External Beam Partial Breast Irradiation Versus Whole Breast Irradiation for in Early Breast Cancer: A Systematic Review and Meta-Analysis

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Research

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Abstract

Purpose

Postoperative radiotherapy can reduce the recurrence of breast cancer. Postoperative radiotherapy is divided into whole breast irradiation (WBI) and partial breast irradiation (PBI) for early breast cancers. Due to the characters of saving time, money, and easy to deliver, external beams PBI (EB PBI) is brought into focus. However, the researches on outcomes, safety, and efficacy between EB PBI and WBI are still insufficient. We concluded a meta-analysis for LRR, regional node recurrence, contralateral breast cancer, distant recurrence, mortality, less acute skin toxicity (≤ 1 grade), late skin toxicity and the cosmetic score of external beam partial breast irradiation (EB PBI) and whole breast irradiation (WBI) to develop a radiotherapy plan for early low recurrence risk breast cancer patients.

Method

We searched Pubmed®Embase®Cochrane Library®Clinicaltrials. Study eligibility criteria are as below: (1) RCTs for EB PBI vs WBI; (2) Histologically confirmed breast cancer; (3) AJCC staged Tis-2N0-1M0; (4) ≥ 40 years old; (5) Tumour size ≤ 3 cm; (6) microscopically clear margins ≤ 5 cm; (7) Mean follow-up time ≥ 5 years. All data is used by Cochrane's Review Manager 5.3 (RevMan) to process.

Results

There were 4 RCT studies included in our study with 1999 patients in EB PBI group and 1999 patients in EB PBI. There was no statistic difference between PBI and WBI groups in local recurrence rates (RR = 1.15; 95% CI, 0.76 to 1.74; p = 0.52; I² = 0%), regional node recurrence (RR = 1.00; 95% CI, 0.49 to 2.04, p = 0.99, I² = 0%), contralateral breast cancer (RR = 0.79; 95% CI, 0.54 to 1.16; p = 0.23; I² = 0%), distant recurrence (RR = 1.00; 95% CI, 0.63 to 1.59; p = 1.00; I² = 0%), non-breast second cancer (RR = 1.03; 95% CI, 0.50 to 2.16; p = 0.93; I² = 83%), mortality (RR = 0.96; 95% CI, 0.60 to 1.55; p = 0.88, I² = 54%). EB PBI had worse cosmetic score (RR = 1.56; 95% CI, 1.04 to 2.34; p = 0.003, I² = 84%), less acute skin toxicity (≤ 1 grade) (RR = 0.17; 95% CI, 0.07 to 0.42; p = 0.0001, I² = 87%) and late skin toxicity (RR = 0.65; 95% CI, 0.48 to 0.88; p = 0.005; I² = 27%) than WBI.

Conclusion

EB PBI has similar LRR, regional node recurrence, contralateral breast cancer, distant recurrence, non-breast second cancer and mortality with WBI. But EB PBI has worse cosmetic score, less acute skin toxicity (≤ 1 grade) and late skin toxicity than WBI.

Introduction

Breast cancer ranks second of the world in cancer incidence and mortality, with more than 2.08 million new cases and more than 630,000 deaths in 2018\textsuperscript{1}. Irradiation can significantly reduce ipsilateral tumor
recurrence and mortality of early breast cancer after breast-conserving surgery\textsuperscript{2–4}. The delay of radiotherapy after breast-conserving surgery was associated with a significantly increased risk of local recurrence\textsuperscript{5}. NCCN guideline-recommended radiotherapy after most of the early breast cancer patients after surgery, especially for breast-conserving surgery\textsuperscript{6}. For early breast cancer patients with low recurrence risk, Partial breast irradiation (PBI) is another choice than traditional whole breast irradiation (WBI)\textsuperscript{6–8}. The proposed of PBI were based on that the recurrence and metastasis sites after breast-conserving surgery is usually located around the primary tumor. About 90\% of recurrence was located in 1 cm from the primary tumor\textsuperscript{7,9–12}. PBI just covers the primary tumor and its surrounding 2-3cm area. PBI can shorten the time of radio-chemotherapy, reduce the treatment cost, and easy to deliver for most radiotherapy centers\textsuperscript{13}. The above advantages and fewer irradiation tissues also improved patients' compliance with postoperative radiotherapy, chemotherapy, and life quality\textsuperscript{5,14}. Although PBI had lower mortality without breast cancer recurrence, However, PBI had higher local recurrence and bad cosmetology outcomes than WBI in previous studies\textsuperscript{15,16}. Vaidya J.S et al\textsuperscript{17} found that PBI compared to WBI reduced the overall mortality of without breast cancer patients by 25 percent over five years. Liu et al\textsuperscript{18} found there was no significant difference in lymph node recurrence, systemic recurrence, and overall survival or mortality. Pan XB et al\textsuperscript{19} found that the risk of secondary malignancy was similar for both PBI and WBI. Hickey BE et al\textsuperscript{20} found PBI or accelerated partial breast irradiation (APBI) has worse cosmetic and some late effects, but it has less acute skin toxicity.

PBI can be divided into brachytherapy PBI (implantation, single-cavity, multi-cavity balloons or combined application), intraoperative PBI (IORT), External beam PBI (EB PBI). However, EB PBI has the advantages of non-invasive and easy to deliver in most radiation centers\textsuperscript{21}. It also had the shortest time and economic costs for patients than WBRT-B, WBI, WBI-accelerated, WBRT intensity-modulated, (APBI)-IC, APBI-HDR interstitial\textsuperscript{22}. EB PBI has a huge application prospect of early breast irradiation than brachytherapy PBI and IORT. Up to now, no systematic meta-analysis on outcomes efficacy and toxicity effects of EB PBI and WBI have been reported. Recently, the latest results of the RAPID were published\textsuperscript{23}, we concluded a meta-analysis for EB PBI and WBI for radiotherapy of early low recurrence risk breast cancer patients.

**Methods**

This paper is reported in accordance with the preferred reporting items for systematic reviews and meta-analyses (PRISMA) statement and was registered at the International Prospective Register of Systematic Reviews (number:156882).

We searched Pubmed (January 2021 to February 1995); Embase (January 2021 to July 1994); Cochrane Library (January 2021 to 2005); Clinicaltrials (January 2021 to June 2015). Study eligibility criteria are as below: (1) RCTs for EB PBI vs WBI; (2) Histologically confirmed breast cancer; (3) AJCC stages Tis-2N0-1M0; (4) ≥ 40 years old; (5) Tumor size ≤ 3 cm; (6) microscopically clear margins ≤ 5 cm; (7) Mean follow-up time ≥ 5 years. Search major themes included ‘partial breast irradiation’ OR accelerated partial breast
irradiation’ AND ‘whole breast irradiation’ And ‘Breast Neoplasm’ (randomized or randomized). MeSH Terms and Emrew terms were used. Then we selected articles in EB PBI vs WBI.

Two independent reviewers collected the data independently. Then they determined the data be included together. We tried to contact the authors to get not reported data. For papers that were updated or re-published, the newest data was abstract. Cochrane’s Review Manager 5.3 (RevMan) was used to assess statistical variance, risk ratios, heterogeneity and sensitivity. Homogeneity and sensitivity analysis were not described in detail because that the study we included was limited and small.

At last 4 papers were included (Fig. 1, Table 1), and all of them are RCTs. The outcomes were about the local regional rate (LRR), regional node recurrence, contralateral breast cancer, distant recurrence, and mortality. The results of the cosmetic score, acute skin toxicity (≤ 1 grade), and late skin toxicity also was included in our meta-analysis. P values of less than 0.05 were considered statistically significant.

### Table 1

| Study, year | Median follow-up (years) | Age(years) | Numbers of patients | Irradiation technology | WBI | PBI |
|-------------|--------------------------|------------|---------------------|------------------------|-----|-----|
| Rodriguez N [1],2013 | 5                       | 61         | 1065 vs 1070        | 3D-CRT                 | 48 Gy/24f,QD | 37.5 Gy/10f,BID |
| Livi L [2],2015 | 5                       | ≥ 40       | 260 vs 260          | IMRT                   | 50 Gy/25f, QD+10Gy/5f,QD | 30 Gy/5f,QD |
| Coles CE [3],2017 | 5                       | 68.64 ± 5.8 | 51 vs 51            | IMRT                   | 40 Gy/15f, QD | 40 Gy/15f, QD |
| TJ W [4],2019 | 8.6                     | 61         | 1065 vs 1070        | 3D-CRT/IMRT            | 42.5 Gy/16f, QD | 38.5 Gy/10f,BID |

### Results

#### Local recurrence rates

There are 3 studies reported local recurrence rates of a total of 3998 patients. local recurrence was defined as any recurrence of the ipsilateral breast or overly skin. Meta-analysis show there was no statistic difference between PBI and WBI groups in local recurrence rates (relative risk [RR] = 1.15; 95% confidence interval [CI], 0.76 to 1.74; p = 0.52; I² = 0%). During at least 5 years of follow-up, there was no significant difference between EB PBI and WBI in the local recurrence rate.

#### Regional node recurrence
We defined regional node recurrence as recurrences of the axilla, supra clavicular fossa, and internal mammary chain. Meta-analysis show there was no statistic difference between EB PBI and WBI groups (RR = 1.00; 95% CI, 0.49 to 2.04, p = 0.99, $I^2 = 0\%$).

**Contralateral breast cancer**

There are 3 studies reported on Contralateral breast cancer. No statistic difference between EB PBI and WBI groups (RR = 0.79; 95% CI, 0.54 to 1.16; p = 0.23; $I^2 = 0\%$).

**Distant recurrence**

Any recurrence of distant organs was considered as distant recurrence. There was no statistic difference between EB PBI and WBI groups (RR = 1.00; 95% CI, 0.63 to 1.59; p = 1.00; $I^2 = 0\%$).

**Non-breast seconded cancer**

There was no statistical difference in non-breast second cancer between EB PBI and WBI groups (RR = 1.03; 95% CI, 0.50 to 2.16; p = 0.93; $I^2 = 83\%$).

**Mortality**

Mortality was reported for 3 studies with a total of 3998 patients. there was no statistic difference between EB PBI and WBI groups. There was no statistic difference between EB PBI and WBI groups in mortality (RR = 0.96; 95% CI, 0.60 to 1.55; p = 0.88, $I^2 = 54\%$).

**Cosmetic scores**

Cosmetic outcome was evaluated on the Harvard Breast Cosmesis Scales$^{23-26}$. It included size, shape, texture, edema, color, thickening, scar, and change into the appearance of the breast. It can be used for accurate cosmetic grading for patients that are treated with breast-conservation surgery and ipsilateral breast tumor recurrence. And it was widely used in clinical trials to accurate cosmetic grading, especially in clinical trials about breast irradiation$^{24}$.

Poor and fair were included in the unsatisfied cosmetic outcome. EB PBI group has more risk of unsatisfied cosmetic outcome than WBI group (RR = 1.56; 95% CI, 1.04 to 2.34; p = 0.003, $I^2 = 84\%$).

**acute skin toxicity (≥ 1 grade) and late skin toxicity (≥ 1 grade)**

Acute radiation morbidity scoring criteria and late radiation morbidity scoring scheme were from the Radiation Therapy Oncology Group(RTOG) and the European Organization for Research and Treatment of Cancer(EORTC)$^{23,26-28}$. Acute skin toxicity greater than grade 1 should be treated for clinical. We mainly compared acute skin toxicity (≥ 1 grade) or late skin toxicity such as edema, pigmentation, depigmentation, telangiectasia, skin to atrophy, ulceration, and so on.
EB PBI would cause less acute skin toxicity (0 grade) than WBI (1 grade) (RR = 0.17; 95% CI, 0.07 to 0.42; p < 0.0001, I^2 = 87%). EB PBI has less late skin toxicity in WBI groups (RR = 0.65; 95% CI, 0.48 to 0.88; p = 0.005; I^2 = 27%).

Discussion

Studies about brachytherapy PBI or intraoperative PBI more than EB PBI. Our meta-analysis compared EB PBI and WBI which systemic proved that EB PBI has similar LRR, regional node recurrence, contralateral breast cancer, distant recurrence, non-breast second cancer, and mortality with WBI for early breast patients. Besides, EB PBI has less acute skin toxicity (0 grade), similar late skin toxicity, and worse cosmetic score than WBI. The protection for subcutaneous tissue during radiotherapy plans delivery and limited the dose in EB PBI will contribute to better cosmetic outcomes. In the study we included, the dose of EB PBI was from 30 Gy/5f to 40 Gy/15 f. In the current report, the dose of EB PBI was 30-50Gy/5-10f, QD/BID^3,25,26,28,29. The best optimal dose of EB PBI radiation has not been determined, National Comprehensive Cancer Network(NCCN) recommend EB PBI deliver in the dose of 34 or 38.5 Gv/10f, BID and 40-42.5 Gy/15-16f, QD for WBI-accelerated. Dose of 4 studies is different which might the source of heterogeneity for LRR, cosmetic score, and acute skin toxicity (0 grade). TJ W^23's study has a maximum proportion of patients. It was the main reason for heterogeneity for LRR and cosmetic score. It compared WBI-accelerated vs APBI, and the dose of it was the highest than other studies. Although its dose of WBI-accelerated and APBI was accepted by NCCN, Highest does and the different delivery fraction of irradiation might induce worst cosmetic score, LRR, local recurrence rates, mortality, and distant recurrence.

Yasmin Korzets et al^16 found that EB PBI may have the best control of local recurrence than brachytherapy PBI and IORT. EB PBI has similar local recurrence rate(LRR) with WBI. But they didn't include outcomes of RAPID^23 and didn't evaluate regional recurrence, cosmetic score, acute, and late skin toxicity. Our study is the systematic meta-analysis of EB PBI versus WBI. We found that EB PBI had no significant difference in LRR, regional node recurrence, contralateral breast cancer, distant recurrence and mortality compared with WBI. These conclusions were similar to Yasmin Korzets's^30 and Liu's^18 study. But we found that EB PBI tends to increase the risk of LRR, decrease the risk of regional node recurrence, contralateral breast cancer, distant recurrence, and mortality, although all of them had no statistical difference between EB PBI and WBI groups.

The selection of PBI patients was controversial. NCCN, GEC-ESTRO, and UK guidelines recommend the patients suitable for PBI were as follows (Table 2). Different guidelines have different Selection criteria for low-risk early breast cancer patients. The patients that could deliver PBI were early breast stages with low recurrence risk. More studies were exploring the application and outcomes of PBI. NCCN guidelines...
recommend PBI was considered in patients as follows. (1) Older than 50 years old, T1, Negative margin ≥ 2 mm, no LVI, ER-positive, and BRCA negative. (2) Low/medium nuclear grading. The measured size of DCIS detected by screening is less than 2.5 cm and the margin of the negative cutting edge is more than or equal to 3 mm. GEC-ESTRO is similar to NCCN guidelines. In addition to the above inclusion criteria, UK consensus statements recommend that tumor size ≤ 3 cm and negative her-2 could also be considered for PBI. The inclusion criteria of PBI are still under discussion.

Table 2
NCCN, GEC-ESTRO, and UK guidelines recommend the patients suitable for PBI

| Risk factors          | NCCN(ASTRO) | GEC-ESTRO                  | UK      |
|-----------------------|-------------|---------------------------|---------|
| Lymph nodes           | positive    | positive                  | negative|
| Margin                | ≥ 2 mm      | ≥ 2 mm                    | 1–2 mm  |
| Tumor size            | ≤ 2 cm, Single center, single tumor, no vascular invasion | ≤ 2 cm, Non-central, single tumor | ≤ 3 cm  |
| Garde                 | 1–2 grade   | -                         | 1–2 grade|
| Hormonal state        | ER positive, BRCA negative | -                        | ER positive, HER-2 negative |
| Age                   | ≥ 50 years  | ≥ 50 years                | ≥ 50 years|
| Dose                  | 38.5 Gy/10f, BID | 32 Gy/8f, bid            | 40 Gy/15f|

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Table 2 NCCN, GEC-ESTRO, and UK guidelines recommend the patients suitable for PBI

Radiation can damage DNA, massive DNA damage and accumulation of unrepaired chromosome breaks induce cell death. Recurrence and metastasis were associated with tumor cells that did not die after radiotherapy. DNA damaged to activate not only p16/Rb but also p53/p21 pathway, all of them take importance of cellular senescence by cellular arrest and oncogene-induced senescence\textsuperscript{31}. Irradiation
induced the ATM kinase activated and p38 mitogen-activated protein kinase (MAPK) through reactive oxygen species (ROS), H2AX, and other DNA repair protein repair DNA damage by NHEJ, HR. Then, activating transcription factor nuclear factor kappa-B activates pRb by p16INK4A ultimately induces durable cell-cycle arrest. Upregulation of inflammatory and fibrotic mediating, epigenetic changes, and disruption of normal oxygen metabolism will increase cytotoxicity reactive oxygen species (ROS) that occur within hours to years of exposure to ionizing radiation. Apoptosis and senescence in cells trigger fibrosis through signaling pathways. The key stages in the metastatic are the appearance of circulating tumor cells (CTCs) seeding and colonizing distant tissues and organs. EMT helps the tumor cells deform into a more metastasizing form. EMT is highly correlated with TGF-β. TGF-β takes an important role in the process of preserving normal tissues, sensitizing tumor, pro-oxidant, and pro-fibrosis.

EB PBI has less acute skin toxicity (≦1 grade), similar late skin toxicity, and worse cosmetic score than WBI which similar to Hickey BE et al's study. Breast induration or fibrosis, atrophy, appearance changed, edema, the color of skin, fatty necrosis might affect the cosmetic score of EB PBI. The 4 studies we included in our meta-analysis were EB APBI vs WBI/WBI-accelerated. The dose of fraction of APBI mostly higher than WBI/WBI-accelerated which might induce worse toxicity in breast tissue. Patients in EB APBI groups got less irradiation exposure to normal tissue lead to less acute skin toxicity (≦1 grade). A higher dose of fraction and less irradiation exposure tissue affected cosmetic score and late skin toxicity. Livi L et al's study was the source of heterogeneity for acute skin toxicity because it has a minimum proportion of patients with acute skin toxicity in the EB PBI group. Moreover, all 3 studies show EB PBI would cause less acute skin toxicity (≦1 grade) than WBI.

The position of tumor bed will atrophy after surgery. The accuracy of tumor bed in EB PBI was not similar to branch therapy and IORT. Moreover, incomplete coverage of potential tumor targets and increased exposure to normal tissues in EB PBI also needs to solve. The recommended PBI exposure range is 1-3cm around the tumor bed, it might incomplete coverage of potential target areas. The different center has a different target area delineation standard. Different medical personnel had different delineation target area. But EB PBI has the largest irradiation range than beachy PBI and IORT which bring EB PBI better prognoses than PBI and IORT. IMRT/VMRT can better fit the tumors than other technologies, the way of delivering irradiation will affect the results. Most PBI studies were about beachy PBI and IORT. But EB PBI was More economical, noninvasive, beautiful, easy to popularize.

In fact, increased irradiation exposure can kill potential cancer cells, but increased exposure to normal tissues such as heart, lung, and contralateral breast not necessarily lead to better outcomes. For PBI, the accurate selection of the population, the expansion of irradiation exposure range, and the adjustment of exposure mode still need to be further explored. There needs more study on EB PBI to help more and more early breast patients benefit from PBI.

**Conclusion**
This systematic review and meta-analysis found that EB PBI has similar LRR, regional node recurrence, contralateral breast cancer, distant recurrence, non-breast second cancer, and mortality with WBI. EB PBI has less acute skin toxicity (≤ 1 grade), similar late skin toxicity, and worse cosmetic score than WBI. EB PBI has a huge application prospect of clinical for early breast cancer patients. More studies about EB PBI are needed in the future.

**Declarations**

**Ethical Approval and Consent to participate**

Not applicable. All data are from published papers.

**Consent for publication**

The authors agree to publish the article.

**Availability of supporting data**

All the data are from published papers.

**Competing interests**

The authors declare no conflict of interest.

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**Authors’ contributions**

Peiling Dai and Kai Chen completed the writing and partly design of the article. Lan Li and Li Wang completed literature retrieval and data collection. Yaoxiong Xia, Yu Hou and Lan Zhang completed the revision of the article. Li Chang and Wenhui Li completed the design of the article.

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Synopsis: EB PBI has similar LRR, regional node recurrence, contralateral breast cancer, distant recurrence, non-breast second cancer and mortality with WBI. But EB PBI has worse cosmetic score, less acute skin toxicity (Ⅰ grade) and late skin toxicity than WBI.

Data Availability Statement

all data or used during the study are available in a repository or online in accordance with funder data retention policies (DOIs: 10.1016/j.ijrobp.2013.08.046; 10.1016/j.ejca.2014.12.013; 10.1016/s0140-6736(17)31145-5; 10.1016/s0140-6736(19)32515-2.)

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Figures

Reviewer 1

relevant references of randomized controlled trials were identified through database searching (Pubmed: 1522, Emtree: 972, Cochrane Library: 92, Clinical trials 44)

188 references after duplicates were removed

2442 references were screened

2436 of records excluded

Reviewer 2

6 of full-text articles assessed for eligibility

1 of full-text article without clear data was excluded. 1 of re-published article was excluded

4 of studies included in qualitative synthesis

Figure 1

Study selection scheme.
Figure 2

Forest plots for outcomes. A: Local recurrence rates; B: Regional node recurrence; C: Contralateral breast cancer; D: Distant recurrence; E: Non-breast second cancer; F: Mortality; G: Cosmetic score; H: acute skin toxicity (≥1 grade); I: late skin toxicity.