A gene-diet interaction-based score predicts response to dietary fat in the Women's Health Initiative: Westerman et al.

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Table S1: Sample size necessary to achieve power of 0.8

| GxE variance explained (%) | N (nominal) | N (suggestive) | N (genome-wide) |
|----------------------------|-------------|----------------|-----------------|
| 0.05 | 14046 | 49488 | 70866 |
| 0.10 | 7021 | 24737 | 35423 |
| 0.50 | 1401 | 4936 | 7069 |
| 1.00 | 699 | 2461 | 3524 |

Power calculations were undertaken using the Quanto tool, with parameters set as follows: additive model, SNP main effect of 0.5% of trait variance, binary environment with 50% prevalence, and environmental effect explaining 10% of the trait variance.

Table S2: Responder score effects on CRF changes in DM trial participants across main-effect filter thresholds

| | | All variants | | | Nominal main effect (p<0.05) | | | Suggestive main effect (p<1e-5) | | |
|----------------------|-------|---------------------|---------|---------|------------------------------|---------|---------|---------------------------------|---------|---------|
| CRF | N^1 | # SNPs ² | SES^3 | P-value | # SNPs ² | SES^3 | P-value | # SNPs ² | SES^3 | P-value |
| BMI | 1988 | 158365 | 0.049 | 0.026 | 6042 | 0.027 | 0.221 | 569 | 0.027 | 0.236 |
| FG | 281 | 161906 | -0.078 | 0.495 | 1924 | 0.016 | 0.791 | 7 | 0.006 | 0.92 |
| HDL-C | 150 | 153942 | 0.028 | 0.63 | 1731 | -0.06 | 0.471 | 42 | 0.085 | 0.258 |
| LDL-C | 145 | 156313 | 0.035 | 0.729 | 1760 | -0.179 | 0.026 | 46 | 0.01 | 0.901 |
| SBP | 2004 | 153942 | -0.016 | 0.473 | 1536 | 0.029 | 0.196 | 6 | 0 | 0.999 |
| TG | 150 | 152006 | -0.184 | 0.203 | 1774 | -0.139 | 0.066 | 47 | -0.043 | 0.573 |

¹ Sample size available with 1-year follow-up for each CRF

² Number of SNPs selected by the pruning-and-thresholding algorithm for each CRF-threshold combination

³ Standardized effect size (SES) represents the regression coefficient estimate in terms of CRF standard deviation per responder score standard deviation

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Table S3: Responder score effects on CRF changes in DM trial participants across ancestries

| | All combined | | | Black | | | Hispanic | | |
|----------------------|------------------|---------|---------|---------------------------|---------|---------|---------------------------|---------|---------|
| CRF | $\overline{N^1}$ | SES^2 | P-value | $\overline{\mathrm{N}^1}$ | SES^2 | P-value | $\overline{\mathrm{N}^1}$ | SES^2 | P-value |
| BMI | 3606 | -0.02 | 0.24 | 1214 | 0 | 0.994 | 404 | 0.018 | 0.719 |
| FG | 572 | 0.08 | 0.043 | 214 | -0.004 | 0.955 | 77 | -0.014 | 0.901 |
| HDL-C | 430 | -0.06 | 0.224 | 206 | -0.096 | 0.182 | 74 | -0.057 | 0.66 |
| LDL-C | 422 | 0.084 | 0.066 | 206 | 0.055 | 0.412 | 71 | 0.126 | 0.227 |
| SBP | 3645 | 0.041 | 0.015 | 1230 | 0 | 0.991 | 411 | -0.008 | 0.868 |
| TG | 430 | -0.075 | 0.112 | 206 | -0.023 | 0.743 | 74 | -0.098 | 0.433 |

¹ N = sample size available with 1-year follow-up measurements for the CRF in question

Table S4: LDL-FRS effects on alternate CRF changes in DM trial participants

| Outcome risk factor | # SNPs in risk score | N^1 | Std. effect size ² | P-value |
|---------------------|----------------------|-------|-------------------------------|---------|
| BMI | 1760 | 1988 | -0.02 | 0.36 |
| SBP | 1760 | 2004 | 0.00 | 0.83 |
| HDL-C | 1760 | 150 | -0.07 | 0.39 |
| TG | 1760 | 150 | 0.11 | 0.16 |
| FG | 1760 | 281 | 0.01 | 0.82 |

 $^{^1~\}mathrm{N} = \mathrm{sample}$ size available with 1-year follow-up measurements for the CRF in question

² Std. effect size represents the regression coefficient estimate in terms of CRF standard deviation per responder score standard deviation

² Std. effect size represents the regression coefficient estimate in terms of CRF standard deviation per responder score standard deviation

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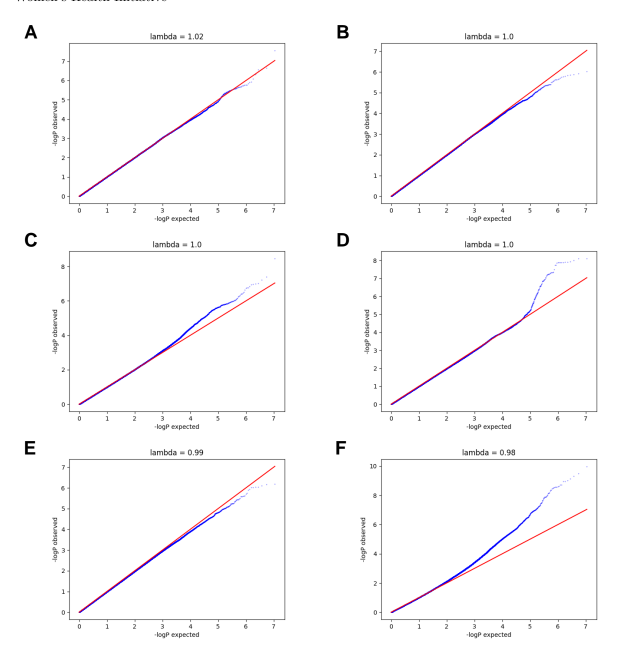


Figure S1: Q-Q plots from individual CRF GWIS. The distribution of p-values from each GWIS is plotted against the expected uniform p-value distribution. Plots correspond to: A) BMI, B) SBP, C) LDL-C, D) HDL-C, E) TG, and F) FG. Lambda values above each plot represent genomic inflation estimates. BMI: body mass index, SBP: systolic blood pressure, LDL-C: LDL cholesterol, HDL-C: HDL cholesterol, TG: triglycerides, FG: fasting glucose.