Trade-Offs Between Harms and Benefits of Different Breast Cancer Screening Intervals Among Low-Risk Women

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Abstract

Background: A paucity of research addresses breast cancer screening strategies for women at lower-than-average breast cancer risk. The aim of this study was to examine screening harms and benefits among women aged 50-74 years at lower-than-average breast cancer risk by breast density. Methods: Three well-established, validated Cancer Intervention and Surveillance Network models were used to estimate the lifetime benefits and harms of different screening scenarios, varying by screening interval (biennial, triennial). Breast cancer deaths averted, life-years and quality-adjusted life-years gained, false-positives, unnecessary biopsies, and the percentage overdiagnosis were assessed by relative risk (RR) level (0.6, 0.7, 0.85, 1 [average risk]) and breast density category, for US women born in 1970. Results: Screening benefits decreased proportionally with decreasing risk and with lower breast density. False-positives, unnecessary biopsies, and the percentage overdiagnosis also varied substantially by breast density category; false-positives and unnecessary biopsies were highest in the heterogeneously dense category. For women with fatty or scattered fibroglandular breast density and a relative risk of no more than 0.85, the additional deaths averted and life-years gained were small with biennial vs triennial screening. For these groups, undergoing 4 additional screens (screening biennially [13 screens] vs triennially [9 screens]) averted no more than 1 additional breast cancer death and gained no more than 16 life-years and no more than 10 quality-adjusted life-years per 1000 women but resulted in up to 232 more false-positives per 1000 women. Conclusion: Triennial screening from age 50 to 74 years may be a reasonable screening strategy for women with lower-than-average breast cancer risk and fatty or scattered fibroglandular breast density.
Table 1. Summary of model features

| Feature | Model E | Model GE | Model W |
|---------|---------|----------|---------|
| Natural history of cancer | Continuous tumor growth | Stage transition | Continuous tumor growth |
| Details on natural history | Variation in growth rates, includes slow- and fast-growing tumors with varying fatal diameters | All lesions begin as DCIS and can evolve through AJCC-6 stages; variation in dwell times in each stage | Variation in growth rates from nonprogressive disease to hyperaggressive tumors |
| Tumors obligated to progress | DCIS nonobligate; invasive obligate | DCIS nonobligate; invasive obligate | DCIS and some small invasive are nonobligate; larger invasive obligate |
| SEER breast cancer data used for model calibration (1975-2010) | Incidence, stage distribution, mortality | Incidence, stage distribution | Incidence and mortality |
| Screen detection conditioned on | Tumor size, modality, age, density, frequency | Modality, age, density, frequency | Tumor size, modality, age, density, frequency |
| Implementation of screening benefit | Smaller tumor size | Younger age and earlier stage | Younger age and smaller tumor size |
| Estimation of overdiagnosis b | Difference screen and no screen | Difference screen and no screen | Difference screen and no screen |
| Implementation of treatment benefit | Cure fraction based on fatal diameter | Hazard reduction | Cure fraction |
| Factors affecting treatment benefit | ER and HER2; age; year of and size at diagnosis | ER and HER2; age; year of and stage at diagnosis | ER and HER2; age; year of and stage at diagnosis |
| Model software program c | Delphi | Schechter et al., 2018 (16) | Alagoz et al., 2018 (17) |
| Detailed model description | van den Broek et al., 2018 (15) | | |

*Adapted from (6). Additional information is available from (18), and at https://resources.cisnet.cancer.gov/registry/site-summary/breast/. AJCC – American Joint Committee on Cancer; DCIS – ductal carcinoma in situ; ER – estrogen receptor.

bOverdiagnosis was defined as screen-detected cancer that would not have been diagnosed in a woman’s lifetime in the absence of screening.

cCombined output from all 3 models was analyzed using SAS (Cary, NC) version 9.4.

be considered. Few studies have assessed the harms and benefits for women at decreased risk (lower than average, i.e., a relative risk [RR] < 1). Women with lower-than-average risk of breast cancer are expected to have a less favorable harm to benefit ratio from untargeted screening, suggesting that less intense screening strategies than biennial screening might be appropriate for this group.

The proportion of women at low risk in the population is substantial; for example, 34% of US women aged 40-74 years have a 5-year risk of developing breast cancer below 1.00% based on the Breast Cancer Surveillance Consortium (BCSC) risk model (8,9). Established factors that are associated with substantially decreased risk for breast cancer include fatty breasts, young age at first birth (younger than 20 years), and young age at menopause (younger than 40 years) with relative risks of 0.6-0.7 (10-12); these factors apply to 8%, 12%, and 13% of US women, respectively (10,11). Factors associated with a more modest decrease in risk, such as 3 or 4 full pregnancies (RR = 0.84) and age at menopause between 45 and 49 years (RR = 0.86) (12,13), are even more common, with 39% and 24% of US women aged 50-79 years reporting those factors, respectively (11).

Breast density has also received attention as an important factor that influences risk of developing breast cancer, as well as affecting the balance between benefits and harms of screening, because low breast density not only leads to a reduced risk for developing disease but also increases the sensitivity of mammography (8,10,14). The aim of this study was to assess the benefits and harms of screening by breast cancer risk, breast density, and screening interval among women aged 50-74 years with lower-than-average risk levels using collaborative modeling. Study results are intended to inform discussions about risk-based screening guidelines and practice.
specific prevalence in the BCSC. Density affected mammography performance (32), whereas mammography performance was assumed to be unaffected by risk. Risk associated with density (Table 3) was combined multiplicatively with the risk of the different risk levels (relative risks). In this way, density and other risk factors were assumed to be independent determinants of breast cancer risk, consistent with observed data. Thus, a 50-year-old woman with heterogeneously dense breasts and a relative risk of 0.7 had a relative risk of 0.875 (0.7*1.25). In each simulation, women were followed until death or a model-specific upper age of 100 or 120 years. To evaluate the efficacy of different screening scenarios, we assumed 100% uptake of screening and treatment. We modeled biennial screening between ages 50 and 74 years (13 screens) and triennial screening between ages 50 and 74 years (9 screens).

### Screening Outcomes

For all screening scenarios, we estimated outcomes per 1000 women alive at age 50 years, including the number of ductal carcinoma in situ (DCIS) and invasive breast cancers detected. Benefits included breast cancer deaths averted, life-years gained, and quality-adjusted life-years (QALYs) gained. To calculate QALYs, we applied health-related quality-of-life utilities by age (33), and we applied quality-of-life decrements by attaching weights to specific health states for women undergoing a mammogram and diagnostics (34) and life-years with breast cancer by stage of disease at diagnosis (35). Harms included overdiagnosis, false-positives, and benign biopsies. Overdiagnosis was defined as screen-detected cancer that would not have been diagnosed in a woman’s lifetime in the absence of screening. In addition, harm to benefit ratios (false-positives per life-year gained and overdiagnosis per breast cancer death averted) were calculated.

### Analysis

We presented all outcomes by subgroups of risk and density for each strategy using the median (minimum, maximum) of the 3 models. Each outcome was compared with a reference value, defined as the model-specific results for biennial screening from age 50 to 74 years, all densities combined (thus, with representative population frequencies of breast density categories), and average risk (RR = 1). We evaluated the differences between screening scenarios by assessing the incremental benefits and incremental harms by dividing the incremental harm by the incremental benefits.

We performed sensitivity analyses on varying utility values for undergoing screening and additional workup and on varying specificity by risk (9) (Supplementary Methods, available online).

### Results

#### Screening Outcomes

Among 1000 women aged 50 years followed over their lifetimes, the number of invasive breast cancers detected when screening biennially between ages 50 and 74 years varied substantially by subgroup; the highest number of invasive breast cancers was a median of 150 (range across models = 150-177) detected in the average-risk (RR = 1) extremely dense group and decreased with decreasing risk and density in all 3 models to 39 (range = 33-52) in the lowest risk-density category (ie, RR = 0.6 and almost...
Table 4. Number of ductal carcinoma in situ (DCIS), invasive breast cancers detected, lifetime benefits, and lifetime harms for biennial screening between ages 50 and 74 years per 1000 women followed over their lifetimes across models

| Breast density at age 50 years | Relative risk | No. of DCIS detected, median (min, max) | No. of invasive BCs detected, median (min, max) | Lifetime benefits, median (min, max) | Lifetime harms, median (min, max) |
|--------------------------------|---------------|----------------------------------------|-----------------------------------------------|--------------------------------------|-----------------------------------|
|                                |               | No. of BC deaths averted | Life-years gained | QALYs gained | False-positives | Biopsies | Overdiagnosis |
| Almost entirely fatty | 0.60 | 16 (11, 24) | 39 (33, 52) | 2.5 (1.6, 2.7) | 40 (38, 44) | 25 (22, 26) | 665 (623, 824) | 102 (92, 128) | 12 (8, 19) |
|                                | 0.70 | 18 (16, 27) | 45 (39, 77) | 2.9 (1.9, 3.1) | 46 (44, 52) | 30 (27, 30) | 663 (620, 821) | 101 (92, 127) | 14 (9, 22) |
|                                | 0.85 | 22 (20, 32) | 54 (47, 101) | 3.5 (2.3, 3.7) | 55 (53, 62) | 38 (33, 38) | 659 (617, 816) | 101 (91, 126) | 17 (11, 26) |
|                                | 1.00 | 25 (22, 37) | 63 (55, 117) | 4.1 (2.7, 4.3) | 64 (62, 73) | 45 (40, 46) | 656 (613, 811) | 100 (91, 126) | 20 (12, 29) |
| Scattered fibroglandular density | 0.60 | 22 (12, 27) | 60 (58, 64) | 3.3 (2.6, 4.3) | 60 (48, 76) | 36 (29, 47) | 1088 (1018, 1267) | 166 (151, 197) | 17 (11, 18) |
|                                | 0.70 | 25 (18, 30) | 74 (67, 89) | 3.8 (3.0, 4.9) | 70 (56, 88) | 43 (35, 56) | 1083 (1011, 1260) | 166 (150, 196) | 20 (12, 21) |
|                                | 0.85 | 31 (23, 36) | 89 (81, 116) | 4.5 (3.6, 6.0) | 84 (67, 106) | 54 (44, 70) | 1074 (1001,1249) | 164 (148, 194) | 24 (15, 24) |
|                                | 1.00 | 36 (25, 41) | 103 (94, 133) | 5.2 (4.2, 7.0) | 98 (77, 124) | 65 (52, 84) | 1065 (991, 1238) | 163 (147, 192) | 27 (17, 27) |
| Heterogeneously dense | 0.60 | 21 (15, 28) | 75 (72, 88) | 4.0 (3.0, 5.2) | 68 (60, 93) | 40 (36, 59) | 1297 (1213, 1495) | 198 (180, 232) | 17 (13, 18) |
|                                | 0.70 | 24 (21, 32) | 102 (87, 106) | 4.6 (3.4, 6.0) | 79 (69, 108) | 49 (44, 71) | 1288 (1202, 1484) | 197 (178, 230) | 19 (15, 20) |
|                                | 0.85 | 29 (26, 38) | 121 (104, 136) | 5.5 (4.2, 7.2) | 95 (82, 130) | 61 (54, 88) | 1274 (1186, 1468) | 195 (176, 228) | 23 (18, 23) |
|                                | 1.00 | 33 (29, 43) | 140 (120, 156) | 6.3 (4.9, 8.4) | 111 (94, 152) | 73 (64, 106) | 1260 (1170, 1452) | 193 (174, 226) | 26 (20, 26) |
| Extremely dense | 0.60 | 18 (17, 30) | 94 (83, 95) | 4.2 (2.9, 6.3) | 65 (63, 113) | 41 (40, 76) | 1023 (961, 1392) | 156 (142, 212) | 15 (14, 17) |
|                                | 0.70 | 24 (21, 34) | 109 (108, 121) | 4.9 (3.3, 7.2) | 75 (73, 130) | 49 (48, 90) | 1014 (952, 1379) | 155 (141, 210) | 17 (15, 19) |
|                                | 0.85 | 30 (26, 40) | 130 (129, 155) | 5.7 (4.0, 8.7) | 90 (86, 157) | 60 (60, 111) | 1001 (938, 1360) | 153 (139, 208) | 20 (18, 22) |
|                                | 1.00 | 33 (30, 45) | 150 (150, 177) | 6.5 (4.7, 10.1) | 106 (98, 182) | 72 (70, 131) | 988 (924, 1342) | 151 (137, 205) | 22 (21, 24) |

*BC = breast cancer; QALYs = quality-adjusted life-years.*
entirely fatty breasts) (Table 4). The trends in lifetime benefits and harms are shown for 1 exemplar model (Figure 1).

### Benefits

The absolute numbers of lifetime benefits decreased with decreasing risk and with decreasing density in all 3 models. For women with lower-than-average risk and fatty breasts, screening led to fewer benefits (breast cancer deaths averted and life-years gained) than for women at average risk and/or with denser breasts (Table 4; Supplementary Table 1, available online). For example, per 1000 women followed over their lifetime, biennial screening from age 50 to 74 years gained 40 (range = 38-44) life-years in low-risk women (RR = 0.6) with fatty breasts, whereas the same strategy gained a median of 64 (range = 62-73) life-years in average-risk women (RR = 1) with fatty breasts and a median of 106 (range = 98-182) in average-risk women (RR = 1) with extremely dense breast (Table 4). The finding that benefits decreased with decreasing risk (approximately linearly) was consistent across models, screening scenarios, density categories, and outcomes (breast cancer deaths averted, life-years gained, QALYs gained). Absolute benefits also increased with increasing density consistently across models, screening scenarios, and risk groups, although the increase was not linear and showed a leveling off for the highest density category (Table 4; Supplementary Table 1, available online). Biennial screening scenarios resulted in more benefits and triennial screening scenarios in all models and for all risk and density subgroups (Figure 1).

### Harms

The number of false-positives were relatively stable over risk given our model assumptions (Table 4; Supplementary Table 2, available online), whereas the number of overdiges increased with decreasing risk (Figure 2). The number of false-positives was highest in breast density category C (heterogeneously dense) (Figure 1). The same trend was found for the number of benign biopsies (Figure 1). The relationship between overdiagnosis and density varied across models: in model E, overdiagnosis increased with increasing density; in model W, overdiagnosis was highest in the 2 middle categories; and in model GE, overdiagnosis slightly decreased with increasing density (Figure 2). When overdiagnosis was expressed as a percentage of all breast cancers detected, the percentage decreased consistently in all models with increasing density from 22.7% (range = 12.1%-31.9%) to 11.6%...
Harm to Benefit Ratios

The ratio between harms and benefits showed diversity across models and measures (Supplementary Table 3, available online). All models predicted a decrease in the number of false-positives per life-year gained with increasing risk and somewhat fewer false-positives per life-year gained in the extremely dense category.

Figure 2. Number of overdiagnosed women per 1000 women aged 40 years followed over their lifetime by density, relative risk (RR), screening scenario, and model: model I (upper part), model GE (middle part), and model W (lower part). Biennial (diamonds): biennial screening between ages 50 and 74 years (13 screens). Triennial (triangles): and triennial screening between ages 50 and 74 years (9 screens). Reference (dotted horizontal line) shows the model-specific values for biennial screening from age 50 to 74 years, all densities combined, average risk (RR = 1).

(range =10.6%-12.5%) for a relative risk of 1 and did not vary by risk.

Screening Scenarios (Biennial vs Triennial)

Biennial vs triennial screening has fewer benefits for the low-risk and low-density subgroups than for average-risk women (Table 5). The additional number of breast cancer deaths averted per 1000 women is 0.4 (range = 0.3-0.6) in women at lowest risk (RR = 0.6) with fatty breasts and 0.6 (range = 0.5-0.7) in women at lowest risk (RR = 0.6) with scattered fibroglandular densities with biennial vs triennial screening. For women with fatty or scattered fibroglandular breast density and a relative risk of 0.6, 0.7, or 0.85, screening biennially (13 screens) vs screening triennially (9 screens) averted less than 1 additional breast cancer death and gained at most 16 life-years and 10 QALYs. For average-risk women with extremely dense breasts, there were 1.5 (range = 1.2-1.5) additional deaths averted, 28 life-years gained, and 19 QALYs gained with biennial vs triennial screening (Table 5).

The number of additional false-positives was highest for the heterogeneously dense category, lowest for the almost entirely fatty category, and did not vary much by risk. For women with fatty or scattered fibroglandular breast density and a relative risk of no more than 0.85, there were up to 232 additional false-positives per 1000 women (Table 5). There were more additional false-positives per additional life-year gained among the low-risk groups, and this ratio decreased with increasing risk in all models (Table 5). The number of additional overdiaagnoses per breast cancer death averted decreased in 2 of the 3 models by risk and density (Table 5). The number of additional screens per additional life-year gained when going from triennial to biennial screening increased with decreasing risk and density consistently across models. In average risk women (RR = 1) with extremely dense breasts, models predicted that 120 (range = 120-145) additional screens were needed to gain 1 life-year when going from triennial to biennial screening, whereas in women at lowest risk (RR = 0.6) with fatty breasts, models predicted a substantially higher number of additional screens needed to gain 1 life-year: 409 (range = 373-644) (Table 5).

Sensitivity Analysis

Varying utility values for undergoing screening and additional workup or varying specificity by risk did not majorly change the ranking and differences between subgroups (Supplementary Tables 4 and 5, available online).

Discussion

This is the first collaborative modeling study of breast cancer screening strategies for women at lower-than-average risk, while considering breast density in this assessment. The results indicate that triennial screening from age 50 to 74 years should be considered for women at lower-than-average risk with low density, because this strategy reduces harms while maintaining a large part of the benefits. This conclusion was robust across models and assumptions about disutility associated with screening and variations in specificity by risk.
**Table 5.** The incremental number of breast cancer deaths averted, life-years gained, quality-adjusted life-years (QALYs) gained, false-positives, additional biopsies, and harm to benefit ratios when moving from triennial to biennial screening between ages 50 and 74 years per 1000 women followed over their lifetime.

| Breast density at age 50 years | Relative risk | No. of additional breast cancer deaths averted | No. of additional life-years gained | QALYs gained | No. of additional false-positives | No. of additional biopsies | No. of additional overdagnosis | Ratio of additional false-positives per additional life-year gained | Ratio of additional overdagnosis per additional breast cancer death averted | Ratio of additional screens per additional life-year gained |
|-------------------------------|--------------|-----------------------------------------------|------------------------------------|-------------|----------------------------------|--------------------------|-----------------------------|---------------------------------|---------------------------------|---------------------------------|
| Almost entirely fatty         | 0.60         | 0.4 (0.3, 0.6)                                | 9 (6, 10)                          | 5 (2, 6)    | 135 (126, 245)                  | 15 (13, 30)              | 1.7 (0.3, 2.1)               | 15 (13, 44)                      | 3.1 (1.1, 5.3)                  | 409 (373, 644)                  |
|                               | 0.70         | 0.5 (0.4, 0.6)                                | 10 (6, 11)                         | 6 (3, 7)    | 135 (125, 244)                  | 15 (12, 30)              | 2.0 (0.4, 2.3)               | 13 (11, 38)                      | 3.2 (1.1, 5.2)                  | 356 (318, 558)                  |
|                               | 0.85         | 0.6 (0.5, 0.8)                                | 12 (8, 14)                         | 8 (4, 9)    | 134 (125, 242)                  | 14 (12, 29)              | 2.4 (0.5, 2.8)               | 11 (9, 30)                       | 3.1 (1.0, 5.0)                  | 290 (259, 437)                  |
|                               | 1.00         | 0.6 (0.5, 0.9)                                | 14 (9, 16)                         | 10 (5, 11)  | 133 (124, 241)                  | 14 (12, 29)              | 2.7 (0.6, 3.3)               | 9 (8, 26)                        | 3.1 (1.1, 5.1)                  | 254 (222, 384)                  |
| Scattered fibroglandular density | 0.60         | 0.6 (0.5, 0.7)                                | 11 (10, 16)                        | 7 (5, 10)   | 232 (216, 375)                  | 26 (22, 46)              | 2.5 (0.4, 3.3)               | 21 (14, 38)                      | 3.4 (0.8, 5.2)                  | 316 (224, 359)                  |
|                               | 0.70         | 0.7 (0.6, 0.8)                                | 13 (12, 18)                        | 8 (6, 12)   | 231 (215, 373)                  | 25 (22, 45)              | 2.9 (0.5, 3.8)               | 17 (12, 32)                      | 3.4 (0.8, 5.3)                  | 267 (194, 307)                  |
|                               | 0.85         | 0.9 (0.8, 1.0)                                | 16 (14, 21)                        | 10 (8, 14)  | 229 (212, 369)                  | 25 (22, 45)              | 3.4 (0.6, 4.5)               | 15 (10, 27)                      | 3.4 (0.8, 5.1)                  | 226 (163, 254)                  |
|                               | 1.00         | 1.0 (0.9, 1.1)                                | 18 (16, 25)                        | 12 (10, 17) | 227 (210, 366)                  | 25 (21, 45)              | 3.8 (0.7, 5.1)               | 13 (8, 22)                       | 3.4 (0.8, 4.9)                  | 196 (138, 213)                  |
| Heterogeneously dense         | 0.60         | 0.8 (0.7, 0.9)                                | 14 (13, 18)                        | 8 (7, 11)   | 283 (265, 444)                  | 15 (11, 38)              | 3.1 (0.6, 3.9)               | 20 (15, 33)                      | 3.5 (0.8, 5.3)                  | 253 (193, 264)                  |
|                               | 0.70         | 0.9 (0.8, 1.0)                                | 16 (16, 21)                        | 10 (9, 14)  | 281 (262, 441)                  | 15 (11, 38)              | 3.5 (0.7, 4.4)               | 18 (12, 28)                      | 3.5 (0.8, 4.9)                  | 219 (165, 226)                  |
|                               | 0.85         | 1.0 (1.0, 1.2)                                | 19 (19, 25)                        | 12 (11, 16) | 277 (258, 435)                  | 14 (11, 37)              | 4.2 (0.8, 5.2)               | 15 (10, 23)                      | 3.5 (0.7, 5.0)                  | 181 (138, 184)                  |
|                               | 1.00         | 1.2 (1.2, 1.4)                                | 22 (22, 29)                        | 15 (14, 20) | 274 (254, 430)                  | 14 (10, 37)              | 4.7 (0.9, 5.9)               | 12 (9, 20)                       | 3.3 (0.8, 4.9)                  | 153 (117, 158)                  |
| Extremely dense               | 0.60         | 1.0 (0.8, 1.0)                                | 18 (15, 18)                        | 10 (10, 12) | 213 (200, 423)                  | 10 (7, 36)               | 3.4 (0.7, 3.5)               | 14 (11, 24)                      | 3.6 (0.8, 4.5)                  | 199 (198, 231)                  |
|                               | 0.70         | 1.1 (0.9, 1.1)                                | 20 (17, 20)                        | 12 (11, 14) | 211 (198, 419)                  | 10 (7, 36)               | 3.8 (0.8, 4.0)               | 12 (10, 21)                      | 3.7 (0.8, 4.4)                  | 173 (170, 203)                  |
|                               | 0.85         | 1.3 (1.0, 1.3)                                | 24 (20, 24)                        | 16 (14, 16) | 208 (195, 413)                  | 9 (7, 35)                | 4.5 (1.0, 4.6)               | 10 (8, 17)                       | 3.6 (0.7, 4.4)                  | 143 (141, 167)                  |
|                               | 1.00         | 1.5 (1.2, 1.5)                                | 28 (23, 28)                        | 19 (16, 20) | 205 (192, 408)                  | 9 (7, 35)                | 5.1 (1.1, 5.2)               | 9 (7, 14)                        | 3.5 (0.7, 4.2)                  | 120 (120, 145)                  |
Our findings are largely in line with previous studies. A previous modeling study, including the same 3 models, focusing on women at increased risk, found that average-risk women with low breast density undergoing triennial screening will maintain a similar or better balance of benefits and harms than average-risk women receiving biennial screening (6). Another modeling study using combined risk-based strategies also found that triennial screening from age 50-74 years was optimal for low-risk and medium-low-risk Spanish women (7) and even investigated less intense strategies (quinquennial screening). Moreover, triennial screening is the currently employed screening frequency in the United Kingdom and has been predicted to lead to a substantial mortality reduction (36). Also, the Canadian Task Force recommends screening with mammography every 2-3 years for women aged 50-69 years (37).

Our results show that for a subgroup of women with a combination of fatty or scattered fibroglandular breast density and low-risk (RR = 0.6, 0.7, 0.85) incremental benefits (deaths averted, life-years gained, and QALYs gained) are small for biennial screening from age 50 to 74 years compared with triennial screening. This is reflected in the higher ratio between additional false-positives and additional life-years gained in the low-risk and low-density subgroups when going from triennial to biennial screening compared to the average-risk population, indicating that there are (relatively) more harms relative to benefits in these subgroups than in the average-risk population.

The models consistently found that the benefits of screening decrease with decreasing risk, whereas the number of false-positives and unnecessary biopsies are mostly stable over categories of low risk. The latter was due to our assumption that mammography performance was unaffected by risk. The benefits also decreased with decreasing density, although the decrease in benefit was not so steep when comparing the highest density category to the next category, indicating that elevated risk among women with high density is a more important determinant of absolute screening benefits than high breast density. With regard to harms, false-positives and unnecessary biopsies were highest in the heterogeneously dense category, whereas the trends in overdiagnosis across density categories varied across models.

These results are useful for informing guidelines and for clinical practice. Because the conditions that result in lower-than-average risk are common, primary care providers could use these results in shared decision-making discussions with women. Most risk factors that lead to a decreased risk are not easily modifiable, but they are relatively straightforward to ascertain. If a subgroup of women can be identified to be at low risk, these women can relatively safely decrease their screening intensity from biennial to triennial.

We acknowledge that breast density is not known in women who have never been screened and is therefore difficult to use to tailor the interval of screening among low-risk women. However, it is possible to tailor the screening interval after a first mammogram based on density, especially because mandated standard reporting of breast density to women after a mammogram has become increasingly more common in the United States. Importantly, the measurement of breast density has become more reliable with automated density measures and has similar accuracy in predicting breast cancers (38-40).

Strengths of this study include consideration of breast density; evaluation of a comprehensive set of outcomes for benefits and harms; and the use of 3 well-established, validated models (19). One of the strengths of collaborative modeling is that the combined results from the different independent modeling groups constitute a sensitivity analysis on model structure. Each model was developed using common data from multiple sources and an elaborate calibration process varying multiple parameters to match population-level breast cancer incidence and mortality data (from Surveillance, Epidemiology, and End Results [SEER]). If models were to include alternative values for standard parameters, they would no longer be calibrated to SEER data, and the resulting predictions could not be viewed as reliable. A strength of our analysis is that each model incorporates different structural assumptions about unobservable natural breast cancer history, including varying assumptions regarding the percent of cancers (invasive and/or DCIS) that do not progress, and sojourn times, which inherently provide a sensitivity analysis on screening benefit. Taken collectively, the cross-model results provide stronger evidence than would any single model varying each parameter individually. In addition, most trends and the ranking of scenarios were very similar across models, except for the overdiagnosis results. We found especially that the trends in overdiagnosis across density categories varied across the models; in model E, the number of overdiagnosed women increases with increasing density, reflecting the higher risk associated with density, whereas in model GE, the number of overdiagnosed women decreased, reflecting the lower sensitivity associated with density, and in model W, overdiagnosis was highest in the 2 middle categories as a result of the 2 opposing causes of higher risk and lower sensitivity.
not include a subset of invasive cancers that do not regress. Second, the models include a range of nonprogressive DCIS, resulting in a wide range of predicted overdiagnosis of DCIS from 34% to 62% (43). Third, the models assume that the benefit of screening arises from either detection at a smaller tumor size or at an earlier stage, and at a younger age. There is a range between these 3 models in predicted mortality reductions of 25%-32% for biennial screening in ages 50-74 years (22). Finally, for sojourn times, model GE includes an age-dependent sojourn time ranging from 2 to 4 years, whereas models E and W simulate continuous tumor growth with certain distributions, resulting in a wide range of distribution of sojourn times, including a subset of tumors with very short sojourn times as well as very long sojourn times. Estimates of mean sojourn times may be biased if they are based on a model that does not allow for non-progressive (overdiagnosed) cancer (44).

Despite the substantial differences between models on these key assumptions, models come to the same conclusion regarding the incremental benefits and harms of biennial vs triennial screening in low-risk women.

Overall, our collaborative modeling study showed that triennial screening from ages 50 to 74 years can be considered for women who have fatty or scattered fibroglandular breast density and average or low risk of developing breast cancer and for women with very low risk at any density level. By undergoing more intense screening, these women are subjected to more harms, with only small added benefits. The results contribute to the growing body of evidence that tailored screening has many advantages over age-based guidelines for average populations (7,45). It will be important to translate our findings, and other results, into clinical practice and test the most effective methods for communication of breast cancer risk and breast density to enhance shared decision making about breast cancer screening.

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**Data Availability**

Input and output data from the models are available by sending a request to the corresponding author of this study (e-mail: n.vanravesteyn@erasmusmc.nl).

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