Results of a nationwide survey on Japanese clinical practice in breast-conserving radiotherapy for breast cancer

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ABSTRACT

The Breast Cancer Group of the Japanese Radiation Oncology Study Group conducted a nationwide questionnaire survey on the clinical practice of postoperative radiotherapy for breast-conserving treatment for breast cancer. This questionnaire consisted of 18 questions pertaining to the annual number of treated patients, planning method, contouring structure, field design, dose-fractionated regimen, application of hypofractionated radiotherapy, boost irradiation, radiotherapy for synchronously bilateral breast cancer, and accelerated partial breast irradiation. The web-based questionnaire had responses from 293 Japanese hospitals. The results indicated the following: treatment planning is performed using relatively similar field designs and delivery methods; the field-in-field technique is used at more than one-third of institutes; the commonest criteria for boost irradiation is based on the surgical margin width (≤5 mm) and the second commonest criteria was age (≤50 years), although some facilities applied a different age criterion (>70 years) for omitting a tumor bed boost; for conventional fractionation, almost all institutes delivered 50 Gy in 25 fractions to the conserved whole breast and 10 Gy in 5 fractions to the tumor bed. This survey revealed that 43% of hospitals offered hypofractionated radiotherapy, and the most common regimens were 42.56 Gy in 16 fractions for whole-breast irradiation and 10 Gy in 5 fractions for boost irradiation. Almost all of the facilities irradiated both breasts simultaneously for synchronously bilateral breast cancer, and the most common regimens were 42.56 Gy in 16 fractions for whole-breast irradiation and 10 Gy in 5 fractions for boost irradiation. Thissurvey revealed that 43% of hospitals offered hypofractionated radiotherapy, conserving treatment nationwide questionnaire survey; Japanese clinical practice; postoperative radiotherapy; breast-conserving treatment

Keywords: nationwide questionnaire survey; Japanese clinical practice; postoperative radiotherapy; breast-conserving treatment
INTRODUCTION
For patients with early-stage breast cancer, radiotherapy to the conserved breast after breast-conserving surgery (post-BCS RT) reduces the risk of local recurrence and cancer-related death, with a potential survival benefit [1, 2]. For these reasons, post-BCS RT has gained widespread use for patients with early-stage breast cancer. Technological advances in radiation therapy and a deeper understanding of clinicopathological features of breast cancer have led to an increase in the variety of methods available for irradiating the conserved breast.

Currently, different options for radiotherapy are available for breast-conserving treatment. An increase in dose delivery methods has resulted from advanced techniques such as 3D conformal radiotherapy (3DCRT), the field-in-field technique (FIF) and intensity-modulated radiation therapy (IMRT). Some studies have shown that boost irradiation (BI) to the tumor bed brings better local control, especially in younger patients, although different clinical trials have used different dose-fractionation regimens [3, 4]. Such evidence may result in the different adaptability and dose-fractionation regimens for BI among different hospitals. Furthermore, hypofractionated radiation schedules have been offered more frequently based on the high levels of evidence for long-term safety and efficacy [5, 6]. A greater understanding of breast cancer biology would allow for optimizing the field design for post-BCS RT. Several studies have reported that accelerated partial breast irradiation (APBI) or partial breast irradiation (PBI) provides satisfactory outcomes in some categories of early-stage breast cancer [7, 8]. It is thus possible that, based on such reports, diverse radiotherapy (dose–fractionation regimens and targeted volumes) may be prevalent in clinical practice.

After the American College of Surgeons Oncology Group (ACOSOG) Z-0011 trial revealed the feasibility of milder surgical axillary management for patients with low-volume axillary disease, axillary dissection is being increasingly omitted in eligible patients (one or two positive sentinel lymph nodes, having undergone breast conservation, scheduled to receive postoperative radiotherapy and systemic adjuvant therapy) [9]. This landmark trial has altered the treatment paradigm for axillary management and has led to the recommendation to omit additional axillary surgery beyond sentinel lymph node biopsy in women meeting the ACOSOG Z0011 inclusion criteria by the National Comprehensive Cancer Network and the American Society of Clinical Oncology [10, 11]. However, the optimal radiation field for use in such patients with low-volume axillary disease treated by mild axillary management is still unclear, although a standard ‘tangent’ or a ‘high tangent’ is generally considered to be appropriate for axillary disease control. Such changes may lead to differences in targeted volumes for post-BCS RT.

Elucidating the details of the variation among hospitals in Japan may help improve medical practices and patient management, and bring to light new obstacles that need to be addressed. This information could also be used for future clinical trials and for helping to inform national health insurance policy. To assess whether variations exist in post-BCS RT practices among different Japanese institutions, the Breast Cancer Group of the Japanese Radiation Oncology Study Group (JROSG-BCG), with the support of the Japanese Society for Radiation Oncology (JASTRO), conducted a nationwide questionnaire survey on the clinical practice of post-BCS RT in Japan. The aim of this study was to report the results of the tendencies in the clinical practice of post-BCS RT in Japan from the nationwide survey.

MATERIALS AND METHODS
The JASTRO approved and supported the initiative to conduct the questionnaire survey. This survey was conducted with institutional ethical approval. The JROSG-BCG formulated a web-based questionnaire system and requested the participation of all Japanese institutes with ≥1 full-time radiation oncologist and with an annual record of treating ≥10 patients with post-BCS RT in clinical practice. Radiation oncologists were requested to answer 18 questions related to post-BCS RT in six categories as follows (see Supplementary Data 1 for details of the questionnaire): (i) the number of the patients treated annually (three questions); (ii) the planning method/the contouring structure (six questions); (iii) the field design (one question); (iv) the dose-fractionated regimen/the practice of hypofractionated radiotherapy (four questions); (v) the criteria for applying BI (two questions); and (vi) the practice of bilateral post-BCS RT/APBI (two questions). Open access to the questionnaire was available during April, May, July and August of 2016, for a total of 4 months. For convenience, university hospitals and hospitals designated for cancer treatment were termed specialized hospitals (SHs) since they specialized in cancer treatment, whereas the other hospitals were termed general hospitals (GHs).

RESULTS
Completed questionnaires were received from 293 institutes. According to the statistical data of 2015 provided by the JASTRO Database Committee, the collection rate of this survey accounted for 55% (293/535) of the total number of institutes (535) that the JASTRO has authorized as facilities specializing in radiotherapy with ≥1 full-time radiation oncologist. Although 66% (192/293) of the institutes were GHs, we obtained responses from 89% (101/113) of SHs.

Number of patients treated annually
Figure 1 shows the number and ratio of patients treated annually with RT or post-BCS RT. In this study, 77% (225/293) of hospitals conducted RT for ≤600 patients; 76% (224/293) of the included facilities conducted post-BCS RT for ≤90 women.

Treatment planning/Contouring for treatment planning
Computed tomography (CT)-based 3DCRT was performed at 98% (288/293) of the included hospitals. In 92% (271/293) of all hospitals, treatment plans were calculated with inhomogeneity correction and with the use of accurate and novel algorithms such as the superposition algorithm. To make homogenous dose distributions, 57% (167/293) of hospitals used physical or dynamic wedge filters and 39% (114/293) employed the FIF technique as the first approach used commonly in each hospital (Fig. 2). The FIF technique was the most commonly used alternative method to homogenize dose distributions (36%, 106/293), if the first approach failed to do it. Among the institutes where no alternative method was generally employed, 47% (45/95) of them used the FIF technique as the first approach. For contouring of the target and the organs at risk (OARs), 66% (192/293) of all institutes formulated a contour of
the converted breast for the clinical target volume (CTV), while 66% (192/293) and 76% (222/293) of all institutes delineated the regions of interest such as the heart and the lung to evaluate the dose distributions, respectively.

Field design of post-breast-conserving surgery radiotherapy
The questionnaire contained questions about the most popular landmarks selected to define the edges of a treatment field for post-BCS RT in each hospital (see Supplementary data 2 for details of the survey result, Figs S1–S6). The middle line was the most common medial edge of the field (71%, 209/293). For the lateral edge, 50% (146/293) applied the boundary from the middle axillary line to posterior, though 4% (11/293) employed the more posterior landmarks such as the posterior axillary line or the anterior aspect of the ipsilateral latissimus. For the cranial edge, the sternal notch was selected by 55% (162/293) of hospitals. For the caudal edge, 64% (188/293) of facilities used the line 1–2 cm below the inframammary fold. The line ≥1 cm anterior to an ipsilateral nipple was the most commonly selected anterior edge for 86% (251/293) of all included institutions, while the posterior edges were based on the central lung distance (CLD) in 69% (202/293) of hospitals; this was the distance between the anterior edge of the lung and the midpoint of the posterior field edge. For the posterior edge, 25% (73/293) of institutes selected a CLD of <2 cm, 33% (96/293) selected a CLD of <2.5 cm, and 11% (33/293) selected a CLD of <3 cm.

Dose and fractionation; criteria for hypofractionated whole-breast irradiation
All participating facilities answered the question about the use of conventional fractionated (CF) and hypofractionated (HF) regimens, but we did not receive the detailed information on the HF dose fractionation from two facilities. Table 1A shows the total dose, dose per fraction, and the total fractions in CF post-BCS RT (dose in fraction <2.5 Gy). With the exception of four institutes that did not employ the CF regimen, 98% (284/289) of all hospitals used a protocol of 50 Gy in 25 fractions for whole-breast-irradiation (WBI) and 92% (266/289) used a BI of 10 Gy in 5 fractions. Table 1B describes the variety of doses and fractions of HF post-BCS RT (dose in fraction ≥2.5 Gy). Compared with CF regimens, there was a wider dispersion in HF schedules among different institutes, especially in those for the dose fractionations of BI. A total of 38% (111/293) of the hospitals employed HF regimens for WBI of 40–43.2 Gy in 16 fractions; the protocol of 42.56 Gy in 16 fractions was used in 30% (89/293) of hospitals and was the most common of all HF regimens. Eleven hospitals did not provide BI in HF schedules. In this study, 43% (127/293) of all hospitals conducted HF post-BCS RT and 11% (33/293) of facilities performed HF post-BCS RT more frequently in clinical practice than CF post-BCS RT (Fig. 3). Regarding the eligibility criteria for HF post-BCS RT, 40% (51/127) of the institutes employed the criteria recommended by the Japanese Breast Cancer Society (JBCS) [12]. The criteria included the following: (i) aged ≥50 years at diagnosis; (ii) pathologic stage of T1–2 N0 and receipt of breast-conserving surgery; (iii) no history of systemic chemotherapy; (iv) within the breast, the minimum dose was ≥93% and the maximum dose was ≤107% of the prescription dose.

Criteria to apply boost irradiation
This survey asked the participating institutes to list the eligibility criteria for offering BI to the tumor bed. All participating hospitals answered the question regarding the eligibility criteria, but in the analysis of surgical margin conditions, we used 89% (262/293) of answers regarding the superficial margin close to the skin or the deep margin close to the muscle and 95% (278/293) of answers regarding the lateral (horizontal) margin, because these answers were clear enough to analyze. Table 2 shows the factors of each institute taken into consideration in determining whether to use BI and the dominant factors that had a strong impact on the decision-making process. Surgical margin was the most commonly used determining factor. Overall, 97% (283/293) of hospitals considered surgical margin as an important factor for decision-making, and 94% (275/293) of the facilities regarded surgical margin as a dominant factor, although the definition of surgical margin as a determining
factor differed between the hospitals in this survey (Table 3). For the
criteria, some institutes used the margin width, whereas others
referred to the margin status (‘close’ or ‘positive’) on the histo-
logical reports. As for the lateral margin, 70% (194/278) of the hos-
pitals regarded a ≤5-mm margin width as a determinant for the
choice of BI. For the superficial margin close to the skin or the
depth margin close to the muscle, the most commonly used marginal
determinant for BI was also the ≤5-mm margin width (45%, 119/
262 or 43%, 113/262, respectively). Age was the second factor used
commonly in the decision about BI. In 38% (110/293) of facilities,
the common thresholds were the analysis of 97 answers reporting clear information on the thresh-
as a dominant factor in 19% (55/293) of hospitals. According to
the analysis of 97 answers reporting clear information on the thresh-
holds of age for BI, the common thresholds were ≤40 and ≤50 years
(36%, 35/97 and 52%, 50/97, respectively), but four facilities used
age >70 years as the age range for omitting BI (see Supplementary
data 2, Table S1).

Method for the bilateral post-BCS RT/practice of APBI
In patients with synchronously diagnosed bilateral breast cancers, 85%
(250/293) of all hospitals delivered the post-BCS WBI to the bilateral
conserved breast without any interval between irradiation to the right
and the left sides. A total of 7% (21/293) of the hospitals conducted
these procedures with an interval of a few weeks between procedures,
4% (13/293) of the facilities offered these procedures sequentially, and
the other institutes used their institutional procedures. Only 2% (5/
293) of the institutes offered APBI as part of clinical practice, and 1%
(3/293) offered this procedure as part of clinical trials. In this study,
97% (284/293) of all hospitals in Japan did not offer APBI.

DISCUSSION
As far as we know, this nationwide web-based questionnaire survey
is the first investigation to reveal the Japanese clinical practice of
post-BCS RT with an analysis of radiation planning. All of the 293
participating institutes responded, and the survey results contain the
opinions of the majority of the SHs (89%, 101/113). Therefore, the
results of this survey likely indicate the tendencies of current
Japanese clinical practice in post-BCS RT.

Almost all hospitals conducted post-BCS RT using 3DCRT planning
based on CT simulation with inhomogeneity correction and with rela-
tively novel calculation algorithms (e.g. superposition). There were no
major discrepancies in field designs between different institutions,
although some minor differences were noted. The contours of the con-
served breast, heart, and lung were delineated for precise dose delivery
in more than half of the included institutes. This survey revealed the popu-
ularity of the FIF technique for achieving homogenous dose delivery. This
result might be a reflection of the prevalence of 3DCRT in Japan, which
might have helped increase awareness of the efficacy of the FIF technique
among clinicians. The importance of ensuring homogeneity of dose deliv-
ery to the conserved breast has been widely recognized and will likely
lead to a more widespread use of the FIF technique. Furthermore, the
updated guideline of an ASTRO, which recommended 3D conformal
treatment planning with the FIF technique, might promote the spread of
the FIF technique in Japan [6].

This investigation revealed the uniformity in the schedules of
CF post-BCS RT and the frequency of HF post-BCS RT use among
Japanese institutes. CF schedules in almost all institutes used 50 Gy
in 25 fractions (5 days per week) for WBI and 10 Gy in 5 fractions
for BI. HF schedules were more varied than those of CF and dif-
f ered in total doses and dose fractionations. The most common
regimens were 42.56 Gy in 16 fractions (5 days per week) for WBI and
10.64 Gy in 4 fractions for BI. Our results showed that 43% of
hospitals employed the HF regimen, and 11% offered the HF sched-
ule more frequently than they did the CF schedule. In the UK, the
HF regimen (40 Gy in 15 fractions) has been adopted widely based
on the long-term follow-up results of randomized controlled trials
[13]. Several studies have revealed the prevalence of the HF regi-
men in Canada and the USA [14, 15]. The HF regimen was not
used commonly in Japan compared with European countries such as
the UK or the Netherlands. One of the reasons might be that the
results of the JCOG 0906, a Japanese multicenter safety trial of HF
post-BCS RT [16], had not commenced at the start of this survey.
We speculate that the HF regimen has likely gained (and will gain
more) popularity in Japan, based on the increased awareness of its efficacy [17, 18], the future announcement of the JCOG0906 result, and the ASTRO’s updated guideline, which contained more flexible eligibility criteria for use of the HF regimen than the JBCS criteria [6].

According to this survey, surgical margin was the most common determining factor for selection of BI, and age was the second factor, although different criteria were adopted in different institutes. Of the Japanese institutes in this survey, 98% considered surgical margin as an important factor, and 94% regarded it as the dominant factor to be used solely in determining the selection of BI. Many of the facilities in this survey used the ≤5-mm margin width in determining the selection of BI. The EORTC 22881–10882 trial clearly revealed the benefit of BI in terms of local control in patients with microscopically complete excision of invasive disease, although BI could lead to worse cosmetic outcomes or severe fibrosis without an apparent impact on survival benefit [3, 4]. The Japanese tendency to base the use of BI on margin width might be related to the hypothesis that a narrower margin is associated with a risk of local recurrence and the concern that BI could cause a worse cosmetic outcome. However, the panel of the Society of Surgical Oncology and ASTRO concluded that choosing BI should not be dependent

Table 1. The list of doses and fractionations for whole-breast irradiation and boost irradiation, and the percentages of hospitals using this approach

| Total dose (Gy) | Does/Fr (Gy) | Total Fr | The number (%) of all hospitals | The number (%) of specialized hospitals |
|-----------------|--------------|----------|---------------------------------|----------------------------------------|
| **A. Conventional fractionation** | | | | |
| WB | 50 | 2 | 25 | 286 (98) | 101 (100) |
| | 50.4 | 1.8 | 28 | 2 (0.7) | 0 (0) |
| | 47.5 | 2.375 | 20 | 1 (0.3) | 0 (0) |
| No regimen | | | | | 4 (1) |
| Total number | | | | | 293 |
| Boost | 10 | 2 | 5 | 267 (92) | 94 (93) |
| | 9 | 3 | 3 | 8 (3) | 3 (3) |
| | 16 | 2 | 8 | 6 (2) | 3 (3) |
| 4–15 | 2–3 | 2–7 | 8 (3) | 1 (1) |
| No regimen | | | | | 4 (1) |
| Total number | | | | | 293 |
| **B. Hypofractionation** | | | | |
| WB | 40.05 | 2.67 | 15 | 5 (2) | 3 (3) |
| | 40–43.2 | 2.5–2.7 | 16 | 111 (38) | 41 (41) |
| | 42.5–50 | 2.5 | 17–20 | 9 (3) | 1 (1) |
| No regimen | | | | | 166 (57) |
| Total number | | | | | 291 |
| Boost | 5–9 | 2.5–3 | 3 | 23 (7.9) | 2 (2) |
| | 10–10.64 | 2.5–2.66 | 4 | 69 (23.7) | 28 (28) |
| | 10–15 | 2–2.66 | 5–6 | 22 (7.6) | 9 (9) |
| No regimen | | | | | 177 (60.8) |
| Total number | | | | | 291 |

Fr = fractionation; WB = whole breast.
on the margin width [19]. Their meta-analysis showed that there was a weak relationship between local recurrence, margin width and dose in contemporary multimodality treatment, and concluded that in patients with negative margins (no ink on the tumor), the use and dose of BI should be based on an a priori estimation of local recurrence risk in the ipsilateral converted breast, and should not be determined, in isolation, by the width of the surgical margin. According to their report, to estimate the a priori risk of local recurrence, not only margin width but also the other factors (age, subtype, the use of systemic therapies, etc.) should be taken into account. This survey showed that some institutes regarded the other factors (age, subtype, etc.) as determining factors for the selection of BI, but (in 2016) many institutes seemed to put the emphasis on margin width in selecting BI in Japan. In the future, the use of all of the criteria related to the a priori risk of local recurrence is likely to spread in Japan, with novel and future evidence for the benefits of optimizing the balance between the benefit and burden of BI. The changes in the Japanese trends in the criteria for selecting BI are worthy topics for future studies.

In regard to age, 38% of Japanese institutes in this survey regarded age as an important factor and 19% regarded it as the dominant factor to use in selection of BI. Many institutes used the age range of ≤40 or ≤50 years as the criterion for applying BI, as young age is associated with local recurrence. The Oxford meta-analysis of breast-conserving surgery demonstrated an inverse relationship between the rate of any first recurrence and age [1]. The EORTC 22881–10882 trial showed that the relative benefit of BI for local control was larger in younger patients than in older patients, and the absolute risk reduction at 20 years was the largest in patients ≤40 years of age (36.0% in the no boost group versus 24.4% in the boost group) [3]. Japanese institutes seemed to use the age range of ≤40 or ≤50 years based on the above evidence.

A majority (85%) of the institutes included in the survey performed bilateral post-BCS RT synchronously in patients with synchronously diagnosed bilateral breast cancers, indicating the high prevalence of this procedure in Japan. This survey also revealed that most of the institutes (98%) did not perform APBI in clinical practice, and thus APBI may not be the method of choice for post-BCS RT in Japan. One of the reasons for this is likely to be that the Japanese Breast Cancer Society Clinical Practice Guideline for Breast Cancer had recommended APBI should be performed in a clinical trial [12]. It is also possible that this tendency may be due to the variation in the outcomes of APBI in different studies and the questionable superiority of APBI over WBI [20]. However, APBI or PBI could be available for selected patients in the near future, based on newly accumulating evidence [8, 21, 22]. Changes in the Japanese trends regarding the use of APBI and PBI would be worthy topics for future studies.

This survey has some limitations. First, this questionnaire was not answered by all eligible institutes in Japan (institutions with ≥1 radiation oncologist working full-time and with ≥10 patients treated with post-BCS RT annually), which implies that the results may not be representative of clinical practices of post-BCS RT in Japan. However, the majority of the SHs (which are the primary cancer treatment centers in Japan) participated in this survey. Therefore, we believe that this survey presents an accurate overview of the tendencies in the clinical practice of post-BCS RT in Japan. Second, the respondents had to select the most appropriate answers from several options in this questionnaire, which may have unintentionally guided them. Third, this survey disclosed the landmarks of field edges for treatment planning, although this information might be insufficient for detecting precise differences in dose distributions on the conserved breasts as applied by the different institutes. Future investigations would need detailed digital imaging data to better understand the variation in prevalent radiation therapy.

The findings of this nationwide survey provided an overview of the current clinical practices of post-BCS RT in Japan and the

| Elements                        | Percentage of hospitals (%) | An important factor | A dominant factor |
|---------------------------------|----------------------------|--------------------|-------------------|
| Age                             | 38                         | 19                 |                   |
| Surgical margin condition       | 97                         | 94                 |                   |
| Nodal metastasis                | 5                          | 2                  |                   |
| Lymphovascular invasion         | 8                          | 0.3                |                   |
| Subtypes of breast cancer       | 6                          | 2                  |                   |
| Others                          | 7                          | 6                  |                   |

*A dominant factor is defined as a factor having a strong impact on the decision to apply BI; i.e. the use of BI could be determined solely by the dominant factor. BI = boost irradiation.
results can be compared with those of future investigations into Japanese practice of post-BCS RT. This study has provided not only the features of current Japanese practice of post-BCS RT in 2016, but also topics for future studies.

SUPPLEMENTARY DATA
Supplementary data are available at Journal of Radiation Research online.

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CONFLICT OF INTEREST
The authors declare that there are no conflicts of interest.

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Table 3. The diversity of the eligibility criteria for choice of boost irradiation: percentage of hospitals arranged according to the margin width or margin status (in each direction)

| Eligibility condition of surgical margin | Percentage of hospitals (%) | | | |
|-----------------------------------------|----------------------------|---|---|---|
|                                         | Superficial margin | Deep margin | Lateral margin |
| Margin widtha ≤1 mm                     | 3                          | 2                          | 2                          |
| Margin widtha ≤2 mm                     | 3                          | 3                          | 3                          |
| Margin widtha ≤5 mm                     | 45                         | 43                         | 70                         |
| Margin widtha = 0 mm                    | 26                         | 29                         | 5                          |
| Margin status (close margin)b           | 5                          | 5                          | 5                          |
| Margin status (positive margin)b        | 13                         | 12                         | 12                         |
| Others                                  | 4                          | 5                          | 2                          |

*The ‘margin width’ was the distance between the area of the tumor invasion and the specimen edge.

*The ‘margin status’ (close margin or positive margin) was provided by the histological report (close margin or positive margin).
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