Angiotensin receptor blockers and risk of myocardial infarction: systematic review

Michael A McDonald, Scot H Simpson, Justin A Ezekowitz, Gabor Gyenes and Ross T Tsuyuki

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Angiotensin receptor blockers and risk of myocardial infarction: systematic review

Michael A McDonald, Scot H Simpson, Justin A Ezekowitz, Gabor Gyenes, Ross T Tsuyuki

Abstract

Objective To evaluate the effect of angiotensin receptor blockers on the risk of myocardial infarction in patients at risk for cardiovascular events.

Design Systematic review of controlled trials of angiotensin receptor blockers.

Data sources Medline, Embase, Cochrane central register of controlled trials, hand search, and contact with investigators.

Selection of studies Predefined criteria were used to select controlled clinical trials comparing use of angiotensin receptor blockers with angiotensin converting enzyme (ACE) inhibitors or placebo in patients at risk for cardiovascular events. Data were extracted for patients' characteristics, interventions, quality of trials, and rates of myocardial infarction.

Results 19 studies with 31 569 patients were included in the analysis. Two studies investigated the use of angiotensin receptor blockers in hypertensive patients, four studies in patients with diabetes and nephropathy, 10 studies in patients with heart failure, and three in patients with recent myocardial infarction or ischaemic syndrome. 11 studies of 21 062 patients allowed for comparison between angiotensin receptor blockers and placebo; nine studies of 10 625 patients allowed for comparison between angiotensin receptor blockers and ACE inhibitors. Use of angiotensin receptor blockers was not associated with increased risk of myocardial infarction compared with placebo (odds ratio 0.94, 95% confidence interval 0.75 to 1.16) nor with increased risk of myocardial infarction compared with ACE inhibitors (1.01, 0.87 to 1.16). Conclusion Treatment with angiotensin receptor blockers was not associated with a significantly increased risk of myocardial infarction. The 95% confidence intervals do, however, not exclude an increase of up to 16% in the risk of myocardial infarction or ischaemic syndrome. Until further analysis, loss to follow-up) to appraise study quality, in addition to predefined criteria (allocation concealment, blinding, intention to treat) resolved by the vote of a third reviewer (JAE). We used standard systematic review techniques, as outlined by the Cochrane Collaboration.6 To evaluate the potential association between use of angiotensin receptor blockers and risk of myocardial infarction, we reviewed the medical literature to identify controlled trials comparing use of angiotensin receptor blockers with placebo therapy and with ACE inhibitors. We searched Medline, Embase, and the Cochrane central register of controlled trials, each from inception to December 2004. The search combined terms related to myocardial infarction ("myocardial infarction", "heart infarction", "death", "mortality") with terms related to angiotensin receptor blockers ("candesartan", "irbesartan", "losartan", " valsartan", " olmesartan", " telmisartan", " eprosartan"), using Boolean operators and database specific syntax. We also hand searched references from review articles and meta-analyses of angiotensin receptor blockers.

Introduction

Evidence is very strong for the use of angiotensin converting enzyme (ACE) inhibitors to reduce morbidity and mortality in patients with left ventricular dysfunction, in patients with recent myocardial infarction, and in patients who are otherwise at high risk for cardiovascular events.5,6 Angiotensin receptor blockers theoretically produce more complete inhibition of angiotensin II and are better tolerated than ACE inhibitors.7-9 Recent trials have, however, not shown their superiority and have been equivocal on their comparative effect.10-14 Verma and Strauss concluded that the use of angiotensin receptor blockers may even confer a risk of harm, specifically through their association with higher rates of myocardial infarction.15 This has caused much concern over using these agents, with many healthcare professionals and patients asking whether angiotensin receptor blockers should be avoided.

Recent, high profile withdrawals of commonly prescribed medications such as rofecoxib have heightened public awareness and concern over adverse drug reactions. In the case of angiotensin receptor blockers, we thought that it would be important to review systematically all available evidence before drawing conclusions on harm. We conducted a systematic review of all published controlled trials to determine the association of angiotensin receptor blockers and myocardial infarction.

Methods

We used standard systematic review techniques, as outlined by the Cochrane Collaboration.6 To evaluate the potential association between use of angiotensin receptor blockers and risk of myocardial infarction, we reviewed the medical literature to identify controlled trials comparing use of angiotensin receptor blockers with placebo therapy and with ACE inhibitors. We searched Medline, Embase, and the Cochrane central register of controlled trials, each from inception to December 2004. The search combined terms related to myocardial infarction ("myocardial infarction", "heart infarction", "death", "mortality") with terms related to angiotensin receptor blockers ("candesartan", "irbesartan", "losartan", " valsartan", " olmesartan", " telmisartan", " eprosartan"), using Boolean operators and database specific syntax. We also hand searched references from review articles and meta-analyses of angiotensin receptor blockers.

According to prespecified criteria, we included all original studies if they were controlled clinical trials, incorporated mono-therapy with angiotensin receptor blockers in at least one of the treatment arms, reported myocardial infarction as either a prespecified outcome or as an adverse event, and were published in English. We excluded all secondary analyses if myocardial infarction events were provided in the parent study. Two reviewers (JAE, GG) independently screened abstracts for eligibility, rejecting those that were not controlled trials, and separate reviewers (MAM, SHS) independently assessed the full text of the remaining articles for final inclusion or exclusion. Disagreements were resolved by the vote of a third reviewer (JAE). We used standard criteria (allocation concealment, blinding, intention to treat analysis, loss to follow-up) to appraise study quality, in addition to quantitative quality assessment by using the scoring system developed by Jadad.17
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Data collection and outcome measures
Two reviewers (MAM, SHS) used standardised data collection forms to extract data from studies for trial design, country of origin, patients’ characteristics, interventions, and outcomes. We verified accuracy of data by comparing collection forms from each reviewer. We documented all reported myocardial infarctions, fatal or non-fatal, according to the definition used by the authors of individual studies and confirmed that cases represented actual numbers of patients with myocardial infarction rather than total numbers of myocardial events. Where data on myocardial infarction were included as a composite end point, we recorded the components of adverse cardiac outcomes that were provided and subsequently contacted study authors for rates of myocardial events. We also contacted authors for complete details on myocardial events for those studies in which only fatal or non-fatal myocardial infarctions were reported, and we ultimately used the best obtainable data in our analysis.

Statistical analysis
We entered data into 2×2 tables and used Cochrane Review Manager software (RevMan 4.2.7, Cochrane, Copenhagen, Denmark) to analyse our results. For studies that found no events in one of the treatment groups, the program uses a correction factor of 0.5 for all cells, to avoid division by 0. For each study, we calculated odds ratios and combined them for the pooled odds ratio with 95% confidence intervals. We used the standard DerSimonian and Laird random effects model for primary analysis,18 we chose this method a priori to account for potential variation between studies owing to differences in study populations. We also performed Mantel-Haenszel fixed effects analyses for estimating pooled odds ratios, to account best for the limited data available from some of the component studies.19 We used the Q statistic for all comparisons to evaluate heterogeneity of effect among studies. If we chose to accept this variation, we performed a sensitivity analysis excluding the study with the highest Q value.20

Results
Figure 1 summarises the process of identifying studies eligible for inclusion in our analysis. We reviewed the full text of 128 articles from 2742 studies identified from our initial literature search and hand search. A total of 24 studies met criteria for inclusion, and we could ascertain data on myocardial infarction for 19.13 14 21–25 27–32 34–37 We excluded four controlled studies identified in the literature search because the control arms were usual care,26 atenolol,27 amiodipine,28 and hydrochlorothiazide.29 Agreement between two investigators for final study inclusion, measured by the κ statistic, was 0.75. All trials had a prospective, parallel design. Nine trials had the maximum Jadad score of 5; four trials scored 4, five scored 3, and one scored 2 (table 1). Allocation concealment was adequate in eight studies (42%), inadequate in two studies (11%), and unclear in the remaining nine studies (47%). With the exception of the study by Di Pasquale et al,29 treatment was assigned in a randomised fashion. Participants were blinded in 19 studies (100%), investigators in 17 studies (89%), and outcome assessors in 18 (95%). Finally, patients were analysed by the intention to treat principle in 15 of the studies (79%).

In our data set, the study by Bakris et al22 compared losartan with both a placebo arm and an enalapril arm; we therefore included the study in both analyses. We did not use information from the amiodipine arm of the irbesartan diabetic nephropathy trial.31 Two studies investigated the use of angiotensin receptor blockers in hypertensive patients,21 22 four studies in patients with diabetes mellitus and nephropathy,21 22 24 25 and three studies in patients with a recent myocardial infarction or ischaemic syndrome.13 36 37 Myocardial infarction, major adverse cardiac events, or cardiac mortality was an adjudicated study end point in nine studies,13 14 21 24 25 28–30 35 whereas myocardial infarction was reported as an adverse event or was reported by the investigators in 10 studies that had primarily physiological or drug tolerability outcomes.22 25 26 27 33 36 36 34–37 Finally, nine studies allowed for comparison between angiotensin receptor blockers and ACE inhibitors,13 14 22 24 25 30–32 34 and 11 studies allowed for comparison between angiotensin receptor blockers and placebo.21 22 25 27–32 Funnel plots for the angiotensin receptor blocker comparisons with placebo studies and angiotensin receptor blockers compared with ACE inhibitor studies are qualitatively symmetrical, indicating the absence of publication bias (fig 2). Other important study characteristics, including number of subjects, mean age, sex, and duration of follow-up, are summarised in the table.

Effect of angiotensin receptor blockers compared with placebo on risk of myocardial infarction
In this analysis we included two hypertension trials, three trials of patients with diabetes and nephropathy, and six heart failure trials, with a total of 10 656 subjects allocated to treatment with angiotensin receptor blockers and 10 406 subjects allocated to placebo. In the group that was treated with angiotensin receptor blockers, 436 myocardial infarctions occurred (4.09%), compared with 450 myocardial infarctions in the placebo group (4.32%). Overall, using angiotensin receptor blockers was not associated with a significant increase in the risk of myocardial infarction, with a pooled odds ratio of 0.94 (95% confidence interval 0.75 to 1.16) from the random effects model (fig 3). Analysis using the fixed effects model similarly showed no significant association of using angiotensin receptor blockers with risk of myocardial infarction (pooled odds ratio 0.95, 0.83 to 1.09).
Effect of angiotensin receptor blockers compared with angiotensin converting enzyme inhibitors on risk of myocardial infarction

Figure 4 shows the results of the comparison of treatment with angiotensin receptor blockers and ACE inhibitors with respect to risk of myocardial infarction in one hypertension study, one diabetes and nephropathy study, four heart failure studies, and three recent studies of myocardial infarction or ischaemic syndrome. Among 5406 patients receiving angiotensin receptor blockers, 435 myocardial events occurred (8.05%), compared with 433 events (8.30%) in 5219 patients receiving ACE inhibitors, resulting in a pooled odds ratio close to unity (1.01, 0.87 to 1.16 by random effects analysis; 1.00, 0.87 to 1.16 by fixed effects analysis). This summary effect size was driven mainly by the OPTIMAAL study, which accounted for 86.8% of the weighted odds ratio in the random effects model, with an individual study odds ratio of 1.01 (0.87 to 1.16).

Discussion

Treatment with angiotensin receptor blockers was not associated with an increased risk of myocardial infarction, according to our systematic review of 19 trials with 31 369 subjects. With a pooled odds ratio very close to unity in our analyses for angiotensin receptor blockers compared with placebo and compared with ACE inhibitor, our results indicate that an aggregate of patients with hypertension, diabetes and nephropathy, heart failure and left ventricular dysfunction, and patients with recent myocardial infarction or ischaemic syndrome were not at greater risk of myocardial infarction when treated with different angiotensin receptor blockers.

Angiotensin receptor blockers versus placebo

In our analysis of treatment with angiotensin receptor blockers compared with placebo including 21 062 patients, we found no significant difference between groups in the incidence of myocardial infarction, although the 95% confidence intervals cannot rule out an increased risk of myocardial infarction of up to 16% or a reduced risk of up to 25%. Our evaluation of angiotensin receptor blockers compared with placebo included the CHARM-alternative trial, which contributed more than 13% to the weighted pooled odds ratio and was the only study to...
show an increase in myocardial infarction rates with use of angiotensin receptor blockers that reached significance. In this study, patients with left ventricular dysfunction and heart failure who were intolerant to ACE inhibitors were randomised to the angiotensin receptor blocker candesartan or placebo. Despite the observed increased incidence of myocardial infarction in the candesartan group, cardiovascular mortality fell overall with treatment with angiotensin receptor blockers. In contrast, among other patients with heart failure and similar background cardiovascular risk, including patients being treated with concomitant ACE inhibitors in the CHARM-added and ValHeFT trials, the point estimates were distributed across the 1.0 odds ratio, implying that using angiotensin receptor blockers is itself not significantly associated with risk of myocardial infarction. Although we have not shown a clear relative benefit on myocardial infarction of treatment with angiotensin receptor blockers compared with placebo, our results indicate that this class of medications is unlikely to be harmful.

### Angiotensin receptor blockers versus ACE inhibitors

As ACE inhibitors have been shown unequivocally to reduce cardiac morbidity and mortality among patients at risk for cardiovascular events, a prespecified comparative analysis with ACE inhibitors was necessary to assess better the safety and rel-

<table>
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<tr>
<th>Study</th>
<th>ARB group (n/N)</th>
<th>Placebo group (n/N)</th>
<th>Odds ratio (random) (95% CI)</th>
<th>Weight (%)</th>
<th>Odds ratio (random) (95% CI)</th>
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<td>Hypertension</td>
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<td>0/58</td>
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<td>Type 2 diabetes and nephropathy</td>
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<td>0/32</td>
<td>0.45 (1.03 to 25.98)</td>
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<td>Haneda et al 2004</td>
<td>1/95</td>
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<td>12.03 (0.94 to 1.44)</td>
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<td>Heart failure</td>
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<td>ARCH-J 2003</td>
<td>0/148</td>
<td>0/144</td>
<td></td>
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<td>CHARM-added 2003</td>
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<td>13.23 (0.62 to 2.92)</td>
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<td>CHARM-alternative 2003</td>
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<td>13.61 (1.11 to 2.34)</td>
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<td>CHARM-preserved 2003</td>
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<td>73/1509</td>
<td>14.18 (0.54 to 1.10)</td>
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<td>SPICE 2000</td>
<td>5/178</td>
<td>5/91</td>
<td>2.63 (0.49 to 1.75)</td>
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<td>Val-HeFT 2001</td>
<td>89/2596</td>
<td>78/2494</td>
<td>15.55 (1.14 to 1.55)</td>
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<tr>
<td>Total (95% CI)</td>
<td>10 656</td>
<td>10 406</td>
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<td>100.00</td>
<td>0.94 (0.75 to 1.16)</td>
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<td>Test for overall effect: z=0.60, P=0.55</td>
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Fig 3 Effect of angiotensin receptor blockers compared with placebo on risk of myocardial infarction

<table>
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<tr>
<th>Study</th>
<th>ARB group (n/N)</th>
<th>ACE Inhibitor (n/N)</th>
<th>Odds ratio (random) (95% CI)</th>
<th>Weight (%)</th>
<th>Odds ratio (random) (95% CI)</th>
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<td>Hypertension</td>
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<td>Bakris et al 2002</td>
<td>1/118</td>
<td>0/113</td>
<td>0.20 (2.90 to 71.88)</td>
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<tr>
<td>Type 2 diabetes and nephropathy</td>
<td>10/120</td>
<td>8/130</td>
<td>2.18 (1.39 to 3.64)</td>
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<tr>
<td>DETAIL 2004</td>
<td>0/148</td>
<td>0/144</td>
<td></td>
<td>Not estimable</td>
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<tr>
<td>Heart failure</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ELITE 1997</td>
<td>4/352</td>
<td>8/370</td>
<td>1.39 (0.52 to 1.74)</td>
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<td>ELITE II 2000</td>
<td>3/1578</td>
<td>28/1574</td>
<td>7.64 (1.11 to 4.85)</td>
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<td>HEAVEN 2002</td>
<td>0/79</td>
<td>2/71</td>
<td>0.22 (0.02 to 1.48)</td>
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<td>REPLACE 2001</td>
<td>0/301</td>
<td>1/77</td>
<td>0.20 (0.08 to 2.10)</td>
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<tr>
<td>Recent myocardial infarction or ischaemic syndrome</td>
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<td>Di Pasquale et al 1999</td>
<td>0/23</td>
<td>3/50</td>
<td>0.23 (0.29 to 5.82)</td>
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<td>OPTIMAAL 2002</td>
<td>384/2744</td>
<td>379/2753</td>
<td>86.83 (1.01 to 5.87)</td>
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<tr>
<td>Spinat et al 2000</td>
<td>5/100</td>
<td>4/101</td>
<td>1.12 (1.28 to 4.90)</td>
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<tr>
<td>Total (95% CI)</td>
<td>5406</td>
<td>5219</td>
<td></td>
<td>100.00</td>
<td>1.01 (0.87 to 1.16)</td>
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<tr>
<td>Test for overall effect: z=0.12, P=0.91</td>
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</table>

Fig 4 Effect of angiotensin receptor blockers compared with angiotensin converting enzyme inhibitors on risk of myocardial infarction
tive effect of angiotensin receptor blockers on the risk of myocardial infarction. In this analysis, we found no difference in this risk between the arms receiving angiotensin receptor blockers and ACE inhibitors. Similar to our analysis comparing angiotensin receptor blockers with placebo, the 95% confidence interval included an up to 16% increased risk of myocardial infarction down to a 13% reduction with angiotensin receptor blockers. Although our evaluation included nine component studies, all of which had 95% confidence intervals crossing 1.0, the overall effect was most heavily influenced by the OPTIMAAL study.30 In this trial, the angiotensin receptor blocker losartan was found to be non-superior in large randomised controlled trials,15-18 our finding supports the notion that they may be a safe and effective alternative for a select group of heart failure patients not taking ACE inhibitors, as shown by a recent meta-analysis.48

Angiotension receptor blockers do not increase risk of myocardial infarction
Our finding that angiotensin receptor blockers are not associated with an increased risk of myocardial infarction stands in contrast to the editorial by Verma and Strauss,33 which drew attention to results from several recent studies, including the VALUE trial,30 the CHARM-alternative36 and CHARMPreserved trials, and the SCOPE,37 LIFE,38 and RENAAL39 trials. Our analysis, which included all but the LIFE38 and VALUE39 trials (excluded from our analysis because they did not have placebo or ACE inhibitor control groups), highlights the importance of assessing all available evidence by using systematic methods, before drawing conclusions.

Limitations
Although we tried to conduct a thorough review of the existing literature, our study has limitations inherent to any systematic review. Firstly, we were unable to obtain data on myocardial infarction events from all studies identified in our literature search. Perhaps most notably, data on myocardial infarction from the valsartan in acute myocardial infarction trial (VALIANT),13 which included 14 703 patients, were not available for our analysis. We informally tested the potential impact of the unavailable data from this trial by assuming a “worst case scenario” in which all 919 patients admitted to hospital for myocardial infarction or heart failure, or both, in the valsartan arm of VALIANT had a myocardial infarction, and 0 patients in the captopril arm were admitted to hospital for myocardial infarction. In this scenario, the pooled odds ratio in our ACE-inhibitor controlled analysis changed from 1.01 (0.87 to 1.16) to 1.40 (0.25 to 7.80). Despite the addition of almost 10 000 patients to our study total, we would still observe no significant association between use of angiotensin receptor blockers and myocardial infarction risk compared with ACE inhibitors. We would further speculate that data from other potential sources not included in our study (from smaller trials, unpublished reports) would not significantly influence our results. Similarly, excluding the non-randomised trial would have little impact on our findings. Other limitations of our study include potential variation in the definition of myocardial infarction between studies, the possibility of effects that are specific to angiotensin receptor blockers and dose dependent, and the potential confounding influence of other treatments.

Conclusions
We conducted a systematic quantitative review of angiotensin receptor blockers and the risk of myocardial infarction. Until information specifically dealing with this issue is available from large, prospectively designed trials, such as the ONTARGET/TRANSCEND trials,40 we must rely on the weight of available evidence to guide decisions on the management of individual patients’ decisions. Our results show that treatment with angiotensin receptor blockers is not associated with increased incidence of myocardial infarction. The authors thank Janice Varney, MLIS, for her help in conducting the literature search. The authors also thank the individual trialists for sharing data from their original studies. Contributors: All authors took part in the planning and design of the study. MAM, SHS, JAE, and GG did the data collection. SHS did the statistical analyses. All participated in the interpretation of the data. MAM and SHS wrote the first draft of the paper. All authors reviewed and revised the paper for important intellectual content. RTT provided leadership for the study, is the senior and corresponding author, and takes responsibility for the content. Funding: None. Competing interests: SHS has received honorariums from Merck Frosst and Sanofi-Aventis. GG has received travel grants and honorariums from Aventis, Pfizer, Servier, Novartis, Boivail, AstraZeneca, Merck-Schering, Merck Frosst, and Procter and Gamble. RTT is supported by the Merck Frosst Chair in Patient Health Management from the University of Alberta. He is a member of the Cardiovascular Advisory Board of Merck Frosst and has received honorariums and research grants from Merck Frosst, Merck-Schering, Pfizer, Sanofi-Aventis, and Novartis. Ethical approval: Not required.

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