Cost-effectiveness, cost-utility and the budget impact of antidepressants versus preventive cognitive therapy with or without tapering of antidepressants

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Background
As depression has a recurrent course, relapse and recurrence prevention is essential.

Aims
In our randomised controlled trial (registered with the Nederlands trial register, identifier: NTR1907), we found that adding preventive cognitive therapy (PCT) to maintenance antidepressants (PCT+AD) yielded substantial protective effects versus antidepressants only in individuals with recurrent depression. Antidepressants were not superior to PCT while tapering antidepressants (PCT−AD). To inform decision-makers on treatment allocation, we present the corresponding cost-effectiveness, cost-utility and budget impact.

Method
Data were analysed (n = 289) using a societal perspective with 24-months of follow-up, with depression-free days and quality-adjusted life years (QALYs) as health outcomes. Incremental cost-effectiveness ratios were calculated and cost-effectiveness planes and cost-effectiveness acceptability curves were derived to provide information about cost-effectiveness. The budget impact was examined with a health economic simulation model.

Results
Mean total costs over 24 months were €6814, €10 264 and €13 282 for AD+PCT, antidepressants only and PCT−AD, respectively. Compared with antidepressants only, PCT+AD resulted in significant improvements in depression-free days but not QALYs. Health gains did not significantly favour antidepressants only versus PCT−AD. High probabilities were found that PCT+AD versus antidepressants only and antidepressants only versus PCT−AD were dominant with low willingness-to-pay thresholds. The budget impact analysis showed decreased societal costs for PCT+AD versus antidepressants only and for antidepressants only versus PCT−AD.

Conclusions
Adding PCT to antidepressants is cost-effective over 24 months and PCT with guided tapering of antidepressants in long-term users might result in extra costs. Future studies examining costs and effects of antidepressants versus psychological interventions over a longer period may identify a break-even point where PCT−AD will become cost-effective.

Declaration of interest
C.L.H.B. is co-editor of PLOS One and receives no honorarium for this role. She is also co-developer of the Dutch multidisciplinary clinical guideline for anxiety and depression, for which she receives no remuneration. She is a member of the scientific advisory board of the National Insure Institute, for which she receives an honorarium, although this role has no direct relation to this study. C.L.H.B. has presented keynote addresses at conferences, such as the European Psychiatry Association and the European Conference Association, for which she sometimes receives an honorarium. She has presented clinical training workshops, some including a fee. She receives royalties from her books and co-edited books and she developed preventive cognitive therapy on the basis of the cognitive model of A. T. Beck. W.A.N. has received grants from the Netherlands Organisation for Health Research and Development and the European Union and honoraria and speakers’ fees from Lundbeck and Aristo Pharma, and has served as a consultant for Daleco Pharma.

Keywords
Depressive disorders; antidepressants; cognitive behavioural therapies; cost-effectiveness; economics.

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In the Disrupt the Rhythm of Depression (DRD) trial (registered with the Nederlands trial register, identifier: NTR1907), we found that adding preventive cognitive therapy (PCT) to maintenance antidepressants (PCT+AD) resulted in a statistically significant risk reduction in terms of time-related proportion of individuals with depressive recurrence compared with maintenance antidepressants only. We could not demonstrate that antidepressants reduced the risk of recurrence more compared with PCT with guided tapering of antidepressants (PCT/−AD). The aim of the current study was to perform cost-effectiveness, cost-utility and budget impact analyses alongside the DRD trial.

Method

Study design

The DRD trial is a single-blind multicentre three-arm randomised controlled trial. The economic evaluation was performed according to the Consolidated Health Economic Evaluation Reporting Standards statement. The study was approved by an independent medical ethics committee (METIGG) and described in detail elsewhere.

Participants and procedure

Participants were recruited via general practitioners, pharmacists, secondary mental healthcare and the media. To be included, participants had to have experienced at least two prior major depressive episodes, had been in remission for at least 8 weeks but no longer than 2 years based on DSM-IV criteria as assessed by the Structured Clinical Interview for DSM-IV Axis-I Disorders (SCID-I) and have a score on the Hamilton Rating Scale for Depression of ≤10. Participants must have used maintenance antidepressants for at least 6 months. Exclusion criteria were (hypo)mania or a psychotic disorder/MDD with psychotic features, current alcohol or drug misuse, organic brain damage, a predominant anxiety disorder or psychotherapy more than twice a month. Because individuals with various depressive episodes over a longer time span also benefit from relapse prevention strategies, we discarded our initial criterion that both major depressive episodes must have occurred within the past 5 years. In addition, PCT was initially offered in groups but we extended this to individual sessions during the trial since many participants were not able to attend the group meetings because of practical concerns.

Randomisation

After obtaining informed consent, participants were randomised to PCT+AD, antidepressants only or PCT−AD by an independent research assistant not otherwise involved in the study using computer-generated random numbers (allocation ratio 10:10:8, respectively). The randomisation was stratified by number of previous major depressive episodes (two versus three or more) and baseline care (general practitioner versus secondary mental healthcare). Participants were informed about their treatment allocation by a research assistant not involved in the follow-up interviews. Independent assessors masked to treatment allocation conducted the follow-up interviews.

Interventions

PCT is a treatment strategy that is effective in preventing recurrence in individuals in remission with recurrent depression and targets potential cognitive vulnerability factors of MDD. It consists of eight manualised individual or group sessions performed by trained psychologists. Issues with therapist adherence were discussed and resolved during supervision meetings. An independent research assistant assessed therapist adherence to the treatment manual and adherence was judged to be high (87%, range: 81–95, for details, see Bockting et al). In the two antidepressant continuation groups, general practitioners and psychiatrists were advised to continue prescribing antidepressants at the minimal required adequate dosage or higher (≥20 mg fluoxetine equivalent) and in the tapering group to taper the participants’ antidepressant within 4 weeks. Participants’ adherence to the randomised condition is described in detail elsewhere. In short, adherence to PCT (i.e. completing at least five sessions) was 88% in the PCT+AD group and 90% in the PCT−AD group. Adherence to antidepressants was monitored with the Trimbos and iMTA questionnaire on costs associated with psychiatric illness (TiC-P). Most (60%) individuals in the PCT−AD group tapered their ADs over 6 months. At 6 months, adherence to antidepressants was 58% in the antidepressants-only group, 60% in PCT−AD group and 65% in the PCT+AD group.

Economic evaluation

Costs

In accordance with the Dutch guidelines, we used a societal perspective in this economic evaluation, in which all costs inside and outside the healthcare sector were assessed. We used the TiC-P, which is a reliable and valid self-report instrument for collecting cost data, to prospectively assess the costs. The TiC-P was designed to refer to the past 3 months and was administered at baseline and subsequently at 3-month intervals, with the exception of the last assessment which had a 9-month interval. For the costs in this 9-month interval we therefore took the costs of the past 3 months multiplied by three. In addition, the maximum number of days medicated was only 28 and therefore we extrapolated all medication use to 3 months.

Costs within the healthcare sector were related to a range of healthcare services participants used during the study, including medication and in-patient stays, and out-patient and primary care appointments. Costs related to PCT included training and supervision of therapists, costs of contacts between participants and therapists and costs of the workbooks used by participants. Patient and family costs included informal care (i.e. the monetary valuation of time invested by relatives or acquaintances in assisting the participant), travel expenses associated with healthcare visits and (psychiatric) home care. Informal care was measured as part of the TiC-P, by asking participants to express the number of hours per week they received care from relatives and/or friends. This was then valued using the proxy good method (i.e. time spent on caregiving was valued at the (labour) market price of a close substitute, in this case housekeeping). Productivity losses were quantified and included absence from work (absenteeism), reduced productivity while at work (presenteeism) and productivity losses of unpaid work. We used the friction cost method to estimate costs associated with productivity losses as a result of illness-related absence from work. To promote comparisons with other economic evaluations, Dutch standard prices were used as unit prices. We estimated true costs of used resources when standard prices were not available. All unit prices were based on the price level of the Euro in 2015. Reference prices established for previous years were adjusted to 2015 prices applying the consumer price index.
Outcomes

The health outcome of the cost-effectiveness analysis was the number of depression-free days over a period of 24 months based on DSM-IV criteria assessed with the SCID-I by masked interviewers after 3, 9, 15 and 24 months. Quality-adjusted life-years (QALYS) over 24 months was the health outcome measure in the cost-utility analysis and represent disease burden by combining quality (expressed in utilities) and quantity of life, where one QALY represents 1 year of perfect health. We used utilities, derived from the EQ-SD-3L that measures health-related quality of life, to calculate QALYs with the area under the curve method. These utilities were calculated with Dutch tariffs to obtain utilities for specific health states. Costs and health outcomes were discounted in accordance with current Dutch guidelines (1.5% health outcomes, 4% costs).

Statistical analyses

This study was conducted alongside a clinical trial and the power calculation of the primary outcome is provided elsewhere. This power calculation was based on detecting a difference in the time-related proportion of individuals with depressive recurrence regarding the following comparisons: (a) adding PCT to antidepressants versus antidepressants only; and (b) antidepressants only versus PCT while tapering antidepressants. Given that the study was only powered to detect a difference in a depression-related outcome and not in costs, we used probabilistic and medical decision-making techniques to draw inferences about cost-effectiveness. Conforming to the trial protocol, the analyses in this manuscript were restricted to examine these two comparisons with as primary analysis a cost-effectiveness analysis using depression-free days as the health outcome and as the secondary analysis a cost-utility analysis using QALYS as the health outcome.

We used the intention-to-treat principle, in which all participants were included in the analyses regardless of adherence to the randomised interventions. To deal with missing data, multiple imputations by chained equations with predictive mean matching, which is recommended for dealing with missing data in (cost-effectiveness) trials, were used in our main analysis, incorporating baseline variables predictive of outcome and drop-out. Multiple imputations were used to make optimal use of the available data and reduce possible selection bias because of non-random drop-out. Costs and outcomes associated with each treatment condition were used to calculate the incremental cost-effectiveness ratio (ICER) relative to an alternative. The formula of the ICER is:

\[
\text{ICER} = \frac{(C_{\text{PCT+AD}} - C_{\text{AD}})}{(QALY_{\text{PCT+AD}} - QALY_{\text{AD}})}
\]

where \( C_{\text{PCT+AD}} \) is the mean costs in the PCT+AD group; \( C_{\text{AD}} \) is the mean costs in the antidepressants-only group; \( QALY_{\text{PCT+AD}} \) is the mean QALYS in the PCT+AD group and \( QALY_{\text{AD}} \) is the mean QALYS in the antidepressants-only group.

The bootstrap method was used for information about the uncertainty of the results. To allow correlated residuals and correct for differences in baseline characteristics in sensitivity analyses, seemingly unrelated regression equations were bootstrapped 5000 times. Simulated values of the estimates for cost and outcome differences were displayed in a cost-effectiveness plane to capture the uncertainty in the ICER estimate. Information about the cost-effectiveness plane can be found in supplementary File 2. Cost-effectiveness acceptability curves (CEACs) were derived that inform decision-makers about the probability that an intervention will be cost-effective, depending on the willingness to pay per additional unit of health outcome.

Both complete cost and effect data were available for 105 participants. For 169 participants at least 50% of the cost data and for 175 at least 50% of the QALY data were available during the 24 months. For 197 participants follow-up data on depression-free days were available. We performed several sensitivity analyses to examine the robustness of our results. The intervention could not be started and follow-up data were not available for 43 participants in the PCT groups for practical reasons. In addition, 16 participants in the PCT groups and 21 in the antidepressants-only group dropped out immediately after randomisation for other reasons. Ultimately, 209 participants had follow-up data and were used in our main analysis. We included all 289 participants in a sensitivity analysis.

Furthermore, we repeated the analyses in participants with at least 50% cost data available and in complete cases. We also repeated the main analysis while correcting for baseline costs and utilities, for whether participants filled out additional momentary assessments, and for whether participants in the therapy groups received the individual or group therapy. Finally, our results for the primary outcomes suggested an increased recurrence rate for PCT/AD compared with antidepressants only in the first 140 days. To account for possible withdrawal symptoms in the first months, we performed an analysis taking into account only the final 1.5 years of the study.

Budget impact analysis

We used a health economic simulation model for depression, DEPMOD, to examine the budget impact when offering the different relapse prevention strategies to an estimated 25% of the target population in the Netherlands. Per-person cost differences as estimated by the trial data were applied to 25% of the target group to estimate the order of magnitude of the budget impact. The budget impact was estimated both from a healthcare perspective and a societal perspective, and considered a 2-year time horizon.

Results

Details of the participant flow have been described elsewhere. Between 14 July 2009 and 30 April 2015, 289 participants were assessed for eligibility and randomised to the PCT+AD (n = 104), antidepressants-only (n = 100) or PCT/AD (n = 85) groups. Of those, 209 provided additional data following randomisation. Demographic and clinical variables are displayed in Table 1. The participants in the treatment groups appeared to have similar characteristics, except for slight imbalances in marital status, education and employment status. Since baseline characteristics of all participants and the participants with any follow-up data were similar and equally distributed over the treatment groups, no indication for a systematic bias because of drop-out was found.

Costs

The various types of costs generated by the three groups and information on the use of healthcare services during the 24 months of the study are presented in supplementary Table 1. Costs are based on the data of participants for whom at least one cost measurement was available during follow-up. Mean costs per participant directly related to PCT were €349, €354, and €0 in the PCT+AD, PCT/AD and antidepressants-only groups, respectively. These costs mainly consisted of costs related to training/supervision of therapists and contacts between participants and therapists. Hospital admissions and care provided by mental healthcare institutions contributed considerably to overall costs within the healthcare sector. Costs associated with productivity losses were substantial.
When visually inspecting supplementary Table 1, in most categories costs appear slightly lower for the PCT+AD group compared with the antidepressants-only group and the antidepressants-only group compared with the PCT/–AD group except for larger reductions for PCT+AD compared to antidepressants-only regarding absenteeism and hospital admissions.

An overview of the mean costs per measurement for all 209 individuals is displayed in supplementary Table 2. Mean total costs of the PCT+AD group appear only higher between 3 and 9 months and lower during the other measurements. Accumulating all costs (supplementary Table 2), mean total costs for the antidepressants-only group was not statistically significant (730) for the PCT/–AD group, 607 (range 194–1.95) for the antidepressants-only group, 662 (range 194–1.99) for the antidepressants-only group (i.e. PCT+AD is dominant). The CEACs in Fig. 1 regarding depression-free days shows a high probability that PCT+AD is dominant. Regarding QALYs, PCT+AD was associated with 67.8% of the bootstrapped ICERs appearing in the south-west quadrant, indicating that costs were lower and health outcomes worse compared with antidepressants only (Table 2 and Fig. 1). Interpretation of outcomes located in the south-west quadrant depends on whether decision-makers are willing to accept a cost reduction for a loss in health. The CEAC in Fig. 1 regarding QALYs shows a high probability that PCT+AD dominates antidepressants only when the willingness-to-accept threshold is low and a decreasing probability if the threshold to accept a reduction in health is increased.

Regarding antidepressants only compared with PCT/–AD, Table 2 and the cost-effectiveness planes (Fig. 2) show that most (72.9% for depression-free days and 80.8% for QALYs) of the bootstrapped ICERs were located in the south-east quadrant, indicating a 93.1% probability that costs were lower and health outcomes better for PCT+AD compared with the antidepressants-only group (i.e. PCT+AD is dominant). The CEAC in Fig. 2 show that the probability that antidepressants only are dominant compared with PCT/–AD is approximately 90% for both depression-free days and QALYs when the willingness to pay per additional

**Table 1** Demographics and clinical variables

|                          | PCT+AD group (n = 104) | Antidepressants-only group (n = 100) | PCT/–AD group (n = 85) |
|--------------------------|------------------------|--------------------------------------|------------------------|
| Age, years:¢ mean (s.d.) | 47.0 (9.3)             | 47.2 (10.5)                          | 47.7 (11.1)            |
| Gender: women; n (%)     | 72 (69)                | 64 (64)                              | 53 (62)                |
| Dutch nationality, began | 101 (97)               | 95/99 (96)                           | 82/84 (98)             |
| Marital status, n (%)    | 27/103 (26)            | 32/99 (32)                           | 28/84 (33)             |
| Single                   | 49/103 (67)            | 59/99 (60)                           | 46/84 (55)             |
| Married/cohabiting       | 7/103 (7)              | 8/99 (8)                             | 10/84 (12)             |
| Education, n (%)         | 20 (19)                | 25/99 (25)                           | 12/84 (14)             |
| Primary and/or secondary education | 31 (30) | 28/99 (28) | 23/84 (27) |
| Vocational education     | 53 (51)                | 46/99 (46)                           | 49/84 (58)             |
| Higher education         | 73/103 (71)            | 65/98 (66)                           | 53/84 (63)             |
| Treatment as usual, n (%)| 32 (31)                | 31 (31)                              | 26 (31)                |
| Specialised mental healthcare | 72 (69) | 69 (69) | 59 (69) |
| General practitioner     | 5 (3–6)                | 4 (3–6)                              | 5 (3–6)                |
| Total HRSD, mean (s.d.)  | 3.6 (3.1)              | 3.8 (3.1)                            | 3.6 (3.0)              |
| Total IDS-SR, mean (s.d.)| 20.4 (11.5)            | 18.5 (10.8)                          | 20.6 (12.1)            |
| Type of antidepressant, n (%) | 85/103 (83) | 79/99 (83) | 69/85 (81) |
| SSRI                     | 1/103 (1)              | 8/99 (8)                             | 1/85 (1)               |
| SNRI                     | 7/103 (7)              | 7/99 (7)                             | 10/85 (12)             |
| Tricyclic antidepressant | 5/103 (5)              | 2/99 (2)                             | 3/85 (4)               |
| Atypical antidepressant  | 1/103 (1)              | 0 (0)                                | 0 (0)                  |
| Monoamine oxidase inhibitor | 0 (0)   | 0 (0) | 0 (0) |
| More than one antidepressant | 5/103 (5) | 2/99 (2) | 2/85 (2) |
| EQ-5D-3L, mean (s.d.)    | 0.84 (0.16)            | 0.80 (0.18)                          | 0.79 (0.17)            |
| Baseline costs, ¥ mean (s.d.) | 1533 (5423) | 1695 (3049) | 1778 (3383) |

PCT + AD, preventive cognitive therapy and antidepressants; PCT/–AD, preventive cognitive therapy with guided tapering of antidepressants; IQR, interquartile range; HRSD, Hamilton Rating Scale for Depression; IDS-SR, Inventory of Depressive Symptomatology Self-Report; SSRI, selective serotonin reuptake inhibitor; SNRI, selective serotonin and norepinephrine reuptake inhibitor.

¢ Two participants were older than 65 years at baseline (i.e. 67 and 68 years).

¥ Data not available for all randomised participants.

Imputed data.

Economic evaluation

The results of the main analyses are presented in Table 2. Table 2 and the cost-effectiveness plane (Fig. 1) show that regarding depression-free days, most (93.1%) of the bootstrapped ICERs were located in the south-east quadrant, indicating a 93.1% probability that costs were lower and health outcomes better for PCT+AD compared with the antidepressants-only group (i.e. PCT+AD is dominant). The CEAC in Fig. 1 regarding depression-free days shows a high probability that PCT+AD is dominant. Regarding QALYs, PCT+AD was associated with 67.8% of the bootstrapped ICERs appearing in the south-west quadrant, indicating that costs were lower and health outcomes worse compared with antidepressants only (Table 2 and Fig. 1). Interpretation of outcomes located in the south-west quadrant depends on whether decision-makers are willing to accept a cost reduction for a loss in health. The CEAC in Fig. 1 regarding QALYs shows a high probability that PCT+AD dominates antidepressants only when the willingness-to-accept threshold is low and a decreasing probability if the threshold to accept a reduction in health is increased.

Regarding antidepressants only compared with PCT/–AD, Table 2 and the cost-effectiveness planes (Fig. 2) show that most (72.9% for depression-free days and 80.8% for QALYs) of the bootstrapped ICERs were located in the south-east quadrant, indicating a 72.9% and 80.8% probability that antidepressants only generated lower costs and better outcomes compared with PCT/–AD (i.e. antidepressants only is dominant). The CEACs in Fig. 2 show that the probability that antidepressants only are dominant compared with PCT/–AD is approximately 90% for both depression-free days and QALYs when the willingness to pay per additional
health gain is zero and the probability for depression-free days slightly decreases and for QALYs slightly increases with additional investments. The sensitivity analyses yielded similar results. When only taking into account the last 1.5 years of the study to examine the cost-effectiveness of antidepressants only compared with PCT/−AD, the gain in depression-free days decreased from 21 to 12 but antidepressants only remained cost-effective compared with PCT/−AD.

**Budget impact analysis**

The target population size in the Netherlands in a given year, meaning the yearly prevalence of people with at least two previous episodes of depression, was estimated to be 110 000, which is roughly 1% of the total population of approximately 10 million people aged between 18 and 65 in the Netherlands. Offering PCT+AD instead of antidepressants alone to 25% of the target population is associated with an estimated decrease in costs of 76 million euro from a healthcare perspective and 95 million euro from a societal perspective. Offering antidepressants alone instead of PCT/−AD to 25% of the target population is associated with an estimated decrease in costs of 9.5 million euro from a healthcare perspective and 83 million euro from a societal perspective.

**Discussion**

Adding PCT to antidepressants had the highest number of depression-free days and lowest costs, PCT/−AD had the lowest number of depression-free days and highest costs, and antidepressants only ranked in-between in terms of depression-free days and costs. Adding PCT to antidepressants was dominant compared with antidepressants only regarding depression-free days and resulted in decreased costs at a population level whereas antidepressants only dominated PCT/−AD in the cost-effectiveness, cost-utility and budget impact analyses.

Adding PCT to antidepressants compared with antidepressants only

Although adding PCT to antidepressants was dominant compared with antidepressants only resulting in an increase (statistically significant) in depression-free days and decrease in costs, QALYs did not significantly differ for PCT+AD compared with antidepressants only. The EQ-5D used in our study might lack sensitivity to detect small improvements in individuals in remission with recurrent MDD. Moreover, it only displays the current health state and therefore does not capture all recurrences during the 24 months of the study. Studies indeed suggest that the EQ-5D might be less representative for individuals with psychiatric symptoms. Therefore, also in line with the trial protocol, we consider the cost-effectiveness analysis using depression-free days as the primary outcome.

The finding that PCT+AD dominated antidepressants only in terms of depression-free days contrasts with another Dutch relapse prevention study where adding mindfulness-based cognitive therapy to maintenance antidepressants did not reduce the risk of recurrence compared with antidepressants only. However, it is congruent with studies demonstrating that sequential cognitive therapy after remission is protective of recurrence and with our primary outcomes. Moreover, our finding is partly in line with a cost-effectiveness study of a relapse prevention strategy for partially remitted depression that found cognitive therapy added to antidepressants and clinical management was likely to be cost-effective compared with antidepressants and clinical management only over 17 months when decision-makers were willing to pay £4500 per recurrence prevented, and with a cost-effectiveness study

| Analysis | Incremental effects (95% CI) | Mean ICER, € | North-east quadrant, % | South-east quadrant, % | South-west quadrant, % | North-west quadrant, % | In inferior quadrant, % |
|----------|-----------------------------|--------------|------------------------|-----------------------|-----------------------|------------------------|------------------------|
| PCT+AD (n = 71) versus antidepressants only (n = 79) | Depression-free days: 3409 (−3408 to −3360) 34.1 (33.5 to 34.7) Dominant | Dominant | 1.9 92.1 29.9 4.5 0.4 | Dominant | 7.4 72.9 16.8 3.0 |
| PCT/−AD (n = 59) versus antidepressants only (n = 79) | Depression-free days: 3015 (−3084 to −2947) 21.0 (20.2 to 21.7) Dominant | Dominant | 1.9 92.1 29.9 4.5 0.4 | Dominant | 7.4 72.9 16.8 3.0 |

| Analysis | Incremental effects (95% CI) | Mean ICER, € | North-east quadrant, % | South-east quadrant, % | South-west quadrant, % | North-west quadrant, % | In inferior quadrant, % |
|----------|-----------------------------|--------------|------------------------|-----------------------|------------------------|------------------------|------------------------|
| PCT+AD (n = 71) versus antidepressants only (n = 79) | Quality-adjusted life-years (QALYs): 3409 (−3408 to −3360) −0.018 (−0.019 to −0.017) Dominant | Dominant | 1.9 92.1 29.9 4.5 0.4 | Dominant | 7.4 72.9 16.8 3.0 |
| PCT/−AD (n = 59) versus antidepressants only (n = 79) | Quality-adjusted life-years (QALYs): 3015 (−3084 to −2947) 0.051 (0.050 to 0.053) Dominant | Dominant | 1.9 92.1 29.9 4.5 0.4 | Dominant | 7.4 72.9 16.8 3.0 |

| Analysis | Incremental costs (95% CI), € | Incremental effects (95% CI) | Cost-effectiveness, depression-free days | Cost-utility, QALYs |
|----------|-----------------------------|-----------------------------|----------------------------------------|---------------------|
| PCT+AD (n = 71) versus antidepressants only (n = 79) | −3409 (−3408 to −3360) 34.1 (33.5 to 34.7) Dominant | −0.018 (−0.019 to −0.017) Dominant | 1.9 92.1 29.9 4.5 0.4 | 7.4 72.9 16.8 3.0 |
| PCT/−AD (n = 59) versus antidepressants only (n = 79) | −3015 (−3084 to −2947) 21.0 (20.2 to 21.7) Dominant | 0.051 (0.050 to 0.053) Dominant | 1.9 92.1 29.9 4.5 0.4 | 7.4 72.9 16.8 3.0 |
regarding a relapse prevention strategy for individuals in (partial) remission using maintenance antidepressants that found family psychoeducation added to treatment as usual was highly likely to be cost-effective compared with treatment as usual over 9 months if a depression-free day would be valued at $20 or more. In both studies, only direct healthcare costs were taken into account.

Karyotaki et al performed a systematic review of economic evaluations alongside randomised controlled trials for depression treatments. Three trials were identified comparing the combination of a psychological intervention and antidepressants versus antidepressants only and inconsistent results were found. Overall, Karyotaki et al concluded that studies likely varied widely in results because of differences in study design and study population and that there remain important gaps in knowledge regarding economic evaluations for MDD treatments. Our study is the first to examine the economic consequences of adding a psychological intervention to maintenance antidepressants in individuals who are recurrently depressed and it shows promising results. More studies are needed to substantiate this finding.

**Antidepressants only compared with PCT while tapering antidepressants**

Health outcomes did not significantly favour antidepressants only compared with PCT/AD, which is in line with our primary outcome. However, the cost-effectiveness plane showed that antidepressants only dominated PCT/AD as most of the bootstrapped ICERs were located in the south-east quadrant where costs are lower and health outcomes better. Antidepressants only also resulted in lower societal costs in the budget impact analysis. These findings are only partly consistent with the findings of two studies examining the (cost)effectiveness of a relapse prevention strategy (mindfulness-based cognitive therapy) with support to taper off maintenance antidepressants compared with maintenance antidepressants only in individuals in remission from recurrent depression. In one study, antidepressants dominated mindfulness-based cognitive therapy with tapering support when the willingness-to-pay threshold was zero per additional recurrence prevented but this reversed when the willingness to pay increased to $1000 and above. Results of the other study also suggested that an improvement in terms of recurrence was achieved at higher costs for mindfulness-based cognitive therapy while tapering antidepressants compared with antidepressants only, but that antidepressants dominated mindfulness-based cognitive therapy while tapering antidepressants in terms of QALYs. The probability that mindfulness-based cognitive therapy was cost-effective compared with antidepressants did not rise above 52% regardless of effect measure.

As also stated in the DRD study, it is important to examine why the results were not better for PCT/AD compared with antidepressants only, whereas the addition of PCT to antidepressants
resulted in additional protection. One explanation might be a temporary imbalance caused by withdrawal of antidepressants. The systematic review of Fava et al\(^43\) suggests that withdrawal symptoms can take many forms and that they usually occur within a few days or weeks but can also persist for a longer period (i.e. up to 1 year). This corroborates the DRD study, where we found a higher risk of recurrence during the first 140 days gradually disappearing thereafter in participants tapering antidepressants with PCT compared with antidepressant only.\(^12\) However, when correcting for this possible destabilisation in the current study, the gain in depression-free days for antidepressants only decreased but costs remained lower and effects slightly better for antidepressants only compared with PCT/−AD. It should be noted that most participants (60%) tapered their antidepressants within 6 months, indicating that the advised 4 weeks is not feasible for many and that possible withdrawal effects may have occurred later during the study.

A second explanation might be that participants had difficulty tapering off antidepressants, resulting in a cycling on and off antidepressants that has shown to generate high costs\(^14\) and might result in progressive tolerance to antidepressants.\(^15\) A third explanation might be that prolonged use of antidepressants results in oppositional tolerance, heightening the recurrence risk after discontinuation.\(^46\)

In the long term, we expect costs and outcomes to be more promising for PCT/−AD compared with antidepressants only as costs of antidepressants will be reduced, and outcomes will be enhanced because of the enduring effects of PCT up to 10 years.\(^4,18\) This is also in line with two cost-effective modelling studies in episodic and maintenance MDD that showed maintenance cognitive–behavioural therapy and antidepressants were both cost-effective strategies over 5 years, with maintenance cognitive–behavioural therapy as a favourable option because of their low costs compared with maintenance antidepressants.\(^47,48\) Future studies are needed to examine at what point in time a break-even point will appear where costs and outcomes will improve for PCT/−AD.

Limitations

Some limitations have to be acknowledged. First, we collected the cost and QALY data via online questionnaires that were not mandatory, resulting in missing data. We handled missing data using multiple imputations. Baseline variables predicted whether data were missing, suggesting data were at least partly missing at random and that multiple imputations may have reduced bias associated with complete case analyses. In addition, our sensitivity analyses suggested that missing data had no substantial influence on the results. However, a possibility remains that missing data were not missing at random. Second, this study was not powered to detect a significant difference in QALYs and costs. Yet, the probabilistic and medical decision-making techniques we used allowed us to make estimations of the cost-effectiveness and cost-utility.
Third, whereas a willingness-to-pay threshold for QALYs is available, this is not the case for depression-free days. Therefore, interpretation of the willingness to pay regarding depression-free days remains speculative. Given that in the current study most outcomes were located in the south-east quadrant where costs were lower and effects better, this limitation does not have a major impact on the implications of this study. Fourth, no pill-placebo control group or control group for PCT was used, which would have enabled examination of the specific effects of antidepressants and PCT. Fifth, the data were collected in the Netherlands and the study comprised individuals in remission from at least two previous major depressive episodes using maintenance antidepressants, which might compromise generalisability. Sixth, the estimated budget impact relies on the assumption that the trial results are generalisable to a larger population within the Netherlands, which might not be the case. Nevertheless, we believe that our results can be generalised as participants were recruited via a wide range of resources and our inclusion and exclusion criteria regarding comorbidity were lenient.

**Clinical implications and recommendations**

For individuals willing to stay on antidepressants, the addition of PCT provides a substantial benefit over antidepressants only in terms of cost-effectiveness and the budget impact. For individuals wishing to taper antidepressants, extra investments might be required. Since preventive effects of psychological interventions last up to 10 years and maintenance antidepressants generate long-term costs, future studies should examine the cost-effectiveness of relapse prevention programmes in maintenance antidepressants over a longer time span than the 24 months of the current study. In addition, future studies should examine how withdrawal symptoms and specific patterns of discontinuing antidepressants are associated with cost-effectiveness and whether PCT should be administered before or after tapering antidepressants.

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**Supplementary material**

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