A Polymorphism in the Melanocortin 4 Receptor Gene (*MC4R*: c.92C > T) Is Associated with Diabetes Mellitus in Overweight Domestic Shorthaired Cats

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**Background:** Feline diabetes mellitus (DM) shares many pathophysiologic features with human type 2 DM. Human genome-wide association studies have identified genes associated with obesity and DM, including melanocortin 4 receptor (*MC4R*), which plays an important role in energy balance and appetite regulation.

**Hypothesis/Objectives:** To identify single nucleotide polymorphisms (SNPs) in the feline *MC4R* gene and to determine whether any SNPs are associated with DM or overweight body condition in cats.

**Animals:** Two-hundred forty domestic shorthaired (DSH) cats were recruited for the study. Of these, 120 diabetics were selected (60 overweight, 60 lean), along with 120 nondiabetic controls (60 overweight and 60 lean). Males and females were equally represented.

**Methods:** A prospective case-control study was performed. Genomic DNA was extracted from blood samples and used as template for PCR amplification of the feline *MC4R* gene. The coding region of the gene was sequenced in 10 cats to identify polymorphisms. Subsequently, genotyping by restriction fragment length polymorphism (RFLP) analysis assessed *MC4R*:c.92C > T allele and genotype frequencies in each group of cats.

**Results:** No significant differences in *MC4R*:c.92C > T allele or genotype frequencies were identified between nondiabetic overweight and lean cats. In the overweight diabetic group, 55% were homozygous for the *MC4R*:c.92C allele, compared to 33% of the lean diabetics and 30% of the nondiabetics. The differences between the overweight diabetic and the nondiabetics were significant (*P* < .01).

**Conclusions and Clinical Importance:** We identified a polymorphism in the coding sequence of feline *MC4R* that is associated with DM in overweight DSH cats, similar to the situation in humans.

**Key words:** Endocrinology; Feline; Genetic markers; Genetics; Molecular biology; Pancreas.

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**Abbreviations:**

- AGRP: agouti-related protein
- BCS: body condition score
- DM: diabetes mellitus
- dNTP: deoxyribonucleotide triphosphate
- DSH: domestic shorthaired
- gDNA: genomic DNA
- GWAS: genome-wide association studies
- IGF-1: insulin growth factor-1
- MC4R: melanocortin 4 receptor
- OR: odds ratio
- PCR: polymerase chain reaction
- RFLP: restriction fragment length polymorphism
- RVC: Royal Veterinary College
- SNP: single nucleotide polymorphisms
- UTR: untranslated region
- α-MSH: alpha melanocyte stimulating hormone
and genetic background plays a key role in the development of the disease. Advances in molecular genetic techniques, availability of the complete human genome sequence, and single nucleotide polymorphism (SNP) chips to allow performance of genome-wide association studies (GWAS) have led to identification of genes associated with type 2 DM in humans. To date, there are no studies that have identified specific genes that predispose cats to DM, although given a similar pathophysiology and environmental risk factors, it seems likely that feline DM could share a similar genetic basis with the human disease.

The melanocortin 4 receptor (MC4R) gene, encoding a transmembrane G-protein–coupled receptor, has been implicated as a susceptibility gene in human DM and obesity. This receptor is mainly expressed in the hypothalamus and plays an important role in the regulation of energy balance and appetite. In a situation of positive energy balance, MC4R is stimulated by alpha-melanocyte stimulating hormone (α-MSH), leading to a feeling of satiety. During periods of starvation, the activity of MC4R is inhibited by its inverse antagonist, agouti-related protein (AGRP), leading to a feeling of hunger. Mutations in this gene are the most common single genetic cause of human obesity, accounting for up to 6% of cases. A correlation between MC4R mutations and human type 2 DM has been shown in several studies, including a recent meta-analysis, which showed an odds ratio (OR) of 1.10 (95% CI 1.07–1.13). Given the influence this gene has on susceptibility to obesity and DM in human beings and given the fact that obesity also poses an important risk factor for DM in cats, MC4R was considered to be a logical candidate gene in the search for genetic factors predisposing to DM in cats. The aim of this study was to confirm the sequence of feline MC4R, to identify SNPs in its coding region, and to perform a case-control study to detect a possible gene association with feline DM and obesity.

### Materials and Methods

#### Animals

Blood samples (EDTA or non-anticoagulated) from diabetic cats were recruited via the UK Companion Animal Diabetes Registry, based at the Royal Veterinary College (RVC), from first opinion veterinary practices in the UK, with samples submitted with a standardized sample submission form (http://www.rvc.ac.uk/cic/documents/GHSubmissionformV1328Jan10DL.pdf), that included details of concurrent diseases and clinical signs, body weight and body condition score (BCS) at and before diagnosis, diet and insulin type and dose. The BCS assessment was carried out according to the routinely used 9- and 5-point BCS systems, information about how to assess the BCS of cats according to these systems was included in the submission form. Blood samples from nondiabetic cats were obtained from the RVC Genetic Archive, consisting of retrospective residual samples (after completion of diagnostic testing) from referral cases seen at RVC hospitals. Ethical approval for sample collection had been obtained previously from the local ethical committee. For the SNP discovery phase, 10 cats (5 Burmese: 2 diabetic, 3 nondiabetic; 5 DSH: 3 diabetic, 2 nondiabetic) were assessed. For the genotyping phase, a total of 240 DSH cats were analyzed, consisting of 4 groups (n = 60 each; with males and females equally represented) of overweight diabetic, lean diabetic, overweight nondiabetic, and lean nondiabetic cats.

The diabetic cats were selected on the basis of the presence of increased serum fructosamine concentration, persistent hyperglycemia, requiring insulin therapy, and having been diagnosed as diabetic by the attending veterinarian for longer than 4 weeks. Insulin growth factor-1 (IGF-1) was measured in all samples, those cats with IGF-1 concentration >800 ng/mL were allocated either to the overweight diabetic group (BCS > 4.5 or 5/9) or to the lean diabetic group (BCS ≤ 4.5 or 5/9). Those cats that were overweight before diagnosis and had lost weight before the diagnosis of DM were included in the overweight group, because weight loss is considered one of the landmark clinical signs of DM. Those cats with known concurrent diseases that could have an impact on body weight were excluded from the study. Based on evaluation of medical records and available diagnostic information, nondiabetic cats >9 years of age that had no clinical or biochemical evidence of DM, specifically hyperglycemia or glycosuria, were selected. Cats with clinical or biochemical evidence of other diseases that could lead to changes in body weight (eg, hyperthyroidism, renal disease, gastrointestinal disease, hepatic disease) were excluded from the study. However, because these samples were obtained from an archive of retrospective clinical material, additional prospective diagnostic testing was not performed. All of the nondiabetic cats were seen in a clinic staffed by board-certified veterinary specialists. There were no known familial relationships among any of the cats included in this study.

#### Feline MC4R SNP Discovery

The initial phase focused on resequencing the coding region of the feline MC4R gene to identify SNPs. Genomic DNA (gDNA) was extracted from EDTA blood samples or blood clots using a blood genomic DNA extraction kit, according to the manufacturer's instructions. Feline MC4R-specific primers located in the 5′ untranslated region (UTR; Sense: 5′-CTCAGAATTTCCGCGCAGAC-3′; starting at position -103) and in the 3′ UTR (Anti-sense: 5′- ACCCATGGCTTACACAGGG-3′; starting at position 1078) were designed according to the available feline MC4R sequence (http://www.ensembl.org/Felis_catus/Info/Index; ENSFCAG000000006540). Assembly ASMA18133v1, Genbank Assembly ID: GC_A_000181335.1). Polymerase chain reaction was performed in 25 L volumes using 0.1 L of DNA polymerase (Immolsel DNA polymeraseβ) with 1 L of gDNA as a template. Each reaction also contained 13 L of water, 5 L of 5× PCR-enhancing additive (Hi-Spec additive®), 2.5 L of ammonium sulfate buffer (NH4 bufferβ), 1.25 L MgCl2 (2.5 mM final concentration), 0.25 L 250 μM dNTPs, and 2 L of primer mix (each at 20 pmol/L). Using a PCR cycler, reactions were heated to 95°C for 10 minutes, followed by 30 cycles consisting of 94°C for 40 seconds, 65°C for 30 seconds, and 72°C for 2 minutes, with a final extension step at 72°C for 10 minutes. PCR products were separated by horizontal gel electrophoresis using a 2% agarose gel and analyzed under 590 nm UV light. Subsequently, DNA was extracted using a gel extraction Kit (GeneJel gel extraction Kit®) according to the manufacturer’s instructions. Amplicons then were submitted for sequencing and compared to each other and with the reference feline genome sequence using sequence analysis software.

#### MC4R Genotyping by RFLP Analysis

Genomic DNA was extracted from EDTA blood or blood clots and PCR performed as described in the SNP discovery section. Amplified DNA then was purified using a PCR clean-up kit
(Genelute PCR clean-up Kit\textsuperscript{b}), subjected to digestion using a restriction enzyme (BstOI enzyme\textsuperscript{d}), which recognizes a restriction site (CC\textsuperscript{'(A/T)GG}) incorporating the MC4R:c.92C allele, absent in the MC4R:c.92T allele. The digestion reaction consisted of 0.5 \(\mu\)L of BstOI restriction enzyme, 0.5 \(\mu\)L of BSA, 0.5 \(\mu\)L of Buffer (Buffer \textsuperscript{C})), and 3.5 \(\mu\)L of purified PCR product. Reactions were incubated for 1 hour at 60°C in a water bath and analyzed by agarose gel electrophoresis as described above. In samples in which gel interpretation was potentially ambiguous, genotyping by RFLP was repeated using a higher DNA concentration. If an unambiguous result still was not obtained, manual sequencing was performed.

**Statistical Analysis**

Statistical analysis was performed using SPSS software package.\textsuperscript{1} The Mann–Whitney \(U\)-test was used to analyze non parametric variables. The Pearson’s two-tailed chi-squared test was used to compare MC4R:c.92C>T SNP genotype and allele frequencies among groups, and the OR with 95% confidence interval (CI) was calculated. Bonferroni correction was applied for multiple group comparisons. Significance was considered at \(P < .01\).

**Results**

**Animals**

The median age of the overweight diabetic cats was 12 years (range, 6–18 years); the median weight was 6.6 kg (range, 5.1–10.0 kg). The median age of the lean diabetic cats was 13 years (range, 5–18 years); the median weight was 4.1 kg (range, 2.6–6.0 kg).

For the overweight nondiabetic group (30 male, 30 female), the mean age was 12 years (range, 9.1–20 years) and the median weight 5.7 kg (range, 3.4–8.3 kg). For the lean nondiabetic group (30 male, 30 female), the median age was 13.9 years (range, 9–19 years) and the mean weight 3.9 kg (range, 2.5–5.7 kg).

The age difference between diabetics and nondiabetics was not statistically significant (\(P = .074\)), whereas the obese cats were significantly heavier than the lean cats (\(P < .001\)).

**Polymorphisms in Feline MC4R**

Initial sequencing results showed the presence of 3 polymorphisms. A non synonymous SNP in position 92 of the coding sequence (MC4R:c.92C>T) was found in the DSH cats only and led to an amino acid change from leucine to proline in the protein sequence. Figure 1 shows the sequencing results from the initial SNP discovery phase (10 cats) and represents the fragment of the coding sequence of MC4R where the non synonymous SNP was detected. Two additional synonymous SNPs (MC4R:c.297C>T and MC4R:c.303C>T) also were present in both the DSH and the Burmese cats.

**MC4R:c.92C Homozygous Genotype in Obese Diabetic DSH Cats**

A case-control study was performed, focusing on the MC4R:c.92C>T SNP, to determine whether there was any association with increased body weight or DM. The RFLP assay was found to be a rapid and simple test to identify MC4R genotypes in cats (Fig 2). In the 2 nondiabetic groups, there was no significant difference in allele or genotype frequencies comparing the overweight and lean cats (Table 1), and data from these cats subsequently were combined into a single nondiabetic control group. There were no differences in allele frequencies, comparing the lean and overweight diabetic groups (\(n = 60\) each group) and the nondiabetic control group (\(n = 120\)) (Table 2). However, in the overweight diabetic group (\(n = 60\)), the majority of cats (55%) were homozygous for the MC4R:c.92C allele (Table 2) compared with the nondiabetic control group (\(n = 120\)), where 30% were homozygous C, a difference that was statistically significant (\(P = .002\)). In contrast, there was no difference

**Fig 1.** Initial single nucleotide polymorphism (SNP) discovery phase, fragment of the coding sequence of feline MC4R in which the non synonymous SNP was detected (purple circle). This SNP (MC4R:c.92 C>T) was detected exclusively in domestic shorthaired (DSH) cats and leads to leucine being replaced by proline in the translated protein. DMBURN1, diabetic Burmese cat1; DMBURN2, diabetic Burmese cat2; DMDSH1, diabetic DSH cat1; DMDSH2, diabetic DSH cat2; DMDSH3, diabetic DSH cat3; NONDMBURM1, non-diabetic Burmese cat1; NONDMBURM2, non-diabetic Burmese cat2; NONDMDSH1, non-diabetic DSH cat1; NONDMDSH2, non-diabetic DSH cat2; NONDMDSH3, non-diabetic DSH cat3; A, adenine; C, cytosine; G, guanine; T, thymine; Y, heterozygous cats (C and T alleles).
in genotype distribution when comparing the lean diabetic cats (n = 60) with the nondiabetic cats (n = 120) (P = .7) and when comparing lean diabetics (n = 60) with the obese diabetic cats (n = 60) (P = .02). Overweight cats that were homozygous for the mutation had a statistically significantly higher OR for DM of 3.67 (95% CI, 1.68–7.95, P = .0007) when compared to nonhomozygous obese cats.

**Discussion**

The present study demonstrates that a genetic factor (homozygosity for the MC4R:c.92C allele) may be associated with DM in overweight DSH cats. Three SNPs were identified in the feline MC4R coding sequence, with 2 of these (MC4R:c297C>T and MC4R:c303C>T) representing synonymous changes and therefore less likely to impact protein structure or function than the non synonymous SNP (MC4R:c.92C>T). Additional polymorphisms may occur upstream (ie, in the gene regulatory regions) or downstream of the coding region, but these areas were not investigated in the current study. Because only 10 cats were evaluated in the SNP discovery phase, it is possible that other less common SNPs are present in feline MC4R. However, because only SNPS with a minor allele frequency >0.1 are normally used in case-control association studies, relatively rare SNPs would not be deemed suitable and therefore were excluded from subsequent genetic analysis.
Previous studies in humans have shown that mutations in the \(MC4R\) gene are associated with obesity, which increases the risk of those individuals developing type 2 DM.\(^{26,29-31}\) The current study failed to demonstrate any association between the \(MC4R:c.92C>T\) polymorphism and overweight body condition in non-diabetic DSH cats, but the sample size was relatively small. Rather than having a direct effect on regulating satiety and food intake and consequently influencing body weight, these results suggest that this polymorphism may be associated with progression to overt DM in overweight cats. Alternatively, this polymorphism might be in linkage disequilibrium with another causative polymorphism or mutation elsewhere in the genome.

Under physiologic conditions, \(MC4R\) is constitutively active, leading to an appetite suppression signal until negative energy balance develops. At this time, a decrease in leptin concentration and a subsequent increase in the inverse agonist AGRP occur. AGRP binding inactivates the \(MC4R\) receptor, leading to an increase in appetite.\(^{28}\) Functional studies performed in humans have demonstrated that mutations that influence the amino acid sequence in the N-terminal region of the protein generate receptors that are unable to maintain their constitutive activity, leading to decreased appetite suppression and increased food intake.\(^{33}\) In addition, several studies have evaluated the role of \(MC4R\) in energy metabolism, but conflicting results have been reported.\(^{34-37}\) Notwithstanding discrepancies in the published literature, studies in mice have indicated that mutations of the \(MC4R\) receptor can lead to a reduction in metabolic rate, indicating that this receptor might play additional roles in energy metabolism and there may be interspecies differences in receptor function. The non-synonymous \(MC4R:c.92C>T\) SNP identified in DSH cats causes an amino acid change from leucine to proline at amino acid position 30.

The GWAS of human type 2 DM, which identified \(MC4R\) as a susceptibility gene, involved relatively large numbers of cases and controls.\(^{25,38}\) Unfortunately, such large-scale genetic studies are difficult to replicate in companion animals, with the result that many studies are relatively underpowered. Regardless, GWAS of several diseases of dogs have yielded results, even with relatively low numbers, when performed within breeds.\(^{29-41}\) This is likely to be similar for genetic studies of cats, if performed within relatively inbred pedigree populations, but in the DSH cat population, which is more outbred, larger sample sizes are likely to be required for case-control genotyping studies. Nevertheless, a significant difference was found in the current study, even with relatively small numbers of DSH cats, suggesting that this gene might play a major role in susceptibility to DM in overweight cats.

The current study has a number of limitations. The diabetic cases and controls were breed- and sex-matched, but not age-matched. Older cats were proactively recruited as controls to decrease the chances of including cats that might become diabetic later in life. Previous studies have demonstrated that the risk of developing DM in cats increases with age,\(^{42}\) and therefore, even though the nondiabetic (overweight and lean) populations were older than the diabetic populations, this is likely to be beneficial rather than detrimental. BCS information was obtained from all cats, including information about BCS before diagnosis when available. Because one of the landmark clinical signs of DM is weight loss, some cats that were overweight before developing the disease may have been included in the lean group. Although an attempt was made to exclude cats that had concurrent diseases that could alter body weight, this was not always possible. This could have changed the results of the study. The overall prevalence of the SNP in the diabetic group (lean and obese), however, still is significantly higher than in the nondiabetic group. Given the effect that \(MC4R\) mutations have on appetite regulation in humans, it would have been valuable to have obtained information about each cat in terms of appetite and food intake. Although some of this information was available for some of the cats, it proved difficult to incorporate these data into the analysis or selection criteria because of the subjectivity of evaluating food intake and appetite, particularly in multi-cat households and when different diets and feeding regimes (e.g., ad libitum or not) were being used.

The current findings are of interest not only in advancing our knowledge of the pathogenesis of feline DM but also in terms of clinical relevance. Thus, \(MC4R\) genotyping of overweight cats might be useful in estimating the relative risk of the cat progressing to overt DM, given an OR of 3.67 comparing the overweight diabetic group with the overweight nondiabetic group. Because feline DM is likely to represent a complex genetic disorder, identification of other susceptibility genes is warranted and, if confirmed to be present, a test based on genetic profiling of multiple genes potentially could be used to establish an overall genetic risk for development of DM. This might then be useful for identifying cats with a greater risk of developing the disease, so that preventative measures (e.g., dietary management) could be implemented to decrease other risk factors (e.g., development of obesity) that might precipitate disease in a genetically susceptible individual.

In conclusion, we have demonstrated that a genetic factor may influence susceptibility to DM in cats. The feline \(MC4R:c.92C>T\) polymorphism might be involved in progression to overt DM in overweight cats, but it remains to be established whether or not a causal relationship is present, or if it is simply an association. If the polymorphism does have functional consequences, this does not seem to be directly related to increased susceptibility to overweight body condition and might, instead, play a role in glycemic control in the presence of insulin resistance or influence pancreatic beta cell function in response to chronic hyperglycemia. Functional studies to investigate the physiologic relevance of this polymorphism and GWAS to identify other susceptibility genes for feline DM are warranted.
Footnotes

a GenElute blood genomic DNA extraction kit; Sigma, Poole, UK
b Bioline, London, UK
c Eppendorf Master Cycler; Eppendorf AG, Hamburg, Germany
d ImageMaster VDS; Pharmacia Biotech/GE Healthcare, Buckinghamshire, UK
e Sigma-Aldrich, Dorset, UK
f GATC Biotech Ltd The London BioScience Innovation Centre, London, UK

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