Genetic determinants of obesity-related lipid traits

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Abstract In our ongoing effort to identify genes influencing the biological pathways that underlie the metabolic disturbances associated with obesity, we performed genome-wide scanning in 2,209 individuals distributed over 507 Caucasian families to localize quantitative trait loci (QTLs), which affect variation of plasma lipids. Pedigree-based analysis using a quantitative trait variance component linkage method that localized a QTL on chromosome 7q35-q36, which linked to variation in levels of plasma triglyceride [TG, logarithm of odds (LOD) score = 3.7] and was suggestive of linkage to LDL-cholesterol (LDL-C, LOD = 2.2). Covariates of the TG linkage included waist circumference, fasting insulin, and insulin:glucose, but not body mass index or hip circumference. Plasma HDL-cholesterol (HDL-C) levels were suggestively linked to a second QTL on chromosome 12p12.3 (LOD = 2.6). Five other QTLs with lower LOD scores were identified for plasma levels of LDL-C, HDL-C, and total cholesterol. These newly identified loci likely harbor genetic elements that influence traits underlying lipid adversities associated with obesity.—Sonnenberg, G. E., G. R. Krakower, L. J. Martin, M. Olivier, A. E. Kwitek, A. G. Comuzzie, J. Blangero, and A. H. Kissebah. Genetic determinants of obesity-related lipid traits. J. Lipid Res. 2004. 45: 610–615.

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Obesity, a highly prevalent and chronic disorder, which carries significant health and economic costs, is associated with decreased longevity and increased risks from such health hazards as insulin resistance and diabetes, elevated blood pressure, and heart disease. Obesity, and abdominal obesity in particular, is strongly associated with an adverse dyslipidemic profile and an increased risk for coronary artery disease (1, 2). While this relationship has been described in detail, its pathophysiological basis remains unclear. Several studies have suggested a biologic cascade in which enhanced plasma flux of free fatty acid (FFA) from highly lipolytic, insulin-resistant adipocytes of the visceral-abdominal region in the basal state, along with its poor suppressibility by insulin in the fed state, could lead to overproduction of triglyceride (TG) and apolipoprotein B (apoB), with subsequent channeling into the small, dense subfractions of VLDL and LDL (1, 3–5). Furthermore, the compensatory hyperinsulinemia of insulin resistance may directly influence hepatic production of these lipid-lipoprotein subcomponents to augment this process. Prospective twin and familial segregation studies have also suggested the existence of strong genetic determinants for this abnormal lipid-lipoprotein profile (6, 7).

In our search for the genetic etiology of the obesity-related metabolic adversities, we performed phenotyping and genotyping of individuals from 507 Caucasian families recruited through an obese proband and identified quantitative trait loci (QTLs) on chromosomes 3 (3q27) and 17 (17p12), which were strongly linked to variation in total body adiposity, abdominal fat partitioning, and insulin resistance (8). The present study was undertaken to identify QTLs that affect the obesity-related plasma lipid profile.

METHODS

Study participants

Details of recruitment strategies, ascertainment, and phenotyping characteristics of the individuals and families studied have been presented previously (8). Briefly, families with at least two obese siblings (body mass index [BMI] > 30), availability of one (preferably both) parent, and one or more never-obese siblings (BMI ≤ 27) were ascertained. These were recruited from the TOPS (Take Off Pounds Sensibly, Inc.) membership in 10 states of the midwestern region of the US (WI, IL, MI, IA, MN, OH, WV, MO, KY, and IN). A questionnaire garnered personal data

Abbreviations: apoB, apolipoprotein B; BMI, body mass index; cM, centiMorgan; cNOS, endothelial nitric oxide synthase; HDL-C, HDL-cholesterol; IBD, identity-by-descent; LDL-C, LDL-cholesterol; LOD, logarithm of odds; NO, nitric oxide; QTLs, quantitative trait loci; STRP, short tandem repeat polymorphism; TG, triglyceride; TNF, tumor necrosis factor; total-C, total cholesterol.

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on each individual detailing his/her health, including asthma, kidney or liver disease, hypertension, heart disease, stroke, thyroid disorders, diabetes, medications, menopausal status and hormonal therapy, weight history, and smoking history. Exclusion criteria included pregnancy, type 1 diabetes, history of cancer, renal or hepatic disease, severe coronary artery disease, substance abuse, corticosteroids or thyroid dosages above replacement dose, history of weight loss of more than 10% in the preceding 12 months, as well as individuals receiving lipid-lowering medications. Phenotypic measurements included BMI, waist and hip circumferences, fasting plasma levels of glucose, insulin, insulin/glucose ratio, cholesterol, LDL-cholesterol (LDL-C), HDL-cholesterol (HDL-C), and plasma TG. A total of 2,209 individuals distributed over 507 Caucasian families of predominantly northern ancestry completed the ascertainment and phenotyping procedures. As reported previously (8), the pairwise relationships within this cohort represented a total of 5,399 relative pairs, with the majority being nuclear family (2,177 parent-offspring and 2,198 sibling) relationships. All protocols have been approved by the Institutional Review Board of the Medical College of Wisconsin and follow IRB guidelines for patient safety and confidentiality. All subjects have given informed consent prior to their participation in these studies.

Phenotyping procedures

BMI (kg/m²) was determined from body weight and height. Waist and hip circumferences (cm) were measured according to the World Health Organization criteria. With participants wearing light clothing, waist circumference was the minimal measurement at the navel region and hip circumference the widest measurement at the hip and buttocks. Fasting plasma TG and cholesterol levels were determined spectrophotometrically. TG assay kits were obtained from Stanbio Laboratory, Inc. (San Antonio, TX), and cholesterol kits from Roche-Boehringer (Indianapolis, IN). HDL-C was determined after phosphotungstic acid/MgCl₂ precipitation, and LDL-C was measured directly after immunospecific separation (Diagostics, St. Louis, MO). All determinations were performed in triplicate. Quality controls were performed to assure stability and reliability of the assays. The respective intraassay and interassay coefficients of variation for the lipid analyses were 4.7% and 5.3% for TG, 5.5% and 6.7% for cholesterol, 5.7% and 6.1% for HDL-C, and 2.9% and 7.5% for LDL-C.

Genotyping procedures

Genomic DNA from all samples was prepared from whole blood with commercial reagents (Puregene, Gentra Systems, Minneapolis, MN) using a nonphenol-based approach. DNA was amplified by PCR and screened using fluorescently labeled primers (Weber screening set 9, Research Genetics, Huntsville, AL) at the Marshfield Medical Research Foundation. This set included 387 short tandem repeat polymorphism (STRP) markers, including 366 autosomal, 17 X-chromosomal, and 4 Y-linked markers (9), and yielded an average map density of approximately 10 centiMorgan (cM). Products were analyzed using automated high-throughput scanning fluorescence detectors, each simultaneously detecting three separate dyes as previously described (8). All reported family relationships between individuals were validated by calculating likelihoods of the relationships based upon the autosomal genotype data (10). Errors in the relations between individuals were ascertained, and as a result, data on eight proband families (70 individuals) were discarded. The genotypic data were examined for Mendelian inconsistencies, and those genotypes proven to be inconsistent were removed. The average heterozygosity of the markers used was 0.79 ± 0.06, and the sex-averaged genetic spacing was 9.1 ± 3.8 cM.

Variance components linkage analysis

A variance component model applied to extended family data was used to test for evidence of linkage with phenotypes related to the metabolic syndrome, using a 10 cM genome-wide map of STRP markers. As described previously (8), an extension of the strategy developed by Amos (10) was used to estimate the genetic variance attributable to a specific chromosomal location (11). This approach is based upon specifying the expected genetic covariances between arbitrary relatives as a function of the identity-by-descent (IBD) relationships at a given marker locus.

Basic variance component linkage analysis includes a QTL-specific component, which was used to test for linkage in our analysis. Using a variance component model (12), we tested the null hypothesis that the additive genetic variance due to a QTL ($\sigma_q^2$) equals zero (no linkage) by comparing the likelihood of this restricted model with that of a model in which $\sigma_q^2$ is estimated. The difference between the two log $\lambda$ likelihoods produces a logarithm of odds (LOD) score that is the equivalent of the classical LOD score of linkage analysis. Twice the difference in log likelihoods of these models yields a test statistic that is asymptotically distributed as a 1/2:1/2 mixture of a $\chi^2$ variable and a point mass at zero (13). This quantitative trait linkage method has been implemented in the program package SOLAR (11), which determines whether genetic variation at a specific chromosomal location can explain the variation in the phenotype (10, 12, 14).

Prior to the linkage analysis, values for kurtosis, the distributional attribute most likely to contribute to inflation of type I error, were determined and found to range from 0.2 to 1.8 (8), well within the acceptable range for the analytical methods employed. Genetic heritabilities for all phenotypes were estimated, and linkage scores were calculated after adjusting the observed LOD scores using the estimated correction constants, as described by Blangero et al. (15, 16), and after adjusting for a standard set of covariates found to influence the phenotypes (8). The set of covariates for these analyses, incorporated into the SOLAR package, included age, sex, and interactions between sex and age, smoking, menstrual status, intake of estrogen or birth control steroids, and history of asthma or diabetes.

Simulation studies suggested that the likelihood ratio test yields expected nominal $P$ values for a wide variety of trait distributions (17); however, as an added precaution, we also obtained an empirical distribution for the LOD scores for this sample by simulation of over 10,000 replicates (15, 16). This approach allows for the calculation of expected LOD scores that are then regressed on the observed LOD scores to obtain the appropriate correction constant. In addition, genome-wide $P$ values were calculated by using a method that adjusts for variation in chromosome-specific marker densities (18).

The use of the variance component approach requires an estimate of the IBD matrix. For this TOPS cohort, a pairwise maximum likelihood-based procedure was used to estimate IBD probabilities (11). To permit multipoint analysis for QTL mapping, we employed an extension (13) of the technique of Fulker and colleagues (19). Estimates of the IBD probabilities were generated at any point on a chromosome by using a constrained linear function of observed IBD probabilities of markers at known locations within the region. This multipoint procedure, which yields substantially greater power to localize QTLs than two-point methods, enabled direct localization of the QTL and construction of confidence intervals. One-LOD unit support intervals were obtained by identifying the peak for the maximal LOD score on the plot of the multipoint results, dropping down one LOD unit, and finding the chromosomal region defined by the shoulders of the curve. For the current data set, a LOD-score
evaluation was performed every centiMorgan along the chromosome, the distances between markers having been determined using CRI-MAP (20).

**RESULTS**

In our study cohort of 507 Caucasian families recruited through an obese proband, we aimed to localize QTLs that affect the complex alterations in plasma lipid profiles associated with obesity. Our data support the notion of a strong genetic determinant influencing variation in plasma lipid levels in our cohort. Table 1 shows the means, ranges, and additive genetic heritabilities of the lipid-lipoprotein phenotypes measured. The additive genetic heritabilities (h²), estimated after allowing for the effects of the covariates listed in Methods, ranged from 0.30 to 0.47.

Table 2 lists adjusted results of the linkage analysis of the total genome scan for fasting plasma TG, total cholesterol (total-C), LDL-C, and HDL-C. Seven QTLs with a LOD score of 1.9 or higher were localized. Most prominent among these is the QTL on chromosome 7 at position q35-q36 that is associated with both plasma TG and LDL-C. Figure 1 shows the multipoint linkage analysis plot of the TG phenotype on chromosome 7 (LOD = 3.7). The 1-LOD-unit support interval surrounding the QTL peak at 174 cM (near marker D7S3058) spanned from 166 cM to 182 cM. In an analysis performed independently from TG, plasma LDL-C level was suggestively linked (LOD = 2.2) at the same QTL.

**TABLE 1.** Plasma concentration and genetic heritability for the lipid traits determined

<table>
<thead>
<tr>
<th>Trait</th>
<th>n</th>
<th>Mean</th>
<th>Range</th>
<th>Additive Genetic Heritability (h²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TG, mg/dl</td>
<td>2,136</td>
<td>124.3 ± 62.1</td>
<td>40–357</td>
<td>0.30 ± 0.04</td>
</tr>
<tr>
<td>Total-C, mg/dl</td>
<td>2,136</td>
<td>197.2 ± 39.4</td>
<td>100–320</td>
<td>0.47 ± 0.06</td>
</tr>
<tr>
<td>HDL-C, mg/dl</td>
<td>2,139</td>
<td>38.6 ± 10.3</td>
<td>16–75</td>
<td>0.31 ± 0.05</td>
</tr>
<tr>
<td>LDL-C, mg/dl</td>
<td>2,137</td>
<td>119.1 ± 33.9</td>
<td>41–219</td>
<td>0.38 ± 0.06</td>
</tr>
</tbody>
</table>

TG, triglyceride; HDL-C, HDL-cholesterol; LDL-C, LDL-cholesterol; n, the number of subjects in whom each lipid trait was measured; total-C, total cholesterol. Values are given as mean ± SD.

**QTLs for HDL-C were found at 2p13.3, 12p12.3, and 15q14 (Table 2). Figure 2 shows the multipoint linkage analysis plot for HDL-C on chromosome 12, the locus where HDL-C showed its strongest linkage signal. The 1-LOD-unit support interval surrounding the QTL peak at 42 cM (between markers D12S373 and GATA91H06) spanned from 30 cM to 60 cM. Also, as shown in Table 2, additional linkages were found for total-C at 7q11.23 and 16q13, and for LDL-C on chromosome 6q21.**

Because of the known correlations between the plasma lipid profile and other features associated with obesity, linkage analyses were repeated to evaluate their contributions to the LOD score of the 7q35-q36 TG linkage. Inclusion of BMI or hip circumference measurements as an additional covariate only minimally influenced the linkage

**TABLE 2.** Chromosomal locations of lipid-lipoprotein phenotypes

<table>
<thead>
<tr>
<th>Phenotype</th>
<th>Locus</th>
<th>Location</th>
<th>LOD</th>
<th>Flank Markers</th>
</tr>
</thead>
<tbody>
<tr>
<td>TG</td>
<td>7q35-7q36</td>
<td>174</td>
<td>3.7</td>
<td>D7S3058</td>
</tr>
<tr>
<td>LDL-C</td>
<td>7q35-7q36</td>
<td>174</td>
<td>2.2</td>
<td>D7S3058</td>
</tr>
<tr>
<td></td>
<td>6q21</td>
<td>148</td>
<td>2.2</td>
<td>GATA184A08 D6S2436</td>
</tr>
<tr>
<td>HDL-C</td>
<td>12p12.3</td>
<td>42</td>
<td>2.6</td>
<td>D12S373 GATA91H06</td>
</tr>
<tr>
<td></td>
<td>13q14</td>
<td>39</td>
<td>2.3</td>
<td>ACTC D15S643</td>
</tr>
<tr>
<td></td>
<td>2p13.3</td>
<td>97</td>
<td>2.0</td>
<td>D2S1394</td>
</tr>
<tr>
<td>Total-C</td>
<td>7q11.23</td>
<td>96</td>
<td>2.3</td>
<td>D7S820</td>
</tr>
<tr>
<td></td>
<td>16q13</td>
<td>20</td>
<td>2.1</td>
<td>ATA41E04 D16S748</td>
</tr>
</tbody>
</table>

cM, centiMorgan; LOD logarithm of odds. The multipoint linkage results for the lipid parameters determined were adjusted for the standard set of covariates described in Methods (variance components linkage analysis). The numbers of subjects for each determination are given in Table 1. Locations (cM) are given by peak LOD.

**Fig. 1.** Multipoint linkage map and logarithm of odds (LOD) scores for triglyceride (blue) and LDL-cholesterol (red) on chromosome 7. The multipoint linkage results for the lipid parameters determined were adjusted for the standard set of covariates described in Methods (variance components linkage analysis).

**Fig. 2.** Multipoint linkage map and LOD scores for HDL-cholesterol on chromosome 12. The multipoint linkage results for the lipid parameters determined were adjusted for the standard set of covariates described in Methods.
power for TG at this locus, the LOD score decreasing to 3.2 by the inclusion of either covariate. Inclusion of waist circumference, however, reduced this score to 2.7, while inclusion of insulin reduced it to 2.0 and inclusion of insulin:glucose to 1.7. The location of the maximal LOD score did not change during these evaluations, nor did the inclusion of these additional covariates affect the heritability estimate for TG.

DISCUSSION

We have localized a QTL on human chromosome 7q35-q36 that affects both plasma TG and LDL-C variation in Caucasian families of predominantly northern European ancestry. Several suggestive QTLs were also localized, including a region on chromosome 12p12.3, which affects plasma HDL-C variation. The large cohort of families studied, the selection of families including both obese and never-obese individuals, and the magnitude of the LOD scores identified emphasize the important contributions of these loci to the genetic architecture associated with the obesity-related lipid disorder.

The additive genetic heritability estimates for the lipid-lipoprotein phenotypes analyzed in our cohort range from 0.30 to 0.47 and strongly suggest the existence of genetic factors, which influence expression of this adverse lipid profile. Similar degrees of heritability influencing plasma levels of the same lipid components have previously been reported in twin, adoption, and family studies in nonobese normal subjects and individuals with established coronary artery disease (21–23). Our confidence in the stability of the QTL on chromosome 7 (7q35-q36) is also heightened by its replication in other studies. In a cohort of Mexican-Americans, Duggirala et al. (24) identified a locus harboring susceptibility genes for hypertriglyceridemia in this same region, while analysis of results from Caucasian families in the Framingham Heart Study provided evidence for a gene near this locus that affects the ratio of plasma TG and HDL-C (25). Moreover, a susceptibility locus for hereditary pancreatitis (26), a disorder frequently resulting in hypertriglyceridemia, has been mapped to this location.

Our analyses demonstrate that variation in plasma LDL-C within our population, analyzed independently from TG, is linked to the same QTL on chromosome 7q35-q36. This single QTL may impact variation on both phenotypes. Previous research has demonstrated a strong statistical correlation between plasma TG levels and alterations in the spectrum of LDL particles (27–29). Our linkage analysis provides support for a genetic origin to this correlation. Further studies to evaluate the LDL density spectra in our study cohort and their relationship to this reported linkage are in progress.

Reduced level of plasma HDL-C is a fundamental feature of the obesity-related lipid disorder, particularly that associated with abdominal obesity. The contribution of genetic factors to HDL-C plasma level variation has been reported previously (30, 31). The suggestive QTL on chromosome 12p12.3 that we identified in our analysis is known to include several candidate genes that could influence a broad spectrum of events regulating lipid-lipoprotein metabolism. One candidate gene is LOX-1 (OLR1), which encodes the human oxidized lipoprotein receptor 1 gene, a cell-surface receptor and a potent mediator of endothelial dysfunction (32). This receptor specifically binds, internalizes, and degrades oxidized LDL (33), which is essential to the development of atherosclerotic lesions (34, 35). A second candidate gene is the LDL-related protein 6, which participates in the clearance and intracellular processing of TG-rich lipoproteins (36). The 12p13 region includes the TNFRSF1 and TNFRSF7 genes, members of the tumor necrosis factor (TNF) receptor superfamily, which appear to influence lipoprotein metabolism and cardiovascular integrity (37, 38). Secretion of TNFs and other proinflammatory cytokines is now viewed as an important facet of obesity-related events leading toward the metabolic syndrome (39, 40). Another candidate near this region is the gene for the apoB mRNA editing enzyme (APOBEC1), which is involved in the catabolism of apoB-containing lipoproteins (41, 42).

Our linkage analyses have identified a number of additional suggestive QTLs to plasma variation in LDL-C, HDL-C, and total-C, some of which have been reported previously in other studies. In a Mexican-American population ascertained on a proband with type 2 diabetes mellitus, Duggirala et al. (24, 43) identified loci for susceptibility genes for hypertriglyceridemia on chromosomes 7 and 15. A significant linkage (LOD = 3.9) was identified with TG at 15q near 20 cM, while a suggestive linkage was found on chromosome 7 (near markers D7S506 and D7S653 at 90–100 cM). The latter linkage is close to our suggestive locus at 96 cM for total-C, while the linkage reported on chromosome 15 (between markers GABRB3 and D13S165 around 20–30 cM) is near the locus we report for HDL-C at 39 cM. Our suggestive linkage for HDL-C on chromosome 2 at 97 cM also replicates with a region linked to low HDL-C level at 91 cM in Finnish families (31). Whether these suggestive linkages are false positives or actually represent additional regions contributing to the complexity of the genetic architecture associated with obesity-related disorders remains to be evaluated.

The QTL on chromosome 7q35-q36 affects both TG and LDL-C, and the effect on TG is influenced by covariates representative of abdominal body fat topography (waist circumference) or insulin resistance (insulin and insulin:glucose). This suggests that this QTL harbors gene(s) that not only directly or indirectly affect plasma lipid levels, but also have a significant impact on other phenotypes associated with obesity and progression toward the metabolic syndrome. One potential candidate in this chromosomal region is the gene for endothelial nitric oxide synthase (eNOS). Among its many functions, nitric oxide (NO) mediates monocyte and leukocyte adhesion and smooth muscle cell proliferation, while many of the metabolic derangements associated with obesity and its associated metabolic syndrome, including hyperglycemia, elevated levels of plasma FFA, insulin resistance, and en-
dothelial cell function, are influenced by levels of NO (44). NO production is stimulated through the activation of NF-κB through elevated levels of glucose, FFA, or insulin and by proinflammatory cytokines such as TNFα. In addition to its intrinsic actions, NO is a scavenger for reactive oxygen species and regulates expression of redox-regulated genes (45). Oxidative stress, which has been implicated in the development of insulin resistance and type 2 diabetes (46), may also contribute to the atherosclerotic process through the production of oxidized LDL. Several common polymorphisms in this gene influence susceptibility to atherosclerosis (47, 48), as well as hypertension (49, 50), insulin resistance, and type 2 diabetes (51, 52). To our knowledge, no association studies have been performed between eNOS and the obesity-related lipid profile. Further studies will need to investigate whether eNOS or its metabolites, possibly in conjunction with the products of LOX-1 and TNFRSF1 and/or TNFRSF7, contribute to the etiology of these lipid disturbances.

Our studies strongly suggest that these QTL areas contain gene(s) and/or regulatory elements that affect the adverse lipid profile associated with obesity and may play important roles in the subsequent development of cardiovascular complications.

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