

Packard, C. J., Young, R., Ross, K., Ford, I., Ambegaonkar, B. M., Brudi, P. and McCowan, C. (2017) Modelling total coronary heart disease burden and long-term benefit of cholesterol lowering in middle aged men with and without a history of cardiovascular disease. *European Heart Journal: Quality of Care and Clinical Outcomes*, (doi:10.1093/ehjqcco/qcx012)

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Deposited on: 10 May 2017

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24 25 26	Chris J Packard ¹ , Robin Young ² , Kevin Ross ² , Ian Ford ² , Baishali Ambegaonkar ³ , Philippe Brudi ³ , Colin McCowan ² .
27	
28 29 30	Robertson Centre for Biostatistics, University of Glasgow, Glasgow, Scotland, UK G12 8QQ
31 32	
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34 35	¹ College of Medical, Veterinary and Life Sciences; ² Robertson Centre for Biostatistics,
36 37	University of Glasgow, Glasgow, Scotland; ³ Merck, Sharpe & Dohme Ltd, Kenilworth, New Jersey USA
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46	Corresponding author:-
47 48	Professor Chris J Packard
49	McGregor Building, Room 203, Floor 2
50 51	Western Infirmary University of Glasgow G12 8QQ
52 53	Telephone (44) 7788447576
54	Email: chris.j.packard@gmail.com
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Abstract

Background

Cumulative CHD events over 20 years were examined in men screened for, and in those

randomised to, the West of Scotland Coronary Prevention Study.

10 Methods and Results

Record linkage provided CHD-related events and days in hospital for the 80,230 screenees,

including the randomised cohort of 6,595 men. Risk factors were determined at baseline, and
disease burden assessed for groups defined by cholesterol. Effects of cholesterol lowering were
modelled from differences between groups, and from the treatment arms of the trial.
Over 20 years, those without a history of CHD (n=61,211) had 23.0 events per 100 subjects in
the lowest cholesterol group (mean 4.0mmol/l) and 65.1 per 100 in the highest (8.8mmol/l).
Corresponding days in hospital were 167.2 to 435.4 per 100 subjects. Analogous figures for men
with a CHD history (n=8,570) were 77.3 to 141.7 events per 100 and 526.1 to 936.7 hospital days
per 100.

Lowering cholesterol by about 1.0mmol/l in men with average cholesterol and no CHD was predicted to be associated with 8.9 fewer events and a saving of 56.0 hospital days per 100. In those with CHD this difference gave, depending on starting level, 26.8 to 36.5 fewer events and savings of 158.2 to 247.3 hospital days per 100 subjects.

Comparison of cumulative events in 45-54 versus 55-64 year olds in the trial revealed greater benefit from intervention in the younger decade.

49 Conclusion

Long-term, longitudinal data reveal the considerable CHD burden in middle-aged men and
 indicate substantial clinical benefits from both moderate and aggressive cholesterol lowering.

Keywords – cholesterol, data linkage, myocardial infarction, clinical trial

Introduction

LDL lowering is a cornerstone recommendation in CHD prevention guidelines [1, 2] with metaanalyses [3, 4] and cost-effectiveness studies [5-7] confirming its clinical and economic utility in secondary prevention and high-risk primary prevention. Population benefit assessments are based necessarily on models generated with tools such as the Framingham risk equation [8], and the outcome of clinical trials that are limited (often to 3 to 5 years) in duration relative to the decades-long course of the disease. Also, traditional indices of clinical utility such as numberneeded-to-treat [9] are probabilities based on the first major event and do not take into account the fact that patients often suffer multiple episodes of CHD over a number of years. Overall, as highlighted again in IMPROVE-IT [10] and HOPE-3 [11], these approaches can lead to a substantial underestimate of the full clinical and economic impact of therapy.

A number of trials have reported extended follow up [12-15], and the present study takes advantage of the availability of comprehensive, longitudinal electronic health records not only for the subjects included in the West of Scotland Coronary Prevention Study (WOSCOPS) but also for the population of approximately 80,000 45 to 64-year-old men who were screened [16]. Based on an evaluation of total CHD burden, we modelled the benefits of having lower LDL in a primary prevention setting which is still an area of contention [17-19], and since the screened population included those with a history of CHD, we assessed also the impact of reducing total cholesterol levels in a secondary prevention setting.

Methods

Recruitment for WOSCOPS [16, 20, 21] involved inviting men aged 45 to 64 years from approximately 120 family practices to visit a screening clinic. Between September 1989 and July 8, 1991, 80,230 attended Study Visit 1 at which risk factors were assessed including cholesterol, blood pressure and smoking habit [16]. Personal history of CHD was evaluated by questionnaire based on patient recall (specifically, whether a doctor had informed the subject that they suffered from chest pain due to angina or had had a heart attack). Those with total cholesterol >6.5mmol/1 were evaluated in 3 subsequent visits and 6,595 men with LDL cholesterol (LDLc) 4.0 to 6.0 mmol/1 but no history of myocardial infarction were randomized to pravastatin 40mg/d or placebo. Followup was for a mean of 4.9 years with final trial visits held in May 1995. The outcome was a risk reduction of about one-third in a range of cardiovascular endpoints [21].

Use of lipid-lowering therapy during the first 5 years after the trial ended was monitored by review of GP records. In the original pravastatin and placebo groups respectively, 28.6% and 24.3% at 1 year, 33.6% and 29.4% at 3 years, and 38.7% and 35.2% at 5 years post-trial were on statins [12]. No further information on statin treatment was available for the trial participants after this point (i.e. 10 years post randomisation). For the screened population, no information on use of lipid-lowering treatment was available across the whole of the 20-year observation period.

Individual-level data were extracted from national electronic hospital discharge records and death registries [22] to provide numbers of clinical events and length of hospitalisations for both the WOSCOPS randomised cohort, and the screenees who did not enter the trial. Events were classified using International Classification of Diseases (ICD) codes. Hospital duration for any single event was truncated at 182 days. No limit was placed on the number of events per subject.

The original trial was approved by the Ethics committees of the University of Glasgow and participating health boards and the long-term follow-up and associated record linkage by the Ethics committee of the Royal Infirmary, Glasgow and the Privacy Advisory Committee of the National Health Service for Scotland.

Association of total cholesterol levels with hospital admissions for CHD causes (ICD9 codes
 410-414 and ICD10 codes I20-I25) was assessed over the observation period (i.e. until October
 2011) or until death. We estimated the benefit of cholesterol reduction by comparing differences
 in cumulative events between groups categorised by baseline total cholesterol level.

An analysis was also undertaken to evaluate the relative benefit of initiating cholesterol lowering at age 45 to 54 compared to 55 to 64 years in the WOSCOPS randomised cohort, and separately in screenee groups with total cholesterol levels similar to those seen in the treatment arms of the trial. The 'QRISK' tool [5,23] (www.qrisk.org) was used to estimate 10-year risk of a CVD event for males aged 50 and 60 years (i.e. the mean ages in the two age decades) with the same risk factor levels as the WOSCOPS randomised subjects at baseline [21] (total cholesterol :HDL cholesterol ratio of 5.0, systolic blood pressure 137mmHg, height 173cm, weight 77.5kg, non-smoker or moderate smoker).

Statistical methods

Total CHD burden for each individual was the cumulative number of CHD hospitalisation events and the total number of days spent in hospital attributed to CHD causes. This was analysed separately for (i) the WOSCOPS randomised cohort with its placebo and pravastatin arms, (ii) WOSCOPS screenees without a history of CHD at baseline ('primary prevention' setting), and (iii) WOSCOPS screenees with a history of CHD ('secondary prevention' setting). The last two groups were divided into five categories (P1 to P5 and S1 to S5 respectively) based on total cholesterol at screening - <=4.5, 4.5-5.5, >5.5-6.5, >6.5-8.0 and >8.0 mmol/l. These intervals were chosen so that, where possible, mean cholesterol levels differed by increments of approximately 1.0mmol/l, and hence our findings could be related to the commonly used metric of risk reduction per mmol/l change [3, 4]. Incidence of the first CHD event (death or hospitalisation for a CHD related reason), the cumulative number of hospital admissions for CHD (events and length of stay in days) and the total number of deaths were obtained for each cholesterol group. Risk ratios were derived by quasi-Poisson regression (which allows for overdispersion in the data) using the lowest total cholesterol category as referent and adjusting for other major risk factors (age, blood pressure, smoking status).

total event rates, expressed per 100 subjects over 20 years, were compared between categories of total cholesterol level to obtain relative risks and differences in number of CHD hospitalisations and hospital days attributed to CHD causes per nominal 1.0mmol/l change. The effects of more aggressive cholesterol reductions were estimated by comparing groups that differed by 2.0mmol/l or more. Groups were also characterised by the percentage of subjects experiencing CHD death, multiple CHD events or a single CHD events (see supplementary tables).

39 All statistical analyses were conducted using R version 3.1.2.

Results

A total of 80,230 men attended Study Visit 1 and individual-level record linkage-based follow-up was available for 76,376 (95.2%). Over the observation period (which ranged from 20 to 22 years), there were 35,690 hospitalisations for CHD causes in 15,398 screenees. Once those with incomplete Visit 1 information were excluded, total CHD burden data were available for 61,211 screenees with no prior history of CHD and 8,570 screenees with a history of CHD. (Note that the analyses of screenees does not include subjects randomised to the trial).

Plasma cholesterol and total CHD burden.

Over 20 years follow up, for all screenees there was a positive relationship between total plasma cholesterol and incidence of a first event of CHD hospitalisation or CHD death for both age decades (Figure 1A).

In screenees with no history of CHD (Figure 1B) the rate in the older compared to the younger decade was about 29% higher, and when these subjects, using the entire age range, were grouped by total cholesterol (P1 to P5) there was a 2.7-fold increase in adjusted risk comparing the lowest to highest levels (Table 1). The cumulative number of events per 100 subjects rose from 23.0 to 65.1, and the corresponding number of days of hospitalisation for CHD causes increased from 167.2 to 435.4 per 100 subjects.

In screenees with a history of CHD there was a positive association between first incidence of a
subsequent event and total cholesterol but little difference between the two age decades (Figure
1C). In Table 1, the adjusted risk ratio virtually doubled across the cholesterol categories S1 to
S4 but did not appear to increase further in the S5 group. In general, cumulative event rates were
2-3 fold higher compared to those seen in screenees without a CHD history. Across the

cholesterol range total CHD burden was 77.3 to 141.7 events per 100 subjects, and the number of days in hospital for CHD causes was 526 to about 1,000 per 100 subjects.

Benefit of cholesterol lowering in a 'primary prevention' setting

Differences in cumulative events and hospital days attributable to CHD for groups of interest are presented in Table 2. In taking this approach to modelling the impact of lowering cholesterol, it is assumed that any difference is attributable solely to variation in LDLc, and that the result is a reasonable estimate of the benefit of using pharmacological intervention.

Reduction in total CHD burden was predicted to be substantial in men with the highest cholesterol levels (P5) who were subject to moderate (25%) or intensive (54%) LDL lowering (Table 2). Group P4 was moderately hypercholesterolaemic with a mean cholesterol similar to that in the placebo arm of WOSCOPS (i.e 7.0mmol/l [21]). Comparing event rates in P4 with those for P3 (with its mean cholesterol of 6.0mmol/l) it can be seen that a nominal 1.0mmol/l difference was associated with a 19.4% lower risk of CHD, 10.2 fewer events and 61.3 fewer days in hospital per 100 subjects. The analogous data from the placebo and pravastatin arms in the trial (where there was a 1.1 mmol/l difference in LDLc during the formal 5-year intervention) were a relative risk reduction of 21% over 20 years of follow up [Table 2 in ref 14] and a difference in cumulative CHD events of 10.8 per 100 subjects [for non-day case admissions; Table 3 in ref 14]. The compatibility, for approximately the same degree of cholesterol lowering, of the trial results with those from modelling screenees lends support to our general approach.

In other scenarios presented in Table 2, it can be seen that even at the population average total cholesterol (in group P3), over 20 years, achievable decreases in cholesterol of the order of 0.9 to 2.0 mmol/l were predicted to prevent respectively 8.9 to 19.4 events and 56.0 to 121.1 days in hospital per 100 subjects. The impact of having lower cholesterol expressed per subject rather

than total number of CHD events is depicted schematically in Figure 2A and B. Here is can be
seen that comparing groups P1 with P3 there was a notable reduction in the numbers
experiencing single and multiple CHD events over the 20 years.
Figure 3A shows the cumulative incidence for CHD hospitalisations in 45 to 54-year-old and 55
to 64-year-old men within the WOSCOPS randomised cohort (n=6544; 51 men with incomplete
Visit 1 information were not included in this analysis), and Figure 3B the difference plot with
time. It is clear from consideration of Figure 3B that the cumulative benefit of cholesterol
lowering in primary prevention is greater if intervention is started in the first compared to the
second age decade. The former group (average age 50 years) in the placebo arm of the trial
experienced about 75 events per 100 subjects by the end of the 20-year observation period
(Figure 3A). The observed accrued benefit at age 65 and 70 years respectively was a decrease of
about 5.1 to 6.2 events per 100 subjects (Figure 3B). On the other hand, waiting until men were
55 to 64 years (average age 60 years) before intervention led to a lower accrued benefit of 2.7
events prevented per 100 subjects by age 65, and 4.7 events prevented by age 70.
The potential impact of starting cholesterol lowering earlier is even clearer if the screenee data
are used to model the outcome for each age decade (Figure 3C). Cumulative differences between
the P4 and P3 groups in Table 1 were determined, and for 45-54 year old men by age 70 years the
number of events prevented by being at a 1.0mmol/l lower total cholesterol was over 10 per 100
subjects, whereas for 55 to 64 year olds by age 70 the same difference in cholesterol saved only 6
events per 100 subjects.
Applying the QRISK2 calculation to the younger decade of the WOSCOPS randomised cohort
(Figure 3B) gave a 10-year cardiovascular disease risk of 5.2% to 9.1% depending on smoking

habit which does not meet the threshold of 10% required before drug treatment is recommended [5]. Finally, it was reassuring again that estimated rates of accrued benefit per 100 subjects were similar in the modelled outcome to those seen in the trial cohort.

Benefit of cholesterol lowering in a 'secondary prevention' setting

The predicted benefit of lowering cholesterol in the screenees with a history of CHD (Table 1), as expected, exceeded that seen in the primary prevention setting. Men in group S3 with mean total cholesterol levels close to that seen in the whole screened population (5.9mmol/l; [16]) had on average 140.6 CHD events per 100 over 20 years (Table1) and utilised 931.7 hospital days for CHD causes. In this 'secondary prevention' setting it was estimated that 26.8 events per 100 men would be prevented by lowering cholesterol by approximately 1.0mmol/l from the mean level, and about 63.2 events per 100 by reducing cholesterol by 2.0mmol/l (Table 2). The impact of this shows the substantial number of subjects experiencing CHD death and multiple CHD events and the predicted reduction when LDLc is 50% lower. The number of hospital days saved amounted to 158.2 to 405.5 per 100 subjects respectively (Table 2). Even comparing the two groups with the lowest cholesterol levels (S1 and S2) a 1.1 mmol/l difference was associated potentially with 36.5 fewer events and a saving of 247.3 hospital days per 100 subjects.

Discussion

For many subjects the 20-year observation period was an estimate of 'lifelong' benefit since the
mean age rose from 55 to 75 years and life expectancy for men in Scotland is 76 years (see
Registrar General's report at www.nrscotland.gov.uk). Our main findings were that total CHD
burden over this time was considerable and strongly related to plasma cholesterol levels both in
men without ('primary prevention' setting) or with ('secondary prevention') a history of CHD at
screening. Predicted benefits of intervention were large in that a decrease of around 1.0mmol/1
would lead 8.9 fewer CHD events per 100 subjects in primary prevention at starting total
cholesterol levels of 6.0mmol/l (close to the population average), and 26.8 to 36.5 fewer events
per 100 in secondary prevention when the initial total cholesterol was 6.0 and 5.0mmol/l
respectively. If the estimated LDLc was reduced by about 50% then the potential decrease in
0 CHD events was 19.4 to 31.5 per 100 subjects in primary prevention and 63.2 CHD events per
100 subjects in secondary prevention (Table 2). The corresponding savings in hospital days were
also substantial.

The need to take a long-term view in a disease with a decades-long trajectory is recognised increasingly. In the IMPROVE-IT [10, 24] and HOPE-3 studies [11] LDL lowering decreased risk of a second or third cardiovascular event to a similar extent to that seen for the first event. Comparing findings in these trials with the total CHD burden modelled here it can be seen that in HOPE-3 the number of co-primary endpoints prevented by a 1.0 mmol/l decrease in a cohort with population average LDLc (3.31 mmol/l) was 1.1 per 100 subjects over 5.6 years (the study duration) [11] whereas the cumulative events over 20 years predicted to be prevented in a similar group by an equivalent cholesterol reduction was 8.9 per 100 with a saving of 56 days in hospital per 100 subjects. Likewise, in IMPROVE-IT [24] the difference in LDLc between the two

treatment arms was 0.4 mmol/l and the number of primary endpoint events prevented was 1.87 per 100 subjects. The same cholesterol decrease (taken as a proportion of the difference between the levels seen in groups S2 and S1 in Table 2) would give 13.3 fewer CHD events and save 95.4 hospital days per 100 subjects over 20 years. With respect to European guidelines [2], patients at very high risk have a recommended LDLc goal of <1.8mmol/l and a desired reduction of >=50% if the starting LDLc is 1.8 to 3.5mmol/l. The present study predicts that the long-term (lifelong) benefit of LDL lowering of this magnitude would be considerable as can be seen by comparing groups S3 and S1 for total events in Table 2, and on a per-subject basis in Figure 2.

Even with the demonstration that primary prevention subjects have a useful risk reduction [19,
25] in an era when the costs of intervention are modest, there is a perceived reluctance to begin
therapy, possibly because the benefits are difficult to quantify in terms understood by patients
and occur in the future [17-19]. Presentation of the total CHD burden that an individual may
experience, and the considerable advantages of treatment, may help deliver better strategies and
compliance. Further, our findings in the two age decades add to conclusion from genetic studies
[26] that earlier intervention is better. Greater emphasis on benefit rather than risk (as
exemplified here in the QRISK2 results in the younger decade) may help further refine
widely in the population [17-19]. This concern, including the increased risk of developing type 2
diabetes, has been addressed in meta-analyses of trials [19], and also in the previously published
long-term followup of the WOSCOPS trial cohort [12,14].

Advantages of the present study were the size of the screened population and the longitudinal nature of the data sets. Also, it was notable that the modelled scenarios for an approximate 1.0mmol/l difference in cholesterol were associated with about a 20% lower relative risk, a value

in line with the 22% seen in meta-analysis of outcome trials [3, 4]. Likewise, the approximately 3-fold higher event rates seen in the 'secondary' versus 'primary' groups were in accord with the relative rates for similar CHD endpoints in 4S (a trial in secondary prevention) and WOSCOPS, contemporaneous studies in subjects with similar high LDLc levels [21, 27]. Limitations were that events were not adjudicated, no women were included, the modelling approach estimated treatment benefit from a difference in cholesterol levels in an epidemiological survey, and we had no information on which screenees were prescribed lipid-lowering medication during followup. Arguably, since statins once they were introduced would be used mainly in those with higher cholesterol levels, this would have the effect of compressing observed differences in event rates between the groups, and so the data presented may be a conservative estimate of benefit. On the other hand, population rates of CHD in Scotland fell substantially over the period of followup, so the impact of lowering cholesterol today may be quantitatively less than the values quoted in the current study (although the majority of events reported for our subjects occurred in the latter 10 years of observation).

In conclusion, this study used electronic health records to create a disease trajectory for each subject and permitted the modelling of clinical benefit of an intervention based on long-term observational data. Our findings support the view that primary CHD prevention is more effective if initiated in early mid-adult life and that the benefits of secondary prevention, given the very high total burden of disease, are substantial.

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Figure legends

Figure 1: Relationship between plasma cholesterol and first CHD event (death or hospitalisation) in (A) the whole group of screenees, (B) men with no self-reported history of CHD at screening, and (C) men who reported a history of CHD at screening.

Figure 2: CHD impact over 20 years for groups P1 vs P3 and S1vs S3 expressed as percentages of subjects experiencing a fatal CHD event, multiple non-fatal CHD events or a single CHD event. These are depicted schematically per 50 subjects for the purposes of clarity. Note, if a subject has both a non-fatal and then a fatal event, he is counted for both in order to show the total CHD burden. Complete data are provided in supplementary tables 1 and 2.

Figure 3: Cumulative events over 20 years for WOSCOPS randomized cohort divided by age
decade and treatment allocation (panel A). Panel B is the difference plot (placebo minus
pravastatin) for each age decade for cumulative events in the two treatment arms. Panel C is the
difference plot by age decade for cumulative CHD events in groups P4 and P3 in Table 1. Groups
P4 and P3 comprise men who were not included in the WOSCOPS randomised trial cohort but
had the same cholesterol levels as the two treatment arms. In panel B 10-year risk estimates
obtained using the QRISK-2 algorithm are provided for the average WOSCOPS trial participant
for non-smokers and moderate smokers.

Acknowledgments

Ian Ford and Robin Young had full access to all the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis.

All authors from the academic institution contributed to the study design, analysis plan,

interpretation of the data, and preparation of the manuscript. Baishali Ambegaonkar and Philippe
Brudi (employees of the funder) contributed to the conceptual design and analytical plan, but had
no role in the conduct of the study or in the analysis of the data. The 'stick-man' schematic in
Figure 2 is borrowed from a diagram generated originally by Prof. Ulrich Laufs.

Funding Sources

The study was supported by a grant from Merck, Sharp & Dohme, Kenilworth, New Jersey as part of an Investigator Initiated Programme. The funder was provided with a draft for comment but editorial control remained with Drs Packard and Ford.

We would also acknowledge the support from The Farr Institute @ Scotland. The Farr Institute
@ Scotland is supported by a 10-funder consortium: Arthritis Research UK, the British Heart
Foundation, Cancer Research UK, the Economic and Social Research Council, the Engineering
and Physical Sciences Research Council, the Medical Research Council, the National Institute of
Health Research, the National Institute for Social Care and Health Research (Welsh Assembly
Government), the Chief Scientist Office (Scottish Government Health Directorates), the
Wellcome Trust, (MRC Grant No: MR/K007017/1).

Conflict of interest disclosures

 Authors Ian Ford, Robin Young and Colin McCowan have no conflict of interest to disclose other than research funding for this project from Merck, Sharpe and Dohme, as declared above. Chris Packard has research grants from Roche and honoraria from Merck, Sharpe and Dohme, Pfizer, and Sanofi. Baishali Ambegaonkar and Philippe Brudi are employees of Merck, Sharpe

12 and Dohme, Kenilworth, NJ, USA.

Table 1: Cumulative CHD events and hazard ratios for screened population by cholesterol level.

Cholesterol Group (limits in mmol/l)	Mean <mark>total</mark> cholesterol <i>[LDL]</i> (mmol/l)	Mean Age (years)	Smokers (current % / former %)	Mean systolic/diastolic BP (mmHg)	No. of subjects (% died)	Observed CHD hospitalisation events per 100 subjects over 20 years	Observed CHD hospital days per 100 subjects over 20 years	Adjusted hazard ratio ^a (95% CI)
]	No CHD history – '	primary prevention'		
P5 (>8.0) ^b	8.77 <i>[6.8]^c</i>	54.1	46.0%/ 29.3% ^d	140.4/86.6	1790 (42.0%)	65.1	435.4	2.7(2.4-3.1)
P4 (>6.5-8.0)	7.05 [5.0]	54.5	42.9%/ 27.4%	139.6/86.2	10767 (39.7%)	52.7	349.6	2.2(2.0-2.5)
P3 (>5.5-6.5)	5.98 [4.0]	54.7	42.0%/ 26.4%	137.7/85.1	22288 (37.3%)	42.4	288.3	1.8(1.7-2.0)
P2 (>4.5-5.5)	5.06 [3.1]	54.6	41.7%/ 25.8%	136.3/84.3	18952 (36.9%)	33.6	232.2	1.5(1.3-1.6)
P1 (<=4.5)	4.00 [2.0]	54.6	43.7%/ 23.9%	135.2/83.4	7414 (41.2%)	23.0	167.2	1.0(referent)
				W	ith CHD history – '	secondary prevention	n'	
S5 (>8.0)	8.81 [6.8]	56.8	47.0%/ 36.1%	138.0/84.8	477 (64.8%)	141.7	936.7	1.8(1.4-2.3)
S4 (>6.5-8.0)	7.10 [5.1]	57.3	40.8%/ 40.6%	137.7/84.8	2210 (64.3%)	152.9	1089.9	1.9(1.6-2.4)
S3 (>5.5-6.5)	6.02 [4.0]	57.6	40.2% 39.1%	138.1/84.7	3105 (61.3%)	140.6	931.7	1.8(1.5-2.2)
S2 (>4.5-5.5)	5.08 [3.1]	57.7	43.5%/ 36.1%	137.2/83.9	2037 (61.8%)	113.8	773.4	1.5(1.2-1.8)
S1 (<=4.5)	3.98 [2.0]	57.7	50.6% 30.4%	135.9/83.1	741 (67.9%)	77.3	526.1	1.0(referent)

^a Hazard ratio from quasi-Poisson regression adjusted for age, smoking and blood pressure.

^b Subjects were categorised into groups according to cholesterol level and CHD history at screening; S1 to S5 had a history of CHD and P1 to P5 did not.

^c LDL cholesterol values were assumed to be 2.0 mmol/l less that the total cholesterol as in [16]

^d Percentages are given for current and former smokers, never smokers comprised the remainder.

Table 2: Modelling prevention scenarios using cumulative CHD event and hospitalization rates in the WOSCOPS screened population.

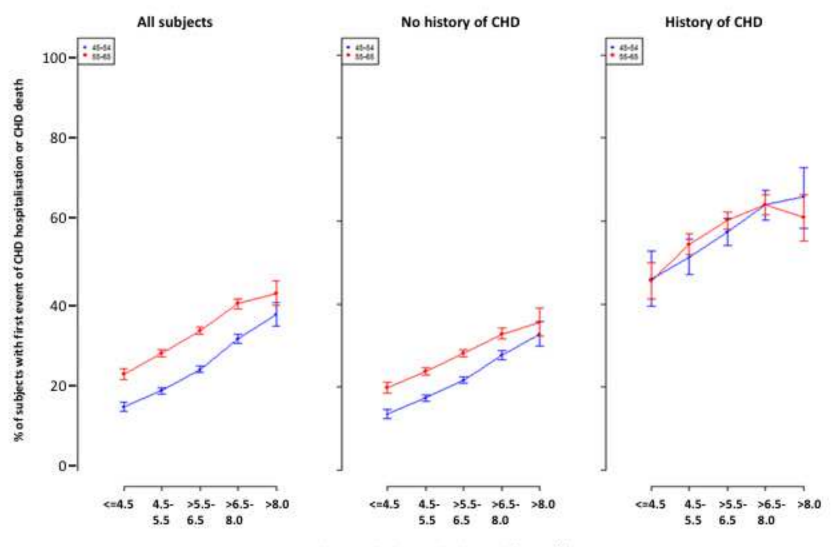
Cholesterol difference	Relative risk reduction ^a	Total CVD events difference over 20 years per 100 subjects	Total CVD hospital days difference over 20 years per 100 subjects
No	CHD history (p	rimary prevention setting)	
Severe hypercholesterolaemia – P5; st	arting total choleste	erol of 8.8 mmo/l (LDL approx 6.8 i	nmol/l) ^b
1.7 mmol/l (25% LDL) ^c difference (change to level seen in P4) 3.7 mmol/l (54% LDL) difference	19.1%	12.4	85.8
(change to level seen in P2)	48.4%	31.5	203.2
Moderate hypercholesterolaemia – P4, WOSCOPS equivalent	; starting total chol	esterol of 7.0 mmol/l (LDL approx.	5.0 mmol/l)
1.0 mmol/l (20% LDL) difference (change to level seen in P3)	19.4%	10.2	61.3
Population average –P3; starting total	cholesterol of 6.0 n	nmol/l (LDL approx 4.0 mmol/l)	
0.9 mmol/l (23% LDL) difference (change to level seen in P2)	20.9%	8.9	56.0
2.0 mmol/l (50% LDL) difference (change to level seen in P1)	45.7%	19.4	121.1
Previou	IS CHD history	(secondary prevention settin	g)
Population average – S3; starting cho	lesterol of 6.0 mmol	l/l (LDL approx 4.0 mmol/l)	-
1.0 mmol/l (23% LDL) difference (change to level seen in S2)	19.1%	26.8	158.2
2.0 mmol/l (50% LDL) difference (change to level seen in S1)	45.0%	63.2	405.5
Low – S2; starting total cholesterol of	5.0 mmol/l (LDL ap	pprox 3.0 mmol/l)	
1.1 mmol/l (35% LDL) difference (change to level seen in S1)	32.0%	36.5	247.3

^a Cholesterol lowering scenarios were tested for a range of starting total cholesterol levels using the groups defined in Table 1. Relative risk reduction compared the CHD event rate in the lower versus higher cholesterol group.

^bEstimated LDL cholesterol was 2.0mmol/l less than the total plasma cholesterol level; see [16].

^c The percent difference in parentheses refers to the predicted difference in LDL cholesterol between the two groups examined. Since they are based on estimates the % change is only an approximate guide.

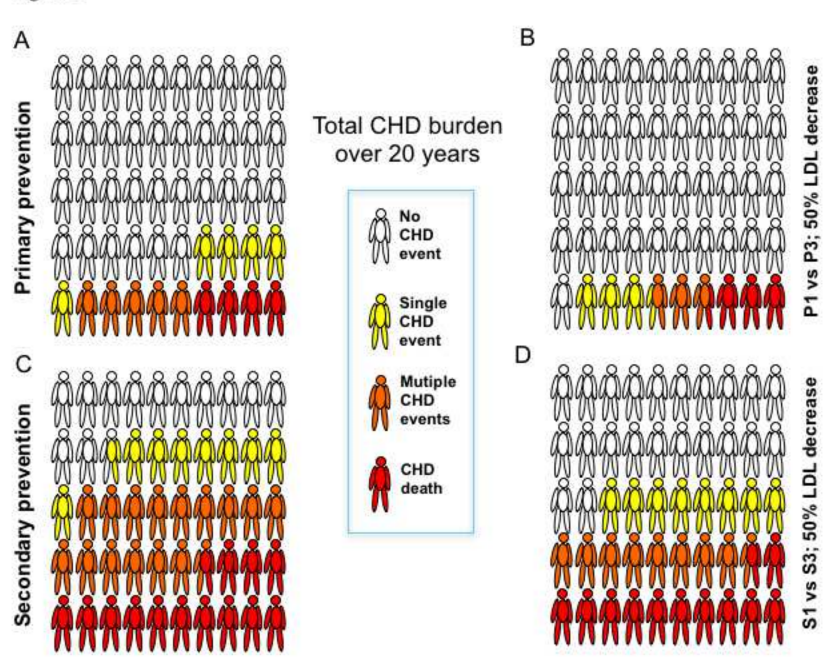
Figure 1.



Baseline total plasma cholesterol (mmol/l)

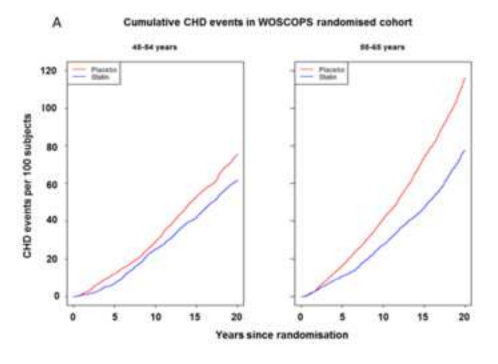
<u>±</u>

Figure 2



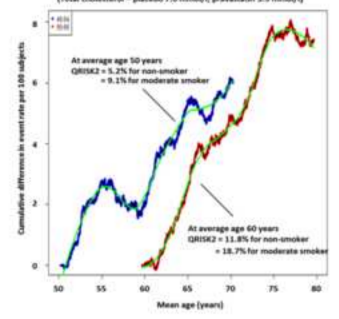
<u>±</u>

Figure 3



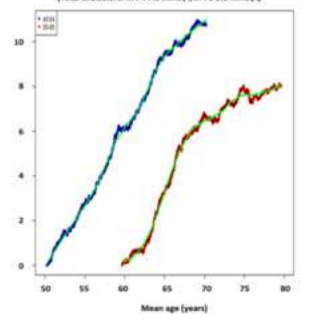


Cumulative difference in CHD events between placebo and pravastatin arms in WOSCOPS randomised cohort. (Total cholesterol – placebo 7.0 mmol/l) praventatin 5.9 mmol/l)





Cumulative difference in CHD events between groups P4 and P3 in screened population. (Total chalesteroi in P4 7.0 mmol/l; in P3 6.0 mmol/l)



Supplementary files new

Click here to access/download Supplementary files Modelling prevention Supplementary tables.docx Modelling total CHD burden and long-term benefit of cholesterol lowering in middle aged men with and without a history of cardiovascular disease.

Word counts

Abstract = 250 words

Main body of text (inc references, tables) = 4880

Permissions information

Permissions

Not applicable

Modelling total CHD burden and long-term benefit of cholesterol lowering in middle aged men with and without a history of cardiovascular disease.

Chris J Packard¹, Robin Young², Kevin Ross², Ian Ford², Baishali Ambegaonkar³, Philippe Brudi³, Colin McCowan².

Robertson Centre for Biostatistics, University of Glasgow, Glasgow, Scotland, UK G12 8QQ

¹College of Medical, Veterinary and Life Sciences; ²Robertson Centre for Biostatistics, University of Glasgow, Glasgow, Scotland; ³Merck, Sharpe & Dohme Ltd, Kenilworth, New Jersey USA

Corresponding author:-

Professor Chris J Packard McGregor Building, Room 203, Floor 2 Western Infirmary University of Glasgow G12 8QQ Telephone (44) 7788447576 Email: chris.j.packard@gmail.com

Abstract

Background

Cumulative CHD events over 20 years were examined in men screened for, and in those randomised to, the West of Scotland Coronary Prevention Study.

Methods and Results

Record linkage provided CHD-related events and days in hospital for the 80,230 screenees, including the randomised cohort of 6,595 men. Risk factors were determined at baseline, and disease burden assessed for groups defined by cholesterol. Effects of cholesterol lowering were modelled from differences between groups, and from the treatment arms of the trial. Over 20 years, those without a history of CHD (n=61,211) had 23.0 events per 100 subjects in the lowest cholesterol group (mean 4.0mmol/l) and 65.1 per 100 in the highest (8.8mmol/l). Corresponding days in hospital were 167.2 to 435.4 per 100 subjects. Analogous figures for men with a CHD history (n=8,570) were 77.3 to 141.7 events per 100 and 526.1 to 936.7 hospital days per 100.

Lowering cholesterol by about 1.0mmol/l in men with average cholesterol and no CHD was predicted to be associated with 8.9 fewer events and a saving of 56.0 hospital days per 100. In those with CHD this difference gave, depending on starting level, 26.8 to 36.5 fewer events and savings of 158.2 to 247.3 hospital days per 100 subjects.

Comparison of cumulative events in 45-54 versus 55-64 year olds in the trial revealed greater benefit from intervention in the younger decade.

Conclusion

Long-term, longitudinal data reveal the considerable CHD burden in middle-aged men and indicate substantial clinical benefits from both moderate and aggressive cholesterol lowering.