INTRODUCTION

The increasing prevalence of type 2 diabetes (T2D) is one of the greatest challenges in public health, in both developed and developing countries. In 2012, approximately 8.3% of the world’s adult population was living with diabetes. Diabetes is a leading cause of cardiovascular disease, renal disease, blindness, and limb amputation. T2D, accounting for 80–90% of all diabetes in Europe, decreases life expectancy by 5–10 years.

Because the onset of T2D can be postponed or partially prevented by changes in the lifestyles of high-risk subjects, the cost-effectiveness of lifestyle (and other) interventions can be increased by improving the precision of risk prediction, thereby enabling targeting of the individuals at highest risk.

Although obesity is the strongest predictor of T2D, it is also known that heritability of T2D is 26–69%, depending on age of onset, thus motivating the search for genetic predictors for T2D. However, despite the large number of published genome-wide association studies (GWAS) of T2D, there is still some skepticism regarding the practical value of identified single-nucleotide polymorphisms (SNPs) in personalized risk prediction for the disease. The main reason is that the effect of individual SNPs on complex common disease phenotypes is relatively weak and/or adds little to predictions based on lifestyle, demographic, and clinical factors.

In GWAS, SNPs need to meet the stringent genome-wide threshold, usually set to $P < 5 \times 10^{-8}$, to be significantly associated with the trait. Even though the sample sizes in GWAS have been increasing steadily over the years, they are still insufficient for SNPs with small effects to pass that threshold. This could explain why it has been shown that all common variants across the genome actually explain a much higher proportion of heritability (50% or more) in many complex traits than one could see based on a small subset of significant SNPs only.

To explain a meaningful proportion of variability in a complex trait and, more importantly, to use this knowledge in risk assessment at an individual level, one needs to construct a numeric summary measure of genetic risk—a genetic (polygenic) risk score (GRS) based on a large number of genotyped variants. Our aim is to develop a GRS with the best possible predictive power and investigate its potential to improve T2D risk stratification in the general population. For this purpose, two sources of data will be used: (i) results of large-scale meta-analyses of GWAS to obtain effect estimates for individual SNPs and (ii) individual-level data of a relatively large population-based cohort from the Estonian Biobank.
to compare versions of the GRS and decide on applicability of the best GRS in practical risk prediction. The best-fitting GRS for prevalent cases is then further validated in the analysis of incident T2D (data obtained by linking the Estonian Biobank cohort database to electronic health records of the participants).

MATERIALS AND METHODS

Estonian Biobank cohort and genotyping

The Estonian Biobank (Estonian Genome Center, University of Tartu) was established with the long-term purpose of implementation of research results in public health and medicine in Estonia. Between 2002 and 2011, the Estonian Biobank recruited a cohort of 51,380 participants, including adults from all counties in Estonia and accounting for approximately 5% of the Estonian adult population during the recruitment period. Broad informed consent signed by participants enabled use of the data for various health research purposes as well as linkage of the data with other health-related databases and registries. An extensive phenotype questionnaire and measurement panel, together with follow-up data from linkage with national health-related registries and electronic health records (the Estonian Health Insurance database), allows assessment of the effects of classic epidemiological risk factors on the incidence of common complex diseases, such as T2D. The research project at the Estonian Genome Centre was approved by the Ethics Committee of Human Studies, University of Tartu, Estonia.

In the present study, a genotyped subset of 10,273 individuals (including 1,181 prevalent T2D cases) from the cohort was analyzed. The DNA samples of this subset were genotyped using either Illumina Human OmniExpress (a sample of 8,085 individuals) or Illumina Cardio-MetaboChip (a case–control sample of 942 T2D cases, 680 cases of coronary artery disease, and 903 random controls) genome-wide arrays. For 337 individuals (including 169 T2D cases) genotyped by both arrays, genotype data from the Cardio-MetaboChip array were used.

During the average follow-up time of 5.63 years, 386 incident T2D cases were observed (in 9,092 individuals free of T2D at recruitment) by 1 April 2014. A total of 1,994 individuals had died by 1 September 2015 (including 1,069 deaths due to cardiovascular causes).

The baseline phenotype data (Table 1) used for this study consist of age, gender, BMI and prevalent T2D status, history of hypertension or high blood glucose level, physical activity level (active versus inactive), and consumption of fresh fruit. For a subset (n = 6,064), data on plasma glucose level and lipid-protein profiles (low- and high-density lipoprotein (LDL and HDL) cholesterol, triglycerides, and total cholesterol) obtained by nuclear magnetic resonance (NMR) profiling were available (nonfasting measurements, with information on the time of last meal available for adjustment).

The genetic data used in this study were selected as follows. First, the Estonian Cardio-Metabochip sample was used in the large-scale GWAS meta-analysis for T2D susceptibility from the DIAGRAM Consortium. We therefore reran the meta-analysis to remove the effects of the Estonian sample because we intended to use the Cardio-Metabochip for developing the optimal GRS score. Second, only SNPs with P < 0.5 for association with T2D in the meta-analysis were chosen for further analysis. A set of independent SNPs with r² ≤ 0.05 was then obtained via the LD-based clumping procedure of PLINK. Finally, clumped SNPs were retrieved from the Estonian Biobank database and filtered for genotyping and imputation quality and minor allele frequency, resulting in a set of 7,502 SNPs for further analysis and GRS construction (Supplementary Table S1 online). Full details on selection and weighting of SNPs can be found in the Supplementary Methods and Figures online.

Statistical analysis

Statistical analysis was performed using R version 3.1.0. The doubly weighted GRS

Suppose there are K independent markers with allele dosages X₁, ..., Xᵦ available for a study, with estimated (linear or logistic) regression parameters from a GWAS meta-analysis β₁, ..., βᵦ and corresponding P values P₁, ..., Pᵦ. We define the general GRS as

$$\text{GRS} = \sum_{i=1}^{K} w_i X_i$$

and different versions of GRS can be defined by varying the choice of weights wᵢ.

The conventionally used GRS (also referred to as the single-weighted GRS) is defined by choosing wᵢ = βᵢ for all markers with pᵢ < p* and wᵢ = 0 otherwise, where p* is a P value threshold (often 5 × 10⁻⁸ or 5 × 10⁻⁶). Let GRSᵦ denote the GRS where pᵢ* is chosen so that exactly k (k < K) markers with the smallest P values have nonzero weight in the score. For any choice of k, GRSᵦ suffers from a phenomenon called “winner’s curse”—by selecting only SNPs with estimated P values below a certain threshold, one systematically selects SNPs with effect overestimated by chance. To correct for the resulting bias, we propose a doubly weighted GRS, denoted by dGRSᵦ, defined by selecting wᵢ = βᵢ πᵢ with πᵢ defined as an estimated probability that the i-th marker belongs to the set of top k SNPs with the strongest effect on the phenotype (“strongest” defined as having the smallest P value on average of all possible studies). We estimate such probability by simulating new values of potential parameter estimates based on the observed estimates and their standard errors (more details are provided in the Supplementary Methods and Figures online). We have conducted simulation studies (not presented in this article) that demonstrated that dGRSᵦ indeed decreases the bias caused by the “winner’s curse” in the single-weighted GRSᵦ. Although in practice the algorithm requires a large number of SNPs, choosing a small value of k will result in near-zero weights for most of the SNPs used.
Comparison of different versions of the GRS in the association with prevalent T2D status

Using the data of the genotyped subsets of the Estonian Biobank cohort, we calculated both GRS and dGRS, by varying k from 1 to all 7,502 of the initially selected SNPs. The effect of each GRS was assessed using age-, sex-, and genotype platform–adjusted logistic regression models for prevalent T2D status. Both BMI-adjusted and unadjusted models were fitted. The fit of (nonnested) models using a different version of the GRS as a covariate was compared using the Cox likelihood ratio test. The GRS producing the highest log-likelihood for both BMI-adjusted and unadjusted models was selected for further validation.

Because the GRS is a continuous measure, one also needs to specify a set of threshold values to classify individuals as being at “high,” “average,” or “low” genetic risk, for instance, to simplify the risk assessment in clinical practice. We investigated whether the quintiles of GRS can be used for that purpose, by using bar charts to compare the observed T2D prevalence in individuals aged 40–79 years across quintiles of the GRS and BMI category (<25, 25–30, 30–35, >35). In addition, bar charts were produced to study the distribution of individuals across GRS quintiles within the subset with prevalent T2D and in the subset of obese (BMI >35) T2D-free individuals aged 60 and older.

Validation of the GRS in the analysis of incident conditions

The GRS was further assessed for its effect on incident T2D in individuals without prevalent T2D at baseline and all-cause and cardiovascular mortality (in all individuals) using Cox proportional hazards modeling with age as the time scale. All models are adjusted for BMI category, smoking level (although smoking is not considered a standard predictor, we included it because of the significant effects of smoking level on T2D risk in our cohort), waist-to-hip ratio (WHR), waist circumference, physical activity level, history of high blood glucose, history of hypertension, fruit and vegetable consumption, and sex.

The analysis was restricted to the subset of 6,280 individuals aged 35–79 at recruitment (302 incident T2D cases) while censoring all T2D diagnoses beyond age 80 because diagnoses in the elderly are often related to significant risk-altering comorbidities (cancer or cardiovascular diseases). In addition, the Cox proportional hazards model was fitted in the subset with available glucose and lipid measurements (3,776 individuals, including 158 incident T2D cases) by adjusting for glucose, triglyceride, and HDL-cholesterol levels in addition to the covariates mentioned previously. The Kaplan-Meier graph of cumulative incidence of T2D was obtained for the subset with BMI >23.

Association of the GRS with other known T2D risk factors

The effects of GRS on BMI, WHR, plasma glucose, total cholesterol, HDL- and LDL-cholesterol, and triglyceride levels were estimated using age- and sex-adjusted linear regression analysis in individuals without prevalent T2D diagnosis. The effects of GRS on BMI and WHR were similarly assessed in individuals with prevalent T2D diagnosis.

Analysis of incremental value of GRS

For prevalent T2D, the area under the receiver operating characteristic (ROC) curve (AUC) was obtained from logistic regression fitted for individuals in the 40–79 age group who were genotyped on the OmniExpress platform. For incident T2D, Harrell’s c-statistic (concordance index) from the Cox proportional hazards models for individuals aged 35–79 with no prevalent T2D diagnosis was obtained.

To study reclassification and 5-year T2D risk predictions, Cox proportional hazards models with and without accounting for GRS were fitted. Improvement in the predictions was assessed using continuous net reclassification improvement (NRI) and integrated discrimination improvement (IDI). Confidence intervals for reclassification indices and c-statistics were estimated with bootstrapping.

Table 1 Baseline characteristics of the two genotyped subsets of the Estonian Biobank cohort

|                          | Illumina Human OmniExpress | Illumina Cardio-Metabochip |
|--------------------------|---------------------------|---------------------------|
| % (n) of females          | 55.8% (8,085)             | 59.5% (2,525)             |
| Age: mean (range)        | 51.1 (1 to 79)           | 58.48 (1 to 80)          |
| Mean follow-up time (range) in years for surviving individuals | 8.0 (7 to 11) | 7.79 (7 to 11) |
| Prevalence of T2D        | 5.0% (372)               | 37.3% (60)               |
| Incident T2D during follow-up (n) | 332                      | 60                        |
| BMI (mean, SE)           | 26.7 (1.2)               | 28.2 (1.3)               |
| Prevalence of overweight (BMI = 25–30) | 33.6% (332)              | 24.6% (60)               |
| Obesity grade I (BMI = 30–35) | 16.5% (332)              | 22.0% (60)               |
| Obesity grade II (BMI >35) | 7.0% (332)               | 15.2% (60)               |
| Current smokers          | 27.6% (332)              | 24.0% (60)               |
| Ex-smokers               | 16.0% (332)              | 18.3% (60)               |

Descriptive statistics are provided separately for both subsets of the Estonian Biobank cohort, depending on the genotyping platform.
RESULTS

Comparison of different versions of the GRS

Results of the model fit for prevalent T2D status with GRS_k and dGRS_k for selected values of k are shown in Table 2. (More detailed results for values of k varied between 1 and 7,502 are provided in Supplementary Table S2 online and a corresponding plot of likelihood ratio statistics is shown in Supplementary Figure S1 online) Compared with the GRS_k (similar to ref. 16), the fit was considerably improved using a GRS_k with a larger number of markers: the highest log-likelihood was achieved with GRS_2100 (BMI-unadjusted models) and GRS_400 (BMI-adjusted models). However, when dGRS_k was used instead, with k = 300 or larger (up to 3,500), the fit became significantly better than that with any GRS_k. The highest log-likelihood was achieved with dGRS_1400 (BMI-unadjusted models) and dGRS_800 (BMI-adjusted models); regardless of whether the analysis was adjusted for BMI, dGRS_1000 provided a fit that was not significantly different from the best-fitting GRS (Cox test \( P > 0.05 \)). Therefore, we used dGRS_1000 in all subsequent analyses (weights shown in Supplementary Figure S2 online).

Association of dGRS_1000 with prevalent T2D

The estimated odds ratio (OR) corresponding to 1 standard deviation (SD) difference in dGRS_1000 was 1.56 (95% CI: 1.45–1.68) in the BMI-unadjusted model and 1.59 (95% CI: 1.46–1.72) in the BMI-adjusted model. The prevalence of T2D by BMI category and quintiles of dGRS_1000 in the subset of the cohort with an age range of 45–79 is shown in Figure 1a (see Supplementary Figure S3 online for a more detailed plot). Although there was no significant interaction (\( P = 0.359 \)) between BMI and dGRS_1000, the association was strongest in overweight or moderately obese individuals (25 < BMI < 35), where the number of T2D cases in the highest GRS quintile was approximately comparable to the total number of cases in the three lowest quintiles.

Figure 1b indicates that approximately one-third of all prevalent T2D cases belong to the highest GRS quintile, but the differences in the observed frequencies of GRS quintiles were greatest in those with BMI <35. However, as indicated by Figure 1c, the majority of severely obese (BMI >35) but T2D-free individuals aged 60 and older belonged to the two lowest GRS quintiles.

Validation of the GRS in the analysis of incident conditions

As shown in Table 3, dGRS_1000 had a highly significant effect (HR = 1.48, 95% CI: 1.32–1.66 per 1 SD; \( P = 1.5 \times 10^{-11} \)) on the risk of developing T2D during follow-up and accounted for a large set of known covariates. According to the likelihood ratio test, the \( P \) value corresponding to the combined effect of the obesity-related parameters in this fitted model for incident T2D is 2.0 \times 10^{-21}, followed by the effect of dGRS_1000. The difference in risks between lowest and highest GRS quintiles was more than threefold (HR = 3.45, 95% CI: 2.31–5.17). It is also important to note that the risk in the highest dGRS_1000 quintile was approximately twice that in the rest of the sample (HR = 1.95, 95% CI: 1.52–2.51), indicating that this subset could be targeted for risk-reducing interventions. This is additionally supported by the fact that the highest dGRS_1000 quintile was associated with 14% higher risk for all-cause mortality (\( P = 0.019 \)) and 27% higher risk for cardiovascular mortality (\( P = 0.001 \)).

The differences in cumulative T2D incidence across dGRS_1000 quintiles can also be seen in Figure 2, which shows that the cumulative risk differences are already obvious by the end of the first year of follow-up.

In the analysis of the effect of dGRS_1000 on incident T2D while additionally adjusting for glucose, triglycerides, and HDL cholesterol, the HR corresponding to 1 SD difference in dGRS_1000 was estimated to be 1.49 (95% CI: 1.27–1.76; \( P = 1.7 \times 10^{-6} \)), indicating that dGRS_1000 has a significant effect on T2D risk that is independent of all available well-known risk factors.

The coefficients of all parameters in the model for incident T2D in the entire cohort of 30,094 initially T2D-free individuals (without genetic effects) and in the genotyped subset of 6,280 individuals (including the effect of dGRS_1000) are shown in Supplementary Tables S3 and S4 online.

Association of dGRS_1000 with known T2D risk factors

The estimated regression coefficients (\( \beta \), SE, \( P \) value) from the analysis of the effect of dGRS_1000 on BMI, WHR, plasma glucose,
In individuals without prevalent T2D, a significant positive association of dGRS_{1000} with plasma glucose ($P = 4.7 \times 10^{-8}$) and triglyceride levels ($P = 8.8 \times 10^{-4}$) and a negative association with HDL-cholesterol level ($P = 8.1 \times 10^{-5}$) were found, suggesting that some individuals at high genetic risk may already have a clearly increased disease risk according to other criteria (or even prediabetes or undiagnosed T2D). No significant association of dGRS_{1000} with BMI was found in T2D-free individuals ($P = 0.42$), and there was only a weak positive association with WHR ($P = 0.012$). Thus, it is unlikely that dGRS_{1000} affected T2D risk through its possible effect on obesity.

The association between BMI and dGRS_{1000} in individuals with existing T2D diagnosis was found to be negative ($P = 0.0039$), suggesting that individuals at high genetic risk are more likely to develop T2D at a BMI lower than that of those with low genetic risk, as supported by the models for both prevalent and incident T2D, where the effects of BMI and GRS appeared to be additive. However, a cautious interpretation of the estimated effect size is needed because the T2D treatment may have affected the BMI of diseased individuals.

**Analysis of incremental predictive ability of the dGRS1000**

Incremental predictive ability of dGRS_{1000} was studied for both prevalent and incident T2D models. Including dGRS_{1000} in the logistic regression model for prevalent T2D improved AUC, irrespective of BMI adjustment (Supplementary Figure S4 online). Harrell’s c-statistic increased by 0.012 (95% CI: 0.004–0.023) after dGRS_{1000} was added to the model for incident T2D. More detailed results for c-statistics and likelihood ratio tests are shown in Supplementary Table S6 online.

Comparing 5-year predictions from models with and without dGRS_{1000} (Supplementary Figure S5 online) resulted in a continuous NRI of 0.324 (95% CI: 0.211–0.444). Further investigation of components of the continuous NRI was undertaken as suggested: the continuous NRI for events was 0.115 (95% CI: 0.02–0.23) and for nonevents was 0.209 (95% CI: 0.182–0.23). The IDI was 0.013 (95% CI: 0.007–0.019).
Table 3 Analysis of the effect of dGRS_{1000} on incident T2D and on all-cause and cardiovascular mortality

| Incident condition (cases/total) | GRS covariate | HR (95% CI) | P value |
|----------------------------------|---------------|-------------|---------|
| Incident T2D (302/6,280), age 35–79 | dGRS_{1000} (effect per 1 SD) | 1.48 (1.32, 1.66) | 1.5 × 10^{-11} |
| dGRS_{1000} quintiles: | 1 (≤20%) | 1 (ref) | | |
| | 2 (20–40%) | 1.55 (0.99, 2.42) | 0.054 |
| | 3 (40–60%) | 2.18 (1.43, 3.31) | 0.0003 |
| | 4 (60–80%) | 2.49 (1.64, 3.78) | 1.9 × 10^{-3} |
| | 5 (≥80%) | 3.45 (2.31, 5.17) | 1.7 × 10^{-3} |
| dGRS_{1000} quintile 5 vs. quintiles 1–4 | 1.95 (1.52, 2.51) | 1.8 × 10^{-4} |
| All-cause mortality (1,994/10,273) | dGRS_{1000} (effect per 1 SD) | 1.03 (0.99, 1.08) | 0.14 |
| dGRS_{1000} quintile 5 vs. quintiles 1–4 | 1.14 (1.02, 1.27) | 0.019 |
| Cardiovascular mortality (1,069/10,273) | dGRS_{1000} (effect per 1 SD) | 1.06 (1.00, 1.13) | 0.046 |
| dGRS_{1000} quintile 5 vs. quintiles 1–4 | 1.27 (1.10, 1.46) | 0.0013 |

Hazard ratios (HR) with 95% confidence intervals (95% CI) from Cox proportional hazard modeling of the effect of dGRS_{1000} on incident T2D (in individuals aged 35–79 and free of T2D at recruitment) and on all-cause and cardiovascular mortality. All models are adjusted for BMI category, smoking level, waist-to-hip ratio, waist circumference, physical activity level, history of high blood glucose, fruit and vegetable consumption, and sex. Age is used as the time scale.

DISCUSSION

We have shown that the combined effect of a large number of genetic markers on T2D risk is of similar strength to (or even stronger than) that of some other known T2D risk factors. To use genetic predictors in practice, the effect of multiple markers needs to be summarized as a GRS. Our analysis shows that the strength of the GRS–disease association can vary greatly depending on how the GRS is calculated and how the SNPs (coded as effect allele count) are selected and weighted.

We propose a methodology called doubly weighting to increase the efficiency of the GRS and reduce the bias created by the “winner’s curse” (SNPs with an effect overestimated by chance in GWAS are oversampled in a list of top hits). A doubly weighted score, dGRSk, captures the effect of k “top” SNPs, but because it is uncertain whether a particular SNP belongs to the “top” set, probability weights for all available independent SNPs (7,502 in our case) are used to account for this uncertainty.

We found that when k ranges from 200 to 3,500 (5,200 in BMI-unadjusted models), the association between dGRSk and prevalent T2D in the Estonian Biobank cohort was clearly stronger than that of the best-fitting conventionally weighted GRS (Supplementary Table S2 online and Supplementary Figure S1 online). The best predictive accuracy was achieved with k = 1,000. This is an indicator that it is likely that the number of independent T2D-associated genomic loci is much larger than currently established by most recent GWAS studies.\footnote{11,17} However, adding too many SNPs to the score will introduce too much random noise, diluting any possible signals. It is possible that an even larger-scale GWAS meta-analysis could reduce the amount of noise (smaller standard errors for parameter estimates); therefore, one could add more markers with weaker signals to the score, leading to further improvements in predictive accuracy.

Our analysis suggests that large-scale genome-wide studies have the potential to explain considerably more variability in the target phenotypes than currently estimated. We have shown that even a partial correction of some common biases in GRS (caused by the “winner’s curse”) may lead to a reduced amount of “missing heritability” (the proportion of genetic variability that is still unexplained by measurable genetic markers).

To demonstrate the potential of the GRS for practical use in personalized risk assessment, it is important to address the added value once the other known predictors are already accounted for. We assessed the effect of GRS when added to other predictors in a model for incident disease and found that the hazards of incident T2D in the top and bottom dGRS_{1000} quintiles differ by approximately 3.5 times. Although obesity is the strongest predictor for T2D, our data suggest that the effects of BMI and GRS on the T2D risk are additive; therefore, an individual with relatively high BMI but with low genetic risk can have the same overall T2D risk level as someone at high genetic risk but average BMI. Despite the observed significant association of the GRS with blood glucose, triglycerides, and HDL-cholesterol levels in T2D-free individuals, the effect of dGRS_{1000} on disease incidence remained practically unchanged after adjustment for these factors, thus suggesting the additive effect of genetic risk while accounting for these parameters.
In conclusion, our analysis suggests that the genetic risk factors for T2D that are captured in the dGRS\textsubscript{100} would improve the predictive accuracy when added to either a (nongenetic) T2D risk score (such as FINDRISC\textsuperscript{18}) or a clinical risk assessment algorithm.

To quantify the potential improvement in risk assessment efficacy, ROC analysis (c-statistics) is an obvious methodological choice. Because age and BMI are already very strong predictors for T2D, the additional incremental value of GRS may seem small\textsuperscript{15} (as also shown in previous studies\textsuperscript{7}). However, the clinically relevant quantity is not the relative contribution of GRS in comparison with age or BMI, for instance, but its ability to distinguish between different individual risk levels in subjects of the same age and BMI. Although the overall improvement in c-statistic for the incident T2D in individuals aged 35–79 is 1.2\%, this increased to 2.1\% in a subset with BMI of 25–35.

It has been debated whether genetic risk estimates based on DNA markers would add any meaningful information in cases where the family history of T2D is known, and it has been shown that complete family history provides better prediction than that achieved using 21 SNPs.\textsuperscript{19} However, the highly polygenic nature of T2D suggests that parent and offspring genetic risk may actually differ considerably because, on average, only half of the risk-affecting alleles are transmitted from each parent to the child. Moreover, in current family structures, it is increasingly difficult to obtain detailed information, if any, from both parents. Therefore, our study suggests that as the number of SNPs included in the GRS increases, the accuracy of GRS-based risk estimate improves in comparison to that based on family history.

One of the limitations of our study was that we have demonstrated the effect of GRS in one ethnically homogenous cohort only. We believe that the conclusions would be similar for other ethnicities as well, because trans-ethnic analyses of T2D have suggested that the effect sizes for common SNPs are relatively homogenous across ethnicities.\textsuperscript{15} However, one should be cautious about extrapolating our results to other populations without further validation.

We also see room for further improvements in the GRS methodology because our method offers only partial correction of the “winner’s curse” bias (SNPs with effects overestimated by chance still tend to obtain larger weights). One could also consider allowing for possible inclusion of correlated SNPs by taking the correlation structure (linkage disequilibrium between SNPs) into account.

In conclusion, we have shown that the proposed GRS is predictive for T2D risk in the long term (from birth on), as reflected by the sex- and age-adjusted comparisons of prevalent cases and controls. We have also demonstrated good predictive ability in the short term when other risk factors such as obesity, diet, physical activity, HDL cholesterol, and blood glucose level have already accounted for (analysis of T2D incidence). Although a longer follow-up time with a greater number of incident T2D cases would enable more precise effect estimation, our results indicate that a GRS with high accuracy, such as dGRS\textsubscript{100}, would significantly improve the best existing risk assessment algorithms for T2D, encouraging its implementation in the practice of personalized medicine.
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