Introduction

Breast cancer is the most common female malignancy in the world, affecting more than 2 million people worldwide every year. A multimodality approach for its management is almost always required with patients undergoing a combination of radiotherapy, chemotherapy and surgery. Radiation therapy is the cornerstone of breast cancer care for definitive treatment approaches. As per existing standard guidelines in post axillary dissection, node positive breast cancer patients are given radiotherapy to lymph nodal area. This is done to improve loco regional control and survival [1-4]. Hypo fractionated radiation schedule has been proven safe and effective for tumour control with added advantage of less treatment time, supported by American Society For Radiation Oncology (ASTRO) group [5-7]. The concern with this dose schedule is a potential increase in acute and late term toxicities [8]. Late toxicities in breast and axilla irradiation are effects on heart, lungs, axilla (lymphedema), brachial plexus, and general cosmesis.

Materials and Methods

From January 2018 – January 2019- 100 breast cancer patients were treated with mastectomy, axillary node dissection (AND) and loco-regional hypo fractionated RT (2.67 Gy per Fraction, in 15 fractions -Total dose 40 Gy) which is in agreement with several randomized trials which have shown to improve local control with mild side effects and acceptable cosmetic outcome [5]. The median age at diagnosis was 60 years (range 40–81). Histology in all patients was ductal carcinoma. Axillary lymph nodes metastasis was confirmed in all women. All patients received systemic chemotherapy either neo adjuvant or adjuvant with schedule of adriamycin and cyclophosphamide. All the patients were assessed priorly to rule out any post-operative complications that
might mimic late toxicities. Follow up of patients was done at 12 months post completion of radiation therapy. LENT-SOMA scale were used to assess lymphedema (0-IV), pain severity (1-IV). RTOG Late toxicity assessment was done for arm stiffness (0-V).

**Results**

Age distribution (in years) of the patient in study was 40-49 – (17%), 50-59 – (32%), 60-69- (41%), 70-79 – (8%), 80-89 – (2%) (Table 1). In 17 patients (17%) lymphedema was recorded during follow up: LENT SOMA grading was as follows; grade I-11 , grade II- 5 and grade III-1 patient (Figure 1). In 12 (12%) of patients symptoms started before radiation therapy and can most likely be attributed to surgery and systemic therapy. 5 (5%) patients developed lymphedema which was likely radiation induced. The time of onset of lymphedema in the latter 5 cases was very variable (from 1 to 11 months from the end of radiotherapy). In 4 patients (grade I-2, grade II-2) it was as a result of disease progression to the lymph nodes. Most common complaint was of pain seen in 25 patients (25%).Severity (LENT SOMA) was-Grade I- 12 (12%), grade II - 9 (9%), grade III- 4 (4%), and grade IV-0 (0%) (Figure 2). Frequent complaint was pain aggravation upon arm movement/exercise, and relieved by rest and analgesics if patients was not able to tolerate it. Arm stiffness was present in 14 (14%), grade I- 9 (9%), grade 2-4 (4%), grade 3- 1 (1%) (Figure 3). Major difficulty faced by patients was restricted abduction of arm. Age group of patients with most symptoms of late toxicities was in 60-69 years of age.

**Discussion**

Hypo fractionated radiotherapy in breast cancer has become a common approach to manage breast cancer. This study investigated the late effects in patients who received this regime to chest wall and nodal regions. Clinical results achieved were similar to those achieved by standard regimens.

As reported in the literature, the irradiation of the lymph node area may increase the risk of side effects such as lymphedema and brachial plexopathy. Lymphedema may represent a standard complication after any axillary surgery. It is difficult to record as no clear parameter to measure lymphedema exists. Also it can be due to many contributing factors such as obesity, chemotherapy, extent of surgery [9-13]. Shah et al., provided the incidence of lymphedema according to the extent of RT after dissection, reporting a rate from 2–35 % after breast irradiation and an increased incidence to 9–65 % in the case of loco-regional irradiation [14]. In this study we used LENT SOMA questionnaire estimate clinical extent of the effects.

Shorter schedule has advantages of being safe, with good outcomes and better cosmesis compared to the standard radiation as it has been studied in several trials [6, 15]. A recent update of START A and START B evaluated the loco-regional RT in a limited group of patients and neither the 5 week nor the 3 week treatment developed significantly worse normal tissue impacts: the assessments of arm and shoulder effects showed no evidence of a detrimental effect for the hypo fractionated schedules [6].

Yarnolds et al. observed no radiation induced brachial plexus toxicity after hypo fractionated irradiation of the axilla and/or supraclavicular fossa. It was stated that the START B regimen (40 Gy in 15 fractions/3 weeks) is equivalent to 47 Gy in 2.0-Gy fractions if the $\alpha/\beta$ value for brachial plexus is 2.0 Gy or to 49 Gy in 2.0-Gy fractions, if $\alpha/\beta=1.0$ Gy [15]. Haffty and Buchholz

![Table 1. Age Distribution of Patients Enrolled in Study](image)

| Age (in Years) | Number of Patients |
|---------------|--------------------|
| 40-49         | 17                 |
| 50-59         | 32                 |
| 60-69         | 41                 |
| 70-79         | 8                  |
| 80-89         | 2                  |

![Figure 1. Graphical Representation of Patients with Lymphedema](image)
commented on the absence of side effects in the small group of patients (116pts of 2215, 7 %) enrolled in the
START B trial and receiving regional hypo fractionated RT: they confirmed that these results are consistent with
modelling of normal tissue effects, which predicts that 40 Gy in 15 fractions should be as safe as the standard
scheme for all normal tissues [16]. Badiyan N. et al. noted that RNI with standard fractionation is associated with
increased toxicity compared to WBI alone but current data do not support this observation [14]. The outcomes
of our investigation seem to confirm what emerges from the literature: patients treated with hypo fractionated RT
(40Gy in 15 fractions with 5 times a week) to the whole breast and infra-supraclavicular fossa did not show a
higher rate of side effects, in 5 patients the lymphedema was radiotherapy induced, of which only 1 was severe.
The favourable results obtained with this fractionation can be the basis for investigating new radiation schemes with
a smaller number of fractions administered in reduced times.

Another concern of hypo fractionation in RNI is toxicity to brachial plexus. Patients can suffer pain, impaired arm movement, paraesthesia and in general poor quality of life due to this. As it depends on total dose, volume of irradiation and dose per fraction, assuming low a/b ratios of 1.0–2.0 for the brachial plexus, hypo fractionated treatment regimens such as 40 Gy in 15 fractions would deliver a lower EQD2 to the brachial plexus than 50 Gy in 25 fractions [17] which significantly increases the risk of late toxicity. Standard RNI has rate of brachial plexopathy of 5% and paraesthesia in 20 % [18-21] but START B found no such case in the study [6, 22]. A retrospective review examining hypo fractionated PMRT with a subset receiving hypo fractionated RNI found no increase in rates of brachial plexopathy [23].

Delanian et al. evaluated the risk of plexopathy with hypo fractionated RNI schedules; incidence being more than 50% in >4 Gy/ fraction, though these studies used outdated techniques. These techniques have been found to have high rates of late effects in a retrospective data review done in Sweden [27, 28]. It is observed that brachial

![Graphical Representation of Patients with Localised Pain](image1)

![Graphical Representation of Patients with Arm Stiffness](image2)
plexopathy has been found to be less than 5% in doses per fraction below 3 Gy [24-26]. Major difficulty faced to record plexopathy is longer follow up requirement as major effects may occur beyond 5 years [29].

In conclusion, this study examined hypofractionated RNI in accordance with previous published literature and evaluates long term toxicities at irradiated sites. These effects were found to be tolerable and mild. But, further review and longer follow ups are required for better understanding of effects at various organ sites; this is also supported by published data.

Demonstrating its safety and efficacy will allow more women to complete adjuvant radiation in shortened treatment duration. Further, from a healthcare economics standpoint, shortened schedules reduces the costs to the healthcare system and allows for increased availability of and improved access to expensive radiation therapy delivery platforms.

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