sequence homology between the *UGT* genes, a Real-Time (RT) PCR method for the *UGT1A4*3* variant (142T>G, *rs2011425*) has not yet been available. Therefore, a novel RT-PCR method was developed at Center for Psychopharmacology using a mixture of allele-specific primers and allele-specific TaqMan probes. The RT-PCR was designed with a common reverse primer (ACC CTT GAG TGT AGC CCA GC) and two forward primers specific to the **1*-allele (TCA GCA TGC GGG AGG CCT) and the **3*-allele (TCA GCA TGC GGG AGG CCG) respectively, in addition to allele specific TaqMan probes (VIC – CAT GGA GCT CCC GCA AG - *MGB* and 6-FAM – CAT GGA GCT CCC GCA CG - *MGB*). The allele specific nucleotides are marked in boldface. The RT-PCR assay was performed by QuantStudio 12K Flex RT-PCR Instrument with QS12K Software, Version 1.2.2 (Thermo Fisher Scientific, Waltham, USA) using 384-well PCR plates. The wells contained 8 μ L of PCR mix, which included 1X TaqMan Universal Genotyping Master Mix (Thermo Fisher), 250 nM of the **1*-allele-specific forward primer, 150 nM of the **3*-allele-specific forward primer, 400 nM of the common reverse primer, 200 nM of the **1*-specific VIC-labelled probe, 120 nM of the **3*-specific FAM-labelled probe and around 30 ng of genomic DNA. Samples were heated95°C for 10 min to initiate Taq DNA polymerase, followed by 50 cycles of amplification (95°C for 15 s and 60°C for 60 s). The RT-PCR method was validated by comparing 92 unique samples with the established restriction fragment length polymorphism (PCR-RFLP) method, described elsewhere [21]. All sample results gave 100% compliance between the methods.

TaqMan assays for rs1670747G>A and rs2926038G>C genotyping were not available at Thermo Fisher Scientific, therefore assays for two SNPs that were in complete linkage disequilibrium with rs1670747 and rs2926038 were purchased (R²=1 in Europeans) [22] i.e. rs1670747/rs416593 (cat. nr: C_771051_10) and rs2926038/rs1513559 (Catnr: C_1509939_10). The RT-PCRs for rs1670747/rs416593 and rs2926038/rs1513559genotyping were performed as recommended by Thermo Fisher Scientific in 8 µL volumes and same temperature cycles as for UGT1A4*3.

Combined haplotypes (UGT2B) from rs1513559 and rs416593 were generated using PHASE 2.0 [23]. These two SNPs are located closely (~41kbp; chr4) and are correlated in a European population (R²=0.31) [22].

2.4 Statistical Analysis

To evaluate the effect of UGT1A4*3 allele and UGT2B haplotype variants on dose-adjusted serum concentration of CLZ 5*N*-glucuronide, CLZ N^+ -glucuronide *and* the various metabolite-to-parent (MPRs) ratios, linear mixed model analyses were used to allow multiple samples per patient. In these analyses age, sex, blood sampling time and smoking habits were included as covariates. The drug exposure was defined as the dose-adjusted serum concentration (concentration/dose ratio, CD ratio). We used ln-transformed values to ensure normal distributions. Undetectable levels of CLZ glucuronides were truncated to the lowest observed value within each CLZ glucuronide respectively to ensure proper calculation of MPRs. All statistical analyses were performed in SPSS®, version 21.0 (IBM® SPSS® Statistics, Armonk, NY, USA). GraphPad version 4 was used for graphical presentations (GraphPad Software, San Diego, CA).

Resident

3. RESULTS

In total, 79 patients with 216 serum concentrations were included of which 61% of the patients were smokers from information on the requisition forms, which was confirmed by measuring high levels of cotinine (table 1). The age of the population was 43.3 years (SD: 14.7y) comprising of a male majority (62%). In the population, three different *UGT2B* haplotypes were identified. The haplotype frequencies were 10.1%, 11.4% and 78.5% for the *UGT2B:AA*, *UGT2B:GA* and *UGT2B:AT* haplotypes, respectively (*UGT2B:AA: rs1513559A*>*G* and *rs416593T*>*A*; *UGT2B:GA* and *rs416593T*>*A*; *UGT2B:AT rs1513559A*>*G* and *rs416593T*>*A*; *UGT2B:AF rs1513559A*>*G* and *rs416593T*>*A*; *UGT2B:AF* haplotype variant, respectively. No significant differences in the Hardy-Weinberg equilibrium were observed between the investigated SNPs (p>0.05). In the population, the average prescribed CLZ dose was 416 mg (table 1). The average absolute serum concentrations of CLZ, *N*-DMC, CLZ *N*-oxide, CLZ *5N*-glucuronide and CLZ*N*⁺-glucuronide were 1496, 1011, 131, 83 and 71 nM, respectively. Smokers had an average cotinine levels 3000-fold higher compared to nonsmokers (table 1 and supplementary figure 1).

In the multivariate analyses adjusting for smoking habits, sex, age and sampling time, UGT1A4*3 carriers were observed with a 1.5-fold higher CD ratio of CLZ compared to noncarriers, but the difference was not statistically significant (UGT1A4*3 carriers vs noncarriers: 5.46 vs 3.69 nM/mg, p=0.064, table 2 and supplementary table 1). Furthermore, UGT2B:GA/GA and UGT2B:GA/AT carriers were associated with decreased CD ratios of N-DMC (UGT2B:GA/GA vs UGT2B:AT/AT: 1.44 vs 2.75 nM/mg, p=0.019; UGT2B:GA/AT vs UGT2B:AT/AT: 2.08 vs 2.75 nM/mg, p=0.062; table 2 and supplementary table 1) as compared to the UGT2B:AT/AT carriers. Compared to UGT2B:AT/AT carriers, significantly reduced N-DMC-to-CLZ MPR ratios were observed in UGT2B:GA/GA and UGT2B:GA/AT carriers, respectively (UGT2B:GA/GA vs UGT2B:AT/AT: 0.42 vs 0.69, p=0.012; UGT2B:GA/AT vs UGT2B:AT/AT: 0.58 vs 0.69 nM/mg, p=0.106; table 2). The CD ratio of CLZ 5N-glucuronide was reduced by 75% in carriers of UGT2B:GA/GA haplotype vs UGT2B:AT/AT carriers (UGT2B:GA/GA vs UGT2B:AT/AT: 0.038 vs 0.16 nM/mg, p=0.030; table 2). Accordingly, the CLZ 5Nglucuronide-to-CLZ MPR ratio was reduced by the same degree in UGT2B:GA/GA carriers (UGT2B:GA/GA vs UGT2B:AT/AT: 0.011 vs 0.039, p=0.030; table 2) when comparing to noncarriers. Compared to noncarriers, UGT1A4*3 carriers had a nominally increased CD ratio of CLZ 5N-glucuronide by 100%, but this effect was not significant (p=0.14, table 2 and supplementary table 1). Compared to UGT2B:AT/AT carriers, UGT2B:GA/GA and *UGT2B:GA/AT* carriers had 70% and 50% (*UGT2B:GA/GA vs UGT2B:AT/AT*: 0.045 vs 0.15 nM/mg, p=0.021; UGT2B:GA/AT vs UGT2B:AT/AT: 0.076 vs 0.15 nM/mg, p=0.016; table 2 and supplementary table 1) reduced CD ratios of CLZ N^+ -glucuronide levels, respectively. Accordingly, the CLZ N^+ -glucuronide-to-CLZ MPR ratios were reduced in UGT2B:GA/GA and UGT2B:GA/AT carriers vs UGT2B:AT/AT carriers (UGT2B:GA/GA vs UGT2B:AT/AT: 0.012 vs 0.039 nM/mg, p=0.022; UGT2B:GA/AT vs UGT2B:AT/AT: 0.021 vs 0.039 nM/mg, p=0.028; table 2 and supplementary table 1).

In the mixed model, smoking reduced the CD ratio of CLZ by 35% at average (p<0.001; supplementary table 1). No effects on the various MPRs and CD ratios of CLZ and CLZ metabolites/glucuronides were observed in UGT2B:AA haplotype vs. UGT2B:AT/AT carriers (p>0.05; figure 1, table 2 and supplementary table 1). No effect on CLZ *N*-oxide was observed among the various UGT1A4*3 and UGT2B:GA carriers vs. noncarriers (figure 1C).

4. **DISCUSSION**

In the present study, the carriers of the *UGT2B:GA* haplotype determined the variants (*rs1513559* and *rs416593*) had significantly decreased CLZ glucuronidation. Thus, the study confirms the importance of these variants for CLZ metabolism and glucuronidation as identified by the recent GWASs [10, 11]. No significant impact on CLZ exposure was observed in carriers of the *UGT2B:GA* haplotype nor the *UGT1A4*3* allele variant when compared to noncarriers. However, the study demonstrated a potential increased CLZ 5*N*-glucuronidation by the *UGT2B:GA* haplotype and *UGT1A4*3* allele variant as previously shown *in vitro* [19]. The altered CLZ glucuronidation by the *UGT2B:GA* haplotype and *UGT1A4*3* allele variants should be further investigated in future clinical studies to assess them in relation to CLZ tolerability.

The reduced glucuronidation rate of CLZ observed in *UGT2B:GA/GA* carriers did not affect the senum levels of CLZ, and thus the clinical effect is unlikely to be changed. However, altered glucuronidation by the *UGT2B:GA* haplotype and *UGT1A4*3* allele variants may involve a shift in other metabolic pathways relevant for the tolerability of CLZ. One of the suggested toxic metabolites, i.e. the CLZ nitrenium ion, is shown to be involved in CLZ-induced agranulocytosis [24]. Therefore, any alterations of the CLZ metabolic pathway caused by changed UGT-mediated glucuronidation may be of clinical relevance in CLZ treatment. Recently, a positive association was shown between levels of *N*-DMC and neutrophil granulocyte [25], which potentially indicate that a shift in metabolic pathways may have an impact on CLZ tolerability. Therefore, one might hypothesize that carriers of the *UGT2B:GA/GA* diplotype may have increased generation of potential toxic metabolites such as the CLZ nitrenium ion, since they show reduced glucuronidation-mediated trapping of the reactive CLZ metabolite. In fact, carriers of *UGT2B:GA/GA* had decreased *N*-DMC levels and *N*-DMC-to-CLZ MPR ratio, further supporting this hypothesis of a nonprotective role of the *GA* haplotype in clozapine tolerability. The effect size on *N*-DMC-to-CLZ MPR ratio of the *UGT2B:GA* haplotype was approximately the same as previously reported in a larger study [10]. On the other hand, *UGT1A4*3* carriers seem to have an increased glucuronidation phenotype which may in turn protect against accumulation of CLZ nitrenium ions and may have a favourable clozapine tolerability profik.

The potential increased glucuronidation rate in UGT1A4*3 carriers has previously been observed in olanzapine-treated patients, an antipsychotic drug with similarities in the chemical structure of clozapine [18]. Thus, the increased function of the UGT1A4*3 variant allele on glucuronidation of clozapine and olanzapine seems to be analogous. In another study, however, the UGT1A4*3 variant allele was shown to decrease the exposure of olanzapine by approximately 25%, which was like the effect size of male sex and smoking habits in schizophrenic patients [21]. This was not the case in our cohort, since the UGT1A4*3 allele variant seems to be a ssociated with

increased CLZ levels, although the number of UGT1A4*3 carriers is low (n=6). Thus, it is important to evaluate this observation in larger studies.

Many of the limitations of the present study are due to the naturalistic setting of the data collection. The study lacked information about comorbidity, organ functions, body weight and potential use of interacting drugs, except from fluvoxamine, carbamazepine, phenytoin, phenobarbital, lamotrigine and valproic acid, which are regularly written on the requisition forms. On the other hand, we had access to several factors that could affect the glucuronidation rate of CLZ, including sex, age and smoking habits, as confirmed by measuring high or low cotinine levels in the blood samples, which were adjusted for in the statistical analyses. As the various *UGT* haplotypes are relevant for CLZ pharmacokinetics, they may potentially also be relevant for CLZ tolerability. However, the number of patients in present study was low and larger clinical studies are warranted, to determine the importance of the *UGT1A4*3* allele and *UGT:GA* haplotype variants in CLZ tolerability.

CONCLUSION

In conclusion, the present study shows that the UGT2B:GA carriers have reduced CLZ glucuronidation and the UGT1A4*3 carriers have potentially increased CLZ 5N-glucuronidation. As reactive metabolites play a role in CLZ-induced toxicity, altered glucuronidation in UGT2B:GA/GA carriers and UGT1A4*3 carriers may potentially alter the inactivation of reactive CLZ metabolites, and hence be relevant for the treatment tolerability.

ETHICS APPROVAL AND CONSENT TO PARTICIPATE

The study was approved by the Regional Committee for Medical and Health Research Ethics (2014/1185), and did not require informed patient consent as only historical data were applied without the potential to cause any harm.

HUMAN AND ANIMAL RIGHTS

No animals were used for studies that are the basis of this research. All human procedures followed were in accordance with the Helsinki Declaration of 1975.

CONSENT FOR PUBLICATION

Not applicable.

AVAILABILITY OF DATA AND MATERIALS

Not applicable.

FUNDING

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CONFLICT OF INTEREST

The authors declare no conflicts of interest, financial or otherwise.

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FIGURE LEGENDS

Figure **1** Effects of the *UGT1A4*3 Allele*, the *UGT2B:GA* and *UGT2B:AA* Haplotypes on Exposure of Clozapine (CLZ), *N*-desmethylclozapine, CLZ *5N*-glucuronide, CLZ *N*⁺-glucuronide, and CLZ *N*-oxide. The univariate comparisons of the *UGT1A4*3* allele and the *UGT2B:GA* and *UGT2B:AA* haplotypes on dose-adjusted serum concentrations (CD) of clozapine (A; CLZ), *N*-desmethylclozapine (*N*-DMC) (B), CLZ *5N*-glucuronide (C), CLZ *N*⁺-glucuronide (D) and CLZ*N*-oxide (E). The horizontal lines represent the mean of the presented subpopulations. Each dot represents a patient sample (both black and grey). Black dots represent the reference group used for statistical comparisons within each allele variant. All values are ln-transformed to ensure normal distribution. Linear mixed model analyses were used and statistical significance was considered at p-value below 0.05.

Supplementary figure **1** The serum cotinine levels in nonsmokers and smokers. Information a bout smoking habits (non-smoker or smoker) were retrieved from the TDM requisition forms. The dotted horizontal line represents the threshold of exclusion of cotinine samples above for non-smoker and cotinine levels below for smokers, respectively.

TABLES

Table 1 Population Characteristics.

Variables	Values
Baseline characteristics:	
Female/Male, n (samples)	30(66)/49(150)
Age, mean years (SD)	43.3 (14.7)
Sampling time, mean hrs (SD)	14.0(2.59)
Dose, mg (SD)	416(193)
Confirmed smokers (%)	48 (61%)
Cotinine levels in smokers/non-	333x10 ⁶ (215)/
smoker, AUC (SD)	$0.11 \times 10^{6} (0.125)$
Genotype characteristics:	
UGT1A4 allele carriers:	
*1/*3, n (samples)	6(18)
*3/*3, n (samples)	0(0)
UGT2B haplotype carriers:	
<i>UGT2B:AT/AT</i> , n (samples)	49 (122)
UGT2B:AT/AA, n (samples)	15 (57)
UGT2B:GA/AT, n (samples)	11 (30)
UGT2B:GA/AA, n (samples)	1(1)
UGT2B:GA/GA, n (samples)	3(6)
Absolute concentrations:	
CLZ, nM mean (SD)	1496 (920)
<i>N</i> -DMC, nM mean (SD)	1011 (583)
CLZ 5N-glucuronide, nM mean (SD)	83 (99)
$CLZN^+$ - glucuronide, nM mean (SD)	71 (88)
CLZ N-oxide, nM mean (SD)	131 (84)

CLZ, clozapine; *N*-DMC, *N*-desmethylclozapine; P, p-value; SD, Standard deviation; SNP, single nucleotide polymorphism.

UGT2B haplotypes are defined based on SNPs rs1513559A > G and rs416593T > A

Table 2 The Effect of UGT1A4*3 Allele and UGT2B Haplotype Variants on Exposure and Metabolic Ratios of Clozapine, N-desmethylclozapine, CLZ 5N-glucuronide, and CLZ N^+ -glucuronide. The effect of UGT1A4*3 allele and UGT2B haplotype (combined rs1513559A>G and rs416593T>A) variants on dose-adjusted serum concentrations (CD) of clozapine (CLZ), N-desmethylclozapine (N-DMC), clozapine glucuronides and various metabolic ratios adjusted for smoking habits, sex, age and sampling withdra wal time.

	MPR N-DMC	MPR CLZ 5N-	MPR CLZ N^+ -	gluc.	$CDCLZN^+$ -	gluc	CDN-DMC		CDCLZ		CDCLZ5N-gluc			
	Fold change (95%	Р	Fold change	P	Fold change	P	Fold change	P	Fold change	Р	Fold change	Р	Fold change	P
	CI)		(95% CI)		(95% CI)		(95% CI)		(95% CI)		(95% CI)		(95% CI)	
UGT2B haloptype:														
AT/AT(ref)	-		-		-		-		-		-		-	
GA/AT	0.84 (0.68, 1.0)	0.11	1.4 (0.72, 2.6)	0.34	0.55 (0.32,0.94)	0.028	0.50 (0.28, 0.88)	0.016	0.76(0.57,1.0)	0.062	0.90 (0.65, 1.3)	0.54	1.2 (0.58, 2.6)	0.58
GA/GA	0.61 (0.42, 0.90) 0	0.012	0.28 (0.087, 0.88)	0.030	0.32 (0.12, 0.84)	0.022	0.29 (0.10, 0.83)	0.021	0.53 (0.31, 0.90)	0.019	0.86 (0.47, 1.6)	0.63	0.24 (0.062, 0.94))0.041
AA/AT	0.96 (0.79, 1.2)	0.63	0.81 (0.46, 1.4)	0.45	0.82 (0.52, 1.3)	0.42	0.85 (0.52, 1.4)	0.51	1.0 (0.77, 1.3)	0.97	1.0 (0.78, 1.4)	0.78	0.84 (0.44, 1.6)	0.59
GA/AA	0.61 (0.31, 1.2)	0.15	0.59 (0.074, 4.8)	0.62	0.22 (0.037, 1.3)	0.092	0.51 (0.075, 3.5)	0.49	1.4 (0.55, 3.6)	0.46	2.3 (0.79, 6.8)	0.12	1.4 (0.12, 16)	0.80
UGT1A4 genotype:														
*3 vs *1/*1	0.92 (0.71, 1.2)	0.54	1.3 (0.62, 2.9)	0.44	1.0 (0.52, 1.9)	0.99	1.5 (0.73, 2.9)	0.28	1.4 (0.95, 2.0)).094	1.5 (0.98,2.2) ().064	2.0 (0.79, 5.0)	0.14

Linear mixed model analyses were used allowing inclusion of multiple measurements per patients. *UGT2B:AT/AT* genotype was used as reference. The statistical analyses were performed after including a ge, sex, smoking habits and sampling time as covariates.

UGT2B haplotype is based on the combination of SNPs rs1513559A>G and rs416593T>A. UGT2B: **GA**, rs1513559**G** and rs416593A; UGT2B: **AA**, rs1513559**A** and rs416593**A**; UGT2B: **AA**, rs1513559**A** and rs416593**T**.

CD, CD ratio (concentration-to-dose ratio); MPR, metabolite-to-CLZ ratio; *N*-DMC, *N*-desmethylclozapine.

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SUPPLEMENTARY MATERIAL

Supplementary table **1** The Effect of UGT1A4*3 Allele, UGT2B Haplotype Variants, Age, Sex and Withdrawal Time on Exposure and Metabolic Ratios of Clozapine, N-desmethylclozapine, CLZ 5*N*-glucuronide, and CLZ N^+ -glucuronide. The table shows the complete statistical model of UGT1A4*3 allele and UGT2B haplotype (combined rs1513559A > G and rs416593T > A) variants on dose-adjusted serum concentrations of clozapine, N-desmethylclozapine (N-DMC), clozapine glucuronides and various

metabolic ratios adjusted for smoking habits, sex, a ge and sampling withdrawal time.

Variables	CDCLZ CD <i>N</i> -DMC			1C	CDCLZ: glucuron	5 <i>N-</i> ide	CDCLZ <i>l</i> glucuroni	V ⁺ - ide	MPR <i>N</i> -DMC		MPR CLZ 5N- glucuronide		MPR CLZ N ⁺ - glucuronide	
	β value, nM/mg(SE)	Р	β value, nM/mg(SE)	Р	β value, nM/mg(SE)	Р	β value, nM/mg(SE)	Р	βvalue (SE)	Р	βvalue (SE)	Р	βvalue (SE)	Р
Intercept	1.46 (0.26)	<0.001	0.96 (0.23)	<0.001	-2.40 (0.62)	<0.001	-1.71 (0.49)	0.001	-0.49 (0.17)	0.003	-3.86 (0.52)	<0.001	-3.19(0.45)	<0.001
UGT1A4*3 carriers	0.39 (0.21)	0.064	0.31 (0.18)	0.094	0.68 (0.46)	0.14	0.37 (0.34)	0.28	-0.082 (0.13)	0.54	0.30 (0.39)	0.44	0.0036 (0.33)	0.99
UGT2B haplotypes:														
AT/AT (ref)	Ref	Ref	Ref	Ref	Ref	Ref	Ref	Ref	Ref	Ref	Ref	Ref	Ref	Ref
AA/AT	0.041 (0.15)	0.78	-0.0047 (0.13)	0.97	-0.18 (0.33)	0.59	-0.16(0.25)	0.51	-0.046 (0.094)	0.63	-0.21 (0.28)	0.45	-0.19(0.23)	0.42
GA/AT	-0.10(0.17)	0.54	-0.28 (0.15)	0.062	0.21 (0.37)	0.58	-0.69 (0.28)	0.016	-0.17 (0.11)	0.11	0.31 (0.32)	0.34	-0.60 (0.89)	0.028
GA/AA	0.84 (0.54)	0.12	0.35 (0.47)	0.46	0.32(1.24)	0.80	-0.67 (0.97)	0.49	-0.50 (0.34)	0.15	-0.52(1.05)	0.62	-1.52 (0.87)	0.092
GA/GA	-0.15 (0.30)	0.63	-0.64 (0.27)	0.019	-1.42 (0.68)	0.041	-1.23 (0.52)	0.021	-0.49 (0.19)	0.012	-1.28 (0.58)	0.030	-1.15 (0.49)	0.022
Smoker	-0.44 (0.12)	<0.001	-0.29 (0.10)	0.007	-0.36 (0.26)	0.18	-0.15 (0.20)	0.46	0.15 (0.075)	0.052	0.085 (0.22)	0.71	0.29 (0.19)	0.13
Male sex	0.016	0.90	-0.018 (0.11)	0.87	0.32 (0.28)	0.26	-0.25 (0.21)	0.23	-0.034 (0.078)	0.66	0.30 (0.23)	0.21	-0.28 (0.20)	0.16
Age (per year)	0.001 (0.004)	0.73	0.001 (0.003)	0.72	0.011 (0.0084)	0.193	0.0029 (0.006)	0.65	-0.002 (0.002)	0.94	0.010 (0.007)	0.16	0.002 (0.006)	0.70
Blood sampling (per hr)	-0.0043 (0.0095)	0.65	0.0029 (0.008)	0.72	-0.020 (0.024)	0.419	-0.015 (0.021)	0.47	0.0076 (0.0056)	0.178	-0.016 (0.021)	0.44	-0.011 (0.018)	0.54

All CD values are presented as Ln-transformed values to ensure normal distribution. Linear mixed model analyses were used to include multiple measurements per patients. UGT2B AT/AT genotype was used as reference. The statistical analyses were performed after including age, sex, smoking habits and sampling time as covariates.

UGT2B haplotype is based on the combination of SNPs rs1513559A > G and rs416593T > A. UGT2B:GA, rs1513559G and rs416593A; UGT2B:AA, rs1513559A and rs416593A; UGT2B:AA, rs1513559A and rs416593T.

CD, CD ratio (concentration-to-doseratio); MPR, metabolite-to-CLZ ratio; SE, standard error; SNP, single nucleotide polymorphism; N-DMC, N-desmethylclozapine.

Figure 1:



Supplementary figure 1:

