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1	<b>CRTC2</b> polymorphism as a risk factor for the incidence of metabolic
2	syndrome in patients with solid organ transplantation
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4	CRTC2 and Post-transplant metabolic syndrome.
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## 50 ABSTRACT

51 Metabolic syndrome after transplantation is a major concern following solid organ transplantation (SOT). The CREB-regulated transcription coactivator 2 (CRTC2) regulates glucose metabolism. The 52 effect of CRTC2 polymorphisms on new-onset diabetes after transplantation (NODAT) was 53 investigated in a discovery sample of SOT recipients (n<sub>1</sub>=197). Positive results were tested for 54 replication in two samples from the Swiss Transplant Cohort Study (STCS,  $n_2=1294$  and  $n_3=759$ ). 55 Obesity and other metabolic traits were also tested. Associations with metabolic traits in population-56 based samples (n<sub>4</sub>=46'186, n<sub>5</sub>=123'865, n<sub>6</sub>>100,000) were finally analyzed. In the discovery sample, 57 58 CRTC2 rs8450-AA genotype was associated with NODAT, fasting blood glucose and BMI (p<sub>corrected</sub><0.05). CRTC2 rs8450-AA genotype was associated with NODAT in the second STCS 59 replication sample (OR=2.01, p=0.04). In the combined STCS replication samples, the effect of 60 61 rs8450-AA genotype on NODAT was observed in patients having received SOT from a deceased donor and treated with tacrolimus (n=395, OR=2.08, p=0.02) and in non-kidney transplant recipients 62 63 (OR=2.09, p=0.02). Moreover, rs8450-AA genotype was associated with overweight or obesity (n=1215, OR=1.56, p=0.02), new-onset hyperlipidemia (n=1007, OR=1.76, p=0.007), and lower 64 HDL-cholesterol (n=1214,  $\beta$ =-0.08, p=0.001). In the population-based samples, a proxy of 65 rs8450G>A was significantly associated with several metabolic abnormalities. CRTC2 rs8450G>A66 67 appears to play an important role in the high prevalence of metabolic traits observed in patients with SOT. A weak association with metabolic traits was also observed in the population-based samples. 68

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71	Key terms:	Transplantation,	metabolic s	yndrome after	transplantation,	genetic polymorphisms.
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# 75 Introduction

The introduction of calcineurin inhibitors (CNIs) - cyclosporine (CSA) and tacrolimus (TAC) - in solid organ transplantation (SOT) has reduced the incidence of acute rejection episodes and improved short-term graft survival<sup>1, 2</sup>. However, both drugs are not devoid of metabolic complications, such as glucose intolerance, hypertension, and hyperlipidemia which can be very pronounced and have a detrimental impact on patients' life quality, and increase the mortality risk due to cardiovascular events<sup>3</sup>. Indeed, cardiovascular disease is responsible for approximately 20-40% of non-graft-related death after the first post-transplantation year<sup>4-6</sup>.

83 New onset diabetes mellitus after transplantation (NODAT) is a serious complication partly related to the use of CNIs, mainly TAC<sup>7, 8</sup>. It is also associated with increased cardiovascular events, infectious 84 complications, and graft loss<sup>9, 10</sup>. There are several risk factors which increase the risk of NODAT in 85 transplantation such as obesity, increased age, male sex, deceased donor, hepatitis C (HCV) status, 86 acute rejection, and African-American or Hispanic descent<sup>9, 11</sup>. The identification of genetic factors 87 involved in the development of NODAT may greatly benefit from a more detailed understanding of 88 the complex metabolic pathways involved in glucose metabolism. Genome-wide association studies 89 (GWAS) conducted to date explain only 10% of type 2 diabetes heritability and more diabetic 90 susceptibility genes remain to be discovered (reviewed in <sup>12</sup> and <sup>13</sup>). A recent GWAS investigating 91 NODAT identified eight polymorphisms<sup>14</sup>. However, only one of these was replicated in another 92 study<sup>15</sup>. Whereas GWAS have been extremely valuable, other approaches are needed to further 93 understand the pathophysiology of NODAT. 94

95 The cAMP-regulated transcriptional coactivator 2 (CRTC2), a transcriptional coactivator that 96 promotes the transcription of genes targeted by the cAMP response element-binding protein<sup>16</sup>, is an 97 interesting target protein in glucose metabolism. CRTC2 belongs to the CRTC family which 98 comprises 2 other members, CRTC1 and CRTC3<sup>17</sup>. CRTC1 is mainly expressed in the central nervous 99 system and we recently showed an association between *CRTC1* polymorphisms and obesity markers 100 (body mass index (BMI) and fat mass) in psychiatric and population-based samples<sup>18</sup>, CRTC2 is

highly expressed in the thymus, and present in both T and B-lymphocytes<sup>17</sup>. It is also expressed in the 101 liver and it plays a direct role during the fasting state in the induction of gluconeogenic genes<sup>16</sup>, It 102 103 further enhances hepatic insulin signalling by stimulating expression of the *insulin receptor substrate* 2 gene, thus triggering a feedback response that limits glucose output from the liver during fasting<sup>19</sup>, 104 Calcineurin, the target of both TAC and CSA, plays an important role in the activation of CRTC2<sup>20, 21</sup>. 105 106 Overexpression of CRTC2 induced by a mutation at 2 regulatory sites rendered *CRTC2* constitutively 107 active in an animal model and permitted CRTC2-target gene activation even when calcineurin was inhibited by CNIs<sup>22</sup>. The authors of this study also showed that CRTC2 is required for  $\beta$ -cell function 108 and proliferation and promoting this pathway could ameliorate symptoms of NODAT<sup>22</sup>. 109

CRTC2 single nucleotide polymorphisms (SNPs) were previously investigated in two Asian 110 populations and one coding SNP (R379C), with a very low minor allele frequency (MAF), was 111 associated with type II diabetes<sup>23</sup> and with lung cancer<sup>24</sup>. So far CRTC2 SNPs have not been 112 investigated in other ethnic populations or with other phenotypes. In this work, we aimed to study the 113 influence of CRTC2 SNPs on the incidence of NODAT in a sample of Caucasian SOT recipients, and 114 115 positive results were then tested for replication in the Swiss transplant cohort Study (STCS). We also aimed to extend our analysis to obesity and other variables of the metabolic syndrome (MetS) 116 following SOT. Finally, we aimed to test if the associations with MetS components could be found in 117 118 general population-based samples (>100,000 subjects).

119

#### 120 Materials and Methods

# 121 Discovery study sample:

This study aimed initially to investigate the effect of genetic polymorphisms of drug metabolizing enzymes and/or transporters on the incidence of different post-transplant complications and on immunosuppressive doses and blood levels<sup>25, 26</sup>. For this study, a total of 197 patients were enrolled between 2003 and 2005 from the outpatient clinic of the Transplant Center of the University Hospital of Lausanne, Switzerland. Patients with functional graft for more than 12 months after transplantation were eligible to participate in the study. Data regarding patient's age, gender, BMI, ethnic origin, donor's age, HLA mismatch, duration of graft cold ischemia and delayed graft function were collected retrospectively from patients' medical files. Immunosuppressive regimens, doses and blood levels were obtained retrospectively at different time-points during the first year post-transplantation. The study was approved by the ethics committee of the University of Lausanne. Patients gave their written informed consent to participate in this pharmacogenetic study. Once included, venous blood samples were collected for DNA extraction and further genotyping analysis.

For the present study, data were collected between October 2011 and April 2012 as related to the 134 135 development of NODAT during the first 5 years following transplantation. Data regarding fasting blood glucose (FBG), glycated haemoglobin (HbA1c), 2hrs oral glucose tolerance test (OGTT), 136 137 insulin and oral anti-diabetic treatment were collected retrospectively from the medical files at the 138 time of transplantation, at 1, 3, 6, 9 and 12 month post-transplantation and at the yearly follow-up until 139 5-year post-transplantation. NODAT was diagnosed if a patient needed anti-diabetic treatment (either 140 insulin or oral anti-diabetic agents) for at least 6 months following transplantation or had several abnormal glucose profiles during the follow-up period that fulfil the criteria given by the WHO and 141 ADA consensuses<sup>27</sup>, including FBG $\geq$ 7.0 mmol/l (in  $\geq$  two occasions) or 2 hours plasma glucose $\geq$ 11.1 142 mmol/l during OGTT <sup>27</sup> (even if not treated with anti-diabetic drugs). Patients who had diabetes or 143 144 prediabetes before transplantation or were transplanted because of diabetic nephropathy were excluded 145 from the present study.

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#### 147 The Swiss Transplant Cohort Study (STCS):

148 The STCS is an ongoing prospective multicenter cohort project (Basel, Bern, Genève, Lausanne, St. Gallen and Zurich), aiming at a nationwide comprehensive and structured data collection in all SOT 149 150 recipients. All recipients of SOTs in Switzerland are prospectively registered since May 2008. 151 Currently more than 3500 patients are included in the STCS. No particular eligibility or exclusion 152 criteria exist for enrolment. After transplantation, all patients are mandatorily followed in their 153 respective transplant centers. After baseline assessment, STCS follow-up assessments take place at 6-, 12 months and yearly thereafter. Biological samples are collected in relation to the case at baseline, 154 and at 6-, and the 12-month visits. Data regarding patient's age, gender, BMI, blood pressure and lipid 155 profiles (total cholesterol, HDL- and LDL-cholesterol) were available for all patients at the different 156

time-points of the STCS follow-up. Full description of this cohort is published elsewhere<sup>28, 29</sup>. For the 157 first STCS replication sample, patients transplanted from May 2008 to 8th of May 2011 with a 158 159 functional graft for at least 12 months after transplantation were included in the analyses (n=1294). 160 NODAT was diagnosed if a patient needed anti-diabetic treatments following transplantation or if such new metabolic event was reported in their case report forms. For the second STCS replication 161 sample, patients transplanted from 9<sup>th</sup> of May 2011 to May 2013 with a functional graft for at least 12 162 163 months after transplantation were included in the analyses (n=759). For this second STCS replication 164 sample, several random and FBG were available in the database, as well as HbA1c. Therefore, as for the discovery sample, NODAT was diagnosed if a patient needed anti-diabetic treatment (either 165 166 insulin or oral anti-diabetic agents) or fulfilled the criteria given by the WHO and ADA consensuses mentioned earlier. For both STCS samples, patients who were diagnosed with diabetes or were 167 168 prediabetics before transplantation or were transplanted because of diabetic nephropathy were excluded from the NODAT analyses. 169

For the combined STCS sample, new-onset hypertension and new-onset hyperlipidemia were diagnosed if patients needed anti-hypertensive and hypolipidemic treatment post-SOT. Patients with previous hypertension or hyperlipidemia were excluded from the new-onset hypertension and newonset hyperlipidemia analyses, respectively. The study was approved by the ethics committee of their respective centers. Patients gave their written informed consent to participate in this pharmacogenetic study.

Recipients younger than 18 years old and recipients with multiple organ transplantation were excluded
from the whole analyses. If a patient received more than one transplantation during the inclusion
period, only data from the first SOT was included in the analyses.

For both the discovery and STCS samples, as abdominal obesity (waist circumference) was notavailable, BMI was used as a marker of obesity.

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#### **182 Population based samples:**

We aimed to replicate results with MetS traits in several population-based samples: the Meta-Analyses
 of Glucose and Insulin-related traits Consortium (MAGIC)(n=46'186)<sup>30</sup>, Genetic Investigation of

ANtropometric Traits (GIANT) consortium (n=123'865)<sup>31-33</sup>, and Genome Wide Associations Scans
 for Total Cholesterol, HDL-C, LDL-C and triglycerides (n>100'000)<sup>34, 35</sup>. More details are in the
 Supplementary Material.

188

# **189 Polymorphism selection and genotyping:**

190 Two tagging SNPs within *CRTC2* gene (rs8450G>A and rs12117078G>C) were selected using 191 HapMap Genome Browser by limiting the search to SNPs with a MAF>5% in the Caucasian 192 population and  $r^2$  cutoff of 0.8. Genotyping in the discovery study sample was done using Taqman 193 allelic discrimination assays. Genetic analyses for the STCS replication samples were done using the 194 fluorescence-based competitive allele-specific PCR technology (KASPar)<sup>36</sup>. More details are in the 195 Supplementary Material.

196

# 197 Statistical analysis:

198 Detailed statistical methods are presented in the Supplementary Material.

199 Discovery and STCS samples:

We assessed the association of *CRTC2* SNPs with MetS traits 1) by applying a logistic regression for binary outcomes adjusted for recipient's age at transplantation and sex. Other variables identified through the univariate analysis (P<0.10) were also added as covariates in these models, 2) by using linear mixed models for continuous variables, and 3) by fitting a Generalized Additive Mixed Model (GAMM) to include a smooth trend for the response in time (allowing multiple observations for each patient) for non-linear continuous variables (BMI and FBG).

206 For the discovery sample, the p-values of the models were adjusted for multiple comparisons using

- 207 Bonferroni correction and the SNP that survived this correction was investigated in the STCS.
- 208 Data for both the discovery and replications samples were analyzed using Stata 12 (StataCorp, College
- 209 Station TX, USA) and R version 2.13.0 software (http://www.R-project.org).
- 210 *Population-based samples:*

We analyzed the associations of *CRTC2* SNP with different MetS traits using multivariate linear regression with allele dosage in which potential confounding factors such as age, sex, and smoking status were added as covariates.

214

#### 215 **Results:**

# 216 Discovery and STCS replication samples:

General characteristics of the discovery and STCS replication samples are presented in the Supplementary Material and Supplementary Tables 1 and 2. Supplementary Table 2 represents the global STCS sample without the exclusion of patients with previous history of diabetes or other metabolic abnormalities before transplantation.

The distribution of *CRTC2* genotypes in all studies did not deviate from the Hardy-Weinberg
equilibrium (P>0.05) and the MAFs were similar to those reported in HapMap (Supplementary Table
3).

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# 225 *CRTC2 SNPs* and NODAT, FBG, and BMI in the discovery study sample:

Carriers of the *CRTC2 rs8450-AA* genotype showed increased risk of NODAT as compared to wildtype *GG* genotype (odd ratio (OR)=6.91, 95%CI:1.52-31.36,  $p_{corrected}$ =0.02). An increased risk was also observed by analyzing the *rs8450* SNP in a recessive model (Table 1). Neither cumulative prednisone dosages nor calcineurin inhibitors' doses, trough levels or concentration/dose ratios tested at 2 time periods (at 1 month or at 12 months post-SOT) were associated with NODAT in the univariate analysis or when used in the multivariate model including *CRTC2* SNPs and the other covariables (data not shown).

FBG were also available at several time-points after SOT. By applying the GAMM model, increased FBG levels were found in patients with the *CRTC2 rs8450-GA and AA* genotypes (0.24 mmol/L,  $p_{corrected}=0.06$  and 0.47 mmol/L,  $p_{corrected}=0.02$ , respectively) as compared to wild-type *GG* genotype. We observed an equal increased of FBG levels by analyzing the *rs8450* SNP in dominant model (Table 1). Patients with the *CRTC2 rs8450-GA* genotype showed a 1.36 kg/m2 increased BMI compared to the wild-type genotype ( $p_{corrected}=0.02$ ). By analyzing the SNP in a dominant model, we observed a significant association between *CRTC2 rs8450* and BMI ( $\beta=1.17, 95\%$ CI:0.17-2.35,  $p_{corrected}=0.04$ ). We observed no association between *CRTC2 rs12117078G>C* SNP and NODAT, FBG or BMI (Table 1).

#### 243 *CRTC2 SNP* in the STCS replication samples:

The *CRTC2 rs8450G>A* SNP that survived corrections for multiple testing in the discovery study sample was investigated in the STCS. Recessive models for this SNP are presented in the text and in Tables 2 and 3 while additive and dominant models regarding NODAT are presented in Supplementary Table 4.

248

# 249 *CRTC2 SNP* and NODAT:

In the univariate analyses, increased recipient's age (p=0.0003), male sex (p=0.03), positive HCV 250 251 status (p=0.003), deceased donor (p=0.04), treatment with TAC (p=0.001) and baseline BMI (p=0.0002) were among the non-genetic factors associated with an increased risk of NODAT and were 252 used as covariates in the genetic models. In the first STCS replication samples, we observed a non-253 significant association between rs8450G>A and NODAT (Table 2). However, by analyzing the SNP 254 255 in the subgroup of patients having received SOT from a deceased donor and treated with TAC, a significant effect of the SNP was observed (n=281, OR=2.36, 95%CI:1.16-4.78, p=0.01)(Table 2). In 256 the second replication sample (n=438), in which the diagnostic criteria for NODAT was the same as 257 for the discovery sample, recipients carrying the rs8450-AA genotype showed a significant increased 258 risk of NODAT (n=438, OR=2.01, 95%CI:1.03-3.91, p=0.04). In the combined STCS sample, the 259 effect of the CRTC2 genotype was only observed in the subgroup of patients having received SOT 260 from a deceased donor and treated with TAC (n=395, OR=2.08, 95% CI:1.15-3.74, p=0.02) (Table 2). 261

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# 265 *CRTC2 SNP* and BMI, new-onset hypertension/ hyperlipidemia:

We investigated the influence of *CRTC2* rs8450G>A on BMI and on the incidence of new-onset hypertension/hyperlipidemia in the combined STCS sample.

Increased recipient's age (p<0.0001), male sex (p<0.0001) and type of transplanted organ (p<0.0001) 268 were among the non-genetic factors associated with increased BMI in the univariate analyses and were 269 used as covariates in the genetic models. BMI values were also dichotomized into normal versus 270 overweight or obese (BMI≥25 kg/m<sup>2</sup>). At 12-months post-SOT, overweight or obese patients were 271 more prevalent in the CRTC2 rs8450-AA genotype compared to the other genotypes (n=1215, 272 273 OR=1.56, 95%CI:1.08-2.25, p=0.02). We observed the same association but with lower significance 274 for the combined STCS sample at 6-months post-SOT (n=1389, OR=1.41, 95%CI:1.01-1.97, p=0.04), 275 and a non-significant association with baseline BMI (n=1515, OR=1.20, 95%CI:0.87-1.65, p=0.27), 276 suggesting that time post-SOT modulates the effect of the CRTC2 genotypes.

By applying the GAMM model to test the association between BMI over time and *CRTC2 rs8450G>A* SNP (n=1625), patients with the *CRTC2 rs8450-AA* genotype showed a 0.47 kg/m<sup>2</sup> (95%CI:0.11-1.02, p=0.01) increase in BMI compared to the other genotypes.

280 We found no association between the CRTC2 rs8450G>A and the incidence of new-onset 281 hypertension (Supplementary Material). On the other hand, carriers of the CRTC2 rs8450-AA 282 genotype showed increased risk of new-onset hyperlipidemia as compared to the other genotypes (n=1007, OR=1.76, 95%CI:1.16-2.66, p=0.007). Additionally, patients with the CRTC2 rs8450-AA 283 284 genotype also showed a significant decrease of HDL-Cholesterol levels in the combined STCS sample 285 (n=1214,  $\beta$ = -0.08 mmol/L, p=0.001). CRTC2 rs8540G>A SNP was not associated with total cholesterol or with LDL-cholesterol blood levels in the combined STCS sample (data not shown). 286 More details can be found in the Supplementary Material. 287

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#### 289 *CRTC2 SNP* and MetS traits in kidney and non-kidney transplant recipients:

Patients were dichotomized into kidney versus non-kidney transplant recipients (Table 3). The *CRTC2 rs8450-AA* genotype was associated with significantly higher OR for NODAT in non-kidney

transplant recipients (n=409, OR=2.09, 95%CI:1.13-3.86, p=0.02), while the *rs8450 AA*-genotype was
only associated with higher risk of new-onset hyperlipidemia in kidney transplant recipients (n=573,
OR=1.95, 95%CI:1.14-3.15, p=0.01).

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#### 296 **Population-based samples:**

Several glycemic traits were available in the MAGIC study (Supplementary Table 5); each rs1572788*C-allele* (in complete linkage disequilibrium (LD) with rs8450 *A-allele*) increased FBG levels by 0.01 mmol/l (n=46'186, p=0.004). Additionally, each rs1572788 *C-allele* decreased HOMA-B by 0.008% (n=46'186, p=0.03). No association was observed between rs1572788T>C and HOMA-IR or the 2 hour OGTT.

Decreased HDL-cholesterol and increased triglycerides were observed for each *rs1572788 C-allele* in the Genome Wide Associations Scans for Total Cholesterol, HDL-C, LDL-C and triglycerides (n=96'908,  $\beta$ =-0.008, p=0.013 and n=93'562,  $\beta$ =0.009, p=0.004, respectively). We observed nonsignificant associations for *rs1572788T*>*C* with total and LDL-cholesterol, as well as with obesity traits studied in the GIANT study, however, the direction of the effect is consistent with increased blood lipids and obesity with each *rs1572788 C-allele* (Supplementary Table 5).

# 308 Discussions:

The transcriptional co-activator (CRTC) family is implicated with energy expenditure (CRTC1 and 309 CRTC3)<sup>37, 38</sup> and glucose metabolism (CRTC2)<sup>16</sup> in animal models. In the present study, we aimed to 310 311 study the effect of CRTC2 SNPs on the development of diabetes in SOT recipients, a population at 312 high risk of developing metabolic abnormalities, and in population-based samples. We also extended 313 our analyses to investigate obesity and other metabolic traits. The CRTC2 rs8450-AA genotype was associated with increased risk of NODAT, increased FBG, and BMI in a sample of 156 patients with 314 315 SOT. Results on NODAT were replicated only in the second STCS replication sample (n=438), while 316 in the first and the combined STCS replication samples, the SNP only showed significant influence in the population at higher risk of developing NODAT, mainly in recipients having received a graft from 317 a deceased donor and treated with TAC. We also observed a significant association for BMI in the 318

STCS replication sample; overweight and obesity were also more frequent in patients carrying the *rs8450-AA* genotype at 12 months post-SOT. *CRTC2 rs8450G>A* was also associated with increased risk of new-onset hyperlipidemia following SOT and with lower HDL-cholesterol blood levels. Furthermore, we also found the increased risk of the *C-allele* of *rs1572788* (in complete LD with *rs8450 A-allele*) on diabetic and lipidemic traits in two large population-based samples (MAGIC and GWAS for Total Cholesterol, HDL-C, LDL-C and triglycerides, respectively), and a consistent direction even though non-significant between *rs1572788T>C* and obesity traits in the GIANT study.

326 *CRTC2 rs8450G>A* was associated with NODAT in the total discovery sample and only in the second 327 STCS replication sample, but not in the total STCS sample in which a significant association was 328 observed only in the subgroup of patients at higher risk of developing NODAT. This could partly be 329 explained by the difference of NODAT diagnostic criteria, which is only based on anti-diabetic 330 treatment in the first STCS replication sample. Another explanation could be the high percentage of 331 patients receiving an organ from a living donor in the first STCS replication sample (28%, 332 Supplementary Table 2) compared to only 11.5% in the discovery sample (Supplementary Table 1). Deceased donor is a common risk factor of NODAT<sup>11</sup> and the impact of the SNP seems to be more 333 prominent in the groups at higher risk of developing MetS traits. Additionally, when differentiating 334 335 kidney and non-kidney transplant recipients (Table 3), we observed a significant association between 336 the CRTC2 rs8450-AA genotype and NODAT in non-kidney transplant recipients. The percentage of 337 non-kidney transplant recipients was higher in the second STCS replication sample compared to the first STCS sample (Supplementary Table 2) and it should be noted that kidney transplant recipients 338 had higher percentages of living donors compared to non-kidney transplant recipients (43% vs. 2% 339 340 respectively). Moreover, kidney transplant recipients usually have obesity and other metabolic 341 problems already before transplantation compared to other solid organ recipients, which could explain 342 the differential effect of the SNP depending on the type of the transplanted organ. Interestingly, in 343 both the discovery and the combined STCS samples, the effect of the SNP was stronger in NODAT 344 cases developed after the first year post-SOT (data not shown), It maybe that the effect of the CRTC2 345 SNP on NODAT is more important in the long-term, but this remains to be further studied.

346 In the GWAS literature, CRTC2 SNPs were not found to be associated with diabetes, obesity or 347 hyperlipidemia. On the other hand, by testing the individual effect of CRTC2 rs1572788T > C SNP on 348 diabetic and lipidemic traits in two population-based samples, a significant effect was observed, even 349 though this effect is very weak and the variance explained by the SNP is very small (Supplementary Table 5) and could thus not be detected by GWAS. However, patients with SOT is a population at risk 350 351 of developing MetS traits, because of the disease itself and/or the immunosuppressive medications, 352 both can act as an important trigger, unmasking different genetic factors. So in this at risk population, the effect of CRTC2 SNP was more pronounced, even in much smaller sample sizes compared to 353 population-based samples. Additionally, in the GWAS of NODAT<sup>14</sup>, CRTC2 SNPs did not reach 354 genome-wide level of significance. Moreover, most of the SNPs retained in this GWAS were not 355 replicated in another study with NODAT<sup>15</sup>. In fact, both GWAS and candidate gene studies are 356 important gene association approaches and both are subject to the same artifacts of spurious 357 358 association findings. Depending on the methodology of these studies (sample size, definition of the 359 phenotypes, MAF of the SNPs, corrections for multiple testing, etc...), discordances between both 360 approaches could be seen. GWAS relies on indirect association to locate a disease-causing variant 361 while candidate gene approach relies on an a priori hypothesis to identify this disease-causing variant. 362 For all of these reasons, replication of positive finding in either approach is of utmost importance to 363 validate the results.

The functional activity of this SNP, in the regulatory 3'UTR, is unknown and data from 1000 364 Genomes<sup>39</sup> showed that 78 SNPs in the *CRTC2* gene region are in high LD ( $r^2$  threshold $\geq 0.8$ ) with 365 rs8450G>A. We used the RegulomeDB database that annotates SNPs with known and predicted 366 regulatory elements in the intergenic regions of the H. sapiens genome<sup>40</sup>. This database reveals known 367 and predicted regulatory DNA elements including regions of DNAase hypersensitivity, binding sites 368 369 of transcription factors, and promoter regions that have been biochemically characterized to regulation 370 transcription. Source of these data include public datasets from GEO, the ENCODE project, and published literature. Regarding CRTC2 rs8450G>A, several of these proxy SNPs (e.g. rs6680140 with 371  $r^2=1$ ) show cis-eQTL effect on the expression of *CRTC2* and fall into the autoimmune regulatory 372

(AIRE) motif, influence the binding of several proteins (*JUN*, *CREBBP* and *ELF1*), and has several
histone marks (H3k09me3, H4k20me1, etc.). Altogether, these data suggest that the SNP might have a
regulatory function.

376 Interestingly, among the analyzed clinical factors, treatment with corticosteroids was not associated 377 with NODAT, neither in the discovery cohort nor in the STCS sample. However, in our samples most 378 of the patients were treated with corticosteroids so it is difficult to draw major conclusions. On the 379 other hand, the cumulative corticosteroids dosages and the duration of therapy rather than the simple 380 corticosteroids administration could be the most influencing factors. In our discovery sample, the 381 cumulative dosages of corticosteroids at the first year post-transplantation were not associated with 382 NODAT outcome. In the literature, several studies did not find any influence of cumulative corticosteroid dosages on the appearance of NODAT<sup>41-45</sup>. The current lower dosages of corticosteroids 383 384 used in different immunosuppressive regimens could contribute to the negative association with NODAT appearance in our study. 385

This study has several limitations and strengths. Random and FBG levels were only available in the 386 387 second STCS replication sample and the diagnosis of NODAT in the first STCS replication sample 388 was only based on anti-diabetic treatments post-SOT. The genotype of the rs8450G>A SNP of the 389 liver transplant donors is unknown, therefore the significant effect of the SNP on NODAT found in 390 non-kidney transplant recipients, in which nearly half of them are liver transplant recipients, should be 391 interpreted with caution. However, CRTC2 is expressed in different tissues and is implicated in 392 glucose regulation through different mechanisms (gluconeogenesis, insulin signaling and beta-cell survival<sup>16, 19, 22</sup>) which could highlight the importance of recipient *CRTC2* genotype even in liver 393 394 transplant recipients. Additionally, by excluding patients with liver transplantation from the nonkidney transplant recipients' analyses, a significant association between the CRTC2 SNP and NODAT 395 396 is still observed (data not shown). The present results do not allow the determination of whether the 397 rs8450G > A SNP is the causative variant or merely a proxy of one or more yet unidentified variants. Despite the possible regulatory functions proposed by the RegulomeDB database, further studies are 398 399 needed to elucidate which precise mechanisms underlie the observed association. This study included 400 people of Caucasian origin and results cannot be generalized to other ethnic groups. On the other hand, 401 the fact that the results were replicated in two independent samples with SOT and in large population-402 based samples, the latter used as a proof of concept of the effect of the polymorphism, strengthens the 403 validity of our data.

In conclusion, our results suggest that variations in CRTC2 play an important role in the high prevalence of MetS complications observed in patients with SOT. Besides, by studying CRTC2 SNPs in population-based samples, we still observed a weak yet significant association, thereby suggesting that the effect of CRTC2 variations is more important in population at risk of developing different MetS traits than in the general population. This is the first study showing the importance of CRTC2 variations with NODAT and other MetS traits in patients with SOT. The assessment of the major risk gene variants would allow predicting vulnerability for developing Mets phenotypes and thus adapt the immunosuppressive treatment from the beginning by genotyping the patients before transplantation. In the long-term, these patients could benefit from individualized immunosuppressive regimens adapted to their genetics and environment.

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441	
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#### 451 Footnote

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- 470 Supplementary data:
- 471 Supplementary information is available at journal's website

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**Table 1** Association between *CRTC2* SNPs and NODAT, blood glucose levels and BMI in patients with solid organ transplantation from the discovery study sample:

	NODAT		FBG		BMI	
	(n=156)		(n=156)		(n=156)	
SNP	OR (95% CI)	P corrected	Difference (mmol/L) compared to wild-type (ref) (95% Cl)	P corrected	Difference (kg/m2) compared to wild-type (ref) (95% Cl)	P corrected
CRTC2 rs84	50					
GG	ref		ref		ref	
GA	1.81 (0.71 - 4.61)	0.46	0.24 (-0.00 - 0.49)	0.06	1.36 (0.28- 2.42)	0.02
AA	6.91 (1.52 - 31.36)	0.02	0.47 (0.07 - 0.86)	0.02	0.37 (-1.38 - 2.06)	0.76
Dominant	2.32 (0.96 - 5.63)	0.12	0.28 (0.05 - 0.51)	0.02	1.17 (0.17 - 2.35)	0.04
Recessive	5.11 (1.24 - 21.03)	0.04	0.36 (-0.03 - 0.74)	0.06	-0.24 (-2.33 - 1.68)	0.80
CRTC2 rs12	117078					
GG	ref		ref		ref	
GC & CC	0.36 (0.11 - 1.16)	0.18	-0.07 (-0.40 - 0.24)	0.70	0.24 (-1.41 - 1.62)	0.78

NODAT: New-onset diabetes after transplantation, FBG: Fasting blood glucose, OR: odd ratio, CI: confidence interval, BMI: body mass index. Regarding NODAT and blood glucose levels analyses: P values were adjusted for age of recipient at transplantation, sex of the recipient and baseline BMI and type of calcineurin inhibitor.

Regarding BMI analyses: P values were adjusted for age of recipient at transplantation, sex of the recipient and type of calcineurin inhibitor.

**Table 2:** Association between *CRTC2 rs8450* SNP in a recessive model and NODAT in the first, second and combined population of STCS and in different subgroups at risk of NODAT:

CRTC2 rs8450		GG&GA (n)	AA	
	N total		NODAT OR (95% CI)	P value
First STCS replication sample <sup>§</sup> :				
Total	711	Ref (n=623)	1.02 (0.62 - 1.70)	0.93
Treatment with TAC & deceased donor	281	Ref (n=244)	2.36 (1.16 - 4.78)	0.01
Second STCS replication sample <sup>§</sup> :				
Total	438	Ref (n=385)	2.01 (1.03 - 3.91)	0.04
Combined STCS replication sample <sup>§</sup> :				
Total	1149	Ref (n=1008)	1.25 (0.84 - 1.87)	0.26
Treatment with TAC & deceased donor	395	Ref (n=340)	2.08 (1.15 - 3.74)	0.02

<sup>\$</sup>Excluding patients with previous history of diabetes before transplantation

NODAT: New-onset diabetes after transplantation, TAC: tacrolimus, OR: odd ratio, CI: confidence interval

P values were adjusted (when appropriate) for age of recipient at transplantation, sex of the recipient,

hepatitis C status, baseline BMI, type of calcineurin inhibitor and type of donor.

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 Table 3: Association between CRTC2 rs8450 SNP and MetS traits in all kidney and non-kidney transplant

 recipients in the combined STCS sample

	Kic	Iney transplant recip	ients	Nor	n-kidney transplant rec	ipients
	n	OR	p-value	n	OR	p-value
	(95	% CI) for rs8450-AA			(95% CI) for rs8450-AA	
NODAT	740	0.87 (0.50 - 1.52)	0.62	409	2.09 (1.13 - 3.86)	0.02
BMI≥25 kg/m2 at 12months post SOT	761	1.59 (0.98 - 2.58)	0.06	454	1.52 (0.84 - 2.75)	0.17
New-onset hyperlipidemia	573	1.95 (1.14 - 3.15)	0.01	434	1.73 (0.84 - 3.56)	0.14
New-onset hypertension	146	1.64 (0.43 - 6.30)	0.47	345	1.44 (0.75 - 2.78)	0.27

OR: odd ratio, CI: confidence interval

NODAT: New-onset diabetes after transplantation, BMI: body mass index

For NODAT analyses: P values were adjusted for age of recipient at transplantation, sex of the recipient,

baseline BMI, type of calcineurin inhibitor and type of donor.

For BMI analyses: P values were adjusted for age of recipient at transplantation and sex of the recipient.

For new-onset hyperlipidemia analyses: P values were adjusted for age of recipient at transplantation, sex of

the recipient, baseline BMI, the type of calcineurin inhibitor and treatment with corticosteroids.

For new-onset hypertension analyses: P values were adjusted for age of recipient at transplantation and sex of the recipient, treatment with corticosteroids, the type of calcineurin inhibitor and type of donor.

1	Supplementary data:
2	Materials and Methods
3	- Population based samples:
4	> The Meta-Analyses of Glucose and Insulin-related traits Consortium (MAGIC)
5	<ul> <li>Genetic Investigation of ANtropometric Traits (GIANT) consortium</li> </ul>
6	> Genome Wide Associations Scans for Total Cholesterol, HDL-C, LDL-C and
7	triglycerides
8	- Polymorphism selection and genotyping
9	- Statistical analysis
10	
11	Results:
12	- General characteristics of the discovery sample
13	- General characteristics of the replication sample (STCS):
14 15	- CRTC2 SNP and new-onset hypertension in the STCS
16 17	- <i>CRTC2</i> SNP and new-onset hyperlipidemia in the STCS
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19	Supplementary Tables 1-5
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# 27 Materials and Methods

#### 28 **Population based samples:**

# 29 The Meta-Analyses of Glucose and Insulin-related traits Consortium (MAGIC)

MAGIC is a large-scale meta-analyses of genome-wide data for continuous diabetes-related traits in participants without diabetes <sup>1</sup>. Meta-analyses of ~2.5 million directly genotyped or imputed autosomal single nucleotide polymorphisms (SNPs) were performed from genome-wide association studies (GWAS). These cohorts include up to 46'186 non-diabetic participants of European descent informative for FBG, the surrogate estimates of beta-cell function (HOMA-B) and insulin resistance (HOMA-IR) derived from fasting variables by homeostasis model assessment and up to 15'234 nondiabetic individuals informative for 2 hour oral glucose tolerance test (OGTT)<sup>2</sup>.

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# 38 Genetic Investigation of ANtropometric Traits (GIANT) consortium

39 The GIANT consortium performed a meta-analysis of GWAS data with a discovery set of 123'865

40 individuals of European ancestry from 46 studies for height  $^3$ , BMI $^4$  and waist-to hip ratio  $^5$ .

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#### 42 Genome Wide Associations Scans for Total Cholesterol, HDL-C, LDL-C and triglycerides

43 Data on lipid traits have been downloaded from "Genome Wide Associations Scans for Total 44 Cholesterol, HDL-C, LDL-C and triglycerides" website <sup>6, 7</sup> which is a meta-analysis of 46 lipid 45 GWASs. These studies together comprise >100,000 individuals of European descent (maximum 46 sample size 100,184 for Total Cholesterol, 95,454 for LDL-C, 99,900 for HDL-C and 96,598 for 47 triglycerides), ascertained in the United States, Europe or Australia.

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# 55 **Polymorphism selection and genotyping:**

Polymorphisms within *CRTC2* were selected using HapMap Genome Browser (release 28)<sup>8</sup>. Two tagging SNPs were found by limiting the search to SNPs within CRTC2 gene  $\pm$  1kb with a minor allele frequency > 5% in the Caucasian population and  $r^2$  cutoff of 0.8: *rs8450G*>A within the 3'UTR and the intronic *rs12117078G*>C SNP.

Genotyping of these 2 SNPS for the discovery study sample was done using Taqman allelic discrimination assays (ABI PRISM 7000 Sequence Detection System; Applied Biosystems, Rotkreuz, Switzerland) after genomic DNA extraction from whole blood. Life technologies Taqman genotyping assays C\_8722376\_10 and C\_30997814\_10 were used for *CRTC2 rs8450G>A* and *rs12117078G>C SNPs*, respectively. Genotyping were performed in the Unit of Pharmacogenetics and Clinical Psychopharmacology, in Lausanne, Switzerland.

Genetic analyses for the first STCS replication sample were performed by KBioscience Institute in
United Kingdom using the novel fluorescence-based competitive allele-specific PCR technology
(KASP<sup>TM</sup>). Details about this technology are available at
<u>http://www.lgcgenomics.com/genotyping/kasp-genotyping-chemistry/</u>.

Genetic analyses for the second STCS replication sample were also performed using the KASP<sup>TM</sup> 70 71 technology on ABI 7500 Fast real-time thermocycler (Applied Biosystems). Genotyping were 72 performed in the service of infectious disease in Lausanne University Hospital, Lausanne, 73 Switzerland, according to manufacturer's protocols (LGC Genomics, UK). The KASP primers were designed by Kraken<sup>TM</sup> assay design and workflow management software and further validated by 74 manufacturer (LGC Genomics, UK). Automated allele calling was performed using SDS software 75 76 according to standard protocols (Applied Biosystems). The following primers were used for CRTC2 77 rs8450G > A genotyping in this sample: primer allele FAM: ACCCCTAGGCATCCGGAAAAG, 78 primer allele HEX: CACCCCTAGGCATCCGGAAAAA and common primer 79 GGGTAGAGGGGGGGGCCCTGGAA.

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# 83 Statistical analysis:

Quantitative data are presented as median and range unless otherwise mentioned, while qualitative data are presented as frequency and percentage. For association studies, the chi-square (Chi<sup>2</sup>) test or the Fisher exact test for binomial variables were used. Differences in allele and genotype frequencies as well as deviation from Hardy-Weinberg equilibrium were assessed using Chi<sup>2</sup> test.

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## 89 Discovery study sample:

For NODAT analyses, logistic regression was applied adjusted for recipient's age at transplantation, 90 91 sex, BMI at baseline and type of CNIs. Due to nonlinearity of our models and the absence of any linear transformation, the association between CRTC2 SNPs with BMI and FBG levels was assessed 92 by fitting a Generalized Additive Mixed Model (GAMM)<sup>9, 10</sup> to allow a smooth trend for the response 93 in time based on multiple observations for each patient (using a thin plate regression spline basis) 94 adjusting for recipient's age at transplantation, sex, and type of CNIs (antidiabetic treatments for FBG 95 analyses). GAMMs were fitted using the mgcv package of R (settings were fixed at package defaults). 96 97 In order to be more conservative, the uncertainty of estimated parameters were assessed by 1000 bootstraps<sup>11</sup> at the subject level and results were similar with those gained by 10'000 bootstraps. 98 Due to the small number of subjects being homozygous for the variant allele of CRTC2 99

100 rs12117078G>C, this SNP was analyzed in a dominant model.

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# 102 **Replication samples (STCS):**

For the statistical analyses of NODAT, new-onset hypertension, new-onset hyperlipidemia, and overweight or obese, logistic regression models were applied adjusted for recipient's age at transplantation and sex. Other variables identified through the univariate analysis (P<0.10) were also added as covariates in these models. GAMM models were applied for BMI and HDL-cholesterol analyses in the STCS sample, as both models were not linear. Linear mixed models were used for systolic and diastolic blood pressure analyses and for blood total cholesterol and LDL-cholesterol analyses. In addition to recipient's age at transplantation, sex, BMI at baseline and type of CNIs, blood pressure models and lipid models were also adjusted for antihypertensive and hypolipidemic drugsintake, respectively.

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# 113 **Population-based samples:**

The associations of *CRTC2* SNP with different MetS traits were analyzed using multivariate linear regression with allele dosage in which potential confounding factors such as age, sex and smoking status were added as covariates. FBG, OGTT, the surrogate estimates of beta-cell function (HOMA-B) and insulin resistance (HOMA-IR) were analyzed in the MAGIC study. BMI, waist circumference and waist-to hip ratio were the only adiposity traits analyzed by the GIANT Consortium. Triglycerides, total cholesterol, HDL-cholesterol and LDL-cholesterol were analyzed in the "Genome Wide Associations Scans for Total Cholesterol, HDL-C, LDL-C and triglycerides" study.

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#### 122 **Results:**

# 123 General characteristics of the discovery sample

124 Among the 197 patients included in the initial study, 17 patients were excluded because they were diagnosed with diabetes before transplantation, 11 patients had no follow-up data in their medical 125 126 files, 2 patients did not sign consents for MetS-related genetic analysis, 4 patients dropped-out from 127 the study and 7 were non-Caucasians. Finally, 156 Caucasian patients were included in the current 128 study, 65% were kidney transplant recipients. The mean age at transplantation was  $47.8 \pm 11$  years, 61% were male and the mean BMI pre-transplant was  $24 \pm 4$  kg/m<sup>2</sup>. During the five years of follow-up 129 post-transplantation, NODAT occurred in 45 (28.9%) patients. Most NODAT cases (67%) occurred 130 during the 1<sup>st</sup> year after transplantation. Most of the patients were treated with CSA (n=102, 65%), 131 132 while 35% (n=54) were treated with TAC. As expected, NODAT was higher in the TAC-treated group compared to CSA<sup>12</sup> (46.3 vs. 19.6 % respectively). Full characteristics of our population and their 133 134 distribution between NODAT and non-NODAT patients are presented in Supplementary Table 1. Recipient age (p=0.002), male sex (p=0.05), donor age (p=0.04), baseline BMI (p=0.0003) and the 135 136 type of CNIs (p<0.001) were significantly different between patients with and without NODAT (Supplementary Table 1). 137

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# 139 General characteristics of the replication sample (STCS):

140 Overall, 1294 patients from the first STCS replication sample received a SOT from May 2008 until 141 May 2011, among them 1219 patients had both the clinical data and the genetic material available and were included in the current study. By excluding patients younger than 18 years old and patients with 142 multiple organ transplantation, 958 patients remained and included in the analysis. The mean age at 143 144 transplantation was 52.3  $\pm$  13 years, 66% were male and pre-transplant BMI was 24.6 kg/m<sup>2</sup> (range:13.7-41.2). During the follow-up period post-transplantation, NODAT, new-onset HTN and 145 new-onset hyperlipidemia occurred in 27.4%, 67.6% and 33.1% of the patients, respectively. Among 146 the patients included in the second STCS replication sample (n=759) with both clinical and genetic 147 148 data and by excluding patients younger than 18 years old and patients with multiple organ 149 transplantation, 667 remained and were included in the analysis. Lower incidence of NODAT, new-150 onset HTN and new-onset hyperlipidemia were observed in this sample, which could be explained by 151 the shorter follow-up duration compared to the first STCS replication sample (Supplementary Table 152 2). Full description of the first, second and combined STCS samples are presented in Supplementary Table 2. All patients from both STCS samples were of Caucasian origin. 153

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# 155 *CRTC2 SNP* and new-onset hypertension in the STCS:

Among the non-genetic factors, type of calcineurin inhibitor (p<0.0001), type of transplanted organ (p<0.0001), treatment with corticosteroids (p<0.0001) and type of donor (p=0.02) were associated with increased risk of new-onset hypertension after SOT in the univariate analyses and were used as covariates in the genetic models. A non-significant association was observed between *CRTC2 rs8450G>A* and new-onset hypertension, except for patients carrying the *rs8450-GA* genotype which showed a protective effect against new-onset hypertension (n=491, OR=0.62, 95%CI: 0.42 - 0.94, p=0.02) compared to wild type genotype.

163 *CRTC2 rs8450G>A* SNP was not associated with systolic or diastolic blood pressure levels (data not
164 shown).

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167	CRTC2 SNP and new-onset hyperlipidemia in the STCS:
168	Among the non-genetic factors, increased recipient's age (p=0.003), baseline BMI (p=0.006), type of
169	calcineurin inhibitor (p=0.005) and treatment with corticosteroids (p<0.0001) were associated with
170	increased risk of new-onset hyperlipidemia after SOT in the univariate analyses and were used as
171	covariates in the genetic models.
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Characteristic	Total	Non-NODAT	NODAT	p-value
Recipient age at transplantation (years), median (range)	48 (22-68)	46 (22-68)	53 (28-68)	0.002
Recipients sex (Males) [%]	60.9	55.9	73.3	0.05
Living donor [%]	11.5	11.7	11.1	0.92
Donor age (years), median (range)	43.5 (10-73)	41 (10-73)	48 (13-69)	0.04
Donor sex (Males) [%]	56.8	63.2	54.8	0.06
Organ, n [%]				
lung	17 (10.9)	10 (9.0)	7 (15.6)	
kidney	102(65.4)	75 (67.6)	27(60.0)	0.46
liver	37(23.7)	26 (23.4)	11 (24.4)	
BMI pre-transplant (kg/m <sup>2</sup> ), median (range)	23.5 (15.8-37.3)	22.3 (15.8-36.2)	26.4 (18.8-37.3)	0.0003
BMI 5 year follow-up (kg/m <sup>2</sup> ), median (range)	26.0 (16.7-43.6)	25.5 (16.7-43.6)	27.1 (19.9-42.3)	0.10
Recipient CMV infection $(R^{+})$ [%]	49.3	52.5	42.2	0.25
Donor CMV infection ( $D^+$ ) [%]	61.5	59.2	66.7	0.39
Recipient and Donor CMV infection $(R^+/D^+)$ [%]	27.6	27.0	28.9	0.81
Calcineurin inhibitors, n [%]				
TAC	54 (34.6)	29 (26.1)	25 (55.6)	~0.004
CSA	102 (65.4)	82 (73.9)	20 (44.4)	<0.001
First transplantation [%]	80.1	77.5	86.7	0.19
Acute rejection episode during first year	47.4	47.2	47.7	0.96

191 Supplementary Table 1: Clinical characteristics of the discovery study sample:

NODAT: New-onset diabetes after transplantation, BMI: body mass index, CMV: cytomegalovirus, R: recipient, D: Donor, TAC: Tacrolimus, CSA: Cyclosporine.

196	Supplementary	Table 2:	Clinical	characteristics	of STCS	;§
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	First STCS	Second STCS	combined STCS replication
Characteristic	replication sample	replication sample	sample
	(n=958)	(n=667)	(n=1625)
Recipient age at transplantation (years),	FF (10 70)	FF (70)	FF (10, 70)
median (range)	55 (18-79)	55 (79)	55 (18-79)
Recipients sex (Males) [%]	66.2	64.6	65.5
Period of follow-up (months), median (range)	12 (0-60)	6 (0-48)	6 (0-60)
Living donor [%]	28.7	24.0	26.7
Donor age (years)	53 (1-86)	55 (1-88)	53 (1-88)
Donor sex (Males) [%]	53.5	53.4	53.4
Organ, n [%]			
Kidney	624 (65.1)	360 (54.0)	984 (60.5)
Liver	158 (16.5)	142 (21.3)	300 (18.5)
Lung	94 (9.8)	97 (14.5)	191 (11.8)
Heart	65 (6.8)	57 (8.6)	122 (7.5)
Islets and pancreas $^{\pounds}$	17 (1.8)	11 (1.6)	28 (1.7)
BMI pre-transplant (kg/m <sup>2</sup> ), median (range)	24.6 (13.7-41.2)	24.9 (13.4-43.5)	24.8 (13.4-43.5)
BMI 1 year follow-up (kg/m <sup>2</sup> ), median (range)	25.3 (15.4-44.6)	24.8 (13.7-41.7)	25.2 (13.7-44.6)
Recipient CMV infection $(R^{+})$ [%]	57.5	59.5	58.2
Donor CMV infection ( $D^*$ ) [%]	53.4	58.4	56.2
Recipient and Donor CMV infection $(R^*/D^*)$ [%]	33.0	36.3	34.3
Calcineurin inhibitors, [%]			
ТАС	65.5	67.1	66.3
CSA	26.7	26.1	26.4
None	7.8	6.8	7.3
Incidence of NODAT [%]	27.4	18.7	24.4
Incidence of New hypertension [%]	67.6	32.8	53.6
Incidence of New hyperlipidemia [%]	33.1	13.8	26.6

<sup>s</sup>patients included in this table represent the global STCS sample with the clinical data available for the analyses without the exclusion of patients with diabetes, hypertension or hyperlipidemia before transplantation. <sup>f</sup>These patients had type I diabetes mellitus as an indication for transplantation and were excluded from NODAT analysis. Abbreviations: BMI: body mass index, CMV: cytomegalovirus, R: recipient, D : Donor, TAC: Tacrolimus, CSA: Cyclosporine, NODAT: New-onset diabetes after transplantation, New hypertension: new onset hypertension after transplantation, New hyperlipidemia: new onset hyperlipidemia after transplantation.

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Supplementary Table 3: Genotype frequencies in the discovery sample and in the first, the second and the combined STCS samples. Hardy-Weinberg equilibrium and a comparison between the observed minor allele frequency and HapMap minor allele frequency in the Caucasian population.

		De Fraguana	$\log p(\theta/)$	Hardy-Weinberg	MAF		
SNP CRTC2 SNPS Frequencies n (%)				Equilibrium (p-value)	observed	НарМар	
Discovery sam	ple (n=156)						
rs8450	GG	GA	AA	0.87	0.30	0.30	
	78 (50.0)	64 (41.0)	14 (9.0)				
rs12117078	<b>GG</b> 126 (80.8)	<b>GC</b> 28 (17.9)	<b>CC</b> 2 (1.3)	0.75	0.10	0.10	
STCS							
rs8450	GG	GA	AA			0.30	
First STCS sample (n=949) <sup>§</sup>	408 (43.0)	429 (44.2)	112 (11.8)	0.96	0.34		
second STCS sample (n=667)	281 (42.1)	302 (45.3)	84 (12.6)	0.84	0.35		
Combined STCS sample (n=1616)	689 (42.7)	731 (45.2)	196 (12.1)	0.92	0.35		

 $\overline{}^{\$}9$  missing genotypes for this sample.

205 MAF: minor allele frequency.

			Additive model			Dominant model			
CRTC2 rs8450		GG	GA		AA		GG	GA&AA	
	N		NODAT OR	p-value	NODAT OR	p-value		NODAT OR	p-value
			(95% CI)		(95% CI)			(95% CI)	
First STCS replication sar	mple:								
Total	711	ref	0.97 (0.68 - 1.38)	0.85	1.01 (0.59 - 1.73)	0.98	ref	0.97 (0.70 - 1.37)	0.88
Treatment with TAC & deceased donor	281	ref	1.08 (0.63 - 1.85)	0.79	2.45 (1.14 - 5.26)	0.02	ref	1.31 (0.79 - 2.17)	0.30
Second STCS replication sample:									
Total	438	Ref	0.81 (0.47 - 1.40)	0.46	1.80 (0.88 - 3.71)	0.11	ref	0.99 (0.60 - 1.62)	0.96
Combined STCS replication sample:									
Total	1149	ref	0.89 (0.66 - 1.20)	0.44	1.18 (0.76 - 1.81)	0.46	ref	0.95 (0.72 - 1.25)	0.70
Treatment with TAC & deceased donor	395	ref	0.81 (0.51 - 1.30)	0.39	1.86 (0.99 - 3.52)	0.06	ref	1.00 (0.65 - 1.54)	0.99

Supplementary Table 4: Association between CRTC2 rs8450 SNP and NODAT in total population of STCS in different subgroups at risk of NODAT:

TAC: tacrolimus, OR: odd ratio, CI: confidence interval

P values were adjusted (when appropriate) for age of recipient at transplantation, sex of the recipient, hepatitis C status, baseline BMI, type of calcineurin inhibitor and type of donor.

Supplementary Table 5: Associations between *CRTC2 rs1572788 T>C* (in complete linkage disequilibrium (LD) with *rs8450 G>A*) and glycemic, lipidemic and obesity traits in several population-based samples:

	Effect of each rs1572788 C-allele						
	(complete LD with <i>rs8450 A-allele</i> )						
phenotype	n	beta	p-value	Explained variance*			
Glucose <sup>£</sup>	46186	0.01	0.004	0.0002			
2h glucose tolerance test <sup>±</sup>	15234	-0.02	0.26				
HOMA-β <sup>±</sup>	46186	-0.008	0.03	0.0001			
HOMA-IR <sup>*</sup>	46186	0.0002	0.96				
Total cholesterol <sup>9</sup>	97148	0.005	0.16				
HDL-Cholesterol <sup>3</sup>	96908	-0.008	0.013	0.0001			
LDL-Cholesterol <sup>9</sup>	92503	0.006	0.06				
Triglycerides <sup>9</sup>	93562	0.009	0.004	0.0001			
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BMI	113955	0.004	0.37				
Waist-hip ratio	55282	0.002	0.71				
Waist circumference <sup>°</sup>							
Male	38305	0.008	0.32				
Female	47320	0.01	0.19				

explained variance by the polymorphism (only calculated for p<0.05).

<sup>£</sup> This clinical variable was analyzed in the MAGIC study. <sup>§</sup> This clinical variable was analyzed in the "Genome Wide Associations Scans for Total Cholesterol, HDL-C, LDL-C and triglycerides". <sup>§</sup> This clinical variable was analyzed in the GIANT study.

2h: 2 hours, HOMA: homeostatic model assessment,  $\beta$ : beta-cell function, IR: insulin resistance, BMI: Body mass index.

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