**CACNA1E Variants Affect Beta Cell Function in Patients with Newly Diagnosed Type 2 Diabetes. The Verona Newly Diagnosed Type 2 Diabetes Study (VNDS)**

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**Abstract**

**Background:** Genetic variability of the major subunit (CACNA1E) of the voltage-dependent Ca^{2+} channel Ca_{v}2.3 is associated to risk of type 2 diabetes, insulin resistance and impaired insulin secretion in nondiabetic subjects. The aim of the study was to test whether CACNA1E common variability affects beta cell function and/or insulin sensitivity in patients with newly diagnosed type 2 diabetes.

**Methodology/Principal Findings:** In 595 GAD-negative, drug naïve patients (mean±SD; age: 58.5±10.2 yrs; BMI: 29.9±5 kg/m²; HbA1c: 7.0±1.3) with newly diagnosed type 2 diabetes we: 1. genotyped 10 tag SNPs in CACNA1E region reportedly covering ~93% of CACNA1E common variability: rs558994, rs679931, rs2184945, rs10797728, rs3905011, rs12071300, rs175338, rs3753737, rs22253388 and rs4652679; 2. assessed clinical phenotypes, insulin sensitivity by the euglycemic insulin clamp and beta cell function by state-of-art modelling of glucose/C-peptide curves during OGTT. Five CACNA1E tag SNPs (rs10797728, rs175338, rs2184945, rs3905011 and rs4652679) were associated with specific aspects of beta cell function (p<0.05–0.01). Both major alleles of rs2184945 and rs3905011 were each (p<0.01 and p<0.005, respectively) associated to reduced proportional control with a demonstrable additive effect (p<0.005). In contrast, only the major allele of rs22253388 was related weakly to more severe insulin resistance (p<0.05).

**Conclusions/Significance:** In patients with newly diagnosed type 2 diabetes CACNA1E common variability is strongly associated to beta cell function. Genotyping CACNA1E might be of help to infer the beta cell functional phenotype and to select a personalized treatment.

**Introduction**

Ca_{v}2.3 is one non-L-type high-voltage-activated (HVA) channel belonging to the voltage-gated Ca^{2+} channel family, whose members mediate Ca^{2+} entry into cells in response to membrane depolarization [1].

Ca_{v}2.3, which is expressed primarily in neuronal cells [2] and in pancreatic beta cells [1], plays a key role in second-phase insulin release [3]. Silencing Ca_{v}2.3 expression in INS-1 cells blunts glucose-stimulated insulin secretion [4]. Ca_{v}2.3^{+/-} mice display fasting hyperglycemia and impaired glucose tolerance [5,6].

Ca_{v}2.3 is composed of a major pore-forming alpha 1 E subunit (CACNA1E), which determines most of the channel’s properties, and of multiple auxiliary subunits (alpha 2, beta, delta, gamma) [1].

Two earlier studies, performed in two different ethnic groups, reported that CACNA1E variants are associated to type 2 diabetes risk [7,8]. However, their findings differed because in one study the metabolic phenotype associated to CACNA1E variability was insulin secretion [8], in the other study it was insulin sensitivity [7]. Both studies used insulin concentrations after a secretory stimulus to explore beta cell function. However, insulin levels are determined by both beta cell function and insulin clearance, which is in turn influenced by insulin sensitivity. Thus, any insulin based assessment of beta cell function also reflects insulin clearance. Indeed, in the former study [8], the association between CACNA1E variability and beta cell function was sought using either surrogate indexes of glucose-induced insulin secretion during OGTTs, or insulin levels in the late part of IVGTTs. In the latter study [7], only acute insulin response after i.v. glucose challenge was measured, i.e. a facet of beta cell function in which Ca_{v}2.3 should play a nonsignificant role [3]. Finally, in both studies, associations between CACNA1E variability and metabolic...
phenootypes were investigated in nondiabetic individuals [7,8]. We hypothesized that, if common CACNA1E variability were relevant in determining beta cell function, this might be evident also when overt diabetes ensues and such an information would be of potential clinical interest.

We therefore undertook the present study to test whether the genetic variability of CACNA1E affects key metabolic phenotypes (beta cell function and insulin sensitivity, both assessed by state-of-art methods) in patients with newly diagnosed type 2 diabetes.

The CACNA1E region was tested by genotyping a number of tag SNPs, selected to capture most (~93%) of common CACNA1E variability. We used the database of the Verona Newly Diagnosed Type 2 Diabetes Study (VNDS), a cohort in which associations between common genetic variants and metabolic phenotypes have already been shown to be detectable [9,10], possibly because of the absence of the potentially confounding effects of long-lasting antidiabetic treatments and of the limited impact of duration and severity of hyperglycemia per se on the glucose insulin homeostatic system.

**Methods**

**Verona Newly Diagnosed Type 2 Diabetes Study (VNDS)**

The VNDS is an ongoing study aiming at building a biobank of patients with newly diagnosed type 2 diabetes [9,10]. A detailed description of the study population and of the experimental design is found online. In this study we report the data collected in 595 patients, whose characteristics are summarized in Table 1. Each subject gave informed written consent before participating in the research, which was approved by the Human Investigation Committee of the Verona City Hospital.

Standard clinical parameters were assessed in all patients. Metabolic tests were carried out on two separate days in random order. On one day an OGTT (75 g) was performed to assess beta cell function, as previously described [11]. On a separate day, a euglycemic insulin clamp was performed to assess insulin sensitivity [9].

**Analytical procedures**

Plasma glucose concentration was measured in duplicate with a Beckman Glucose Analyzer II (Beckman Instruments, Fullerton, CA, USA) or a YSI 2300 Stat Plus Glucose&Lactate Analyzer (YSI Inc., Yellow Springs, OH, USA), at bedside. Serum C-peptide and insulin concentrations were measured by chemiluminescence as previously described [12]. Glycated hemoglobin and serum lipids were measured by standard in-house methods. GAD-antibodies were measured by immunoradiometry (CentAK, Medipan, Germany), according to manufacturer’s instructions.

**Calculations**

The amount of glucose metabolized during the last 60 min of the clamp (M value, reference insulin sensitivity) was computed with standard formulae [13,14].

The following classical indexes of beta cell function were computed [15]:

- Insulinogenic Index: \((\text{Insulin}_{30} - \text{Insulin}_9)/(\text{Glucose}_{30} - \text{Glucose}_9)\), units: mU/mmol;
- Corrected Insulin Response \(\text{CIR}_{120}\): \(\text{Insulin}_{120}/(\text{Glucose}_{120} - 3.89)\) [15]; units: mU*L/mmol².

Further insights in beta cell function were sought by mathematical modeling (see below and Supporting Information S1 Tables S1, S2).

**Modeling of Beta Cell Function**

The analysis of the glucose and C-peptide curves during the OGTTs of the VNDS was performed as described in previous publications [9,10] and builds upon previous works from other laboratories [16,17]. Insulin secretion rate is described as the sum of two components, one responding to the rate of increase of glucose, the other to glucose concentration itself. Modeling details are found online in the Supporting Information S1.

There are two main physiological outputs of the model:

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### Table 1: Clinical and metabolic features of the VNDS population.

| Variable          | Males       | Females    | All         |
|-------------------|-------------|------------|-------------|
| Number (M/F)      | 409         | 186        | 595         |
| Age (yrs)         | 59 [51–65]  | 61 [56–67] | 60 [52–66]  |
| BMI (kg/m²)       | 28.7 [26.1–32.1] | 30.4 [27.2–34.1] | 29.3 [26.4–32.9] |
| Waist (cm)        | 102 [94–111] | 97 [90–103] | 100 [93–108] |
| Fasting P-glucose (mmol/l) | 7 [6.2–8.0] | 7.2 [6.2–8.1] | 7.1 [6.2–8.0] |
| 2hr P-glucose (mmol/l) | 13.1 [10.8–16.2] | 12.9 [10.1–16.3] | 13.1 [10.6–16.2] |
| HbA1c (%)         | 6.7 [6.1–7.5] | 6.6 [6.2–7.4] | 6.6 [6.1–7.5] |
| Triglycerides (mmol/l) | 1.5 [1.0–2.1] | 1.3 [1.0–2.0] | 1.4 [1.0–2.0] |
| HDL-cholesterol (mmol/l) | 1.1 [0.95–1.3] | 1.2 [1.0–1.4] | 1.1 [1.0–1.3] |
| Cholesterol (mmol/l) | 4.8 [4.2–5.5] | 5.1 [4.5–5.8] | 5.0 [4.3–5.6] |
| SBP (mmHg)        | 134 [120–146] | 140 [130–150] | 136 [122–150] |
| DBP (mmHg)        | 80 [80–90]   | 85 [80–90]  | 82 [80–90]  |
| Insulin Sensitivity (µmol/min/m² BSA) | 602 [365–885] | 596 [408–796] | 599 [384–856] |
| Insulinogenic Index (mU/mmol) | 3.6 [2.0–5.9] | 4.8 [2.5–8.2] | 3.9 [2.1–6.9] |
| CIR120’ (mUxL/mmol²) | 0.4 [0.2–1.1] | 0.6 [0.3–1.4] | 0.5 [0.2–1.3] |

Data are presented as median [I.Q. range].

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1. derivative control \([\text{pmol} \cdot \text{m}^{-2} \cdot \text{BSA}] \cdot [\text{mmol} \cdot \text{l}^{-1} \cdot \text{min}^{-1}]^{-1}\) of beta cell function: it is presented as the amount of insulin secreted in response to a rate of glucose increase of 1 mmol/l per min which lasts for 1 minute;

2. proportional control of beta cell function: it is presented as the stimulus-response curve linking glucose concentration (x axis) to insulin secretion rate (ISR \([\text{pmol} \cdot \text{min}^{-1} \cdot \text{m}^{-2} \cdot \text{BSA}]; y axis) at the pre-selected glucose concentrations of 5.5, 8.0, 11.0, 15.0 and 20.0 mmol/l.

Genotyping

A peripheral blood sample was collected from each patient and the DNA was extracted by standard salting out method.

Ten tag-SNPs (rs558994, rs679931, rs2184945, rs10797728, rs3905011, rs12071300, rs175338, rs3753737, rs2253388 and rs4652679), which capture \(~93\%\) of the common genetic variability of this genomic region (about 320 kbp) (fig. 1), were selected with the software GEVALT (GEnotype Visualization and ALgorithmic Tool) [18]. The LD values in our population, expressed as \(r^2\), are consistent with the values reported by HapMap (fig. 1).

Genotypes were assessed by the high-throughput genotyping Veracode technique (llumina Inc, CA), applying the GoldenGate Genotyping Assay according to manufacturer’s instructions [19].

Statistical Analysis

Data are presented as medians and interquartile range, unless otherwise indicated. Hardy-Weinberg equilibrium was tested by chi-square test. Before comparisons, skewed variables were log-transformed to improve the approximation to the Gaussian distribution. Single variant association analyses were carried out by generalized linear models (GLM) as implemented in SPSS and they were adjusted for a number of potential confounders, including age, gender and BMI. Since the logarithmic transformation was insufficient to improve the distribution of the derivative control of beta cell function, the latter was first analyzed by Kruskal-Wallis test. All statistical analyses were carried out with the SPSS 12.0 software. Statistical significance was declared at \(p<0.05\).

Results

Clinical Features of the Study Population

Anthropometric, clinical and metabolic features of the study subjects are shown in table 1. Further details can be found in the Supporting Information S1.

Effects of CACNA1E variants on clinical and metabolic parameters

Allele distribution was compatible with the Hardy-Weinberg equilibrium in all 10 SNPs (\(p>0.16\) or more).

Anthropometric, clinical and metabolic features of the study subjects, layered according to rs558994, rs679931, rs2184945, rs10797728, rs3905011, rs12071300, rs175338, rs3753737, rs2253388 and rs4652679 alleles, are found in the online material (Supporting Information S1 Tables from S3 to S12).
Effects of CACNA1E variants on derivative and proportional control of beta cell function

Both major alleles of rs10797728 (A) and rs175338 (G), which are in modest linkage disequilibrium with each other (D': 0.9, r²: 0.43) (fig. 1), were associated to a significant impairment in the derivative control of beta cell function, i.e. the response of beta cell to the rate of glucose increase (p = 0.01 and p = 0.03 respectively by Kruskal-Wallis) (Table 2). The major allele of rs10797728 was also associated to lower basal insulin secretion rate (−0.07±0.03 log units, p<0.03) (Supporting Information S1 Table S6).

Patients carrying the major alleles A of rs2184945, under a dominant model, (Supporting Information S2 Fig. S1a) or G of rs3905011 (Supporting Information S2 Fig. S1b) had defective proportional control of beta cell function, i.e. the curve relating glucose (stimulus) to insulin secretion rate (response) (p<0.05 respectively by ANOVA). The effects of rs2184945 and rs3905011 genotypes on the beta cell response to glucose concentration were statistically independent of each other (p<0.02 and p<0.01, respectively, in a multivariate model including both SNPs), consistently with their low degree of linkage disequilibrium (D’: 0.26, r²: 0.06).

We then computed a CACNA1E score in order to investigate the combined effect of rs2184945 and rs3905011 on the proportional control, by assigning a score of 1 to all carriers of the rs2184945 A allele, irrespectively of zygoty, and a score of 1 for each rs3905011 G allele. Hence, the CACNA1E score could range from a minimum of 0 (rs2184945 TT–rs3905011 GG genotype) to a maximum of 3 (genotype rs2184945 AA or AT–rs3905011 GG genotype) (Supporting Information S1 Table S13). The higher was the CACNA1E score, the lower was the insulin secretion rate in response to glucose concentration (Fig. 2) (p<0.005).

Finally, the major G allele of rs4652679 was associated to significantly lower basal insulin secretion rate (−0.06±0.03 log units, p<0.04) and to significantly more efficient insulin clearance (+0.07±0.02 log units, p<0.01) (Supporting Information S1 Table S12).

Table 2. Derivative control of beta cell function in patients of the VNDS according to genotype of CACNA1E variants.

| CACNA1E SNP | n | AA | AB | BB | p value (k-w) |
|-------------|---|----|----|----|-------------|
| rs558994    | 514 | 653.6±47.6 | 609.9±40.4 | 568.2±105.3 | 0.69 |
| rs679931    | 515 | 700.1±50.9 | 578.2±42.3 | 712.6±125.4 | 0.11 |
| rs2184945   | 521 | 707.1±54.6 | 585.5±46.7 | 683.1±71.2 | 0.04 |
| rs10797728  | 523 | 658.8±42.1 | 558.4±47.5 | 1015.2±209.8 | 0.01 |
| rs3905011   | 521 | 580.5±46.3 | 646.9±46.2 | 734.7±96.4 | 0.59 |
| rs12071300  | 529 | 642.9±38.8 | 587.5±57.4 | 962.5±205.1 | 0.07 |
| rs175338    | 507 | 658.8±41.5 | 570.3±49.1 | 1054.3±218.7 | 0.03 |
| rs3753737   | 512 | 636.1±42.4 | 656.6±55 | 620.8±94.2 | 0.91 |
| rs2253388   | 515 | 619.5±43.7 | 641.2±51.4 | 754.1±111.1 | 0.32 |
| rs4652679   | 518 | 629.2±42.7 | 636.5±47.9 | 598.6±118.7 | 0.85 |

Data are presented as mean±SE. A non-parametric test has been performed (Kruskal-Wallis) since the variable was not normally distributed.

A = major allele; B = minor allele.

Discussion

This study explored the potential role(s) of ten CACNA1E variants (fig. 1), purposely selected to capture ~93% of common CACNA1E variability, in determining pathophysiological phenotypes (i.e. beta cell function and insulin sensitivity) in patients with newly diagnosed type 2 diabetes of the VNDS. We also genotyped rs679931 and rs3753737 because of their previously reported role in type 2 diabetes risk and in beta cell function regulation [7,8].

The interesting and novel feature of our study is the use of state-of-art methods [16,17,20,21], instead of surrogate markers, to assess beta cell function and insulin sensitivity in a sample of patients with newly diagnosed type 2 diabetes. This population is presumably not yet influenced by long-standing hyperglycemia and/or pharmacological antidiabetic treatment, which could modify phenotype-genotype interaction.

Our data show that up to five (rs10797728, rs175338, rs2184945, rs3905011 and rs4652679) out of ten CACNA1E SNPs may exert some influence on different components of beta cell function (derivative control and proportional control), which in some (rs2184945 and rs3905011 genotypes), but not all cases, are also reflected by changes in glucose control (Supporting Information S1 Tables S3 and S7).

The apparent influence exerted by rs10797728 and 175338 on the derivative control of beta cell function (table 2) is somewhat surprising, in the light of the claim that Cav2.3 is involved in the regulation of only second phase insulin secretion [3] and that the derivative control of beta cell function describes first phase insulin secretion in response to intravenous glucose challenges [11,20,21]. This apparent lack of biological plausibility may indicate that this is simply a chance finding. However, at a close inspection, the data reported by Jing et al., possibly the most thorough study on the relationship between Cav2.3 activity and beta cell function, do show a trend also for reduced first phase insulin secretion in perfused pancreata from Cav2.3/−/− mice [3]. Thus, if replicated...
in future studies, this finding may be of pathophysiological relevance in patients with newly diagnosed type 2 diabetes.

Both rs2184945 and rs3905011 genotypes exerted a strong, independent influence on the proportional control of beta cell function (Supporting Information S2 Fig. S1a and fig. S1b, respectively). Furthermore, our finding that homozygous carriers of “poor” beta cell function alleles at both SNPs had a \( \frac{3}{10} \) reduction in the insulin secretory response to glucose, compares well with the 20–30% reduction of second phase insulin secretion reported in perfused pancreata from \( \text{CaV2.3}^{-/-} \) mice [3]. It is noteworthy that the \( p \) value of the \( \text{CACNA1E} \) score was virtually unaffected (\( p = 0.001 \)) by the inclusion in a multivariate model also of the genetic variants previously reported by us to be associated to beta cell function in the VNDS (rs7901695 or rs7903146 in \( \text{TCF7L2} \) and rs6717980 or rs2384628 in \( \text{GCKR} \) [9,10]). Furthermore, the associations of these two SNPs with indicators of glucose control, such as plasma glucose and HbA1c, are consistent with a potentially relevant role in regulating both basal and glucose induced insulin secretion.

There are a number of limitations in our study. i- This is not a population based study, although we found that VNDS subjects are fairly representative of Italian patients with type 2 diabetes, comparing of their features with previously published studies [22]. However, in our cohort there is a higher than expected prevalence of the male gender, possibly reflecting a gender related referral bias. ii- The size of our cohort is not very large, although the direct assessment of the phenotypes of interest, instead of using surrogate indexes, may partially compensate for it. iii- We did not explore many important pathophysiological phenotypes, e.g. incretin role and liver vs peripheral insulin resistance, which are not measured by the state-of-art methods that we used to assess beta cell function and insulin sensitivity [16,17,20,21]. iv- Our findings need to be replicated in an independent sample of patients, as in all association
studies. This may prove difficult, because generating direct measures of beta cell function and insulin sensitivity in large samples requires time and considerable resources. Indeed, to the best of our knowledge, there is no other large cohort of European patients with type 2 diabetes who underwent the same kind of metabolic phenotyping. Our study was performed in Italian patients; extrapolations to other ethnic groups should be made with caution.

In summary, we have reported that common CACNA1E variation is significantly related to beta cell function in patients with newly diagnosed type 2 diabetes. Hence, we speculate that CACNA1E genotype might be a useful tool to refine diagnosis and prognosis of patients with type 2 diabetes.

Supporting Information

Supporting Information S1 Includes text with supplemental information regarding methods and results, and tables from S1 to S13.

(DOC)

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Supporting Information S2 Includes supplemental figures S1a and S1b.

(DOC)

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Author Contributions

Conceived and designed the experiments: RCB. Performed the experiments: SB MT MLB FT. Analyzed the data: SB MT RCB. Contributed reagents/materials/analysis tools: RCB MT SB MLB FT. Wrote the paper: ET GM PFP EB.