Mortality over two centuries in large pedigree with familial hypercholesterolaemia: family tree mortality study

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Mortality over two centuries in large pedigree with familial hypercholesterolaemia: family tree mortality study

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Abstract

Objective To estimate all cause mortality from untreated familial hypercholesterolaemia free from selection for coronary artery disease.

Design Family tree mortality study.

Setting Large pedigree in Netherlands traced back to a single pair of ancestors in the 19th century.

Subjects All members of pedigree aged over 20 years with 0.5 probability of carrying a mutation for familial hypercholesterolaemia.

Main outcome measure All cause mortality.

Results A total of 70 deaths took place among 250 people analysed for 6950 person years. Mortality was not increased in carriers of the mutation during the 19th and early 20th century; it rose after 1915, reached its maximum between 1935 and 1964 (standardised mortality ratio 1.78, 95% confidence interval 1.13 to 2.76; \( P = 0.003 \)), and fell thereafter. Mortality differed significantly between two branches of the pedigree (relative risk 3.26, 95% confidence interval 1.74 to 6.11; \( P = 0.001 \)).

Conclusions Risk of death varies significantly among patients with familial hypercholesterolaemia. This large variability over time and between branches of the pedigree points to a strong interaction with environmental factors. Future research is required to identify patients with familial hypercholesterolaemia who are at extreme risk and need early and vigorous preventive measures.

Introduction

Familial hypercholesterolaemia is associated with premature cardiovascular disease \(^7\&\) and decreased life expectancy.\(^1\) These associations, however, have been described in families that were investigated because the probands—or even multiple family members—had presented with cardiovascular disease at young age.\(^1\)

Because susceptibility to cardiovascular disease is likely to be affected by additional genetic and environmental factors,\(^2\&\) mortality from familial hypercholesterolaemia may have been preferentially studied in patients and families with multiple risk factors for cardiovascular disease.\(^1\) The natural course of the disorder has not been studied without selection for cardiovascular disease, and estimates for carriers in the general population are therefore lacking.

The recent introduction of molecular diagnostic tools allows the disorder to be diagnosed with certainty.\(^10\) and large scale family screening has been shown to be highly effective.\(^11\) We assessed mortality risk in a large pedigree of carriers of the V408M mutation or Afrikaner-2 mutation in exon 9 of the low density lipoprotein receptor gene.\(^12\)

Subjects and methods

Hypercholesterolaemia was detected in probands A and B during routine screening. In proband C, hypercholesterolaemia was detected after myocardial infarction at the age of 51 years. The probands had mean fasting total serum cholesterol concentrations of 10.24, 9.20, and 12.78 mmol/l, respectively. The three probands were carriers of the V408M mutation on an identical haplotype, and this suggested that they were (distantly) related. We performed genealogical searches using official records of births, marriages, and deaths. Dutch official records for previous centuries are virtually complete because they were stored at several places. The authorities performed these registrations irrespective of socioeconomic status.

The genealogical searches had two phases. Firstly, we traced all maternal and paternal ancestors of the three carriers of the mutation throughout as many generations as possible. We found only one pair of ancestors shared by and connecting the three probands. Secondly, we traced all descendants of this pair and screened all living descendants for the V408M mutation. Figure 1 shows a small part of the pedigree. All first degree relatives of people on the transmission lines of the mutation had a mendelian probability of 0.5 of being affected. As a result of the increasing size of the pedigree in recent generations, the information about ancestors on transmission lines is confirmed many times. Our analyses could have been influenced by consanguinity or non-paternity if the genuine fathers were also carriers of the V408M mutation. Deceased parents and their ancestors could then be interpreted incorrectly as affected. We therefore did genealogical searches of the spouses of the pedigree and tested for the V408M mutation in the living descendants of their siblings. We did not detect consanguinity and we did not find any indication for non-paternity.
The molecular diagnosis of familial hypercholesterolaemia was based on the presence of the V408M mutation. This molecular method has been described elsewhere.\textsuperscript{12} The molecular and genealogical studies were approved by the hospitals' review boards, and all family members studied gave informed consent.

**Statistical methods**

The mortality in the pedigree was compared with the mortality in the Dutch population standardised for age, sex, and calendar period as described previously.\textsuperscript{13–15} In brief, the standardised mortality ratio is the ratio of observed to expected number of deaths. We calculated the expected mortality by multiplying the total number of years lived by the people in the pedigree in each calendar period for each age and sex category by the age and sex specific mortality rates of the Dutch population for each calendar period. The probands and the parental years before birth of the proband were excluded from the analyses. We also ignored the first two decades of life for all people in the pedigree because the registration of juvenile mortality in the 19th century may have been incomplete. Moreover, ancestors who were certainly affected may have been missed because premature death could have decreased the chance of passing on their mutation to present generations. This would lead to underestimation of risk of death. Therefore, we did the primary analyses in relatives of complete sibships (all siblings available for analysis, who had 0.5 probability of being affected. As a result the observed standardised mortality ratios exhibit 50% of the excess mortality from the mutation causing familial hypercholesterolaemia. Secondary analyses were done on people who were definitely affected. We ended all analyses in December 1989, when statins became available to our patients.

We compared mortality between subgroups with Poisson regression (relative risk). Cumulative survival was analysed with Cox's regression (relative risk) and the Kaplan-Meier method. We calculated the 95% confidence interval of the standardised mortality ratio assuming a Poisson distribution of the observed number of deaths and using exact limits. We calculated the 95% confidence interval of the relative risk with Poisson or Cox's regression as the exponent of the regression coefficient and its standard error. Significance was assessed at the 5% level of probability.

**Results**

**Genealogical searches**

We traced a total of 412 descendants in eight generations along the lines of transmission of the V408M mutation in the pedigree. Four sibships with incomplete data due to emigration were excluded from the analyses. The complete sibships contained 387 (94%) relatives, of whom 250 survived for 20 years or more.

**Standardised mortality: variance over time**

Between 1830 and 1989, a total of 70 deaths took place among 250 people with a 0.5 probability of carrying the mutation who were analysed over 6950 person years (table). The overall standardised mortality ratio of these people was 1.32 (95% confidence interval 1.03 to 1.67; P = 0.02). Thirty of the 118 people who were definitely affected died, giving a standardised mortality ratio of 1.59 (1.07 to 2.26; P = 0.02). In the 20th century, mortality from familial hypercholesterolaemia rose, peaking between 1935 and 1964 (standardised mortality ratio 1.78, 1.13 to 2.76; P = 0.003) and then falling. The standardised mortality ratios of relatives who were definitely affected followed a similar trend, with a maximum of 2.29 (1.14 to 4.09; P = 0.005) between 1935 and 1964.

Figure 2 shows the point estimates of the standardised mortality ratios for men and women over time. The period had a significant effect on risk of death after differences in the distribution of age and sex were adjusted for (P < 0.001).
Standardised mortality: variance within generations

Analysis by Poisson regression showed that the risk of death in the family members of probands B and C relative to those of proband A was 1.74 (95% confidence interval 0.82 to 3.67; P = 0.1) and 3.26 (95% confidence interval 1.74 to 6.11; P = 0.001) respectively. Analysing the data with a Cox’s proportional-hazards model did not materially alter the results (data not shown). Figure 3 shows the difference in survival between the branches. The standardised mortality ratio of branch A was 1.04 (95% confidence interval 0.66 to 1.43; P = 0.9).

The Poisson regression analysis allowed synchronous estimation of the variance of risk of death in two directions: through the centuries and within generations. Time had a significant effect on risk of death (P = 0.002) after adjustment for age, sex, and position in the pedigree. The deaths took place in seven generations of the pedigree, and consequently the deceased relatives shared only a small fraction of their genome. Statistical adjustment for the exact family ties did not change the results (data not shown).

Discussion

We found that the excess mortality from familial hypercholesterolaemia varied over time. In the 19th century, mortality seemed lower than in the general population. It rose after 1915, reached a maximum during the 1950s, and decreased thereafter. During the decades with excess mortality, survival in the branches of the pedigree differed significantly, ranging from normal life expectancy to severe excess mortality. This large variation of risk suggests that previous studies, with families based on selected patients, may have overestimated mortality. Moreover, such large variation in mortality in two directions (over time and within generations) in a pedigree indicates that the disorder has strong interactions with environmental factors.

Strengths and weaknesses

The strength of our study is that the natural course of the disorder was assessed free from selection for cardiovascular disease. We started with three probands and traced the pair of distant ancestors from whom the disorder originated. All descendants of these distant ancestors were subsequently identified, the only factor affecting selection being the completeness of the Dutch official records. We then identified the transmission lines of the mutation through the pedigree so that all carriers of the mutation could be analysed. We studied all cause mortality rather than cardiovascular mortality because cause of death was often poorly defined in earlier records.

Our findings are based on a large pedigree of carriers of the V408M mutation in the low density lipoprotein receptor gene. This mutation is consistently associated with greatly raised plasma cholesterol concentrations and may be indicative of familial hypercholesterolaemia in general. Moreover, the mortality from different types of mutations may be similar. However, we cannot exclude the existence of rare mutations that cause a higher risk of death from familial hypercholesterolaemia.

Our analyses are unlikely to have been affected by competing risk. In the past, mortality from other
Familial hypercholesterolaemia is associated with excess mortality in families of patients who present with cardiovascular disease.

Population data are lacking.

Many untreated patients with familial hypercholesterolaemia (about 40%) reach a normal life span.

Standardised mortality ratio was normal in the 19th century and rose to a peak in the 1930s to 1960s.

The variation in mortality suggests an interaction between genetic and environmental factors.

with familial hypercholesterolaemia who show different combinations of risk factors.

Contributors: EJGS and RG JW coordinated the work, evaluated the literature, performed analyses, wrote the first draft of the manuscript, and participated in editing and revising the manuscript. JCD collected the data, performed the molecular work, and participated in editing and revising the manuscript. PH EMDM performed genealogical research and participated in revising the manuscript. AHMS participated in revising the manuscript. JJK performed genealogical research, collected data, and participated in editing and revising the manuscript. Jan P Vandenbroucke, provided methodological support and participated in editing and revising the manuscript. EJGS will act as guarantor.

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Biomedical research has been extremely successful in identifying the mutated genes for monogenic diseases—that is, those for which a single gene mutation is considered necessary and sufficient to produce clinical disease. A prime example of this is familial hypercholesterolaemia, which arises from mutations in the low density lipoprotein receptor gene and carries an increased risk of coronary heart disease. Familial hypercholesterolaemia is one of the commonest autosomal dominant disorders, with over 600 known mutations. There is increasing recognition and evidence that even the monogenic diseases are not simple and that the relation between genotype and phenotype is modified both by other genes and environmental effects.

Sijbrands et al examined mortality over two centuries in a large pedigree with familial hypercholesterolaemia from the Netherlands. Interestingly, the mortality in the pedigree did not differ significantly from national rates until early in the 20th century. Pedigree members who were known to be affected showed the same pattern of mortality as those unaffected, although with higher death rates. This indicates that many untreated people with familial hypercholesterolaemia lived a normal life span. That familial hypercholesterolaemia is compatible with a normal life span under different environmental conditions has been reported earlier—for example in Utah pedigrees and in a Finnish pedigree with the North Karelia mutation.

Sijbrands et al suggest that raised low density lipoprotein concentrations may have protected people from infectious diseases that were more common in earlier centuries, but the absence of other risk factors for coronary heart disease (such as widespread cigarette smoking or a high fat diet) in the 19th century may be equally important. Also, the mortality in different branches of the pedigree differed significantly, suggesting that other genes close to the low density lipoprotein receptor may have a role in modifying risk. The different branches may have also differed in social class or urban residence patterns transmitted culturally from one generation to the next.

Thus, we must recognise that even in monogenic disorders, other genes and the environment can be important. Although the main focus of genetic research has moved on to the common, multifactorial diseases such as coronary heart disease—often termed complex disorders—we have much to learn about the relation between genes, environment, and clinical phenotype even from monogenic disorders.

In a pivotal article about the Human Genome Project two years ago, Francis Collins stated: “Largely, but not entirely, at the behest of our genes, we fare better or worse.” Yet, life expectancy has increased greatly in the past century in the richest nations, and clearly this increase could not be due to genetic factors. Perhaps we should declare that largely at the behest of both our genes and our environment, we fare better or worse.

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