β Blockade after myocardial infarction: systematic review and meta regression analysis

Nick Freemantle, John Cleland, Philip Young, James Mason, Jane Harrison

Medicines Evaluation Group, Centre for Health Economics, University of York, York YO10 5DD Nick Freemantle, senior research fellow James Mason, senior research fellow Jane Harrison, information officer

Department of Cardiology, Castle Hill Hospital, University of Hull, Kingston upon Hull, U16 5JQ John Cleland, professor

Department of Health Sciences and Clinical Evaluation, University of York Philip Young, lecturer in applied statistics

Correspondence to: N Freemantle meg@york.ac.uk

BMJ 1999;318:1730-7

Abstract

Objectives To assess the effectiveness of β blockers in short term treatment for acute myocardial infarction and in longer term secondary prevention; to examine predictive factors that may influence outcome and therefore choice of drug; and to examine the clinical importance of the results in the light of current treatment.

Design Systematic review of randomised controlled trials.

Setting Randomised controlled trials. **Subjects** Patients with acute or past myocardial infarction.

Intervention β Blockers compared with control. Main outcome measures All cause mortality and non-fatal reinfarction.

Results Overall, 5477 of 54 234 patients (10.1%) randomised to β blockers or control died. We identified a 23% reduction in the odds of death in long term trials (95% confidence interval 15% to 31%), but only a 4% reduction in the odds of death in short term trials (-8% to 15%). Meta regression in long term trials did not identify a significant reduction in effectiveness in drugs with cardioselectivity but did identify a near significant trend towards decreased benefit in drugs with intrinsic sympathomimetic activity. Most evidence is available for propranolol, timolol, and metoprolol. In long term trials, the number needed to treat for 2 years to avoid a death is 42, which compares favourably with other treatments for patients with acute or past myocardial infarction. **Conclusions** β Blockers are effective in long term secondary prevention after myocardial infarction, but they are underused in such cases and lead to avoidable mortality and morbidity.

Introduction

β Blockade was once heralded as a major advance in the treatment of patients with myocardial infarction, but current evidence suggests that less than half of eligible patients receive it. $^{1-3}$ The effectiveness of β blockers was appraised by Yusuf et al in 1985, 4 but since then there have been nearly 3000 deaths among 23 000 patients randomised in new trials. Trials of β blockers now include a broader group of patients such as those at high risk or with accompanying heart failure, enabling the benefits identified by Yusuf et al in a restricted group of trials to be extended to such patients.

Methods used in systematic reviews have also advanced. The development of regression techniques within meta analysis enables a more robust examination of the importance of factors that may mediate upon the effectiveness of specific drugs.⁵ Two such factors, intrinsic sympathomimetic activity and cardioselectivity, were identified as potentially important,⁴ and intrinsic sympathomimetic activity in particular

seemed to be related to reduced therapeutic action. Given the changing use of drugs after myocardial infarction, the early promise of β blockade in these patients, and the continuing high rates of mortality associated with myocardial infarction, a new overview of these drugs is timely.

Methods

Objective

We reappraised the effectiveness of β blockers for secondary prevention after myocardial infarction. Our main outcome was all cause mortality and the secondary outcomes were non-fatal reinfarction and withdrawal from treatment. We examined the effectiveness of β blockers in the acute phase immediately after myocardial infarction; their role in longer term secondary prevention; the importance of early initiation after the onset of symptoms; the extent to which specific pharmacological features of different β blockers may affect their performance; the magnitude of benefits achieved by β blockers; and the clinical importance of β blockers.

Inclusion criteria

We included randomised trials without crossover, with treatment lasting more than one day, and with follow up that examined the clinical effectiveness of β blockers versus placebo or alternative treatment in patients who had had a myocardial infarction. Treatment may have begun at any stage before or after myocardial infarction and may have been commenced intravenously.

Search strategy

We conducted sensitive electronic searches of Medline (1966-97 through Ovid), Embase (1974-97 through Dialog), Biosis (1985-97 through Edina), Healthstar (1975-97 through Ovid), Sigle (1980-97 through Blaise-line), IHTA (1990-97 through ECRInet), conference papers index (1984-97 through Dialog), Derwent drug file (1992-97 through Dialog), dissertation abstracts (1992-97 through Dialog), Pascal (1992-97 through Dialog), international pharmaceutical abstracts (1992-97 through Dialog), and science citation index (1981-97 through BIDS).

We reviewed the reference list of each identified study. We also examined existing bibliographies and reviews for relevant studies.

Data abstraction and appraisal of study quality

From each study we abstracted data on the total number of patients randomised to active treatment or control, β blocker, route and dose of drug, duration of treatment, loss to follow up, level of blinding, concealment of allocation, 6 specific study inclusion and exclusion criteria, duration of follow up, deaths, reinfarc-

website extra

References for the trials appear on the BMJ's website

www.bmj.com

Table 1 Characteristics of short term trials comparing β blockers with control (see website for references)

| | Average | | | Concealment | Loss to | Outcome or | Heart | Mortality (No/total No) | | |
|---|---|-----------------------------------|---|---------------|---------------|----------------------------|---|-------------------------|-----------|--|
| Trial | follow up | Drug* | Blinding | of allocation | follow up (%) | endpoint | failure (%) | β Blockers | Controls | |
| Azancot 1982 ^{w1} | 1 month | Acebutolol* | No | Unclear | 0 | Mortality | 0 | 0/14 | 0/12 | |
| Balcon 1966 ^{w2} | 28 days | Propranolol | Double | Unclear | 0 | Mortality | 55 | 14/56 | 15/58 | |
| Barber 1976 ^{w3} | 4 weeks | Propranolol | No | Unclear | Unclear | Mortality, reinfarction | Unclear | 10/52 | 12/47 | |
| Campbell 1984 ^{w4} | In hospital | Timolol* | Unclear | Unclear | 0 | Mortality | Unclear | 1/20 | 2/19 | |
| Clausen 1966 ^{w5} | 14 days | Propranolol | Unclear | Unclear | 0 | Mortality | Unclear | 18/66 | 19/64 | |
| CPRG 1981 ^{w6} | 8 weeks | Oxprenolol | Double | Unclear | 0 | Mortality, reinfarction | 0 | 9/177 | 5/136 | |
| Curtis 1991 ^{w7} | 3.4 days | Propranolol | Double | Unclear | 0 | Mortality | Unclear | 0/18 | 0/12 | |
| Dotremont 1968 ^{w8} | 3-6 weeks | Propranolol | No | No | Unclear | Mortality | 68.6 | 4/36 | 5/36 | |
| Evemy 1978 ^{w9} | 7 months | Practolol* | No | No | Unclear | Mortality | Unclear | 9/46 | 6/48 | |
| Federman 1984 ^{w10} | 28 days | Timolol* | Unclear | Unclear | 0 | mortality | 0 | 1/50 | 0/50 | |
| Fuccella 1968 ^{w11} | 21 days | Oxprenolol | Unclear | Unclear | 14 | Mortality | Unclear | 15/106 | 9/114 | |
| Gupta 1982 ^{w12} | Unclear | Propranolol | Unclear | Unclear | 0 | Mortality | Unclear | 0/25 | 3/25 | |
| Gupta 1984 ^{w13} | 72 hours | Propranolol* | No | Unclear | Unclear | Mortality | Unclear | 0/15 | 0/15 | |
| Heber 1987 ^{w14} | 1 year | LabetaloI* | No | Unclear | Unclear | Mortality | Unclear | 12/83 | 7/83 | |
| Hutton 1979 ^{w15} | 2 days | Propranolol | Unclear | Unclear | 0 | Mortality | Unclear | 0/16 | 0/13 | |
| ICSG 1984 ^{w16} | To discharge | Timolol | Double | Unclear | 0 | Mortality | 57 being treated for heart failure | 3/73 | 4/71 | |
| ISIS-1 Collaborative Group 1986 ^{w17} | 1 year | Atenolol* | No | NA | Unclear | Mortality | Unclear | 1071/8037 | 1120/7990 | |
| Johansson 1980 ^{w18} | 6 months | Practolol* then atenolol | Single | No | Unclear | Mortality | Unclear | 7/25 | 7/29 | |
| Kahler 1968 ^{w19} | Up to 35 days | Propranolol | Double | Unclear | Unclear | Mortality, reinfarction | 11 | 3/38 | 6/31 | |
| Ledwich 1968 ^{w20} | 7 days | Propranolol | Double | Unclear | Unclear | Mortality | Unclear | 2/40 | 3/40 | |
| Lloyd 1988 ^{w21} | 72 hours | Sotalol* | No | Unclear | 0 | Mortality | Unclear | 0/15 | 0/15 | |
| Lombardo 1979 ^{w22} | 20 days | Oxprenolol | Double | Unclear | Unclear | Mortality | 0 | 8/133 | 11/127 | |
| Macleod 1980 ^{w23} | 1 week | Practolol* | Unclear | Unclear | 0 | Mortality | Unclear | 1/26 | 0/26 | |
| McMurray 1991 ^{w24} | 7 days | Xamoterol | Double | Unclear | 0 | Mortality | 31 | 0/25 | 0/26 | |
| MIAMI Trial Research Group 1985 ^{w25} | 15 days | Metoprolol* | Double | Yes | 0.04 | Mortality | 23.5 | 123/2877 | 142/2901 | |
| Mueller 1980 ^{w26} | To discharge | Propranolol* | Double | Unclear | Unclear | Mortality | Unclear | 2/35 | 1/35 | |
| Multicentre 1966 ^{w27} | 28 days | Propranolol | Double | Unclear | 1 | Mortality | 11 | 15/100 | 12/95 | |
| Nigam 1983 ^{w28} | 1 week | Propranolol* | Unclear | Unclear | 0 | Mortality | Unclear | 0/20 | 0/20 | |
| Norris 1968 ^{w80} | 3 weeks | Propranolol | Double | Yes | 0 | Mortality | Unclear | 31/226 | 24/228 | |
| Norris 1978 ^{w29} | To discharge | Propranolol* | No | No | Unclear | Mortality | Unclear | 0/20 | 0/23 | |
| Norris 1984 ^{w30} | In hospital | Propranolol* | No | NA | 0 | Mortality | Unclear | 15/364 | 14/371 | |
| Owensby 1984 ^{w31} | 3 days | Pindolol* | No | NA | Unclear | Mortality | Unclear | 1/50 | 1/50 | |
| Peter 1978 ^{w32} | To discharge | Propranolol* | No | Unclear | 0 | Mortality | 0 | 1/47 | 2/48 | |
| Pitt 1976 ^{w33} | 14 days | Propranolol | Double | Unclear | 0 | Mortality | Unclear | 0/9 | 0/8 | |
| Ranganathan 1988 ^{w34} | 28 days | Timolol* | Double intravenously then by open label orally | Unclear | 2 | Mortality | Unclear | 1/45 | 3/49 | |
| Roberts 1984 ^{w35} | 36 months | Propranolol* | Single | Unclear | 0.2 | Mortality | 4.9 | 24/134 | 20/135 | |
| Singh 1985 ^{w36} | 60 hours | Propranolol* | No | Unclear | 0 | Mortality | Unclear | 0/8 | 0/7 | |
| Sloman 1967 ^{w37} | To discharge | Propranolol* | No | Unclear | Unclear | Mortality | Unclear | 3/26 | 4/23 | |
| Snow 1980 ^{w38} | Short term | Practolol | Unclear | Unclear | 0 | Mortality | Unclear | 19/76 | 15/67 | |
| Thompson 1979 ^{w39} | 1 year | Practolol | Double | Unclear | Unclear | Mortality | Unclear | 5/72 | 6/71 | |
| TIMI IIB Study Group 1989 ^{w40} | 5 days | Metoprolol* (15 mg) | No | Unclear | 3.5 | Mortality, reinfarction | 1.1 | 17/696 | 17/694 | |
| Tonkin 1981 ^{w41} | 1 year | Timolol | Double | Unclear | Unclear | Mortality, reinfarction | Unclear | 1/42 | 1/46 | |
| UKCSG 1983 ^{w42} | To discharge | Timolol | Double | Unclear | Unclear | Mortality | Unclear | 4/56 | 5/55 | |
| Van de Werf 1993 ^{w43} | 10-14 days | Atenolol | Double | Unclear | 0 | Mortality, reinfarction | Unclear | 1/100 | 4/94 | |
| Von Essen 1982 ^{w44} | 14 days | Metoprolol* | Double | Unclear | 0 | Mortality | Unclear | 1/25 | 1/26 | |
| Waagstein 1975 ^{w45} | 1 week | Practolol,* H87/07, or metoprolol | Double | Unclear | 0 | Mortality | Unclear | 0/38 | 0/45 | |
| Wilcox 1980b ^{w46} | 6 weeks | Oxprenolol | Double | Yes | 0 | Mortality | 28 withdrawn owing to severe heart failure | 14/157 | 10/158 | |
| Yang 1987 ^{w47} | 14 days | Betaxolol | Double | Unclear | 0 | Mortality | 9.4 | 0/16 | 0/15 | |
| Yusuf 1980 ^{w48} | 10 days for infarction, 1-4 years for mortality | Atenolol* | No | Unclear | Unclear | Mortality, morbidity | 6.5 | 36/244 | 44/233 | |

CPRG=Coronary Prevention Research Group; ICSG=International Collaborative Study Group; ISIS-1=first international study of infarct survival; MIAMI=metoprolol in acute myocardial infarction; TIMI IIB=thrombolysis in myocardial infarction phase II trial; UKCSG=UK Collaborative Study Group. *Initial dose intravenously.

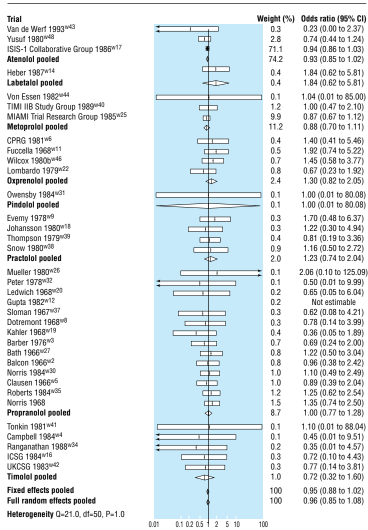


Fig 1 Odds of death and pooled odds ratios in short term trials (arrows indicate 95% confidence intervals exceeding range of plot). ISIS-1=first international study of infarct survival; TIMI IIB=thrombolysis in myocardial infarction phase II trial; MIAMI=metoprolol in acute myocardial infarction; CPRG=Coronary Prevention Research Group; ICSG=International Collaborative Study Group; UKCSG=UK Collaborative Study Group

tions, and withdrawals. Data were checked by a second researcher.

Statistical analysis

We estimated pooled odds ratios for short and long term treatment trials separately using the fixed effects approach of Mantel Haenszel. 78 As we anticipated systematic differences between the results of studies (heterogeneity), we also routinely estimated random effects pooled odds ratios. Standard random effects methods for meta-analysis (pooling the results of studies)9 10 may provide unduly precise estimates of effect, as they assume that the observed distribution of effects is the true treatment distribution—an assumption that may not be valid in sparse data.⁵ 11 12 Therefore, we used the full random effects approach on the basis of the numerical integration techniques using Markov chain Monte Carlo simulation, with appropriate uninformative priors and the "Bugs" software described by Smith et al.⁵ This provides a more robust estimate of the precision of random effects estimates and can account for trial groups that experience no events without resorting to crude fixes such as adding a value to each cell to estimate an individual odds ratio. A further advantage of this approach is that the effects of predictive factors may be examined. Our main treatment related covariates were cardioselectivity and intrinsic sympathomimetic activity, which were examined in the long term trials using a nested random effects logistic regression model (see appendix).

We also made a separate examination of the effects of initial intravenous treatment in long term trials, and the effect of additional treatment options through the proxy variable of publication date before or after the median year (1982). We assessed convergence using the methods described by Geweke¹³ and visual inspection of convergence plots.

We calculated risk differences using standard random effects methods, $^{\rm II}$ and because comparison of risk differences between trials may be affected by different lengths of follow up, we also estimated a pooled incidence risk difference using the approach described by Ioannidis et al. $^{\rm I4}$ This is less robust than the pooled odds ratio but provides a practically interpretable estimate of absolute treatment effect derived directly from the trials. $^{\rm I5}$ For the long term trials, we also calculated pooled estimates of effect for each β blocker using the fixed effects model. $^{\rm 7~8}$

Results

We identified 82 randomised trials that examined the effects of β blockers compared with control and that had data on all cause mortality. Overall, 5477 of 54 234 patients (10.1%) randomised died. Fifty one trials examined acute treatment with β blockers—up to 6 weeks after onset of pain (table 1, and 31 trials examined long term treatment with β blockers—6 to 48 months (table 2).

Short term trials

Overall, 3062 of 29 260 patients (10.5%) randomised in short term trials died. Although 51 trials were identified that examined the effects of short term treatment, only 45 of these had observed deaths in either the intervention or control groups. The major challenge to the quality of this group of trials was that small numbers of patients randomised to treatment or control led to many trials with either no, or only a small number of, deaths.

The pooled random effects odds ratio for the short term trials was 0.96 (95% confidence interval 0.85 to 1.08); that is, a small and non-significant reduction in the odds of death (fig 1). Even if this result is correct it would represent a reduction of only 0.4 deaths in 100 patients, which does not achieve conventional levels of significance (-0.2 to 1) as 250 patients would require treatment to avoid one death (100 to ∞). Analysis of predicted benefit by drug identified no individual drug that differed significantly in effect from the pooled result.

Although most trials were undertaken before the second international study of infarct survival in 1988¹⁶ firmly established the importance of thrombolysis, a large trial of thrombolysis in myocardial infarction in 1989¹⁷ randomised patients who had received recombinant tissue plasminogen activator within 4 hours of

Table 2 Characteristics of long term trials comparing β blockers with control (see website for references)

| | Average | | | Concealment | Loss to follow up | Outcome or | Heart | Mortality (N | lo/total No) | No of reinfarctions | | No of withdrawals | |
|--|--------------------|-----------------------------|---|---------------|----------------------|--|--|---------------------------|-----------------|---------------------|----------|-------------------|---------|
| Trial | follow up | Drug* | Blinding | of allocation | (%) | endpoint | failure (%) | $\beta \ \text{blockers}$ | Controls | β blockers | Controls | β blockers | Control |
| Ahlmark 1974 ^{w49} | 2 years | Alprenolol | Unclear | Unclear | Unclear | Mortality, reinfarction | Unclear | 5/69 | 11/93 | 4 | 15 | 4 | 6 |
| Andersen 1979 ^{w50} | About 1 year | Alprenolol | Double | Unclear | 0 | Mortality | Unclear | 61/238 | 64/242 | _ | _ | 59 | 49 |
| Boissel 1990 ^{w51} | 318 days | Acebutolol | Double | Yes | 0 | Mortality | 49.5 | 17/298 | 34/309 | _ | _ | 102 | 109 |
| Aronow 1997 ^{w52} | 1 year | Propranolol | Unclear | Unclear | Unclear | Mortality, reinfarction | 100 | 44/79 | 60/79 | 3 | 5 | _ | _ |
| Australian and Swedish study 1983 ^{w53} | 2 years | Pindolol | Double | Unclear | Unclear | Mortality, reinfarction | 61 left ventricular dysfunction | 45/263 | 47/266 | 12 | 13 | 76 | 50 |
| Baber 1980 ^{w54} | 9 months | Propranolol | Double | Unclear | Unclear | Mortality, reinfarction | Unclear | 28/355 | 27/365 | 17 | 27 | 82 | 88 |
| Barber 1967 ^{w55} | 2 years | Practolol | Unclear | Unclear | Unclear | Mortality, reinfarction | 26 | 33/207 | 38/213 | _ | _ | _ | |
| Basu 1997 ^{w56} | 6 months | Carvedilol | Double | Unclear | 0 | Mortality, reinfarction | 45 | 2/75 | 3/71 | 4 | 8 | _ | |
| BHAT 1982 ^{w57} | 25 months | Propranolol | Double | Yes | 0.3 | Mortality | 9.2 | 138/1916 | 188/1921 | 103 | 121 | 243 | 179 |
| Darasz 1995 ^{w58} | 6 months | Xamoterol | Double | Unclear | 19 | Mortality, | | 3/23 | 1/24 | _ | _ | 3 | 6 |
| EIS 1984 ^{w59} | 1 year | Oxprenolol | Double | Unclear | Unclear | mortality, | 7.7 | 57/853 | 45/883 | 36 | 38 | 275 | 275 |
| Hansteen | 1 year | Propranolol | Double | Unclear | 0 | Reinfarction Mortality, | 5.9 (taking | 25/278 | 37/282 | 16 | 21 | 70 | 72 |
| Hjalmarson 1981 ^{w61} | 2 years | Metoprolol* | Double (3 months) then open treatment (to 2 years) | Unclear | 1.6 | reinfarction Mortality at 2 years; reinfarction at 3 months | digitalis) | 40/698 | 62/697 | 35 | 54 | 131 | 131 |
| Julian 1982 ^{w62} | 12 | Sotalol | Double | Yes | 0 | Mortality, | 0 | 64/873 | 52/583 | 37 | 38 | 218 | 121 |
| Kaul 1988 ^{w63} | months 6 months | Propranolol | Double | Unclear | 0 | reinfarction Mortality, reinfarction | Unclear | 3/25 | 3/25 | 0 | 4 | 0 | 0 |
| LIT Research Group 1987 ^{w64} | 18 months | (iv) Metoprolol | Double | Unclear | 0.2 | Mortality | 2.1 | 86/1195 | 93/1200 | _ | _ | 381 | 355 |
| Manger Cats 1983 ^{w65} | 1 year | Metoprolol | Double | Unclear | 0 | Mortality | Unclear | 9/273 | 16/280 | _ | _ | _ | |
| Mazur 1984 ^{w66} | 1.5 years | Propranolol | No | Unclear | Unclear | Mortality, reinfarction | Unclear | 5/101 | 11/103 | 5 | 7 | _ | |
| Multicentre international 1975 ^{w67} | Up to 24 months | Practolol | Double | Unclear | 3.4 | Mortality, reinfarction | 0 | 102/1533 | 127/1520 | 69 | 89 | 389 | 382 |
| Norwegian Multicentre Study Group 1981 ^{w68} | 17 months | Timolol | Double | Unclear | Unclear | Mortality | 33 | 98/945 | 152/939 | 88 | 141 | 275 | 219 |
| Rehnqvist 1980 ^{w69} | 1 year | Metroprolol | Unclear | Unclear | 0 | Mortality | Unclear | 4/59 | 6/52 | _ | _ | 12 | 5 |
| Rehnqvist 1983 ^{w70} | 36 months | Metoprolol | Double | Unclear | 0 | Mortality, reinfarction | 24 (taking digitalis) | 25/154 | 31/147 | 18 | 31 | 38 | 35 |
| Reynolds 1972 ^{w71} | 1 year | Alprenolol | Double | Yes | Unclear | Mortality, reinfarction | Unclear | 3/38 | 3/39 | 3 | 2 | 4 | 3 |
| Roqué 1987 ^{w72} | 24 months | Timolol* | Double | Unclear | Unclear | Mortality | Unclear | 7/102 | 12/98 | _ | _ | _ | |
| Salathia 1985 ^{w73} | 1 year | Metoprolol* | Double | Unclear | 0.5 | Mortality, | 10 | 49/416 | 52/348 | _ | _ | 95 | 66 |
| Schwartz 1992 (high risk and low risk) ^{w74} | 22 months | Oxprenolol | High risk† and low risk‡ groups | Unclear | 0 | Mortality, reinfarction | 2 in high risk group; unclear for low risk group | 2/48 15/437 | 12/56 27/432 | 0 | 2 | 11 | 9 |
| SSSD 1993 ^{w75} | 3 years | Metoprolol | No | Unclear | 1.9 | Mortality, reinfarction | 100 | 17/130 | 9/123 | 5 | 6 | _ | |
| Taylor 1982 ^{w76} | 48 months | Oxprenolol | Double | Done | Unclear | Mortality, reinfarction | 0 | 60/632 | 48/471 | 67 | 58 | 185 | 141 |
| Wilcox 1980a ^{w77} | 1 year | Propranolol* or atenolol | Double | Done | 0 | Death | Unclear | 19/127 17/132 | 19/129 | _ | _ | 44 51 | 40 40 |
| Wilhelmsson 1974 ^{w78} | 2 years | Alprenolol | Double | Unclear | 7 | Mortality | Unclear | 7/114 | 14/116 | 16 | 18 | 8 | 8 |
| Yusuf 1979 ^{w79} | 12 months | Atenolol | Double | Unclear | 23 | Death; electrocardiogr aphic signs | Unclear | 1/11 | 1/11 | _ | _ | 2 | 1 |

BHAT=β-blocker heart attack trial; LIT=lopressor intervention. *Initial dose intravenously. †Single blind. ‡Double blind.

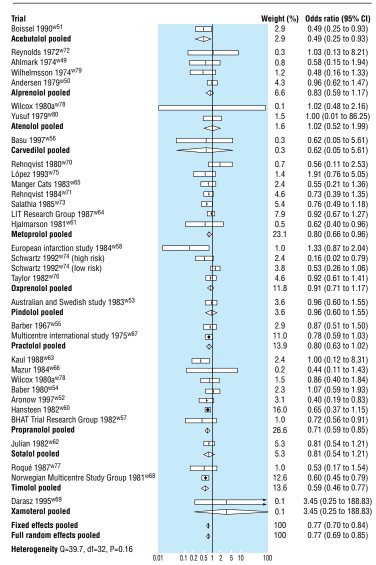


Fig 2 Odds of death and pooled odds ratios in long term trials. LIT=lopressor intervention; BHAT= β -blocker heart attack trial

the onset of pain to early metoprolol or control. During 5 days of follow up, there was no difference in mortality between the two groups. Two subsequent myocardial infarctions were, however, avoided for every 100 patients treated (0.2 to 4).

Long term trials

Overall effects

Overall, 2415 of 24 974 patients (9.7%) randomised in the 31 long term trials died. In general, the quality of studies was reasonably high, with adequate follow up achieved in many trials (table 2), though the proxy quality variable, concealment of allocation, was seldom adequately reported.

Overall, the pooled odds ratio from the full random effects model was 0.77 (0.69 to 0.85). Results from the standard fixed effects model were similar (fig 2).

Because of potential confounding due to the differences in length of study follow up, we used the random effects approach for incidence of risk difference to estimate the normalised annual reduction in mortality across the trials. This approach suggests an annual reduction of 1.2 deaths in 100 patients treated with β blockers after myocardial infarction (0.6 to 1.7); that is, about 84 patients will require treatment for 1 year to avoid one death. A similar approach was used to estimate the effects of treatment on reinfarction, although only 21 of the 34 comparisons provided data on reinfarction, resulting in wider confidence intervals and the potential for reporting bias. This analysis suggests an annual reduction in reinfarction of 0.9 events in every 100 (0.3 to 1.6); that is, about 107 patients would require treatment for 1 year to avoid one non-fatal reinfarction.

Predictors of benefit

Initial intravenous dose—We investigated the extent to which initiation of treatment with an intravenous dose of β blockers predicted mortality in the long term trials. Applying initial intravenous dose as a covariate term in the analysis suggested no additional benefit among patients treated in this manner (odds ratio 0.87, 0.61 to 1.22). Equally, this analysis indicates that there is no reason to delay treatment with a β blocker and that early initiation will lead to a greater period when benefits may be accrued from treatment.

Presence of cardioselectivity or intrinsic sympathomimetic activity—We anticipated that the presence of cardioselectivity and intrinsic sympathomimetic activity would be important predictors of benefit in the trials, a hypothesis examined by Yusuf et al.⁴

Classification of β blockers into those with or without important cardioselective activity or intrinsic sympathomimetic effect is not clear cut, and there is some debate in the literature on the attributes of acebutolol in particular. $^{\text{I8-20}}$ Table 3 describes the attributes of β blockers used in the trials.

The odds ratio for the predictive effect of cardioselectivity on mortality was 1.10 (0.89 to 1.39), showing a non-significant trend towards reduced benefits. The odds ratio for the predictive effect of the presence of intrinsic sympathomimetic activity was 1.19 (0.96 to 1.47), which approaches statistical significance. The results were not sensitive to the classification of acebutolol.

Reduction of benefits over time—We investigated whether benefits were reduced in the trials with additional therapeutic options for treatment intro-

 $\textbf{Table 3} \ \ \text{Classification of attributes of different } \beta \ \ \text{blocker drugs}$

| β Blocker | Cardioselectivity | Intrinsic sympathomimetic activity |
|-------------|-------------------|------------------------------------|
| Acebutolol | - | - |
| Alprenolol | - | + |
| Atenolol | + | - |
| Betaxolol | + | - |
| Carvedilol | - | - |
| Labetalol | - | - |
| Metoprolol | + | - |
| Oxprenolol | - | + |
| Pindolol | - | + |
| Practolol | + | + |
| Propranolol | - | - |
| Sotalol | - | - |
| Timolol | _ | - |
| Xamoterol | + | + |

+=Significant activity; -=no significant activity.

duced, in particular the increasing use of thrombolytic treatment, and aspirin. There is no evidence that treatment in trials after 1982 (the median trial) led to differences in benefit (odds ratio 1.04, 0.82 to 1.28).

Choice of drug—Individually, only four drugs achieved a statistically significant reduction in the odds of death: propranolol (0.71, 0.59 to 0.85); timolol (0.59, 0.46 to 0.77); metoprolol (0.80, 0.66 to 0.96); and acebutolol (0.49, 0.25 to 0.93). The effectiveness of acebutolol is supported by a single moderately sized study, which is open to considerable measurement error. However, trials including propranolol, timolol, and metoprolol include 63% of the available evidence on the effects of long term β blockade in patients who have had a myocardial infarction.

Withdrawal from treatment

Different definitions and reporting made comparison of withdrawal of treatment withdrawal between trials problematic. Similar withdrawal rates between active treatment and placebo groups concealed two opposing effects: more patients are withdrawn from treatment groups because of suspected adverse cardiovascular reactions (most commonly brachycardia and hypotension), whereas in the placebo group withdrawal is more common because of the need for β blockade for hypertension and angina. Trials reports of dizziness, depression, cold extremities, and fatigue were only marginally more common in the treatment than control groups.

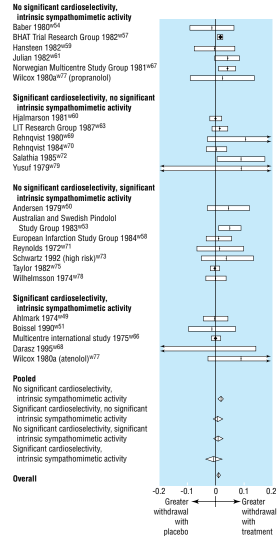
Withdrawal in trials from both treatment and control groups varied from 10% to 30%. No adequate studies have been retrieved to compare directly the comparative tolerability of β blockers with different cardioselectivity or intrinsic sympathomimetic activity.

Overall, 5151 of 21 954 patients (23.5%) withdrew from treatment (table 2). Overall, withdrawal was slightly more common in patients taking β blockers—the difference in the annualised rate of withdrawal compared with placebo being 1.16 in 100 patients treated (0.56 to 1.76, random effects; fig 3). No clinically important differences in withdrawal were observed between β blockers of differing cardioselectivity and intrinsic sympathomimeticity.

Discussion

Considerable evidence supports the routine long term use of β blockers in patients who have had a myocardial infarction, with substantial benefits in terms of reduced mortality and morbidity. Short term β blockade immediately after acute myocardial infarction seems unlikely to be of major benefit unless treatment is continued long term. This finding contradicts recent suggestions that β blockers should be more commonly used intravenously in acute myocardial infarction. 21 In fact, evidence strongly indicates that unless β blockers are continued long term, the benefits suggested by $Owen^{21}$ will not be observed.

The benefits of β blockers on all cause mortality are impressive when compared with other frequently used long term treatments for the same patient group. Table 4 shows the effects of different drugs on the number of patients that would need to be treated for 2 years to avoid one death—for example, after a myocardial infarction 42 patients would need to be treated with β



Pooled incident risk difference=0.0116 (95% CI 0.0056 to 0.0176) Q (combinability for risk difference)=22.7, df=21, P=0.478

Fig 3 Incidence (yearly) of withdrawal from trials

Table 4 Comparison of effect on mortality of different drugs

Drug Number needed to treat*

| Number needed to treat^ |
|--|
| 42 |
| No long term trials in unselected patients |
| 153 |
| 94 |
| ∞ |
| 24 |
| 63 |
| |

^{*}Number needed to avoid death in 2 years of treatment in unselected patients after myocardial infarction.

blockers whereas 292 patients would need to be treated with antiplatelets. The number and length of long term trials showing a consistent benefit for β blockers in unselected patients after myocardial infarction suggest lasting benefits in this comparatively high risk group, and suggest that β blockers should be continued indefinitely.

Have benefits from β blockade declined with availability of new treatments?

Our finding that β blockers benefit a broader group of patients after myocardial infarction supports the findings of the β blocker pooling project. Our finding also agrees with those of the cooperative cardiovascular project, which examined the medical records of 201 752 patients who had had a myocardial infarction. Hat study, mortality was lower in every subgroup of patients treated with β blockade than in untreated patients. The findings of the cooperative cardiovascular project agree with our meta regression analysis, which found no evidence of a reduction in benefits from β blockade in more recent randomised trials. Indeed, rather than being overtaken by newer treatments, β blockers have a comparatively large effect in reducing mortality (table 4).

Which ß blocker?

Cardioselectivity was associated with a non-significant trend towards reduced benefit. The presence of an intrinsic sympathomimetic effect predicted a near significant reduction in benefits and thus drugs with this characteristic should be avoided. We found evidence to support the long term use of propranolol and timolol, the only two drugs indicated for prophylaxis after myocardial infarction in the British National Formulary. The use of either drug led to a substantial reduction in the odds of death, with narrow confidence intervals (fig 2). In contrast, atenolol, which is commonly prescribed in secondary prevention, has been inadequately evaluated in this setting. Although similar efficacy may be achieved-we found no evidence that all β blockers are not equal—it cannot be presumed that the benefits from propranolol, timolol, and metoprolol will be achieved with other drugs.

Have benefits from intravenous β blockers declined over time?

It may be hypothesised that intervention with thrombolytic drugs and antiplatelets reduces the potential for patients to benefit from intravenous β blockade. The first international study of infarct survival25 was completed before the results of the second international study16 became available, and before the use of thrombolytic and antiplatelet treatment was established. In contrast, the comparison of early versus delayed β blockade in a large trial of thrombolysis in myocardial infarction was undertaken in patients who all received thrombolytic and antiplatelet treatment.¹⁷ Although the much larger first international study of infarct survival trial25 achieved a slightly larger reduction in the odds of death with β blockers, measurement error could not be excluded as an explanation for this difference, as indicated by the test for heterogeneity between the trials (Q = 0.025, df = 1, P = 0.87). The thrombolysis in myocardial infarction trial did suggest that early use of intravenous β blockers could reduce the early risk of serious arrhythmias.

Are β blockers underused?

Concern has been voiced that β blockers are used in less than half of eligible patients after myocardial infarction, ¹⁻³ despite substantial benefits and generally low treatment costs. Concern that side effects affect the

Key messages

- The first randomised trials of β blockade in secondary prevention after myocardial infarction were published in the 1960s
- β blockers were once heralded as a major advance, but their use for secondary prevention has declined in recent years
- Firm evidence shows that long term β blockade remains an effective and well tolerated treatment that reduces mortality and morbidity in unselected patients after myocardial infarction
- The benefits from β blockade compare favourably with other drug treatments for this patient group
- Most evidence is for propranolol, timolol, and metoprolol, whereas atenolol, which is commonly used, is inadequately evaluated for long term use

usefulness of β blockers must be tempered by the low yearly withdrawal from β blockers in the long term trials we reviewed. The clinical implications of our results are clear. New is not necessarily better, especially if the aim is to reduce mortality in patients after myocardial infarction. Furthermore, the underuse of β blockers in this group leads to a rate of avoidable death that should not be considered acceptable among those keen to practice evidence based medicine.

Renewed interest in β blockers, particularly in patients with heart failure, ^{26–28} may lead to substantial benefits for a broad range of patients.

We thank Andrew Herxheimer, who assisted in the categorisation of included compounds, and Anne Burton for her diligent help in locating studies and in the preparation of the manuscript.

Contributors: NF developed the protocol for the review, abstracted data, and undertook the majority of statistical analyses. JC conceptualised the review, developed the protocol, and provided clinical interpretation of the included trials and the results. PY developed the meta regression approach and provided methodological support in the review. JM contributed to the development of the protocol, data abstraction, and some of the statistical analyses. JH designed and implemented the electronic search strategies. NF and JC will act as guarantors for the paper.

Funding: SmithKline Beecham Pharmaceuticals UK. The views expressed are those of the authors and not necessarily those of the sponsor.

Competing interests: This study was funded through an unrestricted educational grant by SmithKline Beecham, who supply carvedilol in the United States. JGFC has spoken at many meetings and educational programmes on drugs in heart failure, organised by pharmaceutical and device companies, and received fees and expenses. He has also received research funding from industry as well as the NHS, British Heart Foundation, and US Veterans Administration.

Appendix

Statistical model for random effects regression analysis

$$\log\left(\frac{pt}{1-pt}\right) = \alpha + \delta + \beta I + \gamma S$$

$$\log\left(\frac{pc}{1-pc}\right) = 0$$

Where pt is the probability of an event in the intervention group, pc is the probability of an event in the control group,

I is the presence or absence of significant intrinsic sympathomimetic activity, and S is the presence or absence of significant cardioselectivity. Similarly, α is a constant, δ describes the overall treatment effect, β describes the effect of intrinsic sympathomimetic activity, and γ describes the effect of cardioselectivity.

- Smith J, Channer KS. Increasing prescription of drugs for secondary pre-
- vention after myocardial infarction. *BMJ* 1995;311.917-8. Eccles M, Bradshaw C. Use of secondary prophylaxis against myocardial infarction in the north of England. *BMJ* 1991;302:91-2.
- Viskin S, Barron HV. $\beta\text{-}Blockers$ prevent cardiac death following myocardial infarction: so why are so many infarct survivors discharge without β-blockers? Am J Cardiol 1996;78:821-2.
- Yusuf S, Peto R, Lewis J, Collins R, Sleight P. β-Blockade during and after myocardial infarction: an overview of the randomized trials. Prog Cardiovasc Dis 1985;27:335-71.
- Smith TC, Spiegelhalter DJ, Thomas A. Bayesian approaches to random-effects meta analysis: a comparative study. *Stats Med* 1995;14:2685-99. Schulz KF, Chalmers I, Hayes RJ, Altman DG. Empirical evidence of bias:
- dimensions of methodological quality associated with estimates of treatment effects in controlled trial. *JAMA* 1995;273:408-12. Rothman KJ. *Modern epidemiology*. Boston: MA Little and Brown, 1986.
- Robins J, Breslow N, Greenland S. Estimators of the Mantel-Haenszel variance consistent in both sparse data and large strata models. *Biometrics*
- Fleiss J, Gross AJ. Meta-analysis in epidemiology, with special reference to studies of the association between exposure to environmental tobacco smoke and lung cancer: a critique. *J Clin Epidemiol* 1991;44:127-39.
- DerSimonian R, Laird N. Meta-analysis in clinical trials. Controlled Clin Trials 1986;7:177-88.
- 11 Carlin JB. Meta analysis for 2 β 2 tables: a Bayesian approach. Stats Med 1992;11:141-58.
- 12 Hardy RJ, Thompson SG. A likelihood approach to meta analysis with random effects. Stats Med 1996;15:619-29.
- 13 Spiegelhalter D, Thomas A, Best N, Gilks W. BUGS: Bayesian inference using Gibbs sampling, Version 0.50. Cambridge: Medical Research Council Biostatistics Unit, 1995.
- 14 Ioannidis JPA, Cappelleri JC, Lau J, Skolnik PR, Melville B, Chalmers TC, et al. Early or deferred zidovudine therapy in HIV-infected patients without an AIDS defining illness. Ann Intern Med 1995;122:856-66.
- 15 Freemantle N, Mason JM, Eccles M. Deriving treatment recommenda-tions from evidence within randomised trials: the role and limitation of meta analysis. Intern J Technol Assess Health Care (in press).
- 16 Second International Study of Infarct Survival Collaborative Group. Randomised trial of intravenous streptokinase, oral aspirin, both or neiamong 17 187 cases of suspected acute myocardial infarction: ISIS-2. Lancet 1988;ii:349-60.

- 17 The Thrombolysis in Myocardial Infarction Study Group. Comparison of invasive and conservative strategies after treatment with intravenous tissue plasminogen activator in acute myocardial infarction: results of the thrombolysis in myocardial infarction (TIMI) phase II trial. N Engl J Med 1989;320:618-27.
- 18 Wilhelmsson C, Vedin JA, Wilhelmsen L, Tibblin G. Reduction of sudden deaths after myocardial infarction by treatment with alprenolol: prelimipary results. Lancet 1974:ii:1157-60.
- 19 Feely J, de Vane PJ, Maclean D. β-Blockers and sympathomimetics. BMJ 1983;286:1043-7.
- McDevitt DG. The assessment of β -adrenoceptor-blocking drugs in man. Br J Clin Pharmacol 1977;4:413-25.
- Owen A. Intravenous β-blockade in acute myocardial infarction: should
- be used in combination with thrombolysis. *BMJ* 1998;317:226-7.
 22 Antiplatelet Trialists' Collaboration. Collaborative overview of randomised trials of antiplatelet therapy. I. Prevention of death, myocardial infarction and stroke by prolonged antiplatelet therapy in various categories of patients. BMJ 1994;308:81-106.
- 23 β-Blocker Pooling Project Research Group. The β-blocker pooling project (BBPP): subgroup findings from randomized trials in post infarction patients. Eur Heart J 1988;9:8-16.
- 24 Gottlieb SS, McCarter RJ, Vogel RA. Effect of β-blockade on mortality among high risk and low risk patients after myocardial infarction. N Engl J Med 1998;339:489-97.
- 25 First International Study of Infarct Survival Collaborative Group. Randomised trial of intravenous atenolol among 16 027 cases of suspected acute myocardial infarction: ISIS-1. *Lancet* 1986;ii:57-67.
- 26 Doughty RN, Rodgers A, Sharpe N, MacMahon S. Effects of β-blocker therapy on mortality in patients with heart failure: a systematic overview of randomized controlled trials. Eur Heart J 1997;18:560-5.
- 27 Heidenreich PA, Lee TT, Massie BM. Effect of β-blockade on mortality in patients with heart failure: a meta analysis of randomized clinical trials. J Am Coll Cardiol 1997;30:27-34.
- 28 Cleland JGF, Freemantle N, McGowan J, Clark A. The evidence for β blockers in heart failure. BMJ 1999;318:824-5.
- 29 Latest trials on statins show large benefits? Lancet 1997;350:1525.
- 30 The Multicenter Ditiazem Postinfarction Trial (MDPIT) Research Group. The effect of diltiazem on mortality and reinfarction after myocardial inf- $\ \, {\rm arction.} \, NEnglJ\, Med \,\, 1988; \! \! 319; \! \! 385\text{-}92.$
- Smith P, Arnesen H, Holme I. The effect of warfarin on mortality and reinfarction after myocardial infarction. N Engl J Med 1990;323:147-52.
- Anticoagulants in the Secondary Prevention of Events in Coronary Thrombosis (ASPECT) Research Group. Effect of long term oral anticoagulant treatment on mortality and cardiovascular morbidity after myocardial infarction. Lancet 1994;343:499-503.

(Accepted 6 April 1999)

Risk factors for human hantavirus infection: Franco-Belgian collaborative case-control study during 1995-6 epidemic

N S Crowcroft, A Infuso, D Ilef, B Le Guenno, J-C Desenclos, F Van Loock, J Clement

Puumala hantavirus is the most common human hantavirus infection in Europe.12 It is transmitted to humans by inhalation or contamination of skin breaches by urine or faeces of infected bank voles. Infection ranges from subclinical to a severe influenzalike illness progressing to acute renal failure.³ We carried out a case-control study in an endemic area in France and Belgium to estimate knowledge of hantavirus and identify possible risk factors for infection.

Subjects, methods, and results

National reference laboratories in each country identified cases for the study. A case was defined as someone with laboratory confirmed IgM positive Puumala hantavirus infection between 1 April 1996 and 31 July 1996 in the French departments Nord, Ardennes, and Aisne and Belgian provinces of Hainaut, Namur, and Luxembourg. Controls were matched by sex, community (village), and age group. They were randomly selected from the telephone book. Interviews were carried out by telephone using a standardised questionnaire covering knowledge of hantavirus, distance of the home to a forest, refuse disposal, rodent infestation and control, gardening activities, use of wood for heating or cooking, activities in forests, and entry into rodent infested buildings.

In all, 69/88 (78%) eligible cases were included in the study and 125 controls were recruited. Most cases were in men (51) and those aged 15-65 years (64). Two cases and one control were forestry workers-no others were in occupations thought to be at risk. Forty seven per cent (91/194) of those interviewed had heard of hantavirus infection before becoming ill or being interviewed. Friends were the commonest source of information (44/91, 48%); 63/75 (84%) had heard of the disease in the past 3 years.

The table shows the results of logistic regression. Cases and controls often went walking in forests (odds ratio 0.5, 95% confidence interval 0.1 to 2.7; P = 0.64). Cases were more likely to have entered a building where there might be rodents (1.9, 1.0 to 3.6; P = 0.05)and were more likely to have cleaned (4.2, 1.1 to 15.7;

Epidemiology Scientific Institute of Public Health-Louis Pasteur. Brussels, Belgium N S Crowcroft, fellow, European programme for intervention epidemiology training J-C Desenclos, head of infectious diseases unit

Réseau National de Santé Publique, Saint-Maurice, France A Infuso, fellow, European programme for intervention epidemiology training F Van Loock. epidemiologist

continued over BMJ 1999;318:1737-8