IMPORTANCE For women with a 20% or more familial risk of breast cancer without a known BRCA1/2 (BRCA1, OMIM 113705; and BRCA2, OMIM 114480) or TP53 (OMIM 151623) variant, screening guidelines vary substantially, and cost-effectiveness analyses are scarce.

OBJECTIVE To assess the cost-effectiveness of magnetic resonance imaging (MRI) screening strategies for women with a 20% or more familial risk for breast cancer without a known BRCA1/2 or TP53 variant.

DESIGN, SETTING, AND PARTICIPANTS In this economic evaluation, conducted from February 1, 2019, to May 25, 2020, microsimulation modeling was used to estimate costs and effectiveness on a lifetime horizon from age 25 years until death of MRI screening among a cohort of 10 million Dutch women with a 20% or more familial risk for breast cancer without a known BRCA1/2 or TP53 variant. A Dutch screening setting was modeled. Most data were obtained from the randomized Familial MRI Screening (FaMRIsc) trial, which included Dutch women aged 30 to 55 years. A health care payer perspective was applied.

INTERVENTIONS Several screening protocols with varying ages and intervals including those of the randomized FaMRIsc trial, consisting of the mammography (Mx) protocol (annual mammography and clinical breast examination) and the MRI protocol (annual MRI and clinical breast examination plus biennial mammography).

MAIN OUTCOMES AND MEASURES Costs, life-years, quality-adjusted life-years (QALYs), and incremental cost-effectiveness ratios (ICERs) were calculated and discounted by 3%. A threshold of €22 000 (US $24 795.87) per QALY was applied.

RESULTS This economic evaluation modeling study estimated that, on a lifetime horizon per 1000 women with the Mx protocol of the FaMRIsc trial, 346 breast cancers would be detected, and 49 women were estimated to die from breast cancer, resulting in 22 885 QALYs and total costs of €7 084 767 (US $7 985 134.61). The MRI protocol resulted in 79 additional QALYs and additional €2 657 266 (US $2 994 964.65). Magnetic resonance imaging performed only every 18 months between the ages of 35 and 60 years followed by the national screening program was considered optimal, with an ICER of €21 380 (US $24 097.08) compared with the previous nondominated strategy in the ranking, when applying the National Institute for Health and Care Excellence threshold. Annual screening alternating MRI and mammography between the ages of 35 and 60 years, followed by the national screening program, gave similar outcomes. Higher thresholds would favor annual MRI screening. The ICER was most sensitive to the unit cost of MRI and the utility value for ductal carcinoma in situ and localized breast cancer.

CONCLUSIONS AND RELEVANCE This study suggests that MRI screening every 18 months between the ages of 35 and 60 years for women with a family history of breast cancer is cost-effective within the National Institute for Health and Care Excellence threshold for all densities. Higher thresholds would favor annual MRI screening. These outcomes support a change of current screening guidelines for this specific risk group and support MRI screening.

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Women with a family history of breast cancer have an increased risk of developing breast cancer, and an increased risk of developing it at a relatively young age.\(^1\) In approximately 64\% to 87\% of these women, no causative hereditary gene variant has been found.\(^2\) Because tumor stage at diagnosis is of importance for survival,\(^3\) screening is advised, but guidelines differ substantially.

The American Cancer Society advises additional magnetic resonance imaging (MRI) for women with a lifetime risk of 20\% or more of developing breast cancer,\(^4\) whereas in the Netherlands and the UK, only mammography screening is advised for women at familial risk without a \textit{BRCA1/2} (\textit{BRCA1}, OMIM 113705; and \textit{BRCA2}, OMIM 114480) variant.\(^5\)\(^6\) All guidelines recommend to start screening among women with a familial risk of breast cancer at a younger age than women at average risk.\(^4\)\(^6\)\(^7\) However, younger women often have dense breast tissue,\(^7\) which is associated with decreased mammographic sensitivity.\(^8\) Magnetic resonance imaging screening has a high sensitivity, not affected by breast density.\(^9\)\(^10\) However, MRI leads to more false-positive results and is associated with higher costs.\(^9\)\(^11\) To our knowledge, little is known about the cost-effectiveness of MRI screening for women with a familial risk of breast cancer; one previous study evaluated the cost-effectiveness of MRI screening in women at familial risk of breast cancer without a known gene variant, showing by microsimulation modeling that MRI screening was very costly.\(^11\)

The model was based on data from a nonrandomized study.

Recently, the randomized Familial MRI Screening (FaMRIs) trial showed higher breast cancer detection rates and detection of breast cancer at, on average, an earlier stage when screening with MRI in comparison with mammography in women at increased familial risk without a known \textit{BRCA} or \textit{TP53} (OMIM 151623) variant.\(^9\) In this study, we calculate real-life costs of MRI and mammography in the FaMRIs trial. We estimate the cost-effectiveness by microsimulation modeling, and compare different screening scenarios by varying starting and stopping ages, screening intervals, and combinations of MRI and mammography.

**Methods**

**The FaMRIs Trial**

In the multicenter randomized clinical FaMRIs trial, Dutch women aged 30 to 55 years with a cumulative lifetime breast cancer risk of 20\% or more due to a family history of breast cancer without a known \textit{BRCA1/2} or \textit{TP53} variant were randomly assigned into 2 screening groups after providing written informed consent (trial protocol in Supplement 1). The MRI group received annual MRI plus clinical breast examination (CBE), and mammography every 2 years. The mammography (Mx) group received annual mammography and CBE, in accordance with the Dutch screening protocol.\(^9\) Women refusing randomization could participate in a registration group (Reg-MRI group or Reg-Mx group) by providing consent for registration of their screening results. More details have been described elsewhere.\(^9\)\(^12\) The FaMRIs Study follows the Declaration of Helsinki\(^13\) and was approved by the Erasmus University Medical Center Institutional Review Board (Rotterdam, the Netherlands; reference MEC-2010-292). The FaMRIs trial is registered with the Netherlands Trial Register NL2661.

**The Microsimulation Screening Analysis Model**

In this economic evaluation, conducted from February 1, 2019, to May 25, 2020, we used the Microsimulation Screening Analysis (MISCAN) model, which simulates individual natural histories from birth to death and the natural history of breast cancer. We adjusted the version by Sankatsing et al\(^14\) to extrapolate the findings of the FaMRIs trial. To be able to model the difference in the numbers of detected ductal carcinoma in situ (DCIS) and T1a and T1b tumors between the 2 study groups,\(^9\) 2 additional preclinical states were added to the original MISCAN model: DCIS, \textit{MRI} and T1a/T1b, \textit{MRI} (eFigure 1 in Supplement 2). We assumed that DCIS and T1a and T1b tumors could for some time be detected only by MRI before they could also be detected by mammography or before they become clinically detectable. During all other preclinical states, the tumor could be detected with MRI as well as mammography or clinically diagnosed. Progression through the health states was modeled as a semi-Markov process. The model only takes into account first breast cancers and no contralateral breast cancers.

We assumed the mammographic sensitivity to be 15\% lower than previously used in the model owing to the younger population we modeled.\(^15\) We assumed that CBE would not lead to additional cancer detection.\(^16\) Incidence, dwelling times, stage-specific sensitivities of MRI, and transition probabilities of the additional health state DCIS, \textit{MRI} to DCIS and to T1a/T1b, \textit{MRI} were estimated by calibration using the Nelder-Mead simplex optimization method.\(^17\) We used data from all trial groups (Mx group + Reg-Mx group and MRI group + Reg-MRI group) to increase the amount of data for calibration. Model predictions were calibrated to the number of screening-detected breast cancers per T stage, the number of interval cancers, the number of detected cancers per 10-year age groups, and the number of screening-detected tumors during incident and prevalent rounds, all stratified by screening protocol as observed during the FaMRIs trial. We aimed for all pre-
dicted numbers to fall within 95% Poisson CIs of the observed numbers of tumors.

Probabilities of (false) positive results and diagnostic procedures were obtained from the FaMRIsc trial, stratified by screening modality and by age (<50 and ≥50 years). Both true-positive and false-positive results were associated with diagnostic follow-up and associated costs. For the screening period within the national breast cancer screening program, we applied the same probabilities as for the Mx protocol.

For the situation without screening, we assumed all women with a diagnosed breast cancer would undergo a diagnostic mammogram, CBE, biopsy, or fine needle aspiration, and all women with a diagnosed T2 or higher tumor would undergo an MRI. The percentage of ultrasonographic evaluations performed in diagnosed cases in a situation without screening was assumed to be equal to the percentage of those performed among women with a diagnosed breast cancer within the Mx protocol.

**Screening Strategies**

After calibration, we applied several screening strategies, varying in starting and stopping ages, intervals, and screening modalities. With stopping ages below the age of 75 years, we modeled the women to continue screening within the national screening program until the age of 75 years, consisting of biennial mammography at a local screening unit. Attendance rates were set at 100%.

**Costs**

We applied a health care perspective and considered only direct medical costs (converted to 2018 amounts; eTable 1 in Supplement 2) and costs related to other causes of death. Costs of MRI, mammography in a hospital setting, and ultrasonography were derived from the mean of all published prices.18 The price of mammography in a local screening unit was obtained from the Netherlands Comprehensive Cancer Organisation.19 All other costs were obtained from a study by Saadatmand et al.11 Costs of fine needle aspiration and biopsy were updated and adjusted by adding costs of pathologic examination of the specimen, obtained from the tariff tool.18 Costs of breast-conserving surgery and mastectomy were adjusted assuming 1.5 consecutive hospital days with its price obtained from the Dutch costing manual.20 Costs associated with breast cancer death were assumed to be €19 679 (US $22 179.91) and death due to other causes were assumed to be €15 044 (US $16 955.87).21

We multiplied costs with the resource used during the trial to calculate real-life costs. Mean treatment costs per TN stage were calculated by dividing total treatment costs per TN stage by the number of cancers. Model outcomes were multiplied with aforementioned prices to calculate costs per screening protocol.

**Health State Utilities**

Utility values were obtained from the literature (eTable 2 in Supplement 2). The utility value for the healthy state was based on a study by Versteegh et al.22 Early-stage cancer was associated with disutility of 10%, regional cancer was associated with disutility of 25%, and metastasis was associated with disutility of 40%. A disutility of 0.105 was applied for a positive screening result with a duration of 5 weeks. We did not apply a disutility for screening visits.25

**Statistical Analysis**

We simulated the number of invitations, screening visits, screening-detected cancers, interval cancers, life-years, quality-adjusted life-years (QALYs), deaths from breast cancer, and deaths from other causes, all on a lifetime horizon from age 25 years for a cohort of 10 million Dutch women born in 1980. All results were scaled to 1000 women. Overdiagnosis was defined as detected cancers that would not have been diagnosed in a woman's lifetime in a situation without screening. Incremental cost-effectiveness ratios (ICERs) were calculated by dividing incremental costs by incremental QALYs. We plotted an efficiency frontier representing efficient strategies that are either less costly and more effective, or more costly but more cost-effective than those below the frontier. A cost-effectiveness threshold was based on the National Institute for Health and Care Excellence (NICE) threshold of £20 000 (€22 000 [US $24 795.87]). Average cost-effectiveness ratios were calculated by dividing additional costs by additional QALYs compared with a situation without screening. Costs and effects were discounted by 3%.26

One-way sensitivity analyses were performed for utility values, the price of MRI, and false-positive rates to analyze the association of these parameters with the ICER of the MRI protocol vs the Mx protocol. Utility values were varied ±10% of the base case values and the other parameters were varied ±20% of the base case values. Sensitivity analyses were discounted by 3%.

Five scenario analyses were performed to quantify methodological uncertainty. First, we applied discount rates of 4.0% for costs and 1.5% for effects, according to Dutch guidelines.20 Second, we calculated the ICER without discounting. Third, we applied utility values based on a study by Lidgren et al27 (eTable 3 in Supplement 2). Fourth, we calculated the ICER without costs related to death from other causes. Fifth, we applied a disutility of 0.006 for 1 week for screening participation.24 Scenario analyses were performed for the comparison of the 2 screening protocols of the FaMRIsc trial.

We calculated the risk of radiation-induced breast cancers for an optimal screening strategy with mammography compared with a strategy without mammography. We used the excess absolute risk model28,29 with a glandular dose of a 2-view mammogram of 4.4 mGy.

**Results**

**Real-Life Results During the FaMRIsc Trial**

After a mean follow-up of 4.3 years, 41 tumors were detected in the MRI group, whereas 15 tumors were detected in the Mx group.9 Table 19 shows the number of detected tumors, woman-years at risk, and real-life screening costs according to group, age, and density during the FaMRIsc trial. The MRI protocol
resulted in approximately 2 times higher costs of screening and additional investigation. Mean treatment costs are shown in eTable 4 in Supplement 2.

Model Calibration Results

eFigure 2 in Supplement 2 shows the number of observed breast cancers during the FaMRIsc trial according to T stage and the number of predicted cancers by our calibrated model. All predicted numbers were within the 95% CIs of the observed numbers.

Cost-effectiveness Results

Table 2 and eTable 5 in Supplement 2 display the outcomes of all modeled strategies per 1000 women. With the Mx protocol of the FaMRIsc trial (strategy M), 346 breast cancers would be detected, and 49 women would die from breast cancer, resulting in 22 885 QALYs (discounted by 3%) and total costs of €7 084 767 (US$7 985 134.61) (discounted by 3%) and total costs of €23 497 356 (US$26 483 517.49) (undiscounted). With the MRI protocol of the FaMRIsc trial (strategy V), 377 breast cancers would be detected and 30 breast cancer deaths would occur, resulting in 79 additional QALYs (discounted by 3%) and additional costs of €2 657 266 (US$2 994 964.65) (discounted by 3%) and total costs of €28 024 674 (US$31 866 189.70) (undiscounted). Comparing these 2 protocols resulted in an ICER of €32 277 (US$37 506.01) per QALY gained (discounted).

Both screening protocols of the FaMRIsc trial were dominated by similar screening strategies without CBE (strategy B and U) (Figure 1). Strategies involving MRI resulted in fewer breast cancer deaths, lower numbers of interval cancers, and lower total treatment costs but more overdiagnosed cancers, compared with screening without MRI. The 2 strategies with intervals of 18 months were both on the efficiency frontier (Figure 1). Most strategies on the efficiency frontier consisted of screening from age 35 until 60 years, continued within the national screening program. Switching to screening within the national screening program before age 60 years resulted in higher numbers of clinically diagnosed cancers and breast cancer deaths, and were therefore dominated (eTable 5 in Supplement 2).

Strategy D, consisting of MRI screening every 18 months between ages of 35 and 60 years followed by the national screening program had the highest acceptable ICER, €21 380 (US$24 002.36), when applying the NICE threshold of £20 000 ($22 000 [US$24 795.87]) and was considered optimal. Strategy E, consisting of alternating annual MRI or mammography between the ages of 35 and 60 years, was almost on the efficiency frontier. The effects of this strategy were similar to those of strategy D for somewhat higher cost. Strategies D and E, both followed by screening within the national breast cancer screening program, resulted in a reduction of 98 and 99 breast cancer deaths, respectively, and 65 or 66 overdiagnosed cases, respectively, when compared with a situation without screening.

Sensitivity and Scenario Analyses

Results of the deterministic sensitivity analyses are shown in Figure 2. The ICER was most sensitive to the price of MRI screening and the utility value for DCIS or localized breast cancer.

When applying Dutch discount rates, the ICER of the MRI protocol vs the Mx protocol became lower: €13 108 (US$14 773.83) per QALY gained. The difference in life-years and QALYs between the 2 protocols were 176 and 170, respectively, and the difference in costs was €2 234 665 (US$2 518 657.40). Without discounting, the ICER was €12 376 (US$13 948.80).

In the third scenario analysis, in which we applied a different set of utility values, the difference in QALYs between the MRI protocol and Mx protocol became 71, which was lower compared with the base case. Consequently, the ICER became

<table>
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<th>Characteristic</th>
<th>No. of breast cancers*</th>
<th>Life-years at risk</th>
<th>Costs, € (US $)</th>
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<td>&lt;50 y</td>
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<td>2106</td>
<td>740 188 (834 254.79)</td>
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<td>Total</td>
<td>41*</td>
<td>3218</td>
<td>1 097 766 (1 237 275.59)</td>
<td>230 335 (259 607.12)</td>
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<td>Mx group by age</td>
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<td>7</td>
<td>1215</td>
<td>178 692 (201 401.07)</td>
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<td>3314</td>
<td>520 260 (586 377.24)</td>
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<td>BI-RADS density A-C (0%-75%)</td>
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<td>2743</td>
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<td>176 580 (199 020.67)</td>
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<td>Total</td>
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<td>3249</td>
<td>1 116 397 (1 258 274.31)</td>
<td>236 754 (266 841.88)</td>
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<td>BI-RADS A-C</td>
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<td>105 386 (118 778.98)</td>
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<td>4308</td>
<td>672 531 (757 999.60)</td>
<td>162 635 (183 303.47)</td>
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Abbreviations: BI-RADS, Breast Imaging Reporting and Data System Atlas; FaMRIsc, Familial MRI Screening; MRI, magnetic resonance imaging; Mx, mammography.

* Breast cancers include invasive breast cancers and ductal carcinoma in situ.

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| Abbreviations: ACER, average cost-effectiveness ratio (comparison of a strategy with a situation without screening); FaMRisc, Familial MRI Screening; ICER, incremental cost-effectiveness ratio (comparison of a strategy to the previous nonrandomized strategy in the ranking); LYs, life-years; MRI, magnetic resonance imaging; Mx, mammography; NA, not applicable; QALYs, quality-adjusted life-years.
| a Breast cancers include invasive breast cancers and ductal carcinoma in situ. Results are without discounting. Outcomes contain the effects of both the described strategy and the subsequent national breast cancer screening program.
| b Annual mammography between 40 and 60 years.
| c Alternating annual MRI or mammography between 35 and 60 years.
| d Alternating MRI or mammography every 18 months between 35 and 60 years.

Abbreviations: ACER, average cost-effectiveness ratio (comparison of a strategy with a situation without screening); FaMRisc, Familial MRI Screening; ICER, incremental cost-effectiveness ratio (comparison of a strategy to the previous nonrandomized strategy in the ranking); LYs, life-years; MRI, magnetic resonance imaging; Mx, mammography; NA, not applicable; QALYs, quality-adjusted life-years.
larger: €37 489 (US $42 253.29) per QALY gained (discounted). When not applying costs related to death from other causes, the ICER became €32 712 (US $36 869.20) per QALY gained (discounted), which was similar to the ICER when including these costs. Applying a utility decrement for screening participation hardly affected the ICER, which became €33 534 (US $37 795.67) (discounted).

Radiation Risk
In a situation with additional mammography to the optimal screening strategy (D) consisting of MRI every 18 months between the ages of 35 and 60 years, radiation would induce 0.94 breast cancers and 0.12 breast cancer deaths per 1000 women. In this situation, 3 additional breast cancers would be detected by screening of which 1 would be overdiagnosed, and 2 breast cancer deaths would be prevented (undiscounted) compared with a strategy without additional mammography (strategy D).

Discussion
This economic modeling study of data on Dutch women showed that the detection of more tumors at an early stage and fewer at a late stage by MRI⁹ could be a cost-effective method to reduce breast cancer mortality despite more over-diagnosis and higher costs in comparison with mammography. Yearly MRI seems to bring the largest mortality reduction, but for an ICER higher than allowed by NICE guidelines.³⁰ Neither protocol of the FaMRIsc trial was on the efficiency frontier, mainly owing to the addition of CBE that proved to be inefficient.⁹,¹⁶ Screening with MRI only every 18 months between the ages of 35 and 60 years would be an efficient and cost-effective strategy, with an ICER just below the threshold of £20 000 (€22 000 [US $24 795.87]). We also found that the additional association of mammography with this strategy was limited. Screening consisting of alternating annual MRI and mammography between ages of 35 and 60 years, followed by screening within the national screening program until the age of 75 years was almost on the frontier, with similar effects and more costs as the previously mentioned strategy (MRI only every 18 months between the ages of 35 and 60 years). Most of the efficient strategies consisted of screening from 35 to 60 years of age, with continuation of screening within the national screening program. Furthermore, our
results indicated that the switch to the national screening program should not take place before 60 years of age.

We modeled a Dutch health care setting but we expect the relative difference in health outcomes between our modeled strategies to be similar in other countries. In contrast, unit prices as well as cost-effectiveness thresholds vary substantially per country, which should be taken into account when generalizing our results to other countries.

We simulated one group of women with, on average, the same risk of breast cancer. However, starting screening at 35 years of age may not be beneficial for all women within this group, depending on the youngest age of breast cancer diagnosis of a family member and their individually calculated lifetime risk. Therefore, family history should be taken into account when choosing the starting age for screening.

To our knowledge, one previous study evaluated the cost-effectiveness of additional MRI screening for this group of women. Saadatmand et al calibrated the MISCAN model on data from the 1999-2006 MRI Screening (MRISC) study. The breast cancer incidence in the FaMRIsc Study was higher than that in the MRISC study, and the sensitivity of both MRI and mammography were also higher.

**Strengths and Limitations**

This study has some strengths, including the use of randomized clinical trial data for calibration, which has, to our knowledge, not been done before for this group of women. By using randomized clinical trial data, the model gets more information on the performance of MRI and mammography separately than when these screening modalities are performed simultaneously.

This study also has some limitations. First, the study sample of the FaMRIsc trial was still quite small for calibration. The number of observed cancers stratified by group and stage were small and therefore 95% CIs were large. Therefore, we added the data of the registration groups. However, there may have been a difference in population between women registered and those randomized. A second limitation is the assumption that there is no DCIS that is detectable only by mammography. Third, we were unable to model strategies by breast density categories as the numbers by breast density in the FaMRIsc trial were too small, albeit the associations of MRI with detection seem similar across density categories. A recent study showed the benefit of MRI screening in women with extremely dense breasts. Fourth, we did not measure utility values within our study population. Utility values related to breast cancer vary significantly in the literature and we are aware of the association of these values with the results, as shown in our analyses. Furthermore, we would like to point out the uncertainty of efficiency frontiers as such. Efficiency frontiers are sensitive to changes in underlying data and assumptions, and they do not display uncertainty.

Downsides of MRI are its high costs, more false-positive results, and increased overdiagnosis. Overdiagnosis may be a result of excessive detection of low-grade tumors, but our model cannot distinguish between low-grade or high-grade tumors. Overdiagnosis is captured in our results and the same (dis)utility values were applied to all modeled breast cancer cases because one does not know whether a cancer is overdiagnosed or not.

Applying MRI screening may have some practical implications. Hospitals need to have enough capacity for the screening and for additional diagnostic testing due to more (false) positive results, to prevent waiting lists. In addition, radiologists may need additional training to guarantee good quality, as MRI screening requires expertise.

Currently, abbreviated MRI seems promising, which has shorter acquisition time and reading time while maintaining diagnostic accuracy. A less time-consuming MRI will decrease the price of the test, which has a favorable association with the ICER, as shown in our sensitivity analyses.

**Conclusions**

Based on this cost-effectiveness analysis, MRI screening every 18 months or alternating annual MRI and mammography between the ages of 35 and 60 years may be recommended for women at increased familial risk of breast cancer, both followed by screening within the national screening program, when applying the NICE threshold. Annual MRI was associated with the largest mortality reduction, but for an ICER higher than allowed by NICE guidelines.
Acquisition, analysis, or interpretation of data: Geuzinge, Obdeijn, Rutgers, Saadatmand, Mann, Oosterwijk, Tollenaar, de Roy van Zuidewijn, Lobbes, van’t Riet, Ausems, Loo, Wesseling, Zonderland, Verhoef, Heijnsdijk, Tilanus-Linthorst, de Koning.

Drafting of the manuscript: Geuzinge, Saadatmand, Tollenaar, Lobbes, de Koning.

Critical revision of the manuscript for important intellectual content: Obdeijn, Rutgers, Saadatmand, Mann, Oosterwijk, Tollenaar, de Roy van Zuidewijn, Lobbes, van’t Riet, Hooning, Ausems, Loo, Wesseling, Luiten, Zonderland, Verhoef, Heijnsdijk, Tilanus-Linthorst, de Koning.

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Supervision: Rutgers, Saadatmand, Oosterwijk, Tollenaar, Lobbes, Luiten, Tilanus-Linthorst, de Koning.

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