Genetics of obesity

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Abstract

Objective: The aim was to review and update advances in genetics of obesity.

Design: Analysis and interpretation of recent investigations about regulating the energy balance as well as about gene-nutrient interactions and current nutri-genomic research methods.

Background and main statements: Obesity results from a long-term positive energy balance. However, its rising prevalence in developed and developing societies must reflect lifestyle changes, since genetic susceptibility remains stable over many generations. Like most complex diseases, obesity derives from a failure of adequate homeostasis within the physiological system controlling body weight. The identification of genes that are involved in syndromic, monogenic and polygenic obesity has seriously improved our knowledge of body weight regulation. This disorder may arise from a deregulation at the genetic level (e.g. gene transcription or altered protein function) or environmental exposure (e.g. diet, physical activity, etc.).

Conclusions: In practice, obesity involves the interaction between genetic and environmental factors.

Keywords
Obesity
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Energy expenditure

While it is clear that environmental factors play a significant role in the development of obesity, research work over the last decades has clearly documented a genetic contribution to obesity-related phenotypes as well. Obesity and its related traits represent the archetype of a common complex phenotype. The configuration of multiple genes can range from polygenic (i.e. many genes with a relatively small contribution) to oligogenic (i.e. few genes with large measurable effects often expressed on a residual polygenic background)1. Indeed, it is this oligogenic architecture that justified all current efforts to map genes for complex phenotypes.

Prior to the mid-1990s, most of the work on genetics of human obesity had been limited to demonstrating that variation in a wide range of obesity-related phenotypes could be attributable to varying degrees to the effects of genes2. Quantitative genetics analyses (twins, siblings and family studies) have shown significant heritability, individual’s chances of being obese are increased when having obese relatives, with estimates of heritability generally ranging from 30% to 70%. Different sets of monozygotic twins overfed showed differences in the degree to which excess calories were stores of fat, but the tendency towards increased adiposity within each set of twins were very similar, indicating that genetic factors play a major role in the individual susceptibility to gain weight in a specific environment3–5.

From the late 1980s through the early 1990s, segregation analysis was utilised in an effort to dissect further the genetic component underlying human obesity by attempting to detect the effects of anonymous major genes on obesity-related phenotypes. A number of these studies have reported consistent evidence of an anonymous major gene effect accounting for as much as 40% of the variation in body mass index (BMI) or fat mass in a variety of populations2.

By the early to mid-1990s, researchers began to focus on the goal of identifying the specific genes involved. Initially, such efforts to identify specific genes influencing complex traits such as those associated to obesity relied on the use of a priori selected candidates genes. Such candidates genes are selected on the basis of their perceived role or function in biochemical pathways relevant to the specific phenotype of interest (e.g. the structural gene for a circulating protein). The conservation of hypothalamic pathways in rodents and humans has certainly aided in choosing suitable candidates genes. All the spontaneously occurring obesity mutations in mice either have been found to harbour functionally relevant mutations in humans too or have led to the identification of a system/pathway in which other genes were found to be mutated. The tools recently developed allow easy determination of most gene polymorphisms from a blood sample, especially single-nucleotide polymorphism...
Monogenic obesity (>173 cases in the word)

- LEP, LEPR, POMC, MC4R, MC3R,
- CRHR1-2, GPR24,
- SIM1, PCSK1.

Polygenic obesity (113 candidate genes)

- ADRB1, ADRB2, ADRB3,
- UCP1, UCP2, UCP3,
- PPARG, PPARD,
- ACDC, TNF alpha, IL6,
- AGRP, GHLR, GNB3, etc.

Fig. 1 Genetics of obesity

The mechanisms participating in the effects of leptin and other peptides on food intake and body weight regulation are now becoming clearer. Certain areas of the hypothalamus are rich in specific receptors binding regulatory peptides and triggering central regulatory mechanisms. Factors acting at the central nervous system level include neuropeptide Y (NPY), corticotropin-releasing hormone, proopiomelanocortin, α-melanocyte-stimulating hormone, agouti-related protein, melanin-concentrating hormone, cocaine- and amphetamine-regulated transcript and other molecules. Interaction between them involving complex neuronal mechanisms eventually influence the behaviour and provide important links with neuroendocrine regulation of other vital functions of the organism. Studies in humans have failed to find leptin or any other mutant gene to be the unique ‘obesity gene’. Conversely, multifactorial patterns involving actions of numerous polymorphic gene products now look more likely. Evidence is accumulating that most of these genes encoding central peptide factors as well as their receptors (leptin receptors, melanocortin receptors, NPY receptors) are polymorphic. Dominant inheritance of obesity conferred by missense, nonsense and frameshift mutations in the melanocortin 4 receptor (MC4R) gene has been extensively reported in many populations including French, English, German, American, Italian and Spanish individuals. It has been estimated that 1–6% of extremely obese individuals harbour functionally relevant MC4R mutations. More than 70 mutations of the MC4R, 57 non-synonymous, 5 nonsense and 10 frameshift mutations, have been reported, many of them associated with dominant inheritance of obesity. Functional studies showed that many of the missense mutations also lead to a loss-of-function of the MC4R. Meanwhile, other mutations (i.e., Thr-11-Ser, Arg-18-Cys) and two polymorphisms (Val-103-Ile, Ile-251-Leu) did not modify the function of the MC4R in vitro.

NPY is released from the arcuate hypothalamic nucleus in fasting or in hypoglycaemia situations, its secretion being inhibited after food intake. The Leu7Pro polymorphism in the NPY gene appears to be implicated in lipid metabolism regulation. Some works reported that carriers of the Pro7 allele had higher NPY levels and also body fatness.

A number of peptides synthesised along the gastrointestinal tract also affect food intake. They include ghrelin (orexigenic peptide mainly produced in the stomach), cholecystokinin (produced in the small intestine acting as a short-term satiety signal) and peptide YY3-36 (produced in the colon and suppressing appetite for up to 12 hours). Exploration of these signalling pathways has started and it is becoming clear that polymorphism in relevant genes may have important functional consequences. For the ghrelin receptor gene, two SNPs were reported: Aku204Glu and Phe279Leu, which selectively impair the constitutive activity of the receptor.
in humans leading to short stature and obesity that apparently develop during puberty\textsuperscript{12}.

Moreover, the identification of relevant genes related to food preferences has just started. A novel family of 40–80 human and rodent G protein-coupled receptors expressed in taste receptor cells of tongue and palate epithelia has been identified. Taste 2 receptors (T2Rs) have been shown to function as bitter taste receptor and T1Rs as putative receptor for sweet taste. There is no information on polymorphism in the T1R family genes while some SNPs in T2R have been reported\textsuperscript{13,14}. Rapid progress has been made in this field to elucidate the genetic mechanism controlling formation of food preferences.

**Genes encoding factors implicated in energy expenditure**

The adaptive thermogenesis in humans is closely related to the active mobilisation of lipids from fat tissues and demand special interest in relation to obesity. Central neural pathways responsible for the food intake and energy expenditure regulation are tightly interconnected. The peripheral transmission of central commands to the fat stores is mediated by the sympathetic nervous system. \(\beta\)-adrenoceptor gene families (\(ADRB2\), \(ADRB3\), \(ADRB1\)) are intensively studied candidate genes in the obesity field for their participation in energy expenditure regulation.

The \(\beta2\)-adrenergic receptor gene (\(ADRB2\)) encodes a major lipolytic receptor protein in human fat cells. Two common polymorphisms of the \(ADRB2\) gene, characterised by an amino acid replacement of arginine by glycine in codon 16 (Arg16Gly) and glutamine by glutamic acid in codon 27 (Gln27Glu), have been explored in several diseases such as hypertension and obesity\textsuperscript{2,15–18}. A relationship between the Arg16Gly polymorphism and an altered function of the \(ADRB2\) has been reported leading to a decreased agonist sensitivity. Meanwhile, the Gln27Glu variant was also found to be linked to obesity in some populations. In men, the 27Glu allele has been associated with increased BMI and subcutaneous fat and with elevated leptin and triglycerides levels, while in women, the 27Glu variant was reported to be linked to increased BMI, body fat mass and waist-to-hip ratio\textsuperscript{15–18}. However, other studies in Caucasians (Danish men, Austrian women and German subjects) found no association between the Gln27Glu variant of the \(ADRB2\) gene and obesity\textsuperscript{2,16}.

The \(\beta3\)-adrenergic receptor (\(ADRB3\)) protein plays a role in adipocyte metabolism mediating the rate of lipolysis in response to catecholamines and their agonists have potential anti-diabetes and anti-obesity properties\textsuperscript{2,19–22}. A common polymorphism in this gene, characterised by an amino acid replacement of tryptophan by arginine at position 64 (Trp64Arg), has been identified and may be linked to lower lipolytic activity and account for lipid accumulation in the adipose tissue. A number of articles have reported the relationship between the Trp64Arg variant of the \(ADRB3\) and obesity-related phenotypes. With regard to BMI, more than nine studies have shown a statistically significant association between BMI and the Trp64Arg polymorphism in a variety of populations from 134 to 856 subjects. Besides, two meta-analyses examining the effect of this mutation on BMI have been published for Caucasian populations. One includes 2447 subjects and the significant BMI difference among carriers and non-carriers of the mutation is 0.30\textsuperscript{22}. The second includes 7399 subjects but the results are negative for the association\textsuperscript{22}. This polymorphism has been associated with abdominal/visceral fat obesity in several populations such as Caucasians and Japanese subjects. Similarly, several studies carried out among Mexican American, Japanese and Caucasian women have shown that carriers of the Arg allele had a higher BMI and lower reduction in visceral fat after weight loss\textsuperscript{2,23}. Some authors, however, failed to reproduce the finding on \(\beta\)-adrenoceptors gene variants and further confirmation is required.

More recently, variants of the \(ADRB1\) gene have been studied in relation to obesity. The \(ADRB1\) is also considered a potential candidate gene for obesity because of its role in catecholamine-induced energy homeostasis. Stimulation of \(ADRB1\), a member of G-protein-coupled receptors, mediates energy expenditure and lipolysis in adipose tissue. In the C-terminal intracellular G-protein-coupling domain of \(ADRB1\), a polymorphism Arg389Gly has been identified, with the Arg389 allele vs. the Gly389 allele displaying enhanced adenylate cyclase activity in vitro\textsuperscript{24–25}.

Whereas \(\beta\)-adrenceptors participate in the regulation of adaptive thermogenesis as a component of sympathetic responses, uncoupling proteins (UCPs) are involved in the modulation of heat-generating uncoupled respiration at the mitochondrial level. They represent a family of carrier proteins localised in the inner layer of mitochondrial membranes\textsuperscript{22}. There are different members: UCP1, mostly expressed in brown adipose tissue, has a role in thermogenesis, UCP2 is ubiquitously present in any tissue and UCP3 is mainly expressed in the skeletal muscle and brown adipose tissue. Their putative role as ‘uncoupling proteins’ has been intensively explored; like UCP1, UCP2 mediates mitochondrial proton leak releasing energy stores as heat and thereby affecting energy metabolism efficiency\textsuperscript{22}. The actual functions for UCP2 and UCP3 proteins are still under investigation. It has been proposed that uncoupling proteins act as regulators of energy metabolism, they being fatty acid transmembrane transporters in the mitochondria facilitating proton exchange\textsuperscript{22}.

Moreover, a number of human studies indicated the relationship between UCP polymorphisms and exercise efficiency, resting energy expenditure, substrate oxidation,
energy metabolism, BMI, obesity risk, type 2 diabetes risk, leptin, fat accumulation, body weight changes, physical activity and so on. These observations have led to the consideration of UCP2 and UCP3 as candidate genes for obesity, given their function in the regulation of fuel metabolism.

Several UCP2 gene variants have been described: a G/A mutation in the promoter region −866G/A, a valine for alanine substitution at amino acid 55 in exon 4 (Ala55Val) and a 45 base pair insertion/deletion in the untranslated region of exon 8. The association between these polymorphisms of the UCP2 and various aspects of obesity have been intensively studied. From the literature, it seems that allele G in the promoter region of UCP2 increases obesity risk while it affords relative protection for type 2 diabetes. Meanwhile the Ala55Val polymorphism has shown to be associated with increased exercise efficiency. However, results concerning the exon 8 insertion allele of the UCP2 gene have been inconsistent. While no association with obesity was observed in a number of studies conducted in several populations, significant associations between the exon 8 insertion of the UCP2 gene and BMI or fat mass or presence of obesity were found with P values of 0.01, 0.001, 0.002 and 0.005. A sixth study reported an association of the exon 8 insertion allele with sleeping metabolic rate (P = 0.007).

There are also several UCP3 gene variants. In linkage studies, some of them have been associated to a higher obesity risk. Specifically, the −55C/T polymorphism in the promoter region of this gene has been associated with an elevated BMI, an increased level of adiposity or a greater waist-to-hip ratio. However, other authors have not found any relationship between this polymorphism and a higher risk of obesity or changes in the metabolic rate. Even some studies have reported an inverse correlation with BMI and the presence of −55C/T polymorphism.

Genes encoding factors implicated in adipogenesis

The last group of genes acting in connection with the peripheral regulation of energy expenditure comprises the transcription factors leading to adipogenesis and adipocytes differentiation. The key actor is peroxisome proliferator-activated receptor γ, particularly the adipose-specific isoform PPARγ2. In a meta-analysis examining the Pro12Ala polymorphism in 19 136 subjects, a positive association with BMI was found. In our study, the frequency of the Ala allele, similar to other Caucasian populations, was higher in obese subjects (allelic frequency 0.13) than in controls (0.08), suggesting that this polymorphism was associated with obesity. There is also information on the functional role of PPARγ gene variants. Some mutant proteins appear to have a reduced activity.

Monogenic obesity

The clinical features of human subjects with leptin (or leptin receptor), proopiomelanocortin (POMC), MC4R and proprotein convertase 1 (PC1) deficiency are often associated with an obese phenotype.

Congenital human leptin deficiency has been identified in subjects showing severe early-onset obesity (8 years and 86 kg, or 2 years and 29 kg) with intense hyperphagia and undetectable levels of serum leptin due to a frameshift mutation in the ob gene (deletion G133) in a homozygosis, which resulted in a truncated protein not secreted. Children with leptin deficiency had also profound abnormalities in T-cell number and function consistent with high rates of infection and childhood mortality from infections. Leptin therapy in these subjects has a major effect on appetite with normalisation of hyperphagia and reductions body weight. Leptin receptor-deficient subjects were also found, with the phenotype being similar to those with leptin deficiency. The birth weight was normal, but a rapid weight gain was seen in the first months of life, with severe hyperphagia and aggressive behaviour when food was denied. Basal temperature and resting metabolic rate were normal and they were normoglycemic with mildly elevated plasma insulin as reported for leptin-deficient subjects. But specifically, they also debut with mild growth retardation and impaired basal and stimulated growth hormone secretion.

Homozygous and heterozygous subjects for mutations in POMC have been found. In neonatal life these subjects showed adrenocorticotropic hormone (ACTH) deficiency (the POMC gene encoded ACTH and other peptides), the children have red hair and pale skin due to the lack of melanocyte-stimulating hormone (MSH) action at the melanocortin-1 receptors in skin and hair follicles. The POMC deficiency is associated with hyperphagia and early-onset obesity due to the lack of activation of the melanocortin-4 receptor.

Since 1998 many groups have reported at least 70 mutations in MC4R mostly associated with severe early-onset obesity. Other clinical features of MC4R mutation carriers are hyperphagia, accelerated linear growth in children and marked increase in bone mineral density. Probands with homozygous MC4R mutations show more severe obesity than their heterozygous relatives; thus, the mode of inheritance is codominant. The prevalence of pathogenic MC4R mutations varied, being the highest in obese adults selected for severe childhood obesity, suggesting that both factors, severe obesity and early age of onset, may be markers of MC4R mutations. It has been also shown that pathogenic MC4R mutations are more prevalent in northern European populations than in the Mediterranean or even Asian populations. The figures for prevalence are always quite low, for example 68 obese patients with pathogenic MC4R mutations out of...
2000 or even one mutation in a sample of 3000 obese patients in another published study. Functional analyses of these mutations allow us to classify them on the basis of their effects on receptor signalling. For instance, some authors demonstrated that mutations that caused intracellular retention of the receptor in vitro were associated with earlier age of onset and greater severity of obesity than other mutations.

The extremely low prevalence of pathogenic MC4R mutations in the general population underlies the fact that the obesity epidemic is not an epidemic of new mutations.

Subject carriers of PC1 mutations mainly have severe early-onset obesity, impaired prohormone processing and hypocortisolaemia. Another clinical feature is small intestine dysfunction, which may result from an erroneous maturation of propeptides within the PC1-secreting cells along the gut.

Syndromic forms of obesity

There are about 30 rare syndromes caused by discrete genetics defects or chromosomal abnormalities with obesity as a clinical feature in association with mental retardation, dysmorphic features and organ-specific abnormalities (i.e. pleiotropic syndromes). At least four syndromes show severe hyperphagia and/or other signals of hypothalamic dysfunction, indicating a plausible origin at the level of the central nervous system. The most frequent of these syndromes is Prader-Willi syndrome (prevalence 1/25 000) characterised by obesity, hyperphagia, hypotonia, mental retardation, short stature and hypogonadotropic hypogonism. It is usually caused by lack of the paternal segment 15q11.2-q12, either through deletion of the paternal critical segment (75%) or through loss of the entire paternal chromosome 15 with the presence of two maternal homologues in 22% of patients (uniparental maternal disomy). One suggested mediator of the obesity phenotype is ghrelin, the stomach-secreted peptide that increased appetite by interacting with the product of the gene under observation. The molecular causes that underlie the aetiology of syndromic obesity are far more complex than that for monogenic obesity. Positional genetic strategies have led to the recent identification of several causative mutations responsible for these syndromes; however, in many cases the defective gene product is a ubiquitously expressed protein of unknown function.

Genetic and dietary influences at the present time

The interaction of functional gene polymorphisms with environmental factors (gene–environment interactions) plays a substantial role in obesity risk. Gene–environment interaction implies that in combination the effect of genotype and environment deviates from the additive or multiplicative effects of the two factors. A simple statistical approach to the analysis of gene–environment interaction when dealing with a quantitative trait (e.g. BMD) is to use the statistical test analysis of covariance, with the quantitative trait being entered as the dependent variable, genotype and environmental factors being entered as main effects and with an interaction term between the genotypic and environmental factors. If this term is statistically significant, the implication is that there is a greater or lesser additive effect and an interaction is suggested. However, this outcome could be the result of chance alone, which highlights the importance of confirmatory findings. If the P value for the interaction term is not significant, the implication is that there is a lack of interaction, but it could also simply reflect the lack of power in the study to detect an effect.

From a mechanistic viewpoint, interaction suggests that at the molecular level the effect or byproducts of the two factors. A simple statistical approach to the analysis of gene–environment interaction when dealing with a quantitative trait (e.g. BMD) is to use the statistical test analysis of covariance, with the quantitative trait being entered as the dependent variable, genotype and environmental factors being entered as main effects and with an interaction term between the genotypic and environmental factors. If this term is statistically significant, the implication is that there is a greater or lesser additive effect and an interaction is suggested. However, this outcome could be the result of chance alone, which highlights the importance of confirmatory findings. If the P value for the interaction term is not significant, the implication is that there is a lack of interaction, but it could also simply reflect the lack of power in the study to detect an effect.

From a mechanistic viewpoint, interaction suggests that at the molecular level the effect or byproducts of the environmental insults modifies the molecular function of the product of the gene under observation. The higher obesity risk will come from the co-existence of both genetic and environmental influences at a high scale for a given population. On one hand individuals inherited a number of gene variants in key loci and on the other they adopt a different position on the environmental spectrum of risk by the lifestyle choices they make (e.g. low-fat vs. high-fat diets, high vs. low levels of physical
activity, etc.) Thus, while the environmental factors are modifiable, the genetic factors are not\(^2\). Understanding the gene–environment relationship is one of the big challenges that is being faced.

**Methodological aspects on the study of genetics of obesity**

A complementary strategy to the candidate gene approach for the identification of obesity genes is the use of genome-wide linkage scans. This strategy involves the genotyping of families using polymorphic markers positioned across the whole genome followed by calculation of the degree of linkage of the marker to a disease trait. This approach does not rely on any pre-existing knowledge of the genes related to the disease. For the study of obesity, genome-wide linkage scans have been applied: in families, representative of general population and also in families chosen because they have an around obese proband. This latter approach is very useful for linkage analysis. In fact, the first genome scans for obesity located a quantitative trait locus (QTL) for leptin levels and fat mass at \(2p21\) in Mexican-Americans for the San Antonio Family Heart Study. Subsequently, more than 204 quantitative trait loci for obesity-related phenotypes were found on all but chromosome Y from more than 50 genome-wide scans reported since 1997\(^6\). Most of the genome-wide scans are based primarily on BMI measurements. Several of these QTLs have been confirmed in various studies, at least 38 QTLs showed evidence of linkage in two studies\(^6\).

Nowadays, the availability of OMICs technology represents a major event for studying the genetics of obesity\(^38\)–\(^45\). The different biomarkers at the transcript, protein and metabolite level will hopefully develop into indicative and predictive sets of molecules, their diagnostic power cannot be uncoupled from pre-determined genetic disposition. Human genetic differences appear at the level of SNPs, copy number polymorphisms and the specific combinations of alleles (haplotypes). On top of these levels of sequence variability, epigenetic phenomena, such as DNA methylation, histone acetylation and RNA interference, add to the complexity of individually different gene regulation. Besides, the high throughput offers the opportunity to genotype more than 10,000 samples a day and enables genomewide association studies\(^38\)–\(^45\). DNA hybridisation arrays represent a new tool to study the association of several different polymorphisms with the development of obesity. Gene chips containing 500,000 SNPs are already available allowing the identification of possible mutations in known and unknown genes related to obesity\(^41\). This technology constitutes the appropriate tool for approaching the study of those combinations of genes and mutations that are implicated in the development of obesity in humans, as well as for establishing how environmental factors affects the onset of obesity in individuals with a specific genetic background.

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