

**Hypofractionation in Breast Cancer - A Retrospective Study in a Tribal
Population Based Medical College in West Bengal, India**

ABSTRACT

INTRODUCTION: In a tribal population based area in West Bengal, India though carcinoma cervix is the commonest malignancy in female patients, yet apart from that carcinoma breast is also increasing in number in the recent years. Breast cancer accounts for approximately 26.6% of female malignancy in the radiation oncology out-patient-department of our teaching hospital.

AIMS and OBJECTIVES: To compare conventional RT regimen (50 Gy in 25 fractions over 5 weeks) with one hypofractionated regimen (40Gy in 15 fractions over 3 weeks) in stage II & stage III breast cancer patients as adjuvant radiation therapy in terms of local control, survival and adverse reactions.

MATERIALS and METHODS: It is a retrospective study which has been conducted in the department of Radiotherapy in BSMC (Bankura Sammilani Medical College) spanning from May 2012 to April 2017. A total number of patients included in this study was 302, out of which thirty six patients failed to follow up. So total of 266 patients included in the study were all histologically proved carcinoma breast treated surgically (97.74% by MRM & rest by BCS) with curative intent following which RT was used as adjuvant therapy. In one group (consisting of 133 patients) conventional regimen (50Gy in 25 fractions) was used. In another group (consisting the other 133 patients) dose-schedule used was a hypofractionated one i.e. 40Gy in 15 fractions. Dose per fraction in the 1st group was 2 Gy whereas in 2nd group it was 2.66 Gy. In all patients, RT was given in 5 days a week. Systemic therapy was administered as and when indicated.

RESULT: 4-year disease-free-survival (DFS) in conventional group was 78.94% and in hypofractionated group was 82.70%, (p value >0.05). 4-year overall survival (OS) in

28 conventional group was 81.20% & in hypofractionated group was 85.70%, (p value >0.05).
29 While adverse reactions in terms of both acute & chronic radiation toxicities were
30 considered, there was no significant difference in between the two groups.

31 **CONCLUSION:** There is no significant difference between the conventional regimen and this
32 hypofractionated regimen in terms of OS DFS & adverse reactions in this tribal-based Indian
33 population. Hence, in our institution, we usually prefer Hypofractionated radiotherapy
34 (40Gy/15 fractions) in adjuvant settings for breast cancer patients.

35
36 **Keywords:** Hypofractionation, Breast cancer, Ca Breast.

37 1. INTRODUCTION

38 As we are aware of the fact that radiotherapy is a mandatory modality in the course of
39 treatment for Carcinoma of Breast, various dose prescriptions aside the conventional one
40 had also been tried in particularly adjuvant setting [1]. The goal was to find out an optimum
41 dose prescription by dint of which adequate local control could be achieved respecting the
42 acute and late toxicities. Though breast cancer awareness programs and thorough
43 screening have succeeded enough in developed countries in terms of early diagnosis, in
44 developing countries like India diagnosis at an early stage and early commencement of
45 treatment remain still a challenge [2]. Our practice domain includes a rural-based area i.e.
46 Bankura in West Bengal, India where carcinoma cervix is still the commonest malignancy
47 followed by ca breast as the second commonest malignant entity in the female population.
48 But according to the records of recent years preserved by the Department of Radiation
49 Oncology of Bankura Sammilani Medical College & Hospital, an increase in the incidence of
50 breast cancer is a burning fact. Currently, breast cancer accounts for 26.6% of female
51 malignancies in this area, as recorded, majority of which presented as Locally Advanced
52 Breast Cancer (LABC), with AJCC stage T2 - 4, any N. As recommended, multidisciplinary
53 approach including neoadjuvant chemotherapy (NACT), surgery, adjuvant radiotherapy,
54 adjuvant chemotherapy, hormonal therapy and immunotherapy form the lines of treatment

55 considering all patient factors, disease factors and treatment factors. Modified radical
56 mastectomy (MRM) dominates over Breast Conservation Surgery (BCS) with a statistic of
57 97.74% vs. 2.26% [3]. Due to the belief that removal of the entire diseased breast is
58 mandatory to cure cancer they always opted for MRM even in those favourable cases where
59 BCS might be a better option in term of cosmesis. However our study dealt with adjuvant
60 radiotherapy, which was aimed to compare the so-called conventional breast RT regimen
61 (50 Gy in 25 fractions over 5 weeks) with one hypofractionated regimen (40Gy in 15
62 fractions over 3 weeks) in stage II & stage III breast cancer patients as adjuvant therapy in
63 terms of local control, survival and adverse reactions.

64

65 **2. MATERIALS AND METHODS**

66 2.1 Patients and Methods

67 In this single institutional retrospective study total 302 consecutive patients who got registered
68 between May 2012 and April, 2017 in the outpatient department of Radiotherapy in BSMC(Bankura
69 Sammilani medical college and Hospital) were included. Out of which thirty six patients failed to
70 follow up; so total 266 patients were included in the study finally. After clinical evaluation including
71 local and locoregional examination of bilateral breast and axillae a complete mammogram with
72 proper BIRADS scoring was done. It was followed by a tru-cut biopsy confirming the pathological
73 diagnosis of invasive breast cancer. As fine needle aspiration cytology sample does not suffice to
74 perform immunohistochemistry, tru-cut biopsy was a mandatory inclusion criteria. It was followed
75 by immunohistochemistry stating the oestrogen and progesterone receptor status and HER2 neu
76 amplification status too. Ki 67 was not routinely done in our public hospital before 2014, hence
77 Modified Nottingham Prognostic Index (NPI) Scoring was considered significant to determine the
78 grade of aggressiveness of the infiltrative carcinoma. It was followed by complete metastatic
79 workup including a digital chest X ray sometimes an additional Contrast Enhanced Computed
80 Tomography (CECT) Scan of Thorax , a CECT Scan of the whole abdomen. A Magnetic
81 Resonance Imaging of brain was performed in symptomatic patients with the suspicion of brain
82 metastasis. Patients who were clinical, AJCC anatomic prognostic stage group IIA, IIB, IIIA, IIIB
83 and IIIC were included. Simply, T-stages included were T2- T4 and N-staged included were N0-N3.

84 Significant baseline characteristics used for 1:1 patient matching included history regarding age
85 (<50 years vs. >50 years; no more than 3 years apart), menopausal status (premenopausal vs.
86 postmenopausal), number of relatives affected (1st degree vs. 2nd degree vs. no family history).
87 BRCA 1 and BRCA 2 mutation analysis was not routinely done in our institution. Disease-related
88 factors for patient matching were T-stage, N-stage, AJCC Prognostic stage group, NPI Score,
89 status of post-surgery histopathological examination (HPE) report, ypT and ypN status as patients
90 received Neo Adjuvant Chemotherapy regimens, Hormonal Receptor status, Her-2neu status etc.
91 Other minor factors like age at first child birth (no more than 2 years apart), duration of
92 breastfeeding (obtained from parity), the month that patients received the treatment in question i.e.
93 radiation therapy (no more than 6 months apart) were attempted to match afterwards.

94

95 2.2 Treatment Protocol

96 For selected patients with early breast cancer (EBC) and Large Operable Breast Cancer
97 (LOBC) who were referred for NACT from department of surgery and all LABC patients
98 proper pre-treatment work up including complete blood count, kidney function test, liver
99 function test, diabetic profile, serology and cardiological fitness including echocardiography
100 and electrocardiogram was done. These patients received Taxane based (majority) or
101 Anthracycline Based NACT regimens to achieve downstaging depending on the
102 immunohistochemistry report obtained from true-cut biopsy paraffin blocks. After 14 days
103 following the completion neo-adjuvant chemotherapy the patient was assessed for radical
104 intervention i.e. modified radical mastectomy (MRM) or BCS. After surgery histopathological
105 examination reports were scrutinised for indications for Post Mastectomy Radiation Therapy
106 (PMRT). Finally, adjuvant radiation was planned. All these patients were subdivided into two
107 Groups on the basis of radiation dose-fractionation. The first group was treated with adjuvant
108 Radiation Therapy (RT) with 50Gy in 25 fractions over 5 weeks, i.e. conventional
109 fractionation; while the other group received 40Gy in 15 fraction over 3 weeks, i.e.
110 hypofractionation. Dose per fraction were 2 Gy and 2.66 Gy, respectively. Adjuvant
111 chemotherapy, Hormonal therapy, and Her-2 directed biologic therapy were administered as
112 and when applicable abide by standard evidence-based guidelines. Follow up was done

113 three months according to our institutional protocol. Further treatment included lines of
114 chemotherapies and palliation.

115

116 2.3 Response Assessment

117 After completion of radiation therapy, clinical examination of bilateral breasts and axilla and
118 high-resolution ultrasonography of ipsilateral chest flap, contralateral breast and bilateral
119 axillae was done after 2 months. A chest X-ray and a CECT whole abdomen was done 3
120 monthly. MRI brain was performed on the basis of presenting symptoms as and when
121 required. RECIST v1.1 criteria was used to determine complete response (CR), progressive
122 disease (PD), partial response (PR) or stable disease (SD) in consequent follow ups after
123 completion of treatment. Radiation toxicities (both acute and late) were assessed using
124 RTOG (Radiation Therapy Oncology Group) toxicity grading. Median disease-free survival
125 (DFS) or progression-free survival (mPFS) and overall survival (OS) were analysed using
126 Kaplan-Meier survival over a median follow up of 60 months.

127

128 2.4 Statistical Analysis

129

130 SPSS statistical software version 17 (IBM Corp., Chicago, IL, USA) was used for data
131 analysis. Quantitative data were presented by mean or median as appropriate, and
132 qualitative data were presented as a percentage. OS and PFS/DFS were analysed by the
133 Kaplan-Meier method and compared between both groups by log rank test ($p= 0.05$). The
134 Cox proportional hazards model was used to adjust all prognostic factors. A 2-sided p -value
135 <0.05 was considered statistically significant.

136

137 3. RESULTS

138 In this rural population-based retrospective study a total number of patients included was
139 three hundred two(302). Thirty six patients (36) failed to follow up. Hence, finally two
140 hundred sixty six patients (266) were evaluated for this study ($n = 266$). They have been
141 divided into two groups namely A & B. each containing 133 patients($n 133$). 1:1 patient

142 matching was done considering the criteria mentioned previously. In Group A conventional
 143 fractionation radiation therapy (CFRT) i.e. 50Gy in 25 fractions over 5 weeks was
 144 administered and in Group B hypofractionation radiation therapy (HFRT) i.e. 40Gy in 15
 145 fractions over 3 weeks dose-schedule was used as adjuvant treatment. Electron boost (10 to
 146 15 Gy) was done to the tumour bed where Breast conservation (BCS) performed (though in
 147 2.26% patients only) as primary surgical modality. Acute & chronic reactions were noted and
 148 recorded during & at the completion of radiotherapy & in subsequent follow ups.
 149 Locoregional recurrence (LRR) & Overall survival (OS) & Disease-free survival(DFS) were
 150 also documented. MRM was performed in 96.99% and 97.74 % of patients and BCS was
 151 done in 3.01% and 2.26% followed by boost in **Group A** and **Group B**, respectively. Most
 152 common histopathological variety was Infiltrating duct carcinoma.(84.96% in Group A and
 153 88.72% in Group B). Neoadjuvant chemotherapy was administered in all cases. Taxol based
 154 chemotherapy was used in 90.22% and 90.97% patients in Group A & in Group B,
 155 respectively. **Table 1** depicts patient characteristics and disease-related factors separately
 156 for Group A and Group B.
 157

Table 1. List of patient characteristics attributed during study

| PATIENT CHARACTERISTICS | |
|-------------------------|-----------------------------------|
| | GROUP "A" (CFRT) GROUP "B" (HFRT) |
| MEDIAN AGE | 46 YEARS 50 YEARS |
| TUMOR SIZE | |

| | |
|--|-------------------------|
| T2 | 35(26.3%) 43(32.3%) |
| T3 | 84(63.1%) 82(61.7%) |
| T4 | 14(10.6%) 8(6.01%) |
| LYMPHNODE STATUS | |
| N1 | 40(30.07%) 42(31.57%) |
| N2 | 81(60.90%) 84(63.1%) |
| N3 | 12(9.02%) 7(5.33%) |
| TYPES OF SURGERY | |
| Modified radical mastectomy (MRM) | 129(96.99%) 130(97.74%) |
| Breast Conservation Surgery (BCS) | 4(3.01%) 3(2.26%) |
| HISTOPATHOLOGY | |
| Invasive ductal carcinoma (IDC) | 113(84.96%) 118(88.7%) |

| | |
|----------------------------------|-------------------------|
| Invasive lobular carcinoma (ILC) | 16(12.02%) 12(9.0%) |
| Ductal carcinoma in situ (DCIS) | 4(3.01%) 3(2.2%) |
| NEOADJUVANT CHEMOTHERAPY | |
| TAXOLBASED | 120(90.22%) 121(90.97%) |
| NONTAXOL | 13(9.77%) 12(9.02%) |
| RECEPTOR STATUS | |
| ER+VE | 77(57.89%) 72(54.13%) |
| ER- VE | 56(42.10%) 61(46.86%) |
| PR+ VE | 55(41.35%) 54(40.60%) |
| PR- VE | 78(58.64%) 79(59.39%) |
| HER2NEU +VE | 35(26.31%) 40(30.07%) |
| HER2NEU - VE | 61(45.87%) 54(40.60%) |
| | |

| | |
|--------------------------|-----------------------|
| UNKNOWN/EQUIVOCAL | 37(27.82%) 39(29.33%) |
|--------------------------|-----------------------|

158 There was no significant difference between two Groups regarding radiation toxicity. Most
159 common acute toxicity was skin reactions. RTOG GRADE 1 skin reactions occurred in
160 62.4% patients in Group A & 60.15% patients in Group B. GRADE 2 of the same was
161 evident in 37.59% (for Group A) & 39.85% (for Group B). No grade 3 skin toxicity was
162 noted. (p-value >0.05 i.e. not statistically significant).
163 As recorded, GRADE 1 chronic skin reactions evident in Group A was 51.87% and in Group
164 B it was 53%. GRADE 2 of the same reaction was seen in 42.10% (Group A) & 50.36%
165 (Group B); p value >0.05. (Table 2)

166

167 **Table 2. Percentage of occurrence of skin reaction and chronic reactions in patients**

| SKIN REACTIONS (ACUTE) | GROUP "A"(CFRT) | GROUP "B" (HFRT) |
|-------------------------------|------------------------|-------------------------|
| GRADE 1 | 50(39.59%) | 53(39.8%) |
| GRADE 2 | 83(62.40%) | 80(60.2%) |
| GRADE 3 | 0 | 0 |
| (p>0.05) | | |
| SUBCUTANEOUS TISSUE | GROUP "A"(CFRT) | GROUP "B" (HFRT) |
| GRADE 1 | 71(53.38%) | 69(51.87%) |

| | | |
|----------------------------|------------------------|-------------------------|
| GRADE 2 | 62(46.62%) | 64(48.12%) |
| GRADE 3 | 0 | 0 |
| (p>0.05) | | |
| CHRONIC REACTIONS | | |
| SKIN REACTIONS | GROUP "A"(CFRT) | GROUP "B" (HFRT) |
| GRADE 0 | 5(3.78%) | 8(6.01%) |
| GRADE 1 | 69(51.87%) | 67(50.37%) |
| GRADE 2 | 56(42.10%) | 53(39.84%) |
| GRADE 3 | 3(2.25%) | 5(3.75%) |
| (p>0.05) | | |
| SUBCUTANEOUS TISSUE | GROUP "A"(CFRT) | GROUP "B" (HFRT) |
| GRADE 0 | 6(4.5%) | 5(3.75%) |

| | | |
|--------------------|-----------|------------|
| GRADE 1 | 74(55.6%) | 67(50.3%) |
| GRADE 2 | 50(37.6%) | 53(39.84%) |
| GRADE 3 | 3(2.2%) | 8(6.1%) |
| (p>0.05) | | |

168

169 From the statistical point of view, 4 year local control for the conventional Group (CFRT;
 170 Group A) is 86.46% and for the hypofractionated Group (HFRT; Group B) is 90.6%. (p
 171 value >0.05). 4 year overall survival in group A is 81.20% and in Group B it is 85.70% (p
 172 value >0.05). 4 year Disease-free survival in group A is 78.94 % and in Group B is 82.70%
 173 (p-value >0.05) (Table 3). So on the basis of OS, DFS & locoregional recurrence, there are
 174 no statistically significant differences lies between the two Groups.
 175

| | | |
|--|------------------------|-------------------------|
| Table 3. Data analysis showing significant relationship | | |
| SURVIVAL ANALYSIS (4 YEAR ANALYSIS) | GROUP "A"(CFRT) | GROUP "B" (HFRT) |
| OVERALL SURVIVAL | 108(81.20%) | 114(85.7%) |
| (p>0.05) | | |

| | | |
|--------------------------------------|-----------------|------------------|
| | GROUP "A"(CFRT) | GROUP "B" (HFRT) |
| DISEASE FREE SURVIVAL | 105(78.94%) | 110(82.71%) |
| (p>0.05) | | |
| LOCOREGIONAL CONTROL (4YEARS) | GROUP "A"(CFRT) | GROUP "B" (HFRT) |
| | 105 (78.94%) | 110(82.71%) |
| (p>0.05) | | |

176

177 Figure 1 shows a graphical representation of the probability of subclinical breast tumour
 178 control and normal tissue toxicity with increasing dose in Gy.

179

180

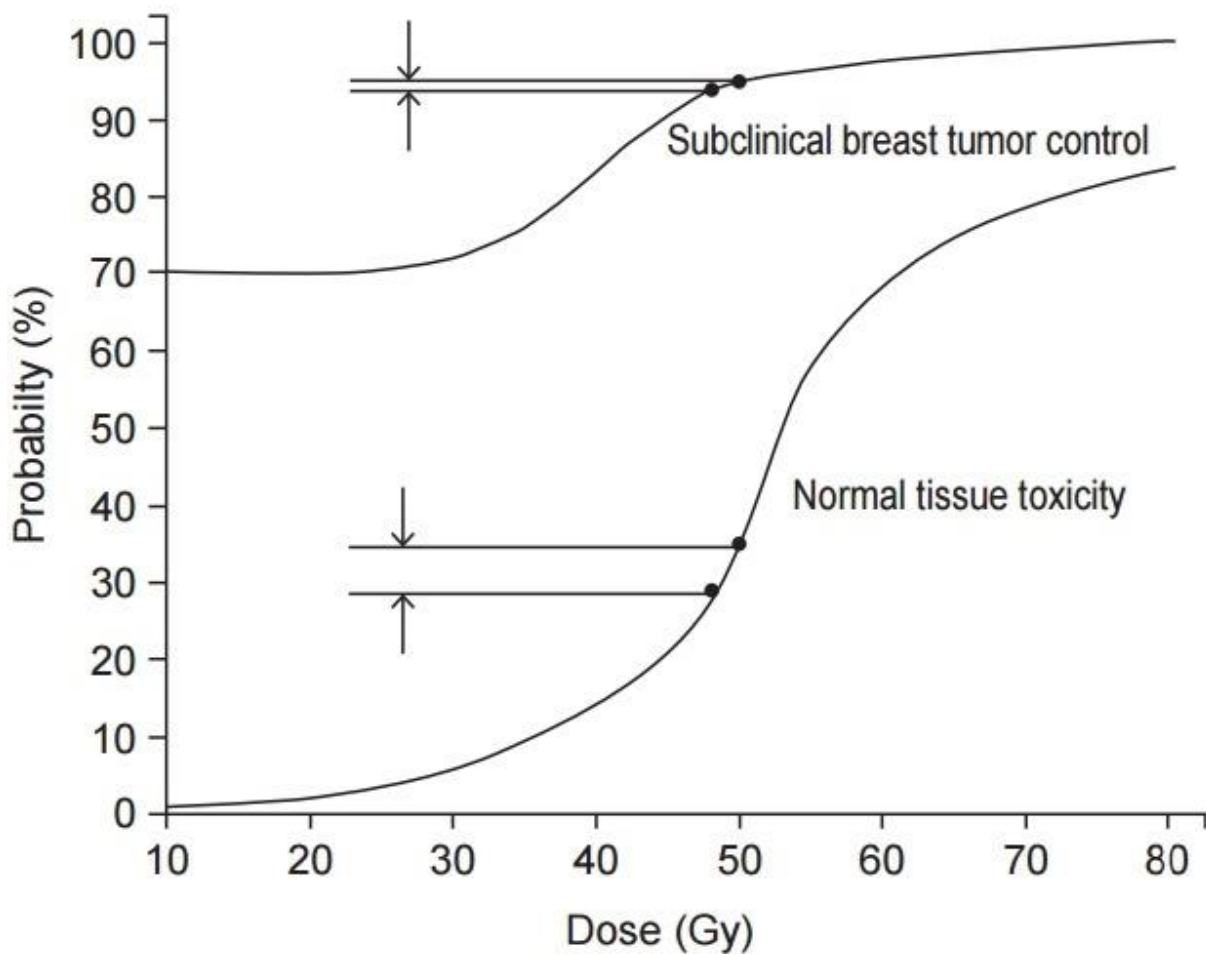


Figure 1

181

182

183

184 4. DISCUSSION

185 Hypofractionation in Carcinoma Breast was cultivated by several study groups from time to
186 time.

187 Whelan et al. [4] conducted Long-Term Results of Hypofractionated Radiation Therapy for
188 Breast Cancer study to determine whether a hypofractionated 3-week schedule of whole-
189 breast irradiation is as effective as a 5-week schedule. Women with invasive breast cancer
190 who had undergone breast-conserving surgery and in whom resection margins were clear
191 and axillary lymph nodes were negative were randomly assigned to receive whole- breast
192 irradiation either at a standard dose of 50.0 Gy in 25 fractions over a period of 35 days (the
193 control group) or at a dose of 42.5 Gy in 16 fractions over a period of 22 days (the
194 hypofractionated-radiation group). The study concluded, at 10 years, 71.3% of women in the

195 control group as compared with 69.8% of the women in the hypofractionated-radiation group
196 had a good or excellent cosmetic outcome (absolute difference, 1.5 percentage points; 95%
197 CI, -6.9 to 9.8) [4].

198

199 Between 1998 and 2002, 2236 women with early breast cancer (pT1-3a pN0-1 M0) at 17
200 centres in the UK were randomly assigned after primary surgery to receive 50 Gy in 25
201 fractions of 2.0 Gy versus 41.6
202 Gy or 39 Gy in 13 fractions of 3.2 Gy or 3.0 Gy over 5 weeks. 749 women were assigned to
203 the 50 Gy group, 750 to the 41.6 Gy group, and 737 to the 39 Gy group. After a median
204 follow up of 5.1 years (IQR 4.4–6.0) the rate of local-regional tumour relapse at 5 years was
205 3.6% (95% CI 2.2–5.1) after 50 Gy, 3.5% (95% CI 2.1– 4.3) after 41.6 Gy, and 5.2% (95%
206 CI 3.5–6.9) after 39 Gy. The estimated absolute differences in 5-year local-regional relapse
207 rates compared with 50 Gy were 0.2% (95% CI -1.3% to 2.6%) after 41.6 Gy and 0.9%
208 (95% CI -0.8% to 3.7%) after 39 Gy. Photographic and patient self-assessments suggested
209 lower rates of late adverse effects after 39 Gy than with 50 Gy, with an HR for the late
210 change in breast appearance (photographic) of 0.69 (95% CI 0.52–0.91, p=0.01). The study
211 concluded the data are consistent with the hypothesis that breast cancer and the dose-
212 limiting normal tissues respond to cancer and the dose-limiting normal tissues respond
213 similarly to change in radiotherapy fraction size. 41.6 Gy in 13 fractions was similar to the
214 control regimen of 50 Gy in 25 fractions in terms of local-regional tumour control [5].

215 Study conducted to test the benefits of radiotherapy schedules using fraction sizes larger
216 than 2.0 Gy in terms of local-regional tumour control, normal tissue responses, quality of life,
217 and economic consequences in women prescribed post-operative radiotherapy. 2215
218 women with early breast cancer (pT1-3a pN0-1 M0) at 23 centres in the UK were randomly
219 assigned after primary surgery to receive 50 Gy in 25 fractions of 2.0 Gy over 5 weeks or 40
220 Gy in 15 fractions of 2.67 Gy over 3 week. 1105 women were assigned to the 50 Gy group
221 and 1110 to the 40 Gy group. After a median follow up of 6.0 years (IQR 5.0-6.2) the rate of
222 local- regional tumour relapse at 5 years was 2.2% (95% CI 1.3-3.1) in the 40 Gy group and
223 3.3% (95% CI 2.2 to 4.5) in the 50 Gy group, representing an absolute difference of -0.7%

224 (95% CI -1.7% to 0.9%)--ie, the absolute difference in local-regional relapse could be up to
225 1.7% better and at most 1% worse after 40 Gy than after 50 Gy. The study interpreted 1105
226 women were assigned to the 50 Gy group and 1110 to the 40 were assigned to the 50 Gy
227 group and 1110 to the 40 Gy group. After a median follow up of 6.0 years (IQR 5.0-6.2) the
228 rate of local-regional tumour relapse at 5 years was 2.2% (95% CI 1.3-3.1) in the 40 Gy
229 group and 3.3% (95% CI 2.2 to 4.5) in the 50 Gy group, representing an absolute difference
230 of -0.7% (95% CI -1.7% to 0.9%)--ie, the absolute difference in local- regional relapse could
231 be up to 1.7% better and at most 1% worse after 40 Gy than after 50 Gy [6].

232 Owen JR in his randomized trial, tested whether fewer, larger fractions were at least as safe
233 and as effective as standard regimens. In this analysis, also assessed the long-term results
234 of tumour control in the same population. In this study 1410 women with invasive breast
235 cancer (tumour stage 1-3 with a maximum of one positive node and no metastasis) who had
236 had local tumour excision of early-stage breast cancer were randomly assigned to receive
237 50 Gy radiotherapy given in 25 fractions, 39 Gy given in 13 fractions, or 42.9 Gy given in 13
238 fractions, all given over 5 weeks. The primary endpoint was a late change in breast
239 appearance, which has been reported elsewhere. 1410 women with invasive breast cancer
240 (tumour stage 1-3 with a maximum of one positive node and no metastasis) who had had
241 local tumour excision of no metastasis) who had had local tumour excision of early stage
242 breast cancer to receive 50 Gy radiotherapy given in 25 fractions, 39 Gy given in 13
243 fractions, or 42.9 Gy given in 13 fractions, all given over 5 weeks. The primary endpoint was
244 late change in breast appearance, which has been reported elsewhere. The study concluded
245 Breast cancer tissue is probably just as sensitive to fraction size as dose-limiting healthy
246 tissues [7].

247

248 Yarnold et al. [8] in his study randomized one thousand four-hundred and ten women with
249 T1-3 N0-1 M0 invasive breast cancer into one of three radiotherapy regimens after local
250 tumour excision of early stage breast cancer; 50 Gy in 25 fractions (F) vs two dose levels of
251 a test schedule giving 39 or 42.9 Gy in 13 F over 5 weeks. Fraction sizes were 2.0, 3.0 and
252 3.3 Gy, respectively. After a minimum 5-year follow up, the risk of scoring any change in

253 breast appearance after 50 Gy/25 F, 39 Gy/13 F and 42.9 Gy/13 F was 39.6, 30.3 and
254 45.7%, from which an alpha/beta value of 3.6 Gy (95% CI 1.8-5.4) is estimated. The
255 alpha/beta value for palpable breast induration was 3.1 Gy (95% CI 1.8-4.4). the study
256 concluded An alpha/beta value of around the study concluded An alpha/beta value of around
257 3 Gy for late normal tissue changes in the breast is derived from the estimated equivalence
258 of 41.6 Gy in 13 fractions and 50 Gy in 25 fractions over 5 weeks, in line with trial predictions
259 [8].

260

261 **Sanz [9] conducted** a study to analyze the results of weekly hypofractionated treatment in
262 486 elderly patients with associated diseases that modify their performance status and do
263 not tolerate long periods of daily irradiation. They were treated with conservative surgery or
264 mastectomy and then adjuvant hypofractionated irradiation, administering 5 Gy or 6.25 Gy in
265 6 fractions, once a week (total dose 30–37.5 Gy) over 6 weeks. The study concluded once-
266 weekly hypo-fractionated radiotherapy is a feasible and convenient option for elderly patients
267 with breast cancer. It is a safe treatment modality with similar survival and local control
268 results compared to standard fractionation, while the side effects are acceptable [9]

269 Sun et al. [10] and Team conducted a phase III noninferior randomized trial to evaluate the
270 efficacy and toxicity of HFRT after mastectomy. In this analysis, 820 high- risk patients
271 mainly with stage III breast cancer were enrolled and followed up for 5 years. Patients were
272 randomly assigned after mastectomy to receive either HFRT (43.5 Gy/15f/3w) or CFRT (50
273 Gy/25f/5w) to the chest wall and supraclavicular nodal region. The primary endpoint was
274 loco-regional recurrence (LRR). The study reported that there were no significant differences
275 in 5-year LRR (8.4% vs. 6.0%, P Z 0.396), DM (21.3% vs. 24.3%, P Z 0.530), DFS (75.1%
276 vs. 74.6%, P Z 0.841), and OS (84.9% vs. 87.1%, P Z 0.562) between HFRT and CFRT
277 group and concluded In patients with high-risk breast cancer after mastectomy, 43.5 Gy
278 delivered in 15 fractions over 3 weeks has comparable efficacy and toxicity at 5 years with
279 standard **fractionation [10].**

280 Randomized controlled trials of altered fraction size versus conventional fractionation for
281 radiation therapy in women with early breast cancer who had undergone breast-conserving

282 surgery. 8228 women in nine studies were analysed. altered fraction size (delivering
283 radiation therapy in larger amounts each day but over fewer days than with conventional
284 fractionation) did not have a clinically meaningful effect on: local recurrence-free survival
285 (Hazard Ratio (HR) 0.94, 95% CI 0.77 to 1.15, 7095 women, four studies, high-quality
286 evidence), cosmetic outcome (Risk ratio (RR) 0.90, 95% CI 0.81 to 1.01, 2103 women, four
287 studies, high- quality evidence) or overall survival (HR 0.91, 95% CI 0.80 to 1.03, 5685
288 women, three studies, high-quality evidence). Acute radiation skin toxicity (RR 0.32, 95% CI
289 0.22 to 0.45, 357 women, two studies) was reduced with altered fraction size. Altered
290 fraction size was associated with less patient-reported ($P < 0.001$) and physician-reported (P
291 $= 0.009$) fatigue at six months (287 women, one study). The review concluded altered
292 fraction size regimens (greater than 2 Gy per fraction) does not have a clinically meaningful
293 effect on local recurrence, is associated with decreased acute toxicity and does not seem to
294 affect breast appearance, late toxicity or patient-reported quality-of- life measures for
295 selected women treated with breast conserving therapy [11].
296

297 The randomized trial was from the MD Anderson Cancer Center, in Houston. The study was
298 conducted in 287 women aged 40 years and older with early- stage breast cancer (stage 0-
299 2), who were randomly assigned to receive either HF-WBI (42.56 Gy in 16 fractions of WBI;
300 $n = 138$) or CF-WBI (50.00 Gy in 25 fractions of WBI; $n = 149$). The rate of physician-
301 assessed toxicity of grade 2 or higher was significantly lower for women receiving HF-WBI
302 (47% vs 78%; $P < .001$), as were acute toxic effects of grade 3 or higher (0% vs 5%; $P = .01$). In particular, rates for physician-
303 assessed fatigue, pruritus, breast pain, and dermatitis were significantly lower for women
304 receiving HF. Although patient-reported quality of life, as reported from the Functional
305 Assessment of Cancer Therapy for Patients with Breast Cancer, was similar for women
306 receiving HF and CF, items associated with lack of energy and trouble meeting family needs
307 favoured women receiving HF. The study concluded treatment with HF-WBI appears to yield
308 lower rates of acute toxic effects than CF-WBI as well as less fatigue and less trouble
309 meeting family needs 6 months after completing radiation therapy [12].
310

311

312 A task force authorized by the American Society for Radiation Oncology weighed evidence
313 from a systematic literature review and produced the recommendations contained herein.
314 The majority of patients in randomized trials were aged 50 years or older, had disease Stage
315 pT1-2 pN0, did not receive chemotherapy, and were treated with a radiation dose
316 homogeneity within $\pm 7\%$ in the central axis plane. Such patients experienced equivalent
317 outcomes with either HF-WBI or CF-WBI. Patients not meeting these criteria were relatively
318 underrepresented, and few of the trials reported subgroup analyses. For patients not
319 receiving a radiation boost, the task force favoured a dose schedule of 42.5 Gy in 16
320 fractions when HF-WBI is planned. The task force also recommended that the heart should
321 be excluded from the primary treatment fields (when HF-WBI is used) due to lingering
322 uncertainty regarding late effects of HF-WBI on cardiac function. Data were sufficient to
323 support the use of HF-WBI for patients with early-stage breast cancer who met all the
324 aforementioned criteria. For other patients, the task force could not reach agreement either
325 for or against the use of HF-WBI, which nevertheless should not be interpreted as a
326 contraindication to its use [13].

327

328 Chan et al. [14] conducted a study to determine if there is an increase in hospital-related
329 morbidity from cardiac causes with HF-WBI relative to CF-WBI. Between 1990 and 1998,
330 5334 women ≤ 80 years of age with early-stage breast cancer were treated with
331 postoperative radiation therapy to the breast or chest wall alone. A population-based
332 database recorded baseline patient, tumour, and treatment factors. The median follow-up
333 was 13.2 years. For left-sided cases, 485 women were treated with CF-WBI, and 2221
334 women were treated with HF-WBI. The 15-year cumulative hospital-related morbidity from
335 cardiac causes (95% confidence interval) was not different between the 2 radiation therapy
336 regimens after propensity-score adjustment: therapy regimens after propensity-score
337 adjustment: 21% (19-22) with HF-WBI and 21% (17-25) with CF-WBI ($P=.93$). For right-
338 sided cases, the 15-year cumulative hospital-related morbidity from cardiac causes was also
339 similar between the radiation therapy groups ($P=.76$). The study concluded there is no
340 difference in morbidity leading to hospitalization from cardiac causes among women with

341 left-sided early-stage breast cancer treated with HF-WBI or CF-WBI at 15- year follow-up
342 [14].

343 Karasawa et al. [15] conducted study to evaluate the efficacy and safety of hypofractionated
344 whole-breast irradiation (HF-WBI) compared with conventionally fractionated (CF) WBI.
345 Patients with early breast cancer (stages 0- II and <3 positive lymph nodes) who had
346 undergone breast-conserving surgery were eligible for the HF- WBI study. HF-WBI was
347 administered at 43.2 Gy in 16 fractions over 3.2 weeks to the whole breast with an additional
348 tumor-bed boost of 8.1 Gy in 3 fractions over 3 days for positive surgical margins or those <5
349 mm. CF-WBI was administered at 50 Gy in 25 fractions over 5 weeks to the whole breast
350 with an additional tumor-bed boost of 16 Gy in 8 fractions over 1.4 weeks to 6 Gy in 3
351 fractions over 3 days, depending on margin status. Grade 2 acute skin reactions were
352 observed
353 for 24 patients (3 %) in the HF-WBI group and 53 for 24 patients (3 %) in the HF-WBI group
354 and 53 patients (14 %) in the CF-WBI ($p < 0.001$) group. The median follow-up period was
355 27 months. Two cases of intrabreast tumor recurrence were observed in each treatment
356 group. Regional lymph node recurrence was observed in 1 HF-WBI patient and 2 CF-WBI
357 patients. The study concluded HF-WBI is superior to CF-WBI in terms of acute skin reaction
358 and has the same short- term efficacy [15].

359
360

361 Kim et al. [16] in phase 2 trial of accelerated, hypofractionated whole-breast irradiation (AH-
362 WBI) delivered as a daily dose of 3 Gy to the whole breast followed by a tumor bed boost.
363 Two hundred seventy-six patients diagnosed with breast cancer (pT1-2 and pN0-1a) who
364 had undergone breast-conserving surgery in which the operative margins were negative
365 were treated with AH-WBI delivered as 39 Gy in 13 fractions of 3 Gy to the whole breast
366 once daily over 5 consecutive working days, and 9 Gy in 3 sequential fractions of 3 Gy to a
367 lumpectomy cavity, all within 3.2 weeks. After a median follow-up period of 57 months
368 (range: 27-75 months), the rate of 5-year locoregional recurrence was 1.4% (n=4), whereas
369 that of disease-free survival was 97.4%. The mean pretreatment percentage breast

370 retraction assessment was 12.00 (95% confidence interval [CI]: 11.14-12.86). The mean
371 value of interval [CI]: 11.14-12.86). The mean value of percentage breast retraction
372 assessment increased to 13.99 (95% CI: 12.17-15.96) after 1 year and decreased to 13.54
373 (95% CI: 11.84-15.46) after 3 years but was not significant ($P > .05$). The study reported AH-
374 WBI consisting of 39 Gy in 13 fractions followed by a tumor bed boost sequentially delivering
375 9 Gy in 3 fractions can be delivered with excellent disease control and tolerable skin toxicity
376 in patients with early-stage breast cancer after breast-conserving surgery [16].

377 Bekelman et al. [17] conducted Retrospective, observational cohort study, in patients with
378 incident early-stage breast cancer treated with lumpectomy and WBI from 2008 and 2013
379 and divided patient into 2 cohorts: (1) the hypofractionation-endorsed cohort ($n = 8924$)
380 included patients aged 50 years or older without prior chemotherapy or axillary lymph node
381 involvement and (2) the hypofractionation-permitted cohort ($n = 6719$) included patients
382 younger than 50 years or those with prior chemotherapy or axillary lymph node involvement.
383 Hypofractionated WBI increased from 10.6% (95% CI, 8.8%-12.5%) in 2008 to 34.5% (95%
384 CI, 32.2%-36.8%) in 2013 in the hypofractionation- endorsed cohort and from 8.1% (95% CI,
385 6.0%-10.2%) in 2008 to 21.2% (95% CI, 18.9%-23.6%) in 2013 in the hypofractionation-
386 permitted cohort. Adjusted mean total health care expenditures in the 1 year after mean total
387 health care expenditures in the 1 year after diagnosis were \$28,747 for hypofractionated and
388 \$31,641 for conventional WBI in the hypofractionation- endorsed cohort (difference, \$2894;
389 95% CI, \$1610- \$4234; $P < .001$) and \$64,273 for hypofractionated and \$72,860 for
390 conventional WBI in the hypofractionation- permitted cohort (difference, \$8587; 95% CI,
391 \$5316- \$12,017; $P < .001$). Adjusted mean total 1-year patient out-of-pocket expenses were
392 not significantly different between hypofractionated vs conventional WBI in either cohort [17].

393

394 Deshmukh et al. [18] constructed a decision-analytic model that followed women who were
395 treated with lumpectomy for early-stage breast cancer. Recurrence, mortality, complication
396 rates, and utilities (five-year radiation-associated quality of life scores), were extracted from
397 RCTs. Costs were based on Medicare reimbursement rates. HF-WBI dominated CF-WBI (ie,
398 resulted in higher quality-adjusted life-years [QALYs] and lower cost) in all scenarios. HF-

399 WBI also had a greater likelihood of cost-effectiveness compared with IORT; under a
400 societal perspective that assumes that radiation-associated disutility persists, HF-WBI
401 results in an ICER of \$17 024 per QALY compared with IORT with a probability of cost-
402 effectiveness of 80% at the \$100 000 per QALY willingness-to-pay of 80% at the \$100 000
403 per QALY willingness-to-pay threshold. If radiation-associated disutility is assumed to
404 discontinue, the ICER is lower (\$11 461/QALY), resulting in an even higher (83%) probability
405 of relative cost-effectiveness. The ICER was most sensitive to the probability of metastasis
406 and treatment cost. The study concluded, for women with early-stage breast cancer
407 requiring adjuvant radiotherapy, HF-WBI is cost-effective compared with CF-WBI and IORT
408 [18].

409 The result of our study clearly suggests that outcome for both dose schedule was equivalent.
410 Hypofractionation is rather cost effective considering the low socio-economic status of our
411 practice domain which reflects a major population of India.

412

413 **5. CONCLUSION**

414 There is no significant difference in between the conventional regimen and this
415 hypofractionated regimen in terms of OS, DFS and adverse reactions. Hence, in our
416 institution, we usually prefer Hypofractionated radiotherapy (40Gy/15 fractions) in adjuvant
417 settings for breast cancer patients.

418

419 **CONSENT**

420 All authors declare that written informed consent was obtained from each patient (or other
421 approved relative).

422 **ETHICAL APPROVAL**

423 All authors hereby declare that all experiments have been examined and approved by the
424 appropriate ethics committee and have therefore been performed in accordance with the
425 ethical standards laid down in the 1964 Declaration of Helsinki.

426

427 **COMPETING INTERESTS**

428 Authors have declared that no competing interests exist.

429

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