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Variation in the plasma membrane monoamine transporter (PMAT, encoded in *SLC29A4*) and organic cation transporter 1 (OCT1, encoded in *SLC22A1*) and gastrointestinal intolerance to metformin in type 2 diabetes: an IMI DIRECT study

Running title: Gut metformin transporters and GI intolerance

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Abstract

OBJECTIVES 20-30% of patients with metformin treated type 2 diabetes experience gastrointestinal side effects leading to premature discontinuation in 5-10% of the cases. Gastrointestinal intolerance may reflect localised high concentrations of metformin in the gut. We hypothesized that reduced transport of metformin via the plasma membrane monoamine transporter (PMAT) and organic cation transporter 1 (OCT1) could increase the risk of severe gastrointestinal side effects.

RESEARCH DESIGN AND METHODS The study included 286 severe metformin intolerant and 1128 tolerant individuals from the IMI DIRECT consortium. We assessed the association of patient characteristics, concomitant medication and the burden of mutations in the *SLC29A4* and *SLC22A1* genes on odds of intolerance.

RESULTS Women (p < 0.001) and older people (p < 0.001) were more likely to develop metformin intolerance. Concomitant use of transporter inhibiting drugs increased the odds of intolerance (OR 1.72, p < 0.001). In an adjusted logistic regression model, the G allele at rs3889348 (*SLC29A4*) was associated with gastrointestinal intolerance (OR 1.34, p = 0.005). rs3889348 is the top cis-eQTL for *SLC29A4* in gut tissue where carriers of the G allele had reduced expression. Homozygous carriers of the G allele treated with transporter inhibiting drugs had over three times higher odds of intolerance compared to carriers of no G allele and not treated with inhibiting drugs (OR 3.23, p < 0.001). Using a genetic risk score derived from rs3889348 and *SLC22A1* variants, the odds of intolerance was more than twice in individuals who carry three or more risk alleles compared with those carrying none (OR 2.15, p = 0.01).

Conclusions: These results suggest that intestinal metformin transporters and concomitant medications play an important role in gastrointestinal side effects of metformin.

Metformin therapy can cause gastrointestinal discomfort that negatively affects quality of life and adherence to prescribed medications. Gastrointestinal side effects usually manifest as nausea, vomiting, diarrhoea, flatulence, indigestion, bloating, abdominal discomfort and stomach ache. 20-30% of metformin treated subjects with type 2 diabetes experience gastrointestinal side effects leading to premature discontinuation in 5-10% of the cases (1, 2). This inhibits adherence to therapy and may lead to a change of treatment, depriving intolerant patients of effective diabetes therapy. Despite its clinical importance, the underlying pathophysiology of metformin intolerance is not yet clear. However multiple possible hypotheses have been proposed including high intestinal metformin concentration (3, 4), its effect on the gut microbiota (5), altered transportation of serotonin or direct serotonergic effects (6), and reduced ileal absorption of bile acid salts (7).

Metformin is not metabolized and is excreted unchanged in the urine. At physiologic pH, it is hydrophilic due to the presence of a quaternary ammonium group that results in a net positive charge. Therefore, Metformin does not efficiently diffuse across the biological membranes and requires carrier-mediated transport. Multiple solute carrier transporters expressed in membranes of the enterocytes, hepatocytes and the kidney are reported to be involved in the absorption, distribution and elimination of metformin. Metformin requires the entire length of the small intestine to be absorbed (8): around 20% of the administered dose is absorbed in the duodenum and 60% in the jejunum and ileum. The remainder reaches the colon and remains unabsorbed. PMAT and OCT1 are reported to play the major role in the intestinal absorption of metformin (9). While PMAT is expressed in the apical (luminal) membrane of the enterocytes, intestinal localization of OCT1 is ambiguous (9-11). An association between reduced function alleles in *SLC22A1* and concomitant use of OCT1 inhibiting drugs with metformin intolerance has been reported (12, 13). An interaction between OCT1 and Serotonin

Transporter (SERT) has also been shown to play an important role in the pathophysiology of metformin intolerance (13).

Whilst PMAT shares extensive substrate and inhibitor overlap with OCTs (14), there are no studies investigating its role in metformin intolerance. Therefore, we hypothesized that reduced transport of metformin by major transporters of metformin, PMAT and/or OCT1, could increase intestinal metformin concentration and subsequently increase the risk of GI side effects. To address this, we used prescribing, biochemistry and clinical data from 286 metformin intolerant and 1128 tolerant individuals from the IMI DIRECT (DIabetes REsearCh on patient straTification) consortium (15). Although OCT3 is expressed in the intestine, there are no common functional variants described, and therefore we do not include OCT3 in this analysis.

Research Design and Methods

Study population

286 metformin intolerant (cases) and 1128 metformin tolerant (controls) subjects were identified from prescribing data in the IMI DIRECT consortium from participating centres across northern Europe (15). Each participant consented to participate in the study and ethical approval was obtained from the medical ethics committees of the respective centres.

All metformin intolerant (cases) and metformin tolerant (controls) had: type 2 diabetes diagnosed clinically, a creatinine clearance ≥ 60 mL/min at metformin exposure, and were white Europeans aged between 18-90 years at recruitment.

Definition of metformin intolerance

The metformin intolerance phenotype was defined in two ways: firstly, individuals who switched to an alternative agent within 6 months of stopping metformin (including modified release metformin) after having had up to 1000 mg daily metformin for up to 6 weeks, who also reported gastrointestinal side effects on the metformin treatment as the reason for switching or where gastrointestinal side effects were clearly documented in the clinical record as a reason for transfer. In an alternative definition, intolerant individuals were defined as those who could not increase their metformin immediate release dose above 500 mg daily despite an HbA1c > 7% (53 mmol/mol) and who either reported gastrointestinal side effects on more than 500 mg, or where gastrointestinal side effects were clearly documented in the clinical record as a reason for transfer.

Where the patient was asked to recall side effects, the intolerant event was limited to be within the last 5 years; if side effects were documented from clinical records then there was no time limit. Participants who did not recall being on metformin or having side effects were excluded (unless clearly documented in clinical records).

Definition of metformin tolerance

Metformin tolerant individuals were defined as those treated with \geq 2000 mg of metformin per day for more than a year (excluding modified release formulations of metformin) and report no side effects.

Clinical covariates

Weight, height and creatinine were defined as the closest measured values within 180 days prior to the index intolerance event (ITE) and BMI was calculated as weight in kg / (height in

m)². The ITE was defined as the date when patients report gastrointestinal symptoms of metformin intolerance for cases, and for controls it is the date when patients start 2000 mg of metformin. Daily dose was the last dose during ITE for cases, and it was determined as the mean dose of prescriptions encashed during the first six months of metformin therapy for controls.

Concomitant medications

Gut metformin transporters have strong substrate and inhibitor overlap (16). Therefore, we identified medications prescribed together with metformin previously reported to inhibit the PMAT and/or OCTs, proteins that mediate transmembrane trafficking of their target molecules, and are required for metformin absorption in the gut. These drugs are selected based on their reported half-maximal inhibitory concentration (IC₅₀) values. Accordingly the use of any of the following medications with metformin were investigated: tricyclic antidepressants (TCAs) (17, 18), proton pump inhibitors (PPIs) (19), citalopram (18), verapamil (17, 18), diltiazem (18), doxazosin (17, 18), spironolactone (17, 18), clopidogrel (20), rosiglitazone (21), quinine (18), tramadol (18, 22), codeine (23), dysopyramide (24), quinidine (21), repaglinide (21), propafenone (17), ketoconazole (17), morphine (22, 23), tropisetron (25), ondasetrone (25), antipsychotic agents (17) and tyrosine kinase inhibitors (26).

Genotyping

DNA samples from participants were genotyped at the University of Oxford (UOXF) using the Illumina Human Core Exome chip v1.0 (HCE24 v1.0). Genotype calling was performed using the GenCall algorithm in the GenomeStudio software supplied by Illumina. Data were subjected to a series of standard quality control analyses in order to highlight poorly performing genetic markers and samples prior to imputation.

Samples were excluded for any of the following reasons: call rate less than 95%, heterozygosity greater than 4 standard deviations (SD) from the mean, high correlation to another sample (pihat ≥ 0.2) or identified as ethnic outlier from constructed axes of genetic variation from principal components analysis implemented in the Genome-wide Complex Trait Analysis (GCTA) software (v1.24.7) (27) using the 1000 Genome as a reference. Further filtration was performed to remove: non-autosomal markers, duplicate markers (sharing the same positions), markers with minor allele frequency (MAF) <1%, Hardy–Weinberg equilibrium (HWE) pvalue < 0.0001 and call rate < 98%. Imputation to the 1000 Genomes Phase 3 CEU reference panel was performed with ShapeIt (v2.r790) (28) and Impute2 (v2.3.2) (29).

Single nucleotide polymorphism selection

As there are no functionally characterised common nonsynonymous SNPs in the *SLC29A4* gene, the tagging intronic SNPs, rs3889348 and rs2685753 ($r^2 = 0.57$, D' = 1) had been previously shown to be associated with trough steady state metformin concentration (30). Therefore, the rs3889348 G>A genotype, was extracted from existing genome-wide data. The frequency of the minor allele (A) of rs3889348 was 38%. Data for previously reported missense *SLC22A1* variants (M420del (18.6%), R61C (7.1%), G401S (3.1%)) were also extracted from the genome-wide data. There was no deviation from HWE for any polymorphism (p > 0.05).

Statistical methods

Categorical data are presented as frequency (percentage) and continuous variables as mean \pm SD if normally distributed or as median and inter quartile range (IQR) otherwise. Students t-test and the Mann-Whitney U test were used to compare differences in quantitative variables distributed normally or not, respectively. Comparison of categorical variables between cases

and controls was done using X^2 test. Logistic regression was used to estimate the association of independent variables with metformin intolerance. Multivariate logistic regression analyses of metformin intolerance were performed with all the covariates included using SNPTEST (version 2.5.2) (31). Association of the intronic rs3889348 G>A in *SLC29A4* was explored assuming additive genetic model. *SLC22A1* variants (M420del, R61C, G401S) were grouped together by summing up the number of risk alleles. Combined unweighted genetic risk score (GRS) was generated as 0, 1, or 2 according to the number of reduced-function alleles in each individual. The combined genotype was then added to the multivariate analyses assuming an additive model. A two-tailed p-value less than 0.025 was considered statistically significant.

Expression quantitative trait locus (eQTL) analyses

We investigated whether rs3889348 is a cis-QTL in the gut using eQTL datasets comprising a total of 246 colon transverse and 122 terminal ilium samples from the Genotype-Tissue expression (GTEx) data release V6 (32). Tissue procurement, gene expression analysis, genotyping and eQTL analysis have been described previously (32-34).

Results

Phenotypic differences between tolerant and intolerant subjects

The characteristics of tolerant and intolerant subjects are presented in Table 1. Women (p < 0.001) and older people at diagnosis or at ITE (p < 0.001) were more likely to be metformin intolerant. Compared to tolerant subjects, metformin intolerant individuals had lower weight (p < 0.001), lower creatinine clearance (p = 0.036) and were treated with a lower metformin dose (p < 0.001).

Concomitant medications and intolerance

This analysis was performed on 237 metformin intolerant and 1128 tolerant subjects who had complete data on history of concomitant medications. Forty percent of metformin intolerant subjects were taking one or more cation transporter inhibitory drugs compared to 24% in tolerant subjects (p < 0.0001) (Table 1). In a logistic regression model adjusted for age, gender, weight and concomitant use of these drugs increased the odds of being intolerant by 70% (OR = 1.72 [1.26-2.32], p < 0.001) (Supplementary Table 1). When the individual drug or drug groups were explored, concomitant use of metformin with either PPIs, TCAs or codeine increased the odds of metformin intolerance significantly (Figure 1). The number of subjects who have been coprescribed metformin with transporter inhibiting drugs are shown in Supplementary Table 2.

Genetic variation in the gut metformin transporters and metformin intolerance

In a logistic regression model, carriers of the G allele had 1.39 [1.15-1.69, p < 0.001] times higher odds of being intolerant to metformin (unadjusted). When rs3889348 was added to a model adjusted for age, sex, weight and genetic substructure, the presence of the G allele was independently associated with metformin intolerance (OR = 1.34[1.09-1.65], p = 0.005) (Supplementary Table 1). No statistically significant difference in any of the baseline phenotypes by genotype was observed (Supplemental Table 3). In addition, no significant interaction between rs3889348, the use of metformin transporter inhibiting drugs and any of the other clinical variables (age, gender) was observed.

We then grouped subjects based on the combination of *SLC29A4* genotype and concomitant use of metformin transporter inhibiting drugs. Taking those with no risk allele and not treated with transporter inhibiting drugs as the reference group, carriers of one and two G alleles treated with transporter inhibiting drugs had more than two (2.44 [1.30-4.78]) and three (3.23 [1.71-

6.39]) fold higher odds of intolerance, respectively, after adjusting for age, sex and weight (Supplementary Table 4).

The association between *SLC22A1* genotypes and metformin intolerance has been previously reported (12, 35). We carried out an analysis on the association between two reduced function (R61C, G401S) and one loss of function (M420del) *SLC22A1* SNPs and metformin intolerance using a combined unweighted GRS. In a logistic regression model adjusted for age, sex, weight, genetic substructure and concomitant use of transporter inhibiting drugs, the *SLC22A1* GRS was not statistically significantly associated with metformin intolerance (OR = 1.35 [0.84-2.12], p = 0.21).

A GRS was then generated from *SLC29A4* and *SLC22A1* variants by summing up the number of risk alleles for each individual. Compared to those with no risk allele, metformin treated subjects with type 2 diabetes having two risk alleles had nearly a two-fold (1.93[1.10-3.65]) increased odds of GI intolerance. Those who carry 3 or more risk alleles had more than twice (2.15[1.20-4.12]) the odds of intolerance (Figure 2).

Sensitivity analysis

There was a big difference in sample size between metformin intolerant and tolerant subjects. In addition, there were significant difference in age and gender between cases and controls. Therefore, we carried out a sensitivity analysis by comparing the intolerant group (n=237) with age and sex matched subgroup of tolerant subjects (n=711). The main findings from the larger metformin tolerant group were confirmed in this sensitivity analysis (Supplementary Table 5 and 6).

rs3889348 is associated with altered PMAT expression in the gut

Given PMAT is one of the major metformin transporters in the gut, we explored the possibility that the intronic SNP, rs3889348 is a cis-eQTL in the intestine utilizing the publicly available

data set from the GTEx portal (Version V6p) (32). The G-allele of rs3889348 (associated with higher risk of intolerance) was significantly associated with lower expression of *SLC29A4* in the terminal ileum of the small intestine ($\beta = -0.42$, $p = 2.1 \times 10^{-04}$) and the transverse colon ($\beta = -0.45$, $p = 1.4 \times 10^{-08}$) (Supplemental Figure 1). rs3889348 is the top cis-eQTL for *SLC29A4* in the transverse colon.

Conclusions

Intestinal absorption of metformin is modulated by the function of cation transporters expressed in the gut. An association between reduced function alleles in the SLC22A1, encoding organic cation transporter 1, and metformin related GI side effects has been previously reported (12, 13, 36). However, the data on intestinal localization of OCT1 is ambiguous; with mixed reports suggesting in the apical (10) and basolateral (11, 37) sides. In addition to OCT1, PMAT also contributes to the intestinal absorption of metformin. PMAT is abundantly expressed in the human intestine and it is concentrated on the tips of the mucosal epithelial layer (38). Carriers of the G allele at this locus (rs3889348) had significantly reduced expression of SLC29A4 in the gut (32). This could lead to higher luminal concentration of metformin. In this current study we demonstrated a significant association of the G allele of an intronic SNP, rs3889348, in SLC29A4 encoding PMAT, with higher odds of GI intolerance after metformin therapy. Each copy of the G allele was associated with 1.34 times higher odds of metformin intolerance. We also show that those who carry two or more variants at either SLC29A4 or SLC22A1, were two-fold more likely to have GI intolerance. Given that PMAT is apically located, this finding suggests that intolerance is driven by increased luminal concentration of metformin, rather than increased enterocyte concentration and direct toxicity to the enterocytes.

There are a number of putative mechanisms whereby increased luminal metformin may increase GI intolerance to metformin (outlined in Figure 3). Firstly, a higher concentration of

metformin in the gut has been shown to inhibit uptake of histamine and serotonin leading to increased luminal concentration of these biogenic amines (13). Metformin is also shown to inhibit diamine oxidase (DAO), an enzyme that degrades histamine, at therapeutic doses (6). Biogenic amines play an important role in the GI pathophysiology. Elevated levels of serotonin and histamine in the GI tract cause GI symptoms such as nausea, vomiting and diarrhea (6, 39). Serotonin is produced mainly in the gut and stored in the enterochromaffin cells of the epithelium. Its release activates gut sensory neurons that will increase intestinal motility, secretion and sensation (39, 40). Increased colon motility and softening of stool consistency has also been observed in serotonin reuptake transporter (SERT) knock-out mice (39, 40). In addition, a recent study from the GoDARTS cohort showed association of a composite SERT genotype, 5-HTTLPR (5-hydroxy tryptamine (serotonin) transporter linked polymorphic region)/rs25531, with intolerance to metformin in subjects with type 2 diabetes (13). In this study, carriers of the low-expressing SERT S* alleles had more than 30% increased odds of metformin intolerance (OR=1.31, 95% CI 1.02-1.67, p = 0.031). Histamine is a monogenic amine stored in the enterochromaffin-like cells within the gastric glands of the stomach. Binding of histamine to the H1, H2 and H4 receptors that are highly expressed in the gut, stimulate gastric acid secretion, increase intestinal motility and smooth muscle inflammation (6).

In addition to the potential role of local concentrations of serotonin and histamine, increased luminal concentrations of metformin could also cause intolerance by other mechanisms that need to be explored. For example, intolerance could be mediated by a reduction in bile acid reabsorption in the ileum leading to elevated bile acid levels in the colon (41), which is known to cause GI disturbances (42). In addition, metformin affects composition and function of the gut microbiota favoring the growth of some species like *Akkermansia* (5, 43-46). Furthermore, increased levels of active and total GLP-1 levels in subjects with type 2 diabetes and without

type 2 diabetes treated with metformin (47) have also been reported and this might increase GI side effects (48) (Figure 3).

In this study we observed increased risk of intolerance with older age, female sex, lower weight and lower creatinine levels. Concomitant use of metformin with the PPIs and TCAs also increase the risk of intolerance. These findings are largely consistent with the results of previous studies, providing further evidence for clinical practice (12, 35). The FDA Adverse Events Reporting System (AERS) suggested that women experience more adverse effects than men (49). Several factors can contribute to these differences. Gender based variability in intestinal expression of drug transporters may result in variability in gut drug concentrations. Women have also slower gastric emptying, altered bile composition and slower intestinal transit time than men (50). These factors could in turn affect the rate and/or extent of absorption of oral medications and hence local drug concentrations in the gut. For a better understanding of the basic mechanisms of sex differences in metformin intolerance, future studies should be designed with a primary focus on this topic.

In summary, we have identified a variant that alters intestinal expression of the cation transporter PMAT (*SLC29A4*) that increases risk of metformin associated gastrointestinal intolerance, and that combined with the previously reported *SLC22A1* variants, this genotype profile can increase odds of metformin intolerance over 2-folds. The apical location of PMAT means that reduced expression will result in increased luminal metformin concentration, suggesting that metformin intolerance is caused by this increased luminal concentration rather than increased enterocyte concentration.

A limitation of this study was the definition for metformin induced GI intolerance. Even though we examined patient reports and clinical records for GI intolerance as a reason for stopping metformin and switching to other medications, there could be other reasons for stopping metformin such as comorbidities that might cause GI disturbance. In addition, initial conclusions drawn from this study need validation and replication in a well-powered independent studies.

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Author contributions

A.D, analysed data, wrote manuscript, performed research. K.Z analysed data, designed research, revised manuscript. N.van L, collected data and revised manuscript. A.M, analysed data and revised manuscript. N.R, analysed data and revised manuscript. R.K, collected data and revised manuscript. P.E, collected data and revised manuscript. S.R, collected data and revised manuscript. A.J, designed research, collected data and revised manuscript. R.H, designed research, collected data and revised manuscript. P.F, designed research, performed research, revised manuscript. M.M, designed research, analysed data and revised manuscript. L.'t H, designed research, collected data and revised manuscript. A.J, analysed data, and revised manuscript. E.R.P designed research, performed research, analysed data, wrote manuscript and revised manuscript

A.D and E.P are guarantors of this work

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Conflict of interests

The authors claim no conflicts of interest

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Tables

Variable	Metformin Tolerant	Metformin Intolerant	р
	(n = 1,128)	(n = 286)	
Age at diabetes diagnosis	55.88 ± 9.44	58.62 ± 10.65	< 0.0001
(years)			
Age at ITE (years)	60.73 ± 9.84	64.63 ± 9.91	< 0.0001
Males/female (male %)	696/432 (61.7%)	117/169 (40.9%)	< 0.0001
Weight (kg)	94.57 ± 18.91	88.84 ± 17.75	< 0.0001
BMI (kg/m ²)	32.11 ± 6.01	31.60 ± 5.95	0.19
Creatinine (µmol/dL)	79.89 ± 16.09	78.41 ± 19.33	0.25
Creatinine clearance (mL/min)	85.17 ± 19.36	82.23±29.44	0.04
Dose (mg)* at diagnosis	1500 (1000-2000)	1000 (500-1000)	< 0.0001
Duration of diabetes (years)	4.0 (1.7-7.0)	4.0 (2.0-9.0)	0.09
Use of metformin transporter	274 (24.29%)	95 (40.08%)	< 0.0001
inhibiting drugs			

Table 1. Baseline characteristics of metformin tolerant and intolerant subjects.

ITE: index intolerance event *Median (IQR). Dose was calculated as the last dose during ITE for cases, and it was determined as the mean dose of prescriptions encashed during the first six months of metformin therapy for controls.

Figure legends

Figure 1. Association of individual intestinal metformin transporter inhibiting drugs with intolerance.

Figure 2. Association of a genetic risk score derived from *SLC29A4* (PMAT) and *SLC22A1* (OCT1) with metformin intolerance. OR: odds ratio; GRS: genetic risk score. Bars indicate standard errors around the mean.

Figure 3. Possible mechanisms for metformin intolerance. A) Metformin is absorbed from the gut lumen via cation transporters such as PMAT, OCT1, SERT and OCT3. B) Increased level of metformin in the gut lumen is observed when metformin is taken with cation transporter inhibiting drugs such as PPIs, TCAs and Codeine. These drugs competitively inhibit metformin uptake by the cation transporters. Metformin is also shown to inhibit diamine oxide, an enzyme that metabolize biogenic amines. In addition, transport capacity of the cation transporters could be reduced in carriers of reduced function (420del, 61C, 401S in *SLC22A1*) or low expressing alleles (rs3889348_G in *SLC29A4*) and hence increased luminal metformin level. Increased level of metformin increases the level of biogenic amines, affect the gut microbiota and elevate bile acid levels. These may cause symptoms of gastrointestinal side effects.

Figures



Figure 1



Figure 2



Figure 3

Supplemental Tables

Supplemental Table 1. Logistic regression model of metformin intolerance

Variable	OR [CI]	р
Age at ITE (years)	1.04[1.03-1.06]	7.44×10 ⁻⁰⁸
Gender	2.31[1.72-3.10]	2.52×10 ⁻⁰⁸
Weight (kg)	1.00[0.99-1.00]	0.20
Use of metformin	1.72[1.26-2.32]	5.17×10 ⁻⁰⁴
transporter inhibiting drugs		
rs3889348	1.34[1.09-1.65]	0.005

Gender coded as women vs men. ITE: index intolerance event

Supplementary Table 2. Number of patients treated concomitantly with metformin transporter inhibiting drugs in the intolerant and tolerant groups.

Drug/ Drug Class	Intolerant group (237)	Tolerant group (1128)	p
PPI	58 (24.5%)	156 (13.8%)	< 0.001
Codeine	22 (9.3%)	35 (3.1%)	< 0.001
ТСА	17 (7.2%)	33 (2.93%)	0.003
Tramadol	10 (4.2%)	18 (1.6%)	0.004
Clopidogrel	5 (2.1%)	14 (1.24%)	0.46
Spironolactone	3 (1.3%)	9 (0.8%)	0.75
Verapamil	3 (1.3%)	8 (0.7%)	0.64

Data are presented as numbers of patients (percentages). TCAs, tricyclic antidepressants; PPIs, proton pump inhibitors.

	rs3889348 genotype			
	AA (n = 205)	GA (n = 681)	GG (n = 532)	р
Age at ITE (years)	61.03 ± 10.50	61.34 ± 10.0	61.94 ± 9.73	0.21
Age at diagnosis (years)	55.97 ± 10.32	56.31 ± 9.67	56.78 ± 9.65	0.27
Gender (women %)	36.6%	42.6%	45.1%	0.11*
Weight (kg)	97.21 ± 19.39	92.53 ± 18.22	93.15 ± 19.43	0.06
BMI (kg/m ²)	32.77 ± 6.02	31.89 ± 6.09	31.88 ± 5.96	0.15
Height (m)	1.72 ± 0.10	1.71 ± 0.10	1.71 ± 0.10	0.29
Creatinine (µmol/dL)	80.51 ± 15.57	79.09 ± 16.29	79.90 ± 17.80	0.95
Creatinine clearance (mL/min)	84.75 ± 18.95	84.98 ± 18.55	84.04 ± 20.80	0.52
HbA1c (%)	8.21 ± 1.59	8.18 ± 1.62	8.39 ± 1.75	0.11
Diabetes duration (years) [†]	4 [1.16-7.00]	4 [2.00-7.19]	4 [1.78-7.00]	0.78 [‡]
Dose (mg) [†]	1000 (1000-1500)	1000 (1000-1500)	1000 (1000-1500)	1.00‡
Drug naïve (%)	44.1%	51.7%	51.4%	0.14*
Use of metformin transporter	22.4%	27.6%	28.8%	0.27^{*}
inhibiting drugs (%)				

Supplemental Table 3. Population characteristics by rs3889348 genotype.

*chi-square test for independence, ‡kruskal-Wallis one-way analysis of variance, †Median (IQR). ITE: index intolerance event

Supplemental Table 4. Joint effect of *SLC29A4* (PMAT) genotype and metformin transporter inhibiting drugs on metformin intolerance.

	Intolerant/Tolerant	OR [95% CI]	р
Carries no risk allele and not treated with metformin	15/141	1	
transporter inhibiting drugs			
Carries one risk allele and not treated with metformin	67/408	1.52 [0.86-2.88]	0.17
transporter inhibiting drugs			
Carries two risk alleles and not treated with metformin	60/305	1.76 [1.01-3.35]	0.04
transporter inhibiting drugs			
Carries no risk allele and treated with metformin	8/37	1.74 [0.64-4.46]	0.25
transporter inhibiting drugs			
Carries one risk allele and treated with metformin	44/137	2.44 [1.30-4.78]	0.007
transporter inhibiting drugs			
Carries two risk alleles and treated with metformin	43/100	3.23 [1.71-6.39]	< 0.001
transporter inhibiting drugs			

Supplementary Table 5. Joint effects of PMAT genotype and metformin transporter inhibiting drugs on intolerance after matching subjects for age and gender.

	OR (95% CI)	р
Weight	0.99[0.98-1.00]	0.08
Use of metformin transporter inhibiting drugs	1.75[1.28-2.38]	< 0.001
rs3889348 G	1.26[1.02-1.59]	0.03

Supplemental Table 6. Joint effect of *SLC29A4* (PMAT) genotype and metformin transporter inhibiting drugs on metformin intolerance after matching patients for age and gender. 237 cases and 711 controls (1 case ~3 controls).

	Intolerant/T	OR [95% CI]	р
	olerant		
Carries no risk allele and not treated with metformin	15/93	-	-
transporter inhibiting drugs			
Carries no risk allele and treated with metformin	8/18	2.76[0.99-7.38]	0.05
transporter inhibiting drugs			
Carries one risk allele and not treated with metformin	67/234	1.78[0.99-3.37]	0.06
transporter inhibiting drugs			
Carries one risk allele and treated with metformin	60/189	1.97[1.09-3.76]	0.03
transporter inhibiting drugs			
Carries two risk allele and not treated with metformin	44/100	2.73[1.45-5.37]	0.003
transporter inhibiting drugs			
Carries two risk alleles and treated with metformin	43/77	3.46[1.82-6.88]	< 0.001
transporter inhibiting drugs			

Supplemental Figures



Supplemental Figure 1. Boxplot of association between rs3889348 genotype and *SLC29A4* (PMAT) expression in the gut, colon transverse (left side) and terminal ilium of the small intestine (right side).