Familial hypercholesterolemia: the Dutch approach
Huijgen, R.

Citation for published version (APA):

General rights
It is not permitted to download or to forward/distribute the text or part of it without the consent of the author(s) and/or copyright holder(s), other than for strictly personal, individual use, unless the work is under an open content license (like Creative Commons).

Disclaimer/Complaints regulations
If you believe that digital publication of certain material infringes any of your rights or (privacy) interests, please let the Library know, stating your reasons. In case of a legitimate complaint, the Library will make the material inaccessible and/or remove it from the website. Please Ask the Library: http://uba.uva.nl/en/contact, or a letter to: Library of the University of Amsterdam, Secretariat, Singel 425, 1012 WP Amsterdam, The Netherlands. You will be contacted as soon as possible.
Chapter 15

Improved access to life insurance after genetic diagnosis of familial hypercholesterolemia: cross-sectional postal questionnaire study

Roeland Huijgen, Sietske J.M. Homsma, Barbara A. Hutten, Iris Kindt, Maud N. Vissers, John J.P. Kastelein, and Jan L.A. van Rijckevorsel

ABSTRACT

Background: A decade ago, in the initial stage of genetic testing for familial hypercholesterolemia (FH) in the Netherlands, it was reported that such screening decreased access to affordable life insurance for mutation carriers. In 2003, in order to improve access to insurance for FH mutation carriers, insurers agreed to underwrite according to a set of guidelines. In this cross-sectional study, we assessed whether access to insurance has improved since the advent of these guidelines.

Methods: We approached 2,825 subjects that had participated in the genetic testing for FH between 1998 and 2003. We compared unconditional acceptance rates before and after FH diagnosis and before and after the guidelines were issued by means of logistic regression analysis.

Results: Our study outcome pertains to 414 FH patients who applied for life insurance. Unconditional acceptance of a policy before DNA diagnosis and before the issue of guidelines occurred in 182 out of 255 (71%) cases, versus 27 out of 35 (77%) cases after DNA diagnosis, but before the issue of guidelines. De facto, 107 out of 124 (86%) patients received unconditional acceptance after DNA diagnosis and after the issue of guidelines ($p$ for trend = 0.002).

Conclusions and implications: Access to life insurance improved for FH patients after molecular diagnosis and it improved even further after the guidelines were issued. Therefore, we argue that limited access to life insurance on the basis of ‘DNA discrimination’ is no longer a valid argument against genetic cascade testing for FH, at least not in our country.
INTRODUCTION

The number of heritable disorders for which a genetic test is available increased from less than 200 in 1993 to more than 1,800 in 2009. DNA diagnostics of a medical condition offers the possibility to start treatment before the disease becomes symptomatic. One of the possible downsides of genetic testing is that a person’s test result may affect one’s ability to obtain life insurance.

Familial hypercholesterolemia (FH) is a condition that meets key criteria for genetic testing. It is a prevalent inherited disorder of lipoprotein metabolism characterized by markedly elevated low-density lipoprotein cholesterol (LDL-C) levels and premature coronary artery disease (CAD). Cholesterol-lowering treatment has been shown to dramatically reduce CAD risk in patients with FH and, as a consequence, early identification by genetic testing for FH followed by effective treatment can reduce the incidence of CAD events substantially.

A publication from 2002 reported, however, that subjects might have impaired access to insurance in the Netherlands after genetic FH diagnosis. On the basis of these findings, our government requested the Association of Insurers to settle these issues with the patient organisation, screening organisation and representatives of the medical profession. Accordingly, they reached agreement in 2003 and, as a result, insurers agreed to underwrite according to a set of specific guidelines. These guidelines stipulate that subjects with genetic FH and free of cardiovascular disease should be offered an unconditional life insurance policy under most conditions. The main requirements were that individuals had LDL-C levels, either treated or untreated, at or below 4.0 mmol/L while other cardiovascular risk factors, such as smoking, hypertension, diabetes or obesity, had to be absent at the time of life insurance application. The effect of these guidelines on access to life insurance for FH patients had never been evaluated.

The purpose of this study is therefore to assess whether FH diagnosis through genetic testing reduced access to life insurance and whether that access improved for subjects with FH after the guidelines were issued.

In the present study, we determined the access to insurance in a large cohort that had been tested for genetic FH. Here we present our results.
Chapter 15

METHODS

Recruitment
Individuals that had participated in the genetic cascade screening program for FH in the Netherlands between January 1998 and June 2003 were eligible for this cross-sectional postal questionnaire survey. We selected those subjects aged between 18 and 55 years in 2008, who had given written informed consent to be approached for scientific studies at the time of genetic testing for FH. In order to include subjects with and without FH in a 2 to 1 ratio (see power calculation below), we approached all selected subjects with genetic FH, whereas we randomly included only one out of three subjects shown not to carry the FH mutation running in their family.

The study was approved by the Medical Ethical Committee of the Academic Medical Center of the University of Amsterdam.

Outcome measures
The main study outcome was the unconditional acceptance rate for life insurance applications.

As all subjects were diagnosed before the guidelines were issued on July 1st 2003, there were three options for the moment of insurance application: before genetic FH diagnosis and before the insurance guidelines were launched (Period 1); after DNA diagnosis and before the guidelines were launched (Period 2); after DNA diagnosis and after the guidelines were launched (Period 3). For each period the unconditional acceptance rate was calculated and stratified for those with and without genetic FH.

Unconditional acceptance was defined as a life insurance application accepted at standard premium and coverage. Processing of the insurance application with additional terms to the contract - from hereon described as “conditional acceptance” - consisted of: i) refusal of the application by the insurance company; ii) acceptance with the obligation to pay a higher premium; acceptance with altered terms of the contract, for example excluding coverage of death by coronary heart disease; or iii) the applicant’s refusal to accept a life insurance policy that would include a higher premium or a clause.

A secondary outcome was the proportion of conditional acceptance of life insurance applications that were unjustified according to the guidelines. The guidelines stipulated that non-normal acceptance was justified in case of LDL-C levels above 4 mmol/L at the time of application, and/or if one or more of the other cardiovascular risk factors i.e. diabetes, hypertension, smoking and obesity were present.7 We defined obesity as having a body mass index (BMI) above 30 kg/m². If the LDL-C level at the
time of application for a life insurance was unknown we estimated the LDL-C level based on the total cholesterol level. We presumed that LDL-C was above 4 mmol/L in cases where the total cholesterol was above 6 mmol/L at the time of application or - if total cholesterol level was also unknown at that time - if the highest ever total cholesterol was above 10 mmol/L.

Data collection
In order to estimate the response rate and to verify the clarity of the questions in the questionnaire, we sent a pilot mailing in January 2008 to the 500 subjects who had been diagnosed with FH most recently. As the questionnaire was not essentially changed based on the experience with the pilot mailings, we included the population from the pilot study in the overall study cohort. The remaining selected subjects were approached for the first time in September 2008. First, a letter explaining the purpose of this study was sent by surface mail with a return form for the participant to indicate whether or not he/she approved of participation (invitation mailing). If the subjects had approved, the questionnaire itself was sent by surface mail (questionnaire mailing). For the mailings that proved undeliverable, we contacted the Municipal Administration of the subjects’ former address and attempted to retrieve their current address. Up to two additional mailings were sent to the subjects that did not respond to the invitation mailing within two months of delay. Similarly, we sent up to two reminders after three months to those who did not return the questionnaire. To assess the reasons for non-response, we called a subset of the group that did not respond to any of the three invitation mailings by telephone. For this purpose, all the non-responders from the pilot study cohort were approached and asked why they did not respond.

Sample size calculation
Under the assumptions that life insurance companies in general accept 97% of the applicants under normal condition and 85% of the FH patients free of CVD, we calculated that 141 subjects with FH and 71 without FH, applying for a life insurance after July 2003, were required to detect a difference in acceptance of 12% with a power of 80% and 0.05 significance level (two-sided). Based on the response rate of a pilot mailing in 500 subjects with FH only, we estimated that we had to approach approximately 2700 subjects.
Statistical analysis

Differences in continuous variables (age, total cholesterol, body mass index) between subgroups were compared using the independent sample t-test. Chi-square tests were applied for comparing distributions of dichotomous data (gender, presence of hypertension or diabetes, smoking).

The unadjusted pairwise association between conditional acceptance of life insurance and clinical variables such as age, sex, hypertension, diabetes, smoking, body mass index, total cholesterol and FH mutation was expressed by odds ratio's estimated by a series of bivariate logistic regression analyses. Subsequently, hierarchical multiple regression analysis with stepwise backward elimination was used to identify independent determinants of conditional acceptance. To handle the entries where the total cholesterol was missing, we assessed the association between total cholesterol and other potential explanatory variables with linear regression of the non-missing entries. Hereafter, the best predictors for total cholesterol were used to impute the missing values for total cholesterol based on nearest neighbour imputation.9 A two sided p-value < 0.05 was considered significant. Statistical analyses were performed with SPSS version 16.

RESULTS

Recruitment and non-response

In total, 6,115 males and females aged between 18 and 55 years had participated in the genetic cascade testing program for FH between January 1998 and June 2003. Of these, 2,558 carried the FH mutation and 3,557 did not. Of the 6,115 tested individuals, 2,825 passed the additional selection steps and were sent an invitation letter. This population consisted of 1,963 (69%) subjects with genetic FH mutation and 862 (31%) without FH.

In total 1,192 (42%) subjects did not respond to our invitation letter (Figure 1). Of the 1,633 (58%) subjects who responded, 1,127 (69%) did consent, 484 (30%) did not consent for various reasons, and in 22 (1%) cases, the invited subjects were deceased. All 1,127 subjects that had consented were sent a questionnaire and 1,040 (92%) returned a completed questionnaire. The 1,040 participants were on average 37 years of age at FH testing, and 527 (50.7%) were male. Baseline demographic and clinical characteristics of patients with and without FH are summarised in Table 1. As expected, those patients with FH had higher untreated total cholesterol levels than those without FH (p<0.001).
An extensive description and analysis of the groups of subjects that did not respond or did not consent was performed (data not shown). In general, the clinical characteristics of the non-participants did not essentially differ from those of the 1,040 participants (data not shown).
Table 1: Demographic and clinical characteristics of the participants

<table>
<thead>
<tr>
<th></th>
<th>With FH</th>
<th>Without FH</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n=765</td>
<td>n=275</td>
<td></td>
</tr>
<tr>
<td>Male gender, n (%)</td>
<td>390 (51)</td>
<td>137 (50)</td>
<td>0.74</td>
</tr>
<tr>
<td>Age years</td>
<td>36.5 ± 9.4</td>
<td>38.4 ± 8.9</td>
<td>0.003</td>
</tr>
<tr>
<td>Hypertension, n (%)</td>
<td>23 (3)</td>
<td>11 (4)</td>
<td>0.43</td>
</tr>
<tr>
<td>Diabetes, n (%)</td>
<td>6 (1)</td>
<td>2 (1)</td>
<td>1.0</td>
</tr>
<tr>
<td>Smoking, n (%)</td>
<td>198 (29)</td>
<td>70 (27)</td>
<td>0.59</td>
</tr>
<tr>
<td>Highest total cholesterol mmol/L</td>
<td>9.5 ± 2.6</td>
<td>5.5 ± 1.4</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Body mass index kg/m²</td>
<td>24.2 ± 3.6</td>
<td>24.7 ± 3.6</td>
<td>0.07</td>
</tr>
</tbody>
</table>

Continuous variables were depicted as mean ± standard deviation. *Percentage based on all individuals for whom information on smoking status was available. FH = genetic diagnosis of familial hypercholesterolemia.

Life insurance

Of the 1,040 participants, 576 (55.4%) had applied for life insurance, on average at the age of 33. Of the 576 who had applied, 327 (57%) were male, and 249 (43%) were female. Other clinical characteristics did not essentially differ between the participants that either had or had not applied for life insurance.

Unconditional acceptance rate of life insurance

Of the 576 applicants for life insurance, 554 (96%) were free of cardiovascular disease at the time of application (Figure 2). Of those, 414 had FH: 255 had applied for a life insurance before genetic FH diagnosis (Period 1) of whom 182 (71.4%) were accepted unconditionally, and a total of 159 subjects had applied for a life insurance after genetic FH diagnosis (Period 2 and 3 together) of whom 134 (84.3%) were normally accepted. So, unconditional acceptance rate for FH patients improved statistically significantly from 71.4% before to 84.3% after genetic diagnosis (p<0.001).

Among the 159 subjects who applied after genetic FH diagnosis, 35 subjects had applied before the guidelines were issued (Period 2) of whom 27 (77.1%) were accepted unconditionally; 124 had applied after the guidelines were issued (Period 3), and of those, 107 (86.3%) were accepted unconditionally. The overall unconditional acceptance rate in individuals with genetic FH improved significantly over the three periods, from 71.4% in Period 1 to 77.1% in Period 2 and to 86.3% in Period 3, with a p for trend of 0.002 (Figure 3).
Figure 2: Flow diagram of participants that applied for life insurance

Questionnaire received
N=1,040

Life insurance (LI) application
N=576

Cardiovascular disease (CVD) before LI
N=22
FH+: N=18 (unconditional acc 5 (18%))
FH-: N=4 (unconditional acc 2 (50%))

Free of CVD at time of LI application
N=554

FH-
N=414

FH+
N=140

FH-
N=45

FH+
N=124

Life insurance application
by subjects free of CVD
Period 3

Period 1
N=255
Unconditional acc 182 (71.4%)

Period 2
N=35
Unconditional acc 27 (77.1%)

Period 1
N=88
Unconditional acc 87 (98.9%)

Period 2
N=7
Unconditional acc 7 (100%)

Period 3 Conditional acceptance
N=1 (2.2%)

Period 3 Unconditional acc
N=44 (97.8%)

acc = acceptance, CVD = cardiovascular disease, FH = with genetic familial hypercholesterolemia diagnosis, LI = life insurance, Period 1 = Application before genetic FH test and before the guidelines were issued, Period 2 = Application after genetic FH test and before the guidelines were issued, Period 3 = Application after genetic FH test and after the guidelines were issued.
Figure 3: Unconditional acceptance rate of life insurance for subjects with genetic FH

FH=familial hypercholesterolemia. The number of applicants accepted under normal conditions and number of overall applicants were: 182 out of 255 in Period 1; 27 out of 35 in Period 2; and 107 out of 124 in Period 3.

Subjects with conditional acceptance of the life insurance application

The life insurance applications of 17 individuals with genetic FH were conditionally accepted after the guidelines were issued in July 2003. No one was refused, 15 (88%) were accepted under the condition that they had to pay a higher premium, one female (#2; 6%) was accepted but cardiovascular mortality was excluded from coverage, and one female (#7; 6%) refused to accept the life insurance because she had to pay a severely increased premium.

In general, these 17 FH patients had a more adverse risk profile than the 107 FH patients who were unconditionally accepted after the guidelines had been issued (Table 2). In fact, 11 of the 17 conditionally accepted subjects had total cholesterol levels before treatment and life style modifications above 10 mmol/L (65%), compared with 13 (25%) of those accepted unconditionally and whose highest total cholesterol level was known (N=52). Of those accepted under specific conditions, 15 (88%) had one or more risk factors that could justify conditional acceptance based on the guidelines (Table 2). We could not identify any reason for conditional acceptance in the two remaining individuals (#2 and #12).

Of the 140 non-FH subjects who applied for life insurance, 138 (98.6%) were accepted at normal premium (Figure 2). Only one 39-year-old male (2.2%) out of the 45 non-FH subjects who applied for life insurance after the guidelines had been issued, had to pay a higher premium. While applying for life insurance, he had hypertension and he smoked. Overall, non-FH subjects were more often accepted unconditionally than subjects with FH when applying for life insurance after the guidelines had been issued (97.8% vs. 86.3%, p=0.045).
Table 2: Characteristics of the FH subjects with conditional acceptance of life insurance (after the guidelines)

<table>
<thead>
<tr>
<th>Acceptance of LI</th>
<th>Age at LI mean ± SD</th>
<th>Male n (%)</th>
<th>HT n (%)</th>
<th>DM n (%)</th>
<th>Smoking n (%)</th>
<th>BMI mean ± SD</th>
<th>Highest TC mean ± SD</th>
<th>Treated at LI</th>
<th>TC at LI</th>
<th>LDL-C* ≤ 4 mmol/L at LI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unconditional (N = 107)</td>
<td>34 ± 8</td>
<td>56 (52)</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>23 (22)</td>
<td>23.2 ± 3.2</td>
<td>8.5 ± 1.9</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Conditional (N = 17)</td>
<td>35 ± 6</td>
<td>12 (71)</td>
<td>1 (6)</td>
<td>1 (6)</td>
<td>6 (35)</td>
<td>25.2 ± 4.6</td>
<td>10.8 ± 3.2</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

Specify per subject with conditional acceptance:

#1 = 41  35  Male  No  No  No  26.5  12  Yes, statin (unspecified)  6  Unknown, TC with treatment 6.0
#2 = 103  37  Female  No  No  No  21.2  10  Yes, atorvastatin 40 mg  4.8  Yes, TC with treatment 4.8
#3 = 141  35  Male  No  Yes  No  22.4  12.9  Yes, statin (unspecified)  8.5  No, TC with treatment 8.5
#4 = 170  28  Male  No  No  Yes  21.5  13  Yes, atorvastatin 40 mg  7.2  No, TC with treatment 7.2
#5 = 195  47  Male  No  Yes  No  34.3  11  Yes, statin (unspecified)  7  No, TC with treatment 7.0
#6 = 578  34  Male  No  Yes  No  21.9  13  Yes, atorvastatin 40 mg  7  No, TC with treatment 7.0
#7 = 625  31  Female  No  No  No  21.6  19  Yes, LDL-apherese  > 6  No, TC with treatment > 6
#8 = 1015  36  Male  No  Yes  No  27.0  9  Yes, atorvastatin 40 mg and ezetimibe 10 mg  5.5  Yes, TC with treatment 5.5
#9 = 1301  23  Female  No  No  Yes  29.4  7.4  Yes, simvastatin 20 mg  5  Yes, TC with treatment 5
#10 = 1372  37  Male  No  No  No  24.8  13.6  Yes, simvastatin 20 mg  6.5  No, TC with treatment 6.5
#11 = 3055  35  Female  Yes  No  No  20.0  11  Yes, atorvastatin 20 mg  4.2  Yes, TC with treatment 4.2
#12 = 3059  32  Female  No  No  No  23.9  8.6  Yes, atorvastatin 40 mg and ezetimibe 10 mg  4  Yes, TC with treatment 4.0
#13 = 3075  34  Male  No  No  No  33.6  10.5  Yes, rosuvastatin 20 mg  7  No, LDL-C with treatment 5.5
#14 = 3225  49  Male  No  No  No  31.7  12  No  12  No, no treatment with high TC
#15 = 3497  41  Male  No  No  Yes  20.8  7.3  Yes, statin (unspecified)  < 6  Yes, TC with treatment < 6
#16 = 3506  36  Male  No  No  No  26.9  5.6  No  < 6  No, LDL-C = 4.1 at application LI
#17 = 3518  30  Male  No  No  Yes  21.7  7.5  Yes, statin (unspecified)  < 6  Yes, TC with treatment < 6

Depicted are the cardiovascular risk factors for the subjects with FH that had applied for a life insurance after the guidelines were issued. *LDL-cholesterol level if it was known at the time of application or, if unknown, an estimate of whether LDL-cholesterol would be ≤ 4.0 mmol/L or not based on known total cholesterol levels. Abbreviations: at LI = at the time of life insurance application; BMI = body mass index (kg/m²); DM = diabetes mellitus; HT = hypertension; LDL-C = low-density lipoprotein cholesterol; mg = milligram; TC = total cholesterol (mmol/L); SD = standard deviation.
Determinants of conditional acceptance of life insurance

Table 3 summarizes the association between conditional acceptance and several clinical variables for all 124 subjects with FH who had applied for life insurance since July 2003. Univariate odds ratio's show that BMI and total cholesterol were positively and statistically significantly associated with conditional acceptance, while age, male gender and smoking showed a statistically non-significant trend. Using multiple regression analysis with stepwise backward elimination, total cholesterol was statistically significantly associated with conditional acceptance. BMI was significantly associated with conditional acceptance only in the model with all 124 subjects. Highest total cholesterol level was the only independent determinant that was statistical significantly associated with conditional acceptance, for patients with FH that had applied in periods 1 and 2 (OR 1.2, 95% CI: 1.1 to 1.3; \( p = 0.006 \)). Thus, the extent of dyslipidemia is an important determinant of conditional acceptance of life insurance applications, even among the subjects that carry an FH mutation and are aware of it.

Similarly, we performed regression analysis on the subjects both with and without FH who had applied for life insurance after the guidelines had been issued. Among these 124 with and 45 without genetic FH, the presence of hypertension, higher untreated total cholesterol levels and increased BMI were all statistically significantly associated with conditional acceptance in univariate analysis. In addition, FH mutation status was associated with a non-significant increased risk of conditional acceptance (OR 7.0, 95% CI: 0.9 to 54; \( p = 0.063 \)). In a multivariate backward model which included total cholesterol levels, total cholesterol remained the only significant determinant of conditional acceptance (OR 1.6, 95% CI: 1.2 to 2.1; \( p = 0.001 \)), whereas the association between FH mutation carriersonhip and conditional acceptance was completely abolished (\( p = 0.99 \)). Thus, the extent of hypercholesterolemia appears to predominate over the presence or absence of FH genotype in determining conditional acceptance.
Table 3: Determinants of conditional acceptance for life insurance after the guidelines for the subjects with FH.

<table>
<thead>
<tr>
<th></th>
<th>Univariate OR [95% CI]</th>
<th>p-value</th>
<th>Model 1 (n=124) OR [95% CI]</th>
<th>p-value</th>
<th>Model 2 (n=69) OR [95% CI]</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age at application years</td>
<td>1.02 [0.96 – 1.09]</td>
<td>0.51</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Male gender</td>
<td>2.19 [0.72 – 6.63]</td>
<td>0.17</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Smoking</td>
<td>2.07 [0.73 – 5.90]</td>
<td>0.17</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Body mass index kg/m²</td>
<td>1.16 [1.02 – 1.33]</td>
<td>0.028</td>
<td>1.16 [1.00 – 1.35]</td>
<td>0.048</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Total cholesterol mmol/L</td>
<td>1.49 [1.13 – 1.95]</td>
<td>0.005</td>
<td>1.74 [1.27 – 2.37]</td>
<td>&lt;0.001</td>
<td>1.48 [1.12 – 1.96]</td>
<td>0.005</td>
</tr>
</tbody>
</table>

Model 1 and 2 resulted from a multiple regression analysis based on stepwise backward elimination with age, sex, smoking, body mass index and total cholesterol as independent variables. For model 1 missing values for total cholesterol and smoking were imputed. For model 2, only those subjects were included for whom all variables in the model were known, which was applicable for 69 (56%) of the 124 subjects. Abbreviations: 95% CI = 95% confidence interval; OR=odds ratio; Total cholesterol= highest level of total cholesterol measured ever.

DISCUSSION

Our findings cautiously suggest that access to life insurance for individuals with genetic FH improved with the introduction of guidelines for insurers. Although the percentage of applicants without FH that obtain insurance on normal terms (98.6%) is significantly greater than for applicants with genetic FH, the vast majority (86.3%) of FH patients are accepted without complications. As such, the unconditional acceptance for life insurance has much improved for patients with FH when compared with a decade ago. Nevertheless, individuals with genetic FH were accepted more often with additional conditions than those without genetic FH. This difference appeared to be determined to a greater extent by variations in cholesterol levels than by mutation carrier ship. Thus, access to life insurance depends more on the actual phenotype of FH than on the genotype.

The findings of a survey that also focussed on life insurance after FH diagnosis in the Netherlands were published in 2002. In that study 46 subjects had applied for life insurance in the period between genetic FH diagnosis and the time of the survey. Of those 46 individuals, 17 (37%) had experienced a broad range of problems with their life insurance application. These problems included outright refusal of the application and conditional acceptance. In that study with its limited sample size, the acceptance of life insurance applications before FH diagnosis was not evaluated. Nevertheless, the authors concluded that patients with FH encountered unanticipated insurance problems as a result of participating in the genetic testing program.
In contrast, our findings suggest that the genetic FH diagnosis itself did not result in the failure to obtain life insurance on normal terms. In fact, in our genetic FH cohort the unconditional acceptance of life insurance improved significantly from 71.4% before molecular diagnosis to 84.3% after FH was identified.

Furthermore, unconditional acceptance of life insurance applications for the cohort with genetic FH increased in successive periods. It is difficult to interpret, however, whether the improvement in access to life insurance after FH diagnosis was primarily attributable to a beneficial effect of the guidelines. In this respect, the elegant studies by Neil and colleagues are of interest. They performed a survey with life insurance applications by fictional FH patients among 41 insurance companies underwriting life insurance in the United Kingdom in 1990 and repeated the study among 26 of these companies in 2002.10, 11 They observed that insurance companies applied a much lower excess mortality rating for FH patients taking statins in 2002 than for the untreated FH patients in 1990, which indicated that insurers recognize that the prognosis of FH patients has improved with more effective treatment. Similarly, the awareness among Dutch insurers of the improved prognosis of treated FH patients made it possible to reach agreement and introduce the guidelines in 2003.7 Since then, several findings have been published which show that the prognosis for treated FH patients has improved in the post statin era.5, 12 These findings may have influenced decision-making among insurers underwriting life insurance even further. Thus, whether improvement in acceptance rate over time is owing to the guidelines or to awareness of improved prognosis of treated FH patients remains difficult to determine. One indication that the guidelines were indeed important was that insurance companies generally adhere to the guidelines. In fact, the vast majority (86.6%) of applicants with FH were accepted unconditionally, and only two of the 17 (12%) conditional acceptances after 2003 could be considered a deviation from the guidelines by the insurers.

Untreated FH patients are at severely increased risk of cardiovascular death at a young age.13 In addition, those with pathogenic mutations that result in higher LDL-C levels have shorter coronary heart disease event free survival than the somewhat milder FH mutations.14 The guidelines for insurers were designed based on that knowledge. Consequently, applicants with FH were required to have LDL-C levels below 4 mmol/L in order to be applicable for unconditional acceptance. In our study cohort, several FH patients applied for life insurance while LDL-C levels were above 4 mmol/L at that time. Among the 17 individuals with FH who were accepted with additional conditions after July 2003, at least 8 had levels above 4 mmol/L. As such, high cholesterol levels proved an important predictor of conditional acceptance.
Of note, in current clinical practice a considerable subset of FH patients does not succeed in achieving LDL-C levels below 4 mmol/L. In a large survey among 1,249 patients that attended outpatient lipid clinics in the Netherlands for more than a year, we observed that only 81% of those achieved LDL-C levels below 4 mmol/L.\textsuperscript{15} Similarly, a survey among 781 subjects who had been diagnosed by the screening program with genetic FH in 2006 showed that in 2008 only 612 (78.4%) had achieved LDL-C levels below 4 mmol/L.\textsuperscript{16} Thus, for some FH patients and physicians the target level stipulated for access to insurance could work as an additional incentive to treat to lower cholesterol levels. But for several others the target level may be discouraging, because it is unachievable for them with currently available treatment options. Even when treated with potent regimens, such patients still present insurance (i.e. actuarial) risk for insurers. Companies underwriting life insurance are private corporations that are not obliged to accept every applicant. Whether insurance companies should make an exception and offer treated FH patients normal access to insurance regardless of their cholesterol levels is beyond the scope of the current article.

One strength of this study is that the patients and control participants were recruited from a clearly defined cohort, i.e. the individuals derived from families harbouring an FH mutation and therefore visited for genetic testing. In addition, we were able to include data of more than 500 individuals that applied for life insurance. However, a potential limitation was non-response. Response rate in this kind of study is lower than in research conducted in a clinical setting and questions about privacy related issues such as health and mortgage may be perceived as sensitive. Bias arises when non-response in a survey is related to the outcome being measured. We assessed the reasons for non-response and non-participation extensively. We identified several reasons for non-response. These reasons in general did not have an impact on the validity of the outcomes, which was confirmed by the reported reasons for non-response which we obtained when we contacted a large subset of individuals that had not responded to the surface mails by telephone.

In conclusion, agreement between insurers and other parties in society to avoid genetic discrimination has proven a viable strategy to improve access to insurances for pre-symptomatic subjects with genetic FH. Accordingly, our findings may also encourage health care workers in other countries to reach similar settlements with insurance companies. Such an approach may also be applied to other treatable genetic conditions in which patient concerns about insurance discourage being tested.\textsuperscript{17, 18} Whether the Dutch approach can be extrapolated to other countries, however, also depends, to some extent on country specific legislation regarding genetic tests.\textsuperscript{19, 20}
If our findings are communicated to individuals from families harbouring an FH mutation, this may reduce the perception of discrimination based on genetic FH. It may enhance the participation rate in the screening program for FH, because it may encourage relatives to be tested, rather than being deterred by concerns about life insurance.
REFERENCES


