Interval breast cancer rates for digital breast tomosynthesis versus digital mammography population screening: An individual participant data meta-analysis

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\textbf{A B S T R A C T}

Background: Digital breast tomosynthesis (DBT) improves breast cancer (BC) detection compared to mammography, however, it is unknown whether this reduces interval cancer rate (ICR) at follow-up.

Methods: Using individual participant data (IPD) from DBT screening studies identified via periodic literature searches July 2016 to November 2019, we performed an IPD meta-analysis. We estimated ICR for DBT-screened participants and the difference in pooled ICR for DBT and mammography-only screening, and compared interval BC characteristics. Two-stage meta-analysis (study-specific estimation, pooled synthesis) of ICR included random-effects, adjusting for study and age, and was estimated in age and density subgroups. Comparative screening sensitivity was calculated using screen-detected and interval BC data.

Findings: Four prospective DBT studies, from European population-based programs, contributed IPD for 66,451 DBT-screened participants: age-adjusted pooled ICR was 13.17/10,000 (95% CI: 8.25–21.02). Pooled ICR was higher in the high-density (21.08/10,000; 95% CI: 6.71–66.27) than the low-density (8.63/10,000; 95% CI: 5.25–14.192) groups (\(P = 0.03\)) however estimates did not differ across age-groups (\(P = 0.32\)). Based on two studies that also provided data for 153,800 mammography screens (age-adjusted ICR 17.69/10,000; 95% CI: 12.22–23.66), DBT’s pooled ICR was 16.83/10,000 (95% CI: 11.89–23.82). Comparative meta-analysis showed a non-significant difference in ICR (\(-0.44/10,000; 95\% \text{CI: } -11.00–10.11\)) and non-significant difference in screening sensitivity (6.79%; 95% CI: -0.73–14.87%) between DBT and DM but a significant pooled difference in cancer detection rate of 33.49/10,000 (95% CI: 23.88–43.10). Distribution of interval BC prognostic characteristics did not differ between screening modalities except that those occurring in DBT-screened participants were significantly more likely to be negative for axillary-node metastases (\(P = 0.005\)).

Interpretation: Although heterogeneity in ICR estimates and few datasets limit recommendations, there was no difference between DBT and mammography in pooled ICR despite DBT increasing cancer detection.

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\textbf{Introduction}

Mammography population screening has been shown to reduce breast cancer (BC) mortality [1–3]; in some screening settings, mammography is being replaced by digital breast tomosynthesis (DBT). This pseudo-3D-mammography technology has been adopted in breast imaging practice as it reduces overlapping parenchyma (relative to mammography) which might reduce false-positive interpretations and enhance visualisation and detection of BC [4–7]. Studies of DBT screening have shown that DBT, used in combination with acquired or synthesized 2D-images, detects an estimated additional 1.6 cancers/1000 screens compared to mammography [6]. However, it remains unknown whether DBT’s incremental BC detection improves outcomes at follow-up, such as reducing the risk of an interval BC, beyond what can be achieved with mammography screening.

BC mortality as an endpoint requires very large studies with long follow-up, so conducting such trials may not be feasible for comparing the effectiveness of DBT and digital mammography (DM) [8]. We
proposed using interval BC rate as a surrogate for screening effectiveness, and described sourcing data on this outcome from prospective studies in a published study plan [8]. Given that interval cancers are not detected at screening and progress to clinical presentation (often due to symptoms), we hypothesised that increased screen-detection of BC would lead to a reduction in interval cancers from DBT relative to DM screening which would be expected to have benefit [8]. This could also inform inferences on whether the additional BC detection from DBT is adding screening benefit, or adding to over-diagnosis (through detection of indolent tumours) or lead-time.

We therefore undertook a collaborative individual participant data (IPD) meta-analysis to assess the impact of DBT screening (versus DM) on interval BC rates in population screening. Because interval cancers are an important but infrequent outcome, pooling datasets gives increased precision in estimation of this endpoint. Using IPD also allows adjustments for differences between studies and study groups and supports subgroup analyses. This paper reports the primary endpoint of the IPD meta-analysis, interval BC rates, based on prospective studies comparing DBT and DM screening.

Methods

An international collaborative meta-analysis using IPD was established in 2016 and detailed in a protocol outlining rationale, study eligibility criteria, literature search method, end-points, and requested data [8]. Study eligibility was based on the following pre-defined criteria: studies that investigated DBT (interpreted alone, or with acquired or synthetic 2D-images) compared to DM screening; used a prospective design for DBT screening; reported data on screen-detected and interval cancers (numbers or rates), or stated that interval cancer data would be obtained at follow-up [8]; and used predominantly biennial screening (aligning with practice in population-based screening programs) [8]. Because some prospective DBT trials used a paired design, comparing DBT and DM screen-reads within-woman, interval BC rates from those studies would represent outcomes for participants screened with DBT (since all had DBT and none had DM-alone). For those studies, interval BC data were requested for a comparison cohort screened with DM only from the same screening program and population.

2.1 Literature searching and collaborative process

Literature searches (MEDLINE, EMBASE and EBM Reviews) were performed at several time-points from July 2016 to November 2019: Fig. 1 describes the literature search, the cumulative search results...
and study identification process in a flow-diagram (including eligible studies [4,9–15] and reasons for excluding studies). Investigators of eligible studies were contacted via email and invited to collaborate (up to three contact attempts were made), and were informed of the study protocol and required data variables. Agreement to provide IPD was documented, and data were accepted for inclusion in the analysis up to June 2020; we anticipate updating the analyses if additional studies provide data in the future.

Ethics: The IPD meta-analysis involved a data-provision plan for which individual studies secured institutional ethics approval, and was also approved by the University of Sydney Ethics Committee for the proposed analyses of IPD sourced from existing studies (Project no. 2017/143) [8].

2.2 Data variables and endpoints

Requested data for screened participants included: age, screen date, mammographic density (as measured in each study), screening round, self-reported symptoms, whether recalled to assessment, outcome (negative, screen-detected or interval cancer), cancer characteristics (histology, size, node status), date of interval BC diagnosis. For each study, variables were collected in a standardised database, including standardised categorisation, to enable pooling of data. Study databases were checked against published reports – differences were discussed with study investigators to achieve resolution or identify the reason for discrepancies (including those imposed by data privacy requirements, such as provision of month/year only for date of birth by some trials).

The primary endpoint was an interval BC (invasive or in-situ), and screening sensitivity (inclusive of BC detection data). Interval BC classification was based on ascertainment of screen-detected participants within 24 months following a negative screen or assessment. Secondary endpoints, comprising cancer detection rates and characteristics, screening recall and predictive value will be reported in future publications.

2.3 Study quality

Eligibility criteria for this IPD meta-analysis required that DBT studies be prospective, to minimise selection of participants to DBT screening. In addition, we describe the characteristics of each study to guide interpretation of quality in the context of population screening, specifically: methods of the prospective DBT study (whether randomisation used); design of comparison groups (whether concurrent, and whether from the same screening program); and the methods used to ascertain interval BCs (whether population cancer registry checked). This information is reported in study-specific summaries and flow-charts (Table 1; Online Appendix-1) supplemented by quality appraisal using the QUADAS-2 tool [16] (online Appendix-1).

2.4 Statistical methods

Study characteristics were summarised descriptively. The number of participants and screens, number of detected and interval cancers, and time to interval cancer diagnosis, were calculated. Screen characteristics were summarised within studies and study-groups (DBT and DM) using the median [interquartile range, IQR] for continuous variables and counts (percent) for categorical variables. Interval cancer characteristics were summarised by study-groups and compared using the Chi-squared test for categorical variables and Kruskal-Wallis test for continuous variables.

Two-stage IPD meta-analysis was performed: study-specific estimates were first calculated using logistic regression, and then synthesised using random-effects meta-analysis. Random-effects models allow for both within-study sampling variability as well as heterogeneity between studies when calculating pooled estimates. If a study had multiple screens from the same participant, a random effect for participant was included within the logistic regression. All meta-analyses were adjusted, by age-group (<50, 50–59, 60+) or by age (continuous) using the median age as reference, and where data were provided also by density (reclassified into low, high). Rate ratios and risk difference with 95% confidence intervals were reported. Sensitivity analyses examined the effect of excluding the small number of participants self-reporting symptoms. P-values <0.05 were considered statistically significant. Analyses were performed using the metafor package [17] (version 2.4–0) for R (version 3.6.3) [18].

Results

We identified eight eligible studies of DBT screening [4,9–15] (Fig. 1), and from these four studies [4,10–12] contributed data according to the IPD meta-analysis protocol inclusive of interval BC data. Collectively these four studies, which have reported outcomes in seven papers [4,10–12,19–21] contributed IPD for 66,451 DBT-screened participants; two of these studies [10,12] also provided data for 153,800 DM-only screens from 101,532 participants (Table 1). Three of the studies were prospective non-randomised trials [4,10,11] one was a prospective cohort study [12]; all four studies investigated DBT in organised population-based screening programs. Study-specific characteristics, including methods and quality evaluation, and data contribution to the meta-analysis are summarised in Table 1, supplemented by study flow-charts and appraisal (online-only Appendix-1); none of the studies were randomised controlled trials (RCT). Median age of DBT-screened populations was similar across studies (Table 1), however it differed between DBT and DM groups for one study [10]. Screening participants in all studies received DBT acquisitions with either DM or synthesised 2D-images, hence for the purpose of interval BC analyses, these will be referred to as DBT-screened participants. However, some studies compared screen-reading (within participants) using DBT alone, DBT with DM, or DBT with synthesised 2D-images (Table 1).

3.1 Interval BC in DBT-screened participants

Across all studies 105 interval BCs were diagnosed in DBT-screened participants at a median 1.3 [IQR 0.9, 1.7] years from screening date (study-specific data shown in Fig. 2). Crude interval BC rate was 15.97/10,000 DBT screens (95%CI: 13.11–19.23), and the age-adjusted (referent 56 years) modelled estimate was 13.17/10,000 (95%CI: 8.25–21.02; Fig. 2) shows there was significant heterogeneity in estimates across studies. Fig. 3 displays study-specific and pooled estimates in age subgroups: although interval BC rate was lower in the 50–59 years age-group (10.08/10,000; 95%CI: 4.88–20.82) relative to younger and older age-groups, there was no evidence that estimates differed significantly by age-groups (P = 0.32).

Fig. 4 shows modelled estimates for density subgroups (three studies contributed data [4,10,11]), indicating significantly higher interval BC rate in the high-density (21.08/10,000; 95%CI: 6.71–66.27) than the low-density (8.63/10,000; 95%CI: 5.25–14.192) group (P = 0.03). The modelled interval BC rate ratio for density (adjusted for study and age) was 2.92 (95%CI: 1.48–5.84) for high versus low density (P = 0.002). Analysis by screen-round was limited to two studies [11,12] of which one was predominantly repeat screening (online-only Appendix-2: Fig. A).

3.2 Interval BC for DM screens & comparative interval BC rates for DBT versus DM

For the two larger studies (Table 1: MBTST, OVVV) [10,12] contributing DM-only screening data, there were 276 interval cancers at a median 1.2 [IQR 0.7, 1.6] years from screening. Crude interval BC rate was 17.58/10,000 DM screens (95%CI: 13.78–22.44) and the age-
| Study [publications reporting outcomes] | Design | Number of screens [N interval cancer analysis]* | Median age [IQR] | Double-reading and recall process | Comparison group* screened with digital mammography (DM) only | Interval cancer ascertainment |
|----------------------------------------|--------|-----------------------------------------------|------------------|-----------------------------------|-------------------------------------------------------------|-----------------------------|
| STORM-1 trial [Ciatto [4]; Houssami [19]] | Prospective trial comparing sequential screen-reading of DM with DBT+DM (paired data) in women ≥ 48 years | 7292 [7235] | 58 [54, 63] | Independent double-read, recall if either reader recalls | NA | NA | Check of regional cancer registry, and pathology and hospital databases; 2 year follow-up |
| MBTST (Malmo trial) [Zackrisson [10]; Johnson [20]] | Prospective trial comparing screen-reading using one-view DBT with 2-view DM (paired data) in women ≥ 40 years | 14,848 [14,711] | 57.4 [48.9, 65.3] | Independent double-read, consensus meeting if score ≥ 3 by either reader | Yes (slightly longer in 2015); yes | 96,037 from 43,769 women [95,497] | 52.3 [45.5, 61.4] Population cancer registry linkage; 2 year follow-up (18 months in subgroup where this screening interval recommended) |
| STORM-2 trial [Bernardi [11]] | Prospective trial comparing in separate arms in women aged ≥ 49 years: screen-reading of DBT with DM (paired data), and screen-reading DBT +DM with DM (paired data); only one set of paired data used in analysis | 9672 [9587] | 58 [53, 63] | Two parallel independent double-reading arms, recall if any reader recalls | NA | NA | Check of regional cancer registry, and pathology and hospital databases; 2 year follow-up |
| OVVV study [Hofvind [12]; Hovda [21]] | Prospective cohort study of DBT* screening vs concurrent (DM-only screened) cohort in women 50–69 years at screening | 34,639 [34,315] | 59 [54, 64] | Independent double-read, consensus meeting if discordant scores | Yes; yes | 57,763 [57,408] | 59 [54, 64] Cancer Registry of Norway Population cancer registry linkage; 2 year follow-up |

STORM = Screening with Tomosynthesis or Mammography; MBTST = Malmo Breast Tomosynthesis Screening Trial; OVVV = Oslo, Vestfold &Vestre Viken study; NA = not applicable (did not provide data).

* Number of screens represents individual participants except for the DM comparison cohort which included repeat screens - number in squared brackets represents number used in the analysis of interval cancer rates using negative screens as the denominator.

* Comparison groups from same population screening programs allowed comparison of interval cancer rates (OVVV compared services within Norwegian program).

* Indicates study used DBT with synthetic 2D-images reconstructed from the DBT acquisitions.
adjusted (referent 56 years) modelled estimate was 17.69/10,000 (95%CI: 13.22–23.66). For these two studies, the age-adjusted modelled estimate of interval BC rate for DBT screening was 16.83/10,000 (95%CI: 11.89–23.82). Temporal distribution showed that a lower proportion of interval BCs was diagnosed in year one of the inter-screening interval for DBT than DM (31.1% for DBT and 40.2% for DM) as shown in online Appendix-2 (Fig. B).

Comparative meta-analyses (Figs. 5 and 6) show an unadjusted interval BC rate difference of 0.08/10,000 (95%CI: −0.44/10,000) indicating a non-significant difference in interval BC rate between DBT and DM screening; Fig. 5 also shows significant heterogeneity in estimates between studies. Fig. 6 shows the rate ratio in subgroup meta-analysis for age, indicating there were no significant differences in interval BC rates between DBT and DM across age-groups (P = 0.86). Sensitivity analyses for these comparative meta-analyses, whereby the small number of screens with self-reported breast symptoms (<0.4% both groups) were excluded, did not alter results (data not shown).
3.3 Cancer detection rate (CDR) and comparative screening sensitivity

Amongst DBT-screened participants across all studies there were 603 screen-detected BCs, a pooled crude CDR of 91.71/10,000 DBT screens; online-only Appendix-2, Fig. C, shows study-specific detection data, and pooled estimates, indicating homogeneous cancer detection rates for DBT. For the two studies [10,12] that also provided data for DM-only screening, there were 895 screen-detected BCs yielding a pooled crude CDR of 58.81/10,000 DM screens (Fig. C, online-Appendix-2, shows study-specific detection data, and age-adjusted pooled estimate). There was a significant pooled difference in CDR of 33.49/10,000 (95%CI: 23.88–43.10) between DBT and DM.

As also shown in Fig. D (online Appendix-2) this significant difference in CDR between DBT and DM screens was evident in comparative meta-analysis across all age-groups.

Meta-analysis comparing screening sensitivity adjusted for age (Fig. 7) estimated a sensitivity difference of 6.79% (95%CI: -0.73–14.87%) between DBT and DM, or a risk ratio of 1.07 (95%CI: 0.99–1.15). As also shown in study-specific estimates [10,12] (Fig. 7), the age-adjusted estimate for one of the studies [MBTST[10]] showed a significant sensitivity difference of 11.2% (95%CI: 3.02–20.01%) between DBT and DM. Sensitivity analyses for the above-reported meta-analyses, that excluded screens with self-reported symptoms (<0.4% both groups), did not alter results.

3.4 Interval cancer characteristics

Table 2 summarises the characteristics of the 366 interval BCs, comprising predominantly (94.3%) invasive cancers, from studies contributing data for DBT and DM screening [10,12]. The median tumour size of invasive interval cancers was 18.0 mm (IQR 13.0, 27.0): median size was 16.0 mm (IQR 12.5, 25.0) in the DBT group vs 19.5 mm (IQR 14.0, 27.0) in the DM group (P = 0.47). Table 2 describes the distribution of histological type, pathological tumour size (pT) category and grade, and lymph node status. Lower proportions of invasive lobular cancer, pT2 tumours, and grade 3 cancers were noted amongst interval cancers in DBT-screened relative to DM-screened groups, although distributions did not significantly differ between modalities (all P > 0.13). There was a significant difference in the distribution of axillary node metastases amongst interval cancers between DBT and DM screening (P = 0.005) — those in DBT-screened participants were more likely to be negative for metastases.

Discussion

Digital breast tomosynthesis (DBT) is being adopted for BC screening in some settings because it increases BC detection rates compared to DM; its effect on recall is variable in European studies however DBT reduces screening recall in studies conducted in the USA[6]. The
A growing body of evidence on DBT’s detection metrics is contrasted by sparse evidence on its effect on outcomes at follow-up of screened populations, yet such evidence is essential to inform breast cancer screening policy decisions. In Europe, for example, DBT has recently received conditional approval for breast cancer screening recommendations that also provided IPD for concurrent cohorts screened with DM. Comparing screening sensitivity showed weak evidence of improved sensitivity from DBT, although DBT significantly increased the rate ratio was almost 3-times in dense (vs low-density) breasts, similar to findings for mammography screening.

For the two larger prospective studies in this meta-analysis that also provided IPD for concurrent cohorts screened with DM alone (153,800 screens) from the same population programs, the age-adjusted pooled interval BC rate for DM was similar to that for DBT. Comparative meta-analysis showed no significant difference in interval BC rates between DBT and DM screening. This finding was consistent across age-groups, so there was no evidence that DBT reduced interval BC rates in any age-group compared to DM. Considering other studies that have evaluated DBT population-based screening, and have reported on interval BCs at follow-up, both Skaane et al. and Bernardi et al. found that there was no significant difference in interval BC rates between DBT-screened groups and DM-screened historical cohorts. Therefore even if these additional studies had been included in the IPD meta-analysis, the findings of our comparative meta-analysis of interval BC rates are unlikely to have changed.

Meta-analysis comparing screening sensitivity showed weak evidence of improved sensitivity from DBT, although DBT significantly improved sensitivity compared to DM. Pooled interval BC rates were significantly higher in dense breasts; adjusting for study and age, the rate ratio was almost 3-times in dense (vs low-density) breasts, similar to findings for mammography screening.

### Table 2

| Variable                      | DBT (n = 90) | DM (n = 276) | Total (n = 366) | P value* |
|-------------------------------|-------------|-------------|----------------|---------|
| **Study**                     |             |             |                |         |
| MBTST                         | 22 (24.4)   | 188 (68.1)  | 210 (57.4)     | 0.005   |
| OVVV                          | 68 (75.6)   | 88 (31.9)   | 156 (42.6)     | 0.005   |
| **Histological type**         |             |             |                |         |
| Ductal carcinoma in-situ      | 7 (7.8)     | 14 (5.1)    | 21 (5.8)       | 0.35    |
| Invasive ductal cancer        | 71 (78.9)   | 206 (75.2)  | 277 (76.1)     |         |
| Invasive lobular cancer        | 9 (10.0)    | 39 (14.2)   | 48 (13.2)      |         |
| Other invasive types          | 3 (3.3)     | 15 (5.5)    | 18 (5.0)       |         |
| Not reported                  | 0           | 2           | 2              |         |
| **Pathological tumour (pT) size** |             |             |                |         |
| pTis                          | 7 (8.3)     | 14 (5.3)    | 21 (6.1)       | 0.43    |
| pT1a-b (<10 mm)               | 15 (17.9)   | 32 (12.2)   | 47 (13.5)      |         |
| pT1c (>10 and ≤20 mm)         | 34 (40.5)   | 106 (40.3)  | 140 (40.3)     |         |
| pT2 (>20 and ≤50 mm)          | 26 (31.0)   | 105 (39.9)  | 131 (37.8)     |         |
| pT3 (>50 mm)                  | 2 (2.4)     | 6 (2.3)     | 8 (2.3)        |         |
| Not reported                  | 6           | 13          | 19             |         |
| **Tumour grade**              |             |             |                |         |
| 1                             | 13 (16.5)   | 35 (14.5)   | 48 (15.0)      | 0.13    |
| 2                             | 44 (55.7)   | 109 (45.0)  | 153 (47.7)     |         |
| 3                             | 22 (27.8)   | 98 (40.5)   | 120 (37.4)     |         |
| Not reported                  | 11          | 34          | 45             | 0.005   |
| **Axillary lymph nodes**      |             |             |                |         |
| Negative                      | 61 (73.5)   | 141 (54.7)  | 202 (59.2)     |         |
| Positive (metastases)         | 20 (24.1)   | 92 (35.7)   | 112 (32.8)     |         |
| Micro-metastases or isolated tumour cells | 2 (2.4) | 25 (9.7) | 27 (7.9) |         |
| Not reported                  | 7           | 18          | 25             |         |

* P value for comparison of the distribution of interval cancer variables between DBT and DM screens.

Note: Not reported: includes missing data or not applicable (these were mostly not applicable in reference to ductal carcinoma in-situ cases); these data were not counted in distribution of proportions.
increased cancer detection rate (and this effect was consistent between studies). Comparison of interval BC characteristics did not find any significant differences between screening modalities, with the exception of axillary-node metastases ($P = 0.005$) – a higher proportion of node-negative and lower proportion of node-positive interval BCs were shown in DBT-screened than DM-screened participants. This finding raises the possibility that DBT screening could reduce the rate of advanced BCs at repeat screening rounds. Temporal distribution of interval BCs, though limited by small numbers at each year, suggests that DBT may shift the time at which interval BC events are diagnosed, so radiological review may provide insights as to whether DBT alters the imaging pattern of interval BCs (for example, fewer false-negatives) [23].

Overall our results indicate that DBT screening, despite increasing BC detection, did not have a measurable impact on interval BC rates, and the increase in screening sensitivity (which was not statistically significant) was due to DBT increasing the number of screen-detected cancers. Our findings should be interpreted with caution despite the strengths of IPD meta-analysis, which enabled more precise analyses of the primary end-point than individual studies and supported statistical adjustments of interval BC rates, because there were limitations that hinder definitive conclusions from our work. The main limitation is that only four from eight eligible studies provided data (some investigators were unable or unwilling to contribute data), and only two studies provided data for concurrent DM screens, so pooled estimates have modest precision. These two studies forming the comparative meta-analysis were from Norway (OVVV) and Sweden (MBTST) [10,12] and were the largest of the four studies contributing data. Although all studies in our meta-analysis had ascertained interval BC events (Table 1), it is noteworthy that both studies contributing to comparative meta-analysis used population cancer registry linkage for this ascertainment. Since both studies were from Nordic nations with highly-organised population screening programs, this could potentially limit generalisability of our findings. Further, we focused on trials conducted in biennial screening programs, so our results may not apply to annual screening practice (where interval BC rates would be relatively lower) [23]. Another limitation is the heterogeneity in interval BC rate estimates, as discussed earlier; we used random-effects meta-analysis to allow for heterogeneity between studies, and explored its possible sources in subgroup analyses. Our findings suggest that there was relatively less heterogeneity in interval BC rates in the 60+ age-group and in the low-density group. Heterogeneity was also evident in comparative estimates based on the MBTST and OVVV studies [10,12]; these two studies used different DBT technology and screen-reading protocols (Table 1) which might account for the relatively divergent estimates of interval BC rates.

Our IPD meta-analysis did not focus on DBT’s initial performance measures (screen-detection and recall rates) because this has been done in study-level meta-analyses from Marinovich et al.[6] Alabousi et al.[27] and Giampietro et al. [28] – however, most DBT screening studies do not report interval BC data so pooled interval BC rates have not been reported in these study-level analyses. We focused on interval BC rates to provide knowledge on an evidence gap in DBT’s effect. Substantially larger IPD datasets that include more studies (including RCTs) would be needed to assess DBT’s effect on interval BC rates in subgroups. We are aware of RCTs that will report other endpoints for screening benefit, such as a reduction in advanced BC rates, and which will report interval BC data potentially supporting larger analyses [29,30].

Using IPD meta-analysis of prospective DBT screening studies, a few studies with heterogeneous results on ICR could be included; our work did not find a significant difference in interval BC rates between DBT and DM screening. Screening sensitivity was not statistically significantly higher for DBT than DM screening, and there was evidence that interval BCs were more frequently node-negative in DBT-screened than DM-screened groups. The findings from our work highlight the need for much larger collaborative analyses of DBT screening to allow comparative evaluation of important but relatively infrequent screening outcomes such as interval BC rates, or advanced breast cancer rates. Our findings can inform future evaluation of DBT screening particularly for population screening programs contemplating potential transition to DBT or retention of standard mammography screening, suggesting that investment in large comparative studies may be a more appropriate strategy than prompt adoption of DBT.

Declaration of Competing Interest

Dr. Houssami reports grants from National Breast Cancer Foundation (NBCF Australia), grants from National Health & Medical Research Council (NHMRC, Australia), during the conduct of the study. Dr. Zackrisson reports grants from the Swedish Cancer Society, grants from Governmental funding (Region Skane), during the conduct of the study; in addition, Dr. Zackrisson has a patent approved, US (application no PCT/EP2014/057,372) issued. The authors have not reported any conflicts of interest.

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Data sharing statement

IPD will not be made available due to privacy regulations that affect IPD provisioning as part of this collaborative study; investigators may be contacted to discuss processes required for potential data access.

Authors’ contributions

Conceptualisation: Houssami, Zackrisson, Hofvind; writing original draft: Houssami; writing (review and edit): all authors; literature search: Hunter, Houssami; data sourcing and collection: Bernardi, Lang, Hunter, Aglen, Johnson, Zackrisson, Hofvind; Project administration: Houssami, Hunter; Statistical analysis: Soerensen, Robledo; access to data: Soerensen, Houssami, Hunter; figures: Soerensen, Hunter; data interpretation: all authors.

Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.eclinm.2021.100804.
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