Addition of a Novel, Protective Family History Category
Allows Better Profiling of Cardiovascular Risk and
Atherosclerotic Burden in the General Population. The
Aklepios Study

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Masterproef voorgedragen in de master in de specialistische geneeskunde
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Abstract

Objectives: Whereas the importance of family history (FH) is widely recognized in cardiovascular risk assessment, its full potential could be underutilized, when applied with its current simple guidelines-based definition (cFH): presence of premature cardiovascular disease (CVD) in a first-degree relative. We tested the added value of a new, extended family history definition (eFH), also taking into account later onset of disease, second-degree relatives and number of affected relatives, on profiling cardiovascular risk and atherosclerotic burden in the general population.

Design: longitudinal population study.

Setting: random, representative population sample from Erpe-Mere and Nieuwerkerken (Belgium, primary care).

Subjects: 2524 male/female volunteers, aged 35–55 years, free from overt CVD.

Main outcome measures: Subjects were extensively phenotyped including presence of atherosclerosis (ultrasound) and a newly developed FH questionnaire (4 generations).

Results: Compared to cFH, eFH was superior in predicting an adverse risk profile (glycemic state, elevated blood pressure, lipid abnormalities, presence of metabolic syndrome components) and presence of atherosclerosis (all age & sex-adjusted p < 0.05). Unlike cFH, eFH remained a significant predictor of subclinical atherosclerosis after adjusting for confounders. Most relations with eFH were not graded but showed clear informational breakpoints, with absence of CVD (including late onset) in any first-degree relative being a negative predictor of atherosclerosis, and a particularly interesting phenotype for further study.

Conclusions: A novel, extended FH definition is superior to the conventional definition in profiling cardiovascular risk and atherosclerotic burden in the general population. There remain clear opportunities to refine and increase the performance and informational content of this simple, readily-available inexpensive tool.

Introduction

Cardiovascular disease (CVD) aggregates in families [1,2,3,4]. Family history (FH) represents the integration of risk within a family from shared genetic susceptibilities and familial clustering of environmental exposures, lifestyles and behaviours [5]. Accurately defining FH of CVD will have increasing importance in the prevention and treatment of CVD in the post-genome era [6,7,8]. Although the term FH is frequently used, there is no common definition [8]. Nearly all definitions are assessments of either “any FH of CVD” or “CVD history in a first-degree relative” and are usually treated as a simple binary variable according to the
occurrence or non-occurrence of disease [9,10]. The current most common definition used in guidelines (cFH) is occurrence of premature CVD (<55 years for men and <65 years for women) in a first-degree relative [11]. Taking into account additional elements could extend the information content of “family history”. Multiple approaches attempting to define which are the key elements of FH have been studied, including age at onset (premature, late occurrence of disease), degree of relationship (first, second-degree), type of relative (sibling, parent), number of affected relatives and lineage (maternal, paternal) [10,12,13,14].

In first-degree relatives coronary heart disease (CHD) risk is greater given younger ages of onset, but -to a lesser extent- also late-onset CHD is associated with early-onset CHD in the proband [10,12,13]. Furthermore, sibling history of CHD might be a stronger risk factor than parental history [10,15,16]. CHD in second-degree relatives is associated with early-onset CHD in the proband, especially with more than one affected relative or with early-onset disease [12]. Increased CHD risk is associated with increasing numbers of first- and second-degree relatives with CHD [10,12]. With regard to lineage the evidence for differential transmission of CHD is far from uniform [10,12,13,17].

Taking into account these key additional elements, we propose a novel, extended FH definition (Asklepios eFH) and define its additional value in describing the risk factor profile and presence of subclinical cardiovascular damage in a large representative population sample.

**Methods**

**Ethics Statement**

The study complies with the declaration of Helsinki, the protocol was approved by the ethical committee of the Ghent University Hospital and all subjects gave written informed consent.

**Study Population**

Subjects were derived from the Asklepios Study, an extensively phenotyped population-representative random sample of 2524 male/female volunteers aged 35–55 years, from the Belgian communities of Erpe-Mere and Nieuwerkerken, free from clinically overt CVD at baseline. An in-depth description of the ASKLEPIOS study protocol has been published [18].

Exclusion criteria were: 1. clinical presence of atherosclerosis/atherothrombosis; 2. major concomitant illness; 3. Diabetes mellitus (DM) type 1, and type 2 if proven macro-vasculopathy or significant renal impairment; 4. conditions precluding accurate haemodynamic assessment (atrial fibrillation, pregnancy); 5. inability to provide informed consent [18].

**Participant Examination: Overview**

After obtaining written informed consent, review of questionnaire data and rest, measurements included: basic clinical data, blood sampling and cardiac and vascular echography. All measurements were single observer. Blood pressure (BP) was recorded using cuff-patient matched bilateral triplicate measurements on a sitting subject using a validated oscillometric device (Omron HEM-907). Body mass index (BMI) was calculated as weight (kg)/height (m)$^2$. Metabolic syndrome (MS) was defined according to the revised ATP-III criteria [19].

**Biochemical Analyses**

All subjects were fasting, had refrained from smoking for at least 6 hours and were screened for intercurrent infection/inflammation before blood sampling (in which case blood sampling was postponed). Serum parameters were measured on a Modular P automated system (Roche Diagnostics, Mannheim, Germany), in an ISO 9002 certified reference laboratory. Impaired fasting glycermia (IFG) denotes a fasting glucose level ≥100 mg/dl and ≤126 mg/dl (diabetes). High-sensitive C-reactive protein (hs-CRP) concentrations were measured by a high-sensitive, particle-enhanced immunoturbidimetric method (Roche Diagnostics, Mannheim, Germany) [20]. Coefficient of variation (CV) of all tests described above was <3.0%. Serum oxidized low-density lipoprotein concentration was measured by a sandwich enzyme-linked immunosorbent assay (Mencodia, Uppsala, Sweden) [21,22]. Total CV was <7.4%.

**Subclinical Cardiovascular Damage**

Carotid and femoral arteries were carefully scanned bilaterally for the presence of plaque (focal protrusion >50% compared to adjacent sites, absolute thickness >1.5 mm). Intima-media thickness (IMT) was defined as the distance from the leading edge of the lumen-intima interface to the leading edge of the media-adventitia interface, measured in end-diastole, at the far wall, 1–2 cm before the bifurcations [23]. Intra-observer coefficient of variation was 5.2% [24]. Atherosclerosis was defined as a carotid or femoral IMT ≥0.9 mm and/or presence of carotid or femoral plaque.

**Family History**

The Asklepios FH Questionnaire (see Questionnaire S1), created for this study, provided data on the occurrence of CVD in 4 generations of the respondent’s genetic family (parents, grandparents, siblings and offspring). As the participants had several days to complete the questionnaire, they could obtain additional information from family members. The study nurse together with the subject reviewed the questionnaire during the visit. For all family members, respondents provided the year of birth and the year and cause of death. The questionnaire further queries for the occurrence of fatal and nonfatal CVD events: myocardial infarction, coronary revascularisation, peripheral vascular intervention of inguinal or lower limb arteries, stroke, carotid revascularisation or sudden cardiac death.

We propose a more comprehensive extended FH construct and divided participants into 3 categories according to their FH: high, moderate and low risk (Fig. 1) [25]. The eFH definition takes into account additional elements such as age at onset of disease (premature, late occurrence), degree of relationship (first, second-degree [grandparents]) and number of affected relatives. The high-risk category was based on literature review and practice guidelines [11,12]. The low-risk category was adapted from literature and a stratification model from Scheuener et al. used to address common chronic diseases in a prenatal setting and subsequently shown to be useful in assessing FH in internal medicine [6,10,12,13,26]. All other subjects where categorized as moderate risk.

**Statistical Analysis**

Statistical analysis was performed using SPSS Statistics 20.0. (SPSS Inc., Ill. USA) and R (R-2.15.3, www.r-project.org). In Table 1, we compared the anthropometric, biochemical, metabolic, lifestyle, and other classic cardiovascular risk factors in the different FH categories. As most of these variables were continuous variables, we used age- and sex-adjusted general linear models (GLM) and data are reported as estimated marginal means (95% confidence interval). For categorical variables, the differences between the FH risk categories were calculated by using chi-square tests.
We used logistic regression to calculate the adjusted odds ratios of having atherosclerosis, according cFH and eFH classes (first age- and sex adjusted and subsequently multivariate adjusted using confounders age, sex, total cholesterol, HDL-cholesterol, systolic BP, smoking, DM and BMI). The level of significance was set at \( p < 0.05 \); we used a \( p < 0.025 \) when comparing the eFH classes to account for multiple testing.

Risk models for presence of atherosclerosis were compared using the PredictABEL package within R (version 1.2–1, July 2012) [27,28,29]. As the high-risk categories are near identical in the Asklepios cFH and eFH definitions, the analysis was only meaningful in the cFH negative group in which we compared a baseline model including classic cardiovascular risk factors to a model to which the novel Asklepios eFH definition was added. The baseline risk model included age, sex, total cholesterol, HDL-cholesterol, smoking, BMI and diabetes mellitus. We assessed continuous net reclassification improvement (NRI) and integrated discrimination improvement (IDI). Since there are no meaningful risk categories for the presence of atherosclerosis, we calculated the continuous NRI, which is the most objective measure of improvement in risk prediction and can be used universally.

**Results**

The FH questionnaire was completed by 2491 out of 2524 Asklepios subjects (Fig. 1). For 2151 out of 2491 subjects (86.4%) all necessary information was available to evaluate FH. We excluded 340 subjects (13.6%) who could not be correctly classified because of insufficiently accurate knowledge of their FH. Basic characteristics and risk factor profile of this unclassifiable group were similar to the overall population, except for a higher BMI (26.5 kg/m\(^2\) versus 25.8 kg/m\(^2\)).

According to the cFH definition, 1706 subjects (79.3%) had a negative FH and 445 (20.7%) a positive FH. The Asklepios eFH classification categorized 419 subjects (19.5%) as low, 1280 (59.5%) as moderate and 452 (21.0%) as high risk.

The new eFH high-risk group is almost identical to the guidelines-defined cFH positives (intraclass correlation coefficient 0.995). Seven subjects, categorized as negative in the cFH were categorized as high risk in the eFH (those having at least two second-degree relatives (grandparents) with premature CVD). The new eFH definition mainly differs from the conventional definition (cFH) by sub-stratifying the cFH negative group into two categories in the eFH: a large moderate-risk subgroup and a smaller low-risk subgroup (Fig. 1).

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* Positive if at least 1 first degree relative with premature CVD

Premature denotes occurrence of CVD < 55 years in men and < 65 years in women

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**Figure 1. Overview of the distribution of subjects according to the conventional and to the proposed new extended family history definitions.** The graph shows the distribution of participants according to the conventional, guidelines-based definition (cFH) and the proposed new extended Asklepios family history definition (eFH). The new eFH high-risk group is almost identical to the guidelines-defined cFH positives. Seven subjects, categorized as negative in the cFH were categorized as high risk in the eFH (those having at least two second-degree relatives (grandparents) with premature CVD). The new eFH definition mainly differs from the conventional definition (cFH) by sub-stratifying the cFH negative group into two categories in the eFH: a large moderate-risk subgroup and a smaller low-risk subgroup.

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<table>
<thead>
<tr>
<th>Variable</th>
<th>New extended family history definition (Asklepios eFH)</th>
<th>Test statistic</th>
<th>Conventional family history definition (cFH)</th>
<th>Test statistic</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Low risk</td>
<td>Moderate risk</td>
<td>High risk</td>
<td>Low risk</td>
</tr>
<tr>
<td></td>
<td>n = 419</td>
<td>n = 1280</td>
<td>n = 452</td>
<td></td>
</tr>
<tr>
<td>Age (years)*</td>
<td>44.2 (43.6–44.8)</td>
<td>46.4 (46.0–46.7)</td>
<td>46.2 (45.7–46.8)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Sex (% male)</td>
<td>47.8 (42.9–52.6)</td>
<td>48.1 (45.3–50.8)</td>
<td>46.6 (42.0–51.2)</td>
<td>0.86</td>
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<tr>
<td>Height (cm)</td>
<td>169.3 (168.7–169.9)</td>
<td>169.1 (168.8–169.4)</td>
<td>169.3 (168.8–169.9)</td>
<td>0.75 0.58 0.94 0.51</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>73.1 (71.9–74.3)</td>
<td>73.7 (73.0–74.4)</td>
<td>74.9 (73.8–76.0)</td>
<td>0.072 0.034 0.038 0.071</td>
</tr>
<tr>
<td>Body Mass Index (kg/m²)</td>
<td>25.3 (25.0–25.7)</td>
<td>25.6 (25.4–25.9)</td>
<td>26.0 (25.6–26.4)</td>
<td>0.048 0.016 0.018 0.095</td>
</tr>
<tr>
<td>Waist circumference (cm)</td>
<td>85.7 (84.7–86.7)</td>
<td>86.4 (85.8–87.0)</td>
<td>87.9 (87.0–88.8)</td>
<td>0.004 0.017 0.002 0.010</td>
</tr>
<tr>
<td>Metabolic, biochemical, inflammatory parameters</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Total cholesterol (mmol/l)</td>
<td>5.5 (5.4–5.6)</td>
<td>5.6 (5.6–5.7)</td>
<td>5.6 (5.6–5.7)</td>
<td>0.062 0.010 0.028 0.96</td>
</tr>
<tr>
<td>LDL-cholesterol (mmol/l)</td>
<td>3.30 (3.21–3.38)</td>
<td>3.31 (3.35–3.46)</td>
<td>3.34 (3.36–3.52)</td>
<td>0.042 0.024 0.052 0.46</td>
</tr>
<tr>
<td>Non-HDL-Cholesterol (mmol/l)</td>
<td>3.84 (3.75–3.93)</td>
<td>3.98 (3.93–4.03)</td>
<td>4.04 (3.95–4.13)</td>
<td>0.005 0.005 0.008 0.26</td>
</tr>
<tr>
<td>HDL-cholesterol (mmol/l)</td>
<td>148.1 (146.6–151.7)</td>
<td>153.6 (151.6–155.7)</td>
<td>155.9 (152.6–159.3)</td>
<td>0.008 0.006 0.008 0.011</td>
</tr>
<tr>
<td>Ln Triglycerides (mmol/l)</td>
<td>4.49 (4.44–4.53)</td>
<td>4.56 (4.53–4.59)</td>
<td>4.60 (4.56–4.65)</td>
<td>0.45 (4.52–4.57)</td>
</tr>
<tr>
<td>Lipid-lowering therapy (%)</td>
<td>5.6 (3.2–8.0)</td>
<td>6.9 (5.6–8.3)</td>
<td>6.9 (4.6–9.1)</td>
<td>0.62 0.36 0.38 0.98</td>
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<tr>
<td>Glycemic state (%)</td>
<td>89.7</td>
<td>84.4</td>
<td>84.1</td>
<td>0.009 0.024 0.048 0.198</td>
</tr>
<tr>
<td>IFG ≥100 mg/dl</td>
<td>9.5</td>
<td>143</td>
<td>13.5</td>
<td></td>
</tr>
<tr>
<td>Type 2 diabetes</td>
<td>0.7</td>
<td>1.3</td>
<td>2.2</td>
<td>1.2</td>
</tr>
<tr>
<td>Number of MS components (%)</td>
<td>0.68</td>
<td>23.4</td>
<td>33.1</td>
<td>38.4</td>
</tr>
<tr>
<td>Systolic BP (mmHg)</td>
<td>124.8 (123.5–126.0)</td>
<td>127.1 (126.3–127.8)</td>
<td>128.5 (127.2–129.7)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>
Table 1.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Low risk</th>
<th>Moderate risk</th>
<th>High risk</th>
<th>Overall</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low vs Moderate</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Test statistic ρ</td>
<td>$&lt;0.001$</td>
<td>$0.005$</td>
<td>$&lt;0.001$</td>
<td>$&lt;0.001$</td>
</tr>
<tr>
<td>p</td>
<td>0.001</td>
<td>0.006</td>
<td>0.003</td>
<td>0.004</td>
</tr>
</tbody>
</table>

**Diastolic BP (mmHg)**
- Low vs Moderate: 78.6 (77.6–79.5) vs 80.0 (79.5–80.6) vs 81.4 (80.5–82.3)
- Test statistic p: 0.001, 0.005, 0.001, 0.008

**Drug-treated hypertension (%)**
- Low vs Moderate: 7.5 (4.6–10.4) vs 11.1 (9.4–12.7) vs 12.0 (9.1–14.7)
- Test statistic p: 0.067, 0.035, 0.019, 0.64

**Physical activity (MET; 3.5×/week, 37 (33–42) vs 36 (34–39) vs 34 (29–38)**
- Test statistic p: 0.51, 0.75, 0.31, 0.30

**Ever smoker (%)**
- Low vs Moderate: 44 (39–49) vs 49 (46–52)
- Test statistic p: 0.173, 0.039, 0.112, 0.75

**Global risk calculation**
- Ln 10-year SCORE risk (Belgium; current age; %)
- Test statistic p: 0.60, 0.52, 0.56, 0.59

**Novel Asklepios eFH Versus Guidelines-defined cFH**

Age and sex-adjusted analyses according to cFH and eFH on anthropometric, biochemical, metabolic, lifestyle, and other classic cardiovascular risk factors are shown in Table 1.

In age and sex-adjusted GLM analyses, a FH of CVD (using either definition) was associated with significantly higher BMI, systolic and diastolic BP, triglycerides, oxidized LDL-cholesterol, 10-year CVD risk (SCORE) and a significantly lower HDL-cholesterol.

Furthermore the novel eFH definition (but not cFH) showed significant associations with glycemic state, non-HDL-cholesterol, LDL-cholesterol, the number of MS components, and it showed borderline significant associations with total cholesterol and fibrinogen.

We subsequently assessed 2×2 comparisons (using age and sex-adjusted GLM) of the three categories of the novel eFH (low, moderate and high risk; Table 1). Most of the additional information when using the novel eFH definition can be explained by the newly defined low risk category (Table 1). The risk profile of the eFH moderate-risk group is not that different from the eFH high-risk group and the near-identical cFH positive group. Therefore, the lower risk profile in the eFH negative group (consisting of eFH low+moderate-risk groups), seems to be mainly driven by the admixture of the eFH low-risk component (Table 1; total, LDL- and non-HDL-cholesterol, triglycerides, glycemic state, number of MS components, oxidized LDL-cholesterol, systolic BP and 10-year CVD risk).

**Subclinical Vascular Damage**

Finally, we assessed the burden of subclinical cardiovascular damage according to both FH definitions. The prevalence of atherosclerosis according to both FH definitions in our study population is presented in Fig. 2.

In age- and sex-adjusted logistic regression analyses, taking the moderate-risk eFH group as the reference category, odds ratios for prevalent atherosclerosis were 0.67 (95% CI 0.51–0.87, p = 0.003) in the low-risk eFH group versus the moderate-risk eFH group. There was no significant increase in prevalent atherosclerosis when comparing eFH high-risk versus moderate-risk categories. For the conventional definition, there was no significant difference on the prevalence of atherosclerosis between eFH positives versus negatives (OR 1.22, 95% CI 0.96–1.54, p = 0.106).

In multivariate adjusted analyses, using classical confounding risk factors (age, sex, total cholesterol, HDL-cholesterol, systolic BP, smoking, BMI and DM), the odds ratios were 0.74 (95% CI 0.56–0.98) in the low-risk eFH group versus the moderate-risk eFH group (Fig. 3). Again, no significant increase in prevalent atherosclerosis was observed when comparing eFH high-risk versus moderate-risk categories or when using the guidelines definition.

We performed sensitivity analyses by 1) analyzing the data for women and men separately, 2) for subjects above and below the age median (45 years) separately, and 3) corrected for educational achievement (as a proxy for social class). We also performed the multivariate analyses by using 1) waist hip ratio and 2) waist circumference instead of BMI as a marker of obesity. The results remained essentially unchanged by these further analyses.

Finally, we tested net reclassification improvement (NRI) and integrated discrimination improvement (IDI) on predicting the presence (or absence) of atherosclerosis by adding the novel eFH definition to a multivariable prediction model including age, sex, systolic blood pressure, total cholesterol, HDL-cholesterol, smoking, BMI and diabetes mellitus in those subjects in the eFH
negative (eFH low and moderate risk) categories. The continuous NRI was significant (0.217 (95% CI 0.120–0.315), p = 0.00001), the IDI borderline significant (p = 0.068).

Discussion

In this study, we designed and tested a novel extended family history definition (Asklepios eFH) which: (1) better correlates with metabolic risk factor burden and; (2) independently predicts the presence of subclinical atherosclerosis beyond conventional risk factor burden, unlike cFH assessments. The eFH definition also shows significant improvement in reclassification for the prediction of prevalent atherosclerosis. The new definition differs from the conventional cFH definition in two aspects. First, it includes key additional elements (identified from literature review) so that the new eFH definition also takes into account later occurrence of disease, disease in second-degree relatives (grandparents) and number of affected relatives. Second, the eFH definition divides participants into 3 rather than 2 categories (Fig. 1): a high-risk group, which is almost identical to the guidelines-defined cFH positive group, and a moderate-risk and novel low-risk group, that has a manifestly lower prevalence of subclinical atherosclerosis.

The new Asklepios eFH definition is superior in detecting adverse CVD risk profiles in the general population. It exposes a significantly greater differential in risk profile than cFH. The eFH definition (but not the cFH definition) was additionally associated with unfavourable glycemic and lipid profiles, more components of the metabolic syndrome and more atherosclerosis. Furthermore, an important novel finding is that most relations with cFH were not graded but showed clear informational breakpoints with the eFH low-risk group being particularly interesting. Much of the additional information extracted by the novel eFH definition can be attributed to the presence of this newly defined low-risk category. The cFH positive group, the eFH high-risk and large eFH moderate-risk groups have quite similar risk factor profiles and preclinical atherosclerotic burdens. Separating out the low-risk category in many cases abolished a large part of the step-up in adverse risk profiles found in the cFH positive group, suggesting that the differences between the guidelines positive and negative groups are (in part) due to the admixture of the eFH low-risk group to the latter (i.e. some differences are due to a significantly better risk profile in the eFH low-risk group).

Most importantly, the new eFH definition (but not the conventional definition) was able to identify presence of atherosclerosis beyond conventional risk factor burden, indicating that FH conveys additional information, not completely characterized by simply measuring a risk factor profile.

A few studies already showed the value of extending FH beyond a simple yes/no question about presence of disease in a first-degree relative [10,12,14]. Scheuner et al. investigated various binary definitions of FH and found significant associations between a personal history of CHD and an additional FH that goes beyond having first-degree relatives with early-onset CHD [12]. In line with their work we took into account key additional elements in our new eFH definition, and we elaborated on their dichotomous definitions by defining a three-tier definition encompassing both the classic high-risk group, as well as a novel and highly interesting low-risk group.

Figure 2. Presence of atherosclerosis according to the proposed new extended family history definition (eFH) and the conventional definition (cFH). Unadjusted data show remarkably less atherosclerosis in the eFH low-risk group (14%) versus the eFH moderate-risk (42%) and eFH high-risk group (43%). There were no large differences observed in prevalence of atherosclerosis when comparing the cFH positive versus negative group (43% versus 39%).

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Asklepios Extended FH

Conventional FH

Figure 2. Presence of atherosclerosis according to the proposed new extended family history definition (eFH) and the conventional definition (cFH). Unadjusted data show remarkably less atherosclerosis in the eFH low-risk group (14%) versus the eFH moderate-risk (42%) and eFH high-risk group (43%). There were no large differences observed in prevalence of atherosclerosis when comparing the cFH positive versus negative group (43% versus 39%).

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That presence of a positive FH is associated with an adverse risk profile is largely in agreement with published reports [30,31]. This familial aggregation of cardiovascular risk factors reflects the genetic and environmental influence on the causal pathways of familial CVD. In line with published data we found an association with fibrinogen (a thrombotic risk factor), but not with hs-CRP, for which literature is inconsistent [32,33,34,35,36].

Regarding the presence of atherosclerosis (a prognostically well-validated non-invasive surrogate endpoint to assess cardiovascular risk), the eFH low-risk group had significantly less atherosclerosis compared to the eFH moderate- and high-risk groups. Although other studies already demonstrated significantly more atherosclerosis in patients with a FH of CVD [37,38]. Our study shows, for the first time, that this is in large part due to significantly less atherosclerosis in a "low-risk" group that can be readily identified by eFH (but not by cFH), separating it from an intermediate group (which would conventionally be classified as having a negative FH). The eFH moderate risk group demonstrates almost identical prevalence of atherosclerosis as the eFH positive group (corresponding to the eFH high-risk group). This large eFH moderate-risk group with a substantial atherosclerotic burden would be overlooked when using the conventional FH definition.

Clinical Relevance

Although FH is an important risk factor for CVD and plays an important role in medical practice, it is underused in CVD prevention efforts [8,39]. Recent data from a randomized controlled trial looking at the added value and feasibility of systematically collecting FH, found that it increases the proportion of persons identified as having high cardiovascular risk for further targeted prevention (many risk factors are amenable to interven-

dction, by lifestyle or pharmacologically) [40,41]. FH also received increased visibility as a risk qualifier in the new European guidelines on CVD prevention, where a positive cFH is considered to increase the 10-year risk of a fatal CV by 1.7-fold in women and by 2.0-fold in men [42]. Conversely, it is suggested that 10-year CVD risk (SCORE) may be lower in those with a FH of longevity. Whilst knowledge of FH may not affect clinical decision making in those at very high or very low predicted risk, it may aid in discriminating risk among the very large group of subjects at intermediate levels of predicted risk [43]. Since low-risk populations for CVD are gaining interest, our eFH low-risk group with a more favourable risk profile and significantly less atherosclerosis, could be an interesting phenotype for further study [44]. Yeboah et al. recently studied novel risk markers, comprising (premature and non-premature) FH of CHD in a first-degree relative, for improvement in cardiovascular risk assessment in intermediate-risk individuals. The authors found that FH was an independent predictor of CHD/CVD in intermediate-risk individuals. Furthermore, besides coronary calcium score, FH performed the best for CHD risk reclassification (NRI = 0.160). Interestingly, most of the correct reclassification was based on subjects reclassified into a lower risk category [45].

Intuitively, it seems likely that shared lifestyle risk factors (smoking, diet, physical inactivity) represent (non-genetic) pathways through which FH influences risk of CVD. The literature is inconsistent [36,46]; we found no clear associations between FH and lifestyle parameters. It is possible that increased perception of familial risk does not automatically lead to changed behaviour; some people may even adopt a fatalistic outlook and make no efforts at all to decrease their risk [47].

Figure 3. Adjusted odds ratios for presence of subclinical atherosclerosis according to the proposed new extended family history definition (eFH) and the conventional definition (cFH). Odds ratios (95% confidence intervals) for the presence of subclinical atherosclerosis were adjusted for age, sex, total cholesterol, HDL-cholesterol, systolic BP, smoking, DM and BMI. Taking the moderate-risk eFH group as the reference category, odds ratios for prevalent atherosclerosis adjusted for classical risk factors mentioned above are 0.74 (95% CI 0.56–0.98) in the low-risk eFH group versus the moderate-risk eFH group. There was no significant increase in prevalent atherosclerosis when comparing cFH positives versus negatives or eFH high-risk versus moderate-risk categories.

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Study Strengths and Limitations

The major limitation of the present study is the absence of outcome data. We used a surrogate measure that is well validated as a prognostic marker: presence of atherosclerosis. Furthermore, it remains to be tested if our findings can be extrapolated to the much younger (where relatives might not have aged sufficiently to have suffered CV events) or older populations. One of the major problems of defining and studying FH is that it is a “moving target”. A 52-year old man today that is eFH low-risk could end up tomorrow in the eFH high-risk category after his brother had a myocardial infarction. Reassuringly, sensitivity analyses in our cohort looking at the older and younger subjects (with correspondingly older/younger family members) showed similar results. A major strength is the population-based nature of the study, combining a well-balanced, representative sample with stringent methodology, and a broad and detailed array of carefully assessed cardiovascular intermediate phenotypes.

Self-reported FH of CVD was not validated through medical records, which is another potential limitation. Many, though not all, previous studies showed that questionnaires considering FH of CVD can be considered as accurate and people can correctly report their FH for CVD [15,48,49,50,51]. Moreover, our assessment of FH through self-report is similar to general practice, thus the present findings can be generalized to the usual clinical setting. We used a categorical definition rather than a (theoretically attractive) continuous FH score, where entry is restricted to more informative (i.e. larger) families and a single affected family member is ruled out [50]. Considering the characteristics of our study population families (average European family size), we followed the recommendations from Silberberg et al., who previously recommended that the use of categorical definitions are more likely to be adequate in smaller families and few affected relatives [52]. Finally, 13.6% of subjects could not be correctly classified because of insufficient accurate knowledge of their FH.

Conclusion

In this study, we designed and tested a novel extended family history definition (Aklepios eFH) which: (1) better correlates with metabolic risk factor burden and (2) independently predicts the presence of subclinical atherosclerosis beyond conventional risk factor burden, unlike cFH assessments and (3) shows significant improvement in reclassification for the prediction of prevalent atherosclerosis. Adding information on non-first degree relatives, late occurrence of disease and number of affected relatives to the FH construct improves the discrimination for cardiovascular risk factors and atherosclerotic burden in order to better target individuals for CVD prevention efforts. The new eFH definition separates the eFH negative group into two categories: a large eFH moderate-risk group, and a smaller eFH low-risk group. The latter is a particularly interesting phenotype for further study, having a more favourable risk profile and significantly less atherosclerosis (odds ratio 0.74). There remain clear opportunities to refine and increase the performance and informational content of this readily available, simple, inexpensive tool.

Supporting Information

Questionnaire S1 The Asklepios Family History (FH) Questionnaire. The Asklepios FH Questionnaire was created specifically for this study. It provides data on the occurrence of cardiovascular disease (CVD) in 4 generations of the respondent’s genetic family (parents, grandparents, siblings and offspring). For all family members, respondents provided the year of birth and the year and cause of death. The questionnaire further queries for the occurrence of fatal and nonfatal CVD events: myocardial infarction, coronary revascularisation, peripheral vascular intervention of inguinal or lower limb arteries, stroke, carotid revascularisation or sudden cardiac death. (PDF)

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Author Contributions

Conceived and designed the experiments: CVd ER MDB TG. Performed the experiments: CVd ER MDB TG. Analyzed the data: CVd ER MDB TG DDB. Contributed reagents/materials/analysis tools: CVd ER MDB TG DDB. Wrote the paper: CVd ER MDB TG PS DDB.

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A Novel Family History Definition
Toevoegen van een nieuwe “protectieve” categorie aan de definitie van familiale voorgeschiedenis omschrijft beter het cardiovasculair risicoprofiel en de afwezigheid van atherosclerose in de algemene bevolking. De Asklepios Studie

Inleiding
Cardiovasculair lijden komt vaak voor binnen families. Naast genetische factoren delen familieleden ook omgevingsfactoren, levensstijl en aangeleerde gedragspatronen. Binnen het domein van de cardiovasculaire preventie is het correct definiëren van familiale voorgeschiedenis belangrijk en dit zal in het post-genoom tijdperk ook steeds belangrijker worden. De term "familiale voorgeschiedenis" wordt alomtegenwoordig gebruikt zowel in de dagelijkse praktijk, als in de literatuur, hoewel er geen eenduidige definitie bestaat. Meestal maakt men gebruik van "een familielid met prematuur cardiovasculair lijden" of "vroegtijdig cardiovasculair lijden bij een eerstegraadsverwant" en wordt het veelal beschreven als een simpele binaire variabele.

In de zoektocht naar een meer performante definitie van “familiale voorgeschiedenis” zijn reeds verschillende sleutelvariabelen bestudeerd: leeftijd van het eerste cardiovasculair event, graad van verwantschap, type verwantschap, aantal aangetaste familieleden en aantasting binnen de maternele of paternele lijn.

Doelstelling
We gaan op zoek naar een nieuwe, bredere definitie (eFH; extended family history) van een positieve familiale voorgeschiedenis voor cardiovasculair lijden rekening houdend met de hierboven beschreven reeds bestudeerde variabelen om beter het cardiovasculair risicoprofiel in te schatten. We zullen de additionele waarde van de nieuwe definitie beschrijven.

Methodologie
Dit onderzoek is onderdeel van de Asklepiosstudie, die een willekeurige steekproef omvat uit de algemene bevolking met 2524 gezonde mannelijke en vrouwelijke vrijwilligers tussen 35-55 jaar, afkomstig uit Erpe-Mere en Nieuwerkerken (een
gedetailleerde beschrijving met de in- en exclusiecriteria van de ASKLEPIOS studie is reeds gepubliceerd).

De vrijwilligers ontvingen een uitgebreide vragenlijst waar, naast persoonlijke gegevens, onder andere ook de familiale voorgeschiedenis werd bevraagd. Bij het studiebezoek werden de vrijwilligers, na het doornemen van de vragenlijst, klinisch onderzocht (bloeddruk, lengte en gewicht) en werd een bloedafname verricht. Daarnaast kreeg elke patiënt ook een vasculaire echografie (carotis en femoralis) en uitgebreide echocardiografie.

De Asklepios vragenlijst (Fig 1) voor familiale voorgeschiedenis (speciaal ontwikkeld voor dit onderzoek), gaf informatie over cardiovasculair lijden binnen 4 generaties (grootouders, ouders, zussen/broers en kinderen). Van alle familieleden werd de geboortedatum en indien van toepassing de doodsoorzaak bevraagd. Daarnaast werd specifiek naar fatale en niet fatale cardiovasculaire events gevraagd: myocardinfarct, coronaire revascularisatie, perifeer vasculair lijden (onderste ledematen of revascularisatie van de carotis), CVA of plotse dood. Aan de hand van deze gegevens wordt op zoek gegaan naar een bredere definitie van familiale voorgeschiedenis van cardiovasculair lijden en wordt de additionele waarde ervan bestudeerd.

We stellen een uitgebreidere FH (family history) definitie (eFH) voor en deelden vrijwilligers in 3 categorieën in (hoog, matig en laag risico), afhankelijk van hun familiale voorgeschiedenis (Fig 2). De nieuwe definitie houdt rekening met bijkomende elementen zoals leeftijd van begin van de ziekte (prematuur of laattijdige ziekte), graad van verwantschap (eerste of tweedegraadsverwant) en het aantal aangetaste familieleden. De hoog risicogroep werd gebaseerd op gegevens uit de literatuur en op de guidelines. De laag risicogroep werd gebaseerd op een stratificatiemodel van Scheuner et al. ontworpen om (prenataal) het risico op het ontwikkelen van prevalente internistische aandoeningen in te schatten. De overige vrijwilligers vielen in de matig risicogroep.

Resultaten en discussie

In de algemene bevolking correleert de nieuwe eFH definitie beter met de aanwezigheid van metabole risicofactoren en voorspelt het daarenboven de aanwezigheid van subklinisch atherosclerotisch vaatlijden bovenop en onafhankelijk van de klassieke
risicofactoren. De nieuwe eFH definitie vertoont een significante verbetering in reclassificatie voor de predictie van aanwezigheid van atherosclerose.

De nieuwe definitie verschilt van de conventionele in twee aspecten: enerzijds houdt het rekening met laattijdige events, tweedegraadsverwanten en het aantal aangetaste familieleden. Anderzijds worden patiënten in drie in plaats van twee categorieën ingedeeld: de hoog risicogroep die bijna identiek is aan de positieve FH, een matig risicogroep en een ‘protectieve’ laag risicogroep waar een manifeste lagere prevalentie van atherosclerose werd vastgesteld.

De eFH definitie is superieur in het detecteren van een slechter cardiovasculair risicoprofiel in de algemene bevolking en was geassocieerd met een hogere glycemie, een ongunstig lipidenprofiel, meer componenten van het metabool syndroom en meer atherosclerose. De cFH (klassieke definitie) positieve FH, de eFH hoog risicogroep en grotendeels de eFH matig risicogroep vertonen een min of meer gelijkaardig risicoprofiel en aanwezigheid van subklinisch atherosclerotisch vaatlijden. De nieuwe eFH definitie was in staat de aanwezigheid van atherosclerose te voorspellen bovenop en onafhankelijk van de klassieke risicofactoren.

De belangrijkste informatie werd echter gehaald uit de laag risicogroep. De eFH stelt ons in staat een laag risicogroep te identificeren (los van de matig FH risicogroep), met significant minder atherosclerose dan de matig risicogroep. De matig risicogroep, die initieel als ‘negatieve FH’ zou bestempeld worden, vertoont een bijna identieke aanwezigheid van atherosclerose als de positieve cFH groep, die groep wordt dus waarschijnlijk onderschat.

**Klinische relevantie**
Hoewel FH een belangrijke risicofactor is in de cardiovasculaire preventie wordt deze in de dagelijkse klinische praktijk te weinig bevraagd en gebruikt. Het navragen van FH identificeert beter het aantal personen met een hoog CV risicoprofiel. FH is daarenboven een ‘risk qualifier’ in de guidelines (het tien jaarsrisico op CVD wordt x1.7 voor vrouwen en x2 voor mannen). Er wordt daarenboven gesuggereerd dat de SCORE gereduceerd zou kunnen worden bij patiënten met een gunstige FH van ‘lang leven’.
Het kennen van de familiale voorgeschiedenis zal geen verschil maken bij patiënten met een zeer hoog of zeer laag CV risicoprofiel maar zou wel kunnen helpen in de zeer grote groep mensen met een intermediair CV risicoprofiel. In een studie van Yeboah et al. was FH een independente predictor van CVD in de intermediair risicogroep, daarenboven was FH, na calciumscore, de beste parameter om patiënten te reclassificeren, opmerkelijk was echter dat de beste reclassificatie gebeurde door patiënten te reclassificeren naar een lagere risico categorie.

Intuitief zou men denken dat het delen van levensstijl (roken, dieet, fysieke activiteit) binnen een familie het risico op CVD zou kunnen beïnvloeden. Wij vonden echter geen associatie tussen FH en levensstijlparameters. Anderzijds betekent dit dat het hebben van een familiale voorgeschiedenis spijtig genoeg niet automatisch leidt naar een gezondere levensstijl.

Besluit

In deze studie ontwikkelden we een nieuwe meer uitgebreide definitie van FH die beter correleert met het metabool risicoprofiel en onafhankelijk van de klassieke risicofactoren de aan- of afwezigheid van subklinische atherosclerose voorspelt.

De nieuwe definitie toont een significante verbetering in reclassificatie voor de aanwezigheid van subklinische atherosclerose. Het toepassen van info over tweedegraadsverwanten, laattijdige ziekte en het aantal aangetaste familieleden aan de definitie van FH verbetert de discriminatie van het cardiovasculair risicoprofiel.

De grootste bevinding is dat de nieuwe definitie de ‘negatieve’ FH groep onderverdeelt in 2 categorieën: een grote matig risicogroep en kleinere laag risicogroep. Belangrijk is dat de matig risicogroep, die in de klassieke definitie als negatief beschouwd zou worden, bijna identiek is qua risicoprofiel aan de hoog risicogroep. De laag risicogroep echter blijkt een interessant fenotype voor verdere studies met een meer gunstig CV risicoprofiel en significant minder atherosclerose.
FIGUUR 1

Uw moeder

E1. Geboortedatum (jaartal)  

E2. Is deze persoon nog in leven?  
- O ja, zij woont in  
- O nee, zij is overleden in het jaar  

E3. Zo deze persoon overleden is, wat was de overlijdensoorzaak?  

E4. Maakte zij ooit het volgende door? (meerdere keuzes mogelijk)  
- O een hartinfarct of hartaanval  
- O een ingreep aan de kranen slagaders (balonondilatatie, overbruggings)  
- O een ingreep aan de slagaders van spieren of been  
- O een herseninfarct, hersenbloeding of beroerte  
- O een ingreep aan de hersenligadegaders  
- O een plots overlijden  
- O geen van bovenstaande  
- O ik weet het niet  

E5. Heeft deze persoon ooit volgende risicofactoren gehad? (meerdere keuzes mogelijk)  
- O een te hoge bloeddruk  
- O een te hoog cholesterol  
- O suikerziekte (diabetes)  
- O overgewicht  
- O heeft deze persoon gerookt (gedurende meer dan een jaar)  
- O geen van bovenstaande  
- O ik weet het niet
FIGUUR 2

ASKLEPIOS Cohort
n = 2024

No questionnaire available
n = 33
Unclassifiable family history
n = 340

Subjects included in analyses
n = 2151

Conventional Family History definition (cFH)

Positive
n = 445

Negative
n = 1706

Asklepios Extended Family History definition (eFH)

High Risk
n = 452
≥ 1 first degree relative with premature CVD
≥ 2 second degree relatives with premature CVD

Moderate Risk
n = 1280
≥ 1 first degree relative with late-onset CVD
≥ 2 second degree relatives with late-onset CVD
1 second degree with premature CVD

Low Risk
n = 419
< 1 second degree relative with late-onset CVD
no known CVD in any first or second degree relatives

* Positive if at least 1 first degree relative with premature CVD
Premature denotes occurrence of CVD < 55 years in men and < 65 years in women