Comparison of Sliding Window and Field-In-Field Techniques For Tangential Whole Breast Irradiation Using The Halcyon and Synergy Agility Systems

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Abstract

**Background:** To develop a tangential treatment technique for whole breast irradiation using the Varian Halcyon and to compare it with Elekta Synergy Agility plans.

**Methods:** For 20 patients two comparable treatment plans with respect to dose coverage and normal tissue sparing were generated. Tangential field-in-field treatment plans (Pinnacle/Synergy) were replanned using the sliding window technique (Eclipse/Halcyon). Plan specific QA was performed using the portal Dosimetry and the ArcCHECK phantom. Imaging and treatment dose were evaluated for treatment delivery on both systems using a modified CIRS Phantom.

**Results:** The mean number of monitor units for a fraction dose of 2.67 Gy was 515 MUs and 260 MUs for Halcyon and Synergy Agility plans, respectively. The homogeneity index and dose coverage were similar for both treatment units. The plan specific QA showed good agreement between measured and calculated plans. All Halcyon plans passed portal dosimetry QA (3%/2 mm) with 100% points passing and ArcCheck QA (3%/2 mm) with 99.5%. Measurement of the treatment and imaging dose with the CIRS phantom resulted in lower dose to the contralateral breast for the Halcyon plans.

**Conclusions:** For the Varian Halcyon a plan quality similar to the Elekta Synergy device was achieved. For the Halcyon plans the dose contribution from the treatment fields to the contralateral breast was even lower due to less interleaf transmission of the Halcyon MLC and a contribution of scattered dose from the collimator system.

Background

Breast cancer is one of the most common malignant tumors in women. Standard treatment for early-stage breast cancer is breast-conserving surgery followed by adjuvant whole breast irradiation [1].

The sparing of the organs at risk is important as the long term survival probability for diagnosed stage I and II patients treated with radiation after breast-conserving therapy is high [2, 3]. Different treatment modalities are available for radiation delivery like three dimensional conformal therapy (3D-CRT) [4, 5], tomotherapy [6–9], static-field intensity-modulated radiation therapy (IMRT) and volumetric modulated arc therapy (VMAT) [9–11].

The Varian Halcyon linac offers 6 MV flattening-filter-free (FFF) photon beam energy with a dual-layer multi-leaf collimator (MLC). Treatment on the Halcyon requires daily image-guidance because there are no light field, optical distance indicator or lasers at the treatment isocenter available.

The aim of this work was to develop a treatment technique suitable for whole breast irradiation using the Halcyon system resulting in a tangential field arrangement. The dosimetric effect of daily imaging on the Halcyon unit was investigated. A comparison of the Halcyon and Synergy Agility plans in terms of imaging dose and deliverability was performed by QA measurements.

Methods

**Patient characteristics**
All patient characteristics are listed in Table 1. The patients were treated via tangential whole breast irradiation without supraclavicular or internal mammary lymph nodes. The size of the planning target volumes ranged from 352 cm³ to 1439 cm³ with a mean target volume of 814 cm³. The cohort consisted of two subgroups. The first ten patients were treated with Elekta Synergy Agility (Elekta Medical Systems, Crawley, UK). Treatment plans with tangential beams were re-planned for Halcyon to compare the plan quality and dose to OARs retrospectively. The second subgroup of 10 patients was scheduled for whole breast irradiation on the Varian Halcyon system (Varian Medical Systems, Palo Alto, CA, USA, version 2.0) while treatment plans for Halcyon and Synergy Agility were created. Finally, two comparable plans were available for all patients.
| Treatment machine | Patient | Tumor site | PTV in cm³ | Energy | Fields | Monitor Units |
|-------------------|---------|------------|------------|--------|--------|---------------|
| Synergy Agility   | 1       | left       | 690        | 6 & 10MV | 6MV FFF | Synergy | 4 | 2 | 245 | 446 |
|                   | 2       | left       | 1163       | 6MV     | 6MV FFF | Synergy | 4 | 2 | 268 | 499 |
|                   | 3       | left       | 832        | 6MV     | 6MV FFF | Synergy | 4 | 2 | 261 | 502 |
|                   | 4       | left       | 946        | 6MV     | 6MV FFF | Synergy | 4 | 2 | 263 | 440 |
|                   | 5       | left       | 435        | 6MV     | 6MV FFF | Synergy | 4 | 2 | 253 | 436 |
|                   | 6       | right      | 790        | 6MV     | 6MV FFF | Synergy | 4 | 2 | 256 | 470 |
|                   | 7       | right      | 509        | 6 & 10MV | 6MV FFF | Synergy | 5 | 2 | 254 | 470 |
|                   | 8       | right      | 934        | 6MV     | 6MV FFF | Synergy | 4 | 2 | 269 | 494 |
|                   | 9       | right      | 351        | 6 & 10MV | 6MV FFF | Synergy | 5 | 2 | 263 | 492 |
|                   | 10      | right      | 1220       | 6MV     | 6MV FFF | Synergy | 4 | 3 | 274 | 700 |
| Halcyon           | 11      | left       | 1016       | 6MV     | 6MV FFF | Synergy | 4 | 2 | 277 | 536 |
|                   | 12      | left       | 798        | 6MV     | 6MV FFF | Synergy | 4 | 2 | 259 | 415 |
|                   | 13      | left       | 589        | 6MV     | 6MV FFF | Synergy | 4 | 2 | 266 | 575 |
|                   | 14      | left       | 861        | 6MV     | 6MV FFF | Synergy | 4 | 2 | 254 | 484 |
|                   | 15      | left       | 1035       | 6MV     | 6MV FFF | Synergy | 4 | 2 | 254 | 641 |
|                   | 16      | right      | 1439       | 6MV     | 6MV FFF | Synergy | 4 | 2 | 266 | 536 |
|                   | 17      | right      | 494        | 6MV     | 6MV FFF | Synergy | 4 | 2 | 243 | 459 |

Patient characteristics and details of the beam configuration, energy and segments used. Stated values indicate mean ± SD. Patient #1–10 were treated on the Synergy Agility machine and patients #11–20 were treated on the Halcyon system.
### Treatment planning

Patients were positioned in supine position with both arms raised. The dose prescription was 40 Gy in 15 fractions followed by a boost of 10 Gy in 5 fractions (mean dose). PTV was delineated closed to the surface. Dose homogeneity was maintained by the standard deviation (SD) in PTV-5 mm of less than 3%. PTV-0.5 mm corresponds to the CTV as defined in the ESTRO Guideline [12, 13].

Pinnacle (Philips Radiation Oncology Systems, Fitchburg, WI, version 16.2) was used for treatment planning of the Elekta Synergy Agility plans. A 3D CRT field in field technique was used with flattened 6 MV and 10 MV beams in forward planning mode [4]. The gantry angle was optimized in the beam's eye view (BEV) for a minimum lung area and beam divergence toward the lung was compensated by adjusting the gantry angles of the beams accordingly. The beams exceeded the patient surface by minimum of 2 cm. One to two segments per field were added to improve target coverage and dose homogeneity and to omit wedges. Final dose was calculated using the collapsed cone algorithm and a grid size of 0.3 cm.

Treatment planning for the Halcyon system was performed in Eclipse (Varian Medical Systems, Palo Alto, CA, USA, version 15.6). Tangential fields were defined using 6 MV FFF beams. The dose distribution was optimized with a sliding window technique. A virtual bolus was added on top of the PTV during the optimization process with a density of 0.26 g/cm³ and a thickness of 2 cm. With a minimum dose objective for the virtual bolus the optimizer was forced to enlarge the field size into the air. An uncertainty analysis was carried out by shifting the isocenter (1 cm) in dorsal and contralateral direction which should result in comparable target coverage for PTV-5 mm. For final dose calculation, the bolus was removed. The same gantry angles were used for the Halcyon and Synergy Agility plans. No couch rotation was utilized in the clinical plans for any of the patients selected. In Eclipse, dose was calculated with a grid size of 0.25 cm using the Acuros External Beam Algorithm, version 15.6.06.

### Treatment plan comparison

We compared the dose-volume histograms (DVHs) of the Halcyon with the Synergy Agility plans. DVH parameters are listed in Table 2. For PTV, the dose received by 2%, 50%, 80%, 95% and 98% volume (D02, D50, D80, D95, D98) were compared. The mean and standard deviation of these parameters were calculated for each patient. The results showed that the Halcyon plans resulted in lower doses to the PTV compared to the Synergy Agility plans, indicating better dose homogeneity and target coverage for the Halcyon system.
D80, D95, D98) were evaluated. The homogeneity index (HI) was defined as HI = (D02 - D98) / D50. For OARs, the maximum and mean doses were assessed. The number of monitor units (MUs) and the correlation between PTV size and number of MU were analysed for all plans. Both machines (Synergy Agility and Varian Halcyon) were calibrated equally: a dose of 1 Gy in 10 cm depth corresponds to 100 MU.

Table 2
DVH comparison between Synergy and Halcyon plans

|                  | Synergy Agility | Halcyon      | p value |
|------------------|-----------------|--------------|---------|
| **PTV**          |                 |              |         |
| D95 in Gy        | 33.9 ± 2.3      | 33.5 ± 2.5   | NS*     |
| D80 in Gy        | 38.0 ± 0.5      | 38.6 ± 0.4   | p < 0.01|
| D_{Max} in Gy    | 43.0 ± 0.6      | 43.3 ± 0.8   | NS      |
| HI               | 0.13 ± 0.01     | 0.12 ± 0.02  | NS      |
| **PTV-0.5**      |                 |              |         |
| D_{Min} in Gy    | 35.3 ± 1.0      | 35.4 ± 0.2   | NS      |
| D95 in Gy        | 38.5 ± 0.2      | 38.8 ± 0.3   | 0.03    |
| D_{Mean} in Gy   | 40.1 ± 0.1      | 40.1 ± 0.1   | NS      |
| SD in Gy         | 1.0 ± 0.1       | 0.8 ± 0.1    | 0.01    |
| **Lung ipsilateral** |           |              |         |
| D_{Mean} in Gy   | 4.7 ± 0.9       | 4.4 ± 1.1    | NS      |
| **Lung contralateral** |       |              |         |
| D_{Mean} in Gy   | 0.3 ± 0.1       | 0.2 ± 0.1    | p < 0.01|
| **Breast contralateral** |       |              |         |
| D_{Mean} in Gy   | 0.5 ± 0.2       | 0.3 ± 0.1    | p < 0.01|
| D01 in Gy        | 1.8 ± 0.7       | 1.0 ± 0.5    | p < 0.01|
| **Heart**        |                 |              |         |
| D_{Max} in Gy    | 31.7 ± 10.5     | 33.0 ± 8.4   | NS      |
| D_{Mean} in Gy   | 1.7 ± 0.4       | 1.5 ± 0.5    | p < 0.01|

* Not significant (NS)

** Only left-sided plan were considered for the evaluation of the heart dose

Dose to target volumes and organs at risk in Synergy Agility plans and Halcyon plans. Stated values indicate mean ± SD and statistical analysis was performed using the Wilcoxon test. The differences were considered statistically significant when p < 0.05.
Dosimetric evaluation

Plan specific QA

In order to test the deliverability of the Halcyon treatment plans, plan specific QA was performed. Plans were delivered to an ArcCHECK Phantom (Sun Nuclear Corporation, Melbourne, FL, USA) with the cavity plug present equipped with a Semiflex 31010 ionization chamber (PTW-Freiburg, Freiburg, Germany) in the center. For point dose measurement, passing criteria were percent dose deviation of less than 3% from the expected dose. The ArcCHECK measurements were evaluated with 3%/2 mm and 2%/2 mm gamma criteria with tolerance level of 96.5% and 95% of pixel passing, global normalization and 10% low dose threshold. Portal dosimetry was evaluated for each field with a gamma criterion of 2%/1 mm and a tolerable passing rate of 94%.

Imaging and treatment dose

To evaluate the imaging and treatment dose a left-sided treatment plan was generated for a dynamic thorax phantom (CIRS Incorporated, Norfolk, Virginia, USA). The phantom was used in static mode. Breast surrogates were attached to simulate a breast treatment. Breast surrogates were created from silicon material with 3 inserts for an ionization chamber for each side (see Fig. 1). A Unidos dosimeter (PTW-Freiburg, Freiburg, Germany) with 0.3 cm³ thimble chamber (type TM31013) was used for dose measurements. The dosimetric influence of different imaging modes (MV planar imaging, volumetric kV CBCT) was compared by measuring the dose in the ipsilateral and contralateral breast. In total, six point dose values were acquired for each imaging setup. For the Halcyon CBCT system, the breast mode with 125 kV and 45 mAs was used followed by iterative image reconstruction. This reconstruction algorithm is designed to reduce noise and to enhance image quality with high resolution. MV portal imaging is limited to 0° and 90° gantry position for the Halcyon system. For the Synergy Agility treatment setup, imaging dose of kV CBCT (120 kV, 70,4 mAs, S20) was compared with tangential MV images at 0° gantry position followed by the tangential treatment angles.

For this anthropomorphic breast phantom, treatment plans were optimized for both the Synergy Agility and Halcyon machines. For our standard protocol, imaging and treatment doses were measured and the accumulated dose in the contralateral breast was determined. A typical imaging procedure on the Synergy Agility system would be to perform imaging with anterior-posterior and tangential MV portals before the first treatment followed by imaging once a week (3 fractions of MV imaging and 15 treatment fractions). In contrast, the treatment on the Halcyon system required imaging before each treatment fraction (15 CBCT imaging fractions and 15 treatment fractions).

Statistical analysis

A statistical comparison of the Halcyon and the Synergy Agility plans was implemented to analyze dosimetric differences between the two machines using the Wilcoxon test for bound comparison. The differences were considered statistically significant when p < 0.05. IBM SPSS Statistics for Windows, version 26 (IBM Corp., Armonk, N.Y., USA) was used for statistical evaluation.

Results

Treatment plan comparison
The beam configuration for Halcyon and Synergy Agility plans are listed Table 1. For most cases, 6 MV and 4 fields were sufficient for the field-in-field technique with the Synergy Agility. In 3 out of 20 cases, a field arrangement of 5 beams was needed. A mixture of 6MV and 10MV was used in 3 cases. For treatment planning on the Halcyon unit, two treatment fields were used for application of a sliding window technique. Just for one case 3 fields were applied.

Table 1 outlines the difference in MUs for the Halcyon vs. Synergy Agility plans. The mean number of MUs used for the Halcyon delivered plans was 515 MUs, whereas the Synergy Agility delivered plans used 260 MUs (p < 0.001). No significant correlation was found between PTV size and number of Mus (R² = 0.5 and R² = 0.1 for Halcyon and Synergy Agility plans, respectively).

Overall, the plan quality of Halcyon plans was comparable to plans delivered on Elekta Synergy linac. Mean dose values in the PTV and OARs are listed in Table 2. For both treatment units the homogeneity index (HI) was similar: Synergy Agility plans achieved HI of 0.13 while the Halcyon plans achieved 0.12. Slightly improved dose homogeneity was observed for the Halcyon plans. Only the D80 in PTV differed significantly by 0.6 Gy on average (p < 0.01). The SD in PTV-5 mm was 1.0 Gy and 0.8 Gy on average for Synergy Agility and Halcyon plans, respectively. Significant differences were observed for D95 and SD in PTV-5 mm. However, given that dose differences are very low, we do not consider them of clinical relevance. Nominal target coverage (D95 and D80) was comparable for Halcyon and Synergy Agility plans.

Mean dose in the lungs and heart was slightly improved for Halcyon plans. Significant differences were found for the mean dose in the contralateral lung (0.1 Gy), the heart (0.2 Gy) and the contralateral breast (0.2). Maximum dose in the heart was 1.3 Gy lower on average for Synergy Agility plans but not statistically significant.

**Plan specific QA**

Phantom measurements with the ArcCHECK and portal dosimetry were performed for the Halcyon plans. All plans were successfully delivered on the Halcyon unit. Table 3 shows the ion chamber measurements and gamma passing rates for all patient plans. For the ArcCHECK, the mean deviation of the ion chamber dose was 0.97% with a maximum deviation of 2.67%. The pass rate of the diode measurements was 99.5% for the 3%/2mm gamma criterion.
### Table 3
Plan specific QA results

| Patient | Ion chamber dose deviation in % | Portal Dosimetry | ArcCHECK |
|---------|----------------------------------|------------------|----------|
|         |                                  | Field #1 | Field #2 | 3%/2mm | 2%/2mm  |
|         |                                  | 2%/1mm   | 2%/1mm   | > 94%  | > 94,5% | > 95%   |
| 1       | -2.67                            | 99.3     | 96.6     | 100    | 100     |
| 2       | -1.57                            | 99.5     | 96.2     | 100    | 98.3    |
| 3       | -1.62                            | 99.3     | 96       | 100    | 98.6    |
| 4       | -1.51                            | 99.5     | 97.1     | 99.8   | 97.2    |
| 5       | -1.90                            | 99.3     | 96.5     | 99.7   | 96.9    |
| 6       | -1.31                            | 96.8     | 98       | 100    | 98.8    |
| 7       | -1.15                            | 95.3     | 97.6     | 99.7   | 96.8    |
| 8       | -0.99                            | 97.7     | 98.1     | 100    | 99.1    |
| 9       | -1.27                            | 96.0     | 97.5     | 100    | 99.6    |
| 10      | -1.27                            | 96.7     | 97.2     | 98.4   | 92.2    |
| 11      | -1.19                            | 98.4     | 97.5     | 97.6   | 95.5    |
| 12      | -1.21                            | 99.5     | 97.3     | 99.5   | 95.1    |
| 13      | -1.03                            | 98.9     | 97.4     | 100    | 98.9    |
| 14      | -0.81                            | 98.9     | 96.6     | 99.8   | 96.5    |
| 15      | -1.06                            | 99.2     | 94.6     | 98.9   | 92      |
| 16      | -1.62                            | 97.8     | 96       | 97.6   | 88.3    |
| 17      | -1.58                            | 94.8     | 97.2     | 99.7   | 97.8    |
| 18      | 2.49                             | 94.6     | 95.8     | 98.5   | 95.8    |
| 19      | 2.39                             | 97.1     | 97.5     | 99.8   | 99      |
| 20      | -0.47                            | 97.2     | 95.8     | 100    | 99.2    |
| Mean    | 0.97                             | 97.8     | 96.7     | 99.5   | 96.8    |
| SD      | 1.25                             | 1.7      | 0.9      | 0.8    | 3.0     |

Results of the plan specific QA for ion chamber measurement, portal dosimetry and the ArcCHECK phantom.

All portal dosimetry measurements passed the 3%/2 mm gamma criterion with 100% of the pixels. The evaluation of the 2%/1 mm gamma criterion resulted in a mean pass rate of 97.8% and 96.7% for field #1 and field #2, respectively.
Imaging and treatment dose

For treatment setup on a Synergy Agility unit, the imaging dose of MV portals was compared with kV CBCT. A summary of all measurements is given in Table 4 and Fig. 2. The lowest imaging dose in the contralateral breast (points 1–3) was measured for the anterior-posterior and tangential MV portals on the Synergy Agility (in total 6 MU). Dose in points 1, 2 and 3 was 0.1 mGy, 0.1 mGy, and 0.4 mGy, respectively. Slightly larger doses were measured for the kV CBCT setup with 0.1 mGy, 0.2 mGy and 0.8 mGy in points 1–3. The cumulated dose of all treatment and imaging fractions for the Synergy Agility plan would result in 195.8 mGy, 291.8 mGy and 541.1 mGy in points 1, 2 and 3 of the contralateral breast, respectively (Table 4). Dose was accumulated for 3 fractions of MV imaging and 15 treatment fractions.

Table 4
Imaging and treatment dose

|                          | Dose in mGy | Contralateral side | Ipsilateral side |
|--------------------------|-------------|--------------------|-----------------|
|                          |             | 1  | 2  | 3  | 4  | 5  | 6  |
| Synergy Agility kV CBCT  | 0.1         | 0.2| 0.8| 1.9| 2.0| 2.9|
| MV portals*              | 0.1         | 0.1| 0.4| 52.5| 46.5| 39.7|
| Treatment                | 13.0        | 19.4| 36.0| 2737.1| 2711.8| 2704.2|
| 15 fx treatment          | 195.0       | 291.3| 540.0| 41056.5| 40677.2| 40563.4|
| 3 fx imaging             | 0.2         | 0.4 | 1.1| 157.5| 139.4| 119.1|
| Total                    | 195.8       | 291.8| 541.1| 41213.9| 40816.6| 40682.5|
| Halcyon kV CBCT*         | 1.0         | 0.6 | 1.0| 2.6 | 2.6 | 2.6 |
| MV portals               | 3.5         | 3.2 | 3.8| 23.9 | 25.8 | 24.5 |
| Treatment                | 5.9         | 8.8 | 16.5| 2649.2| 2634.7| 2645.0|
| 15 fx treatment          | 88.0        | 131.6| 247.5| 39737.7| 39519.8| 39675.4|
| 15 fx imaging            | 14.3        | 9.6 | 14.3| 38.3 | 38.3 | 38.3 |
| Total                    | 102.3       | 141.1| 261.9| 39776.0| 39558.0| 39713.7|

Dose was measured for the ipsilateral and contralateral side in the CIRS phantom for kV and MV imaging procedures and one treatment fraction. Total dose for our standard imaging protocols (marked with *) and all treatment fractions is listed.

For treatment setup on the Halcyon unit, the imaging dose of MV portals was compared with the kV CBCT. The MV portals deliver more dose to the contralateral breast than the kV CBCT. For MV portal imaging from 0° and 90° gantry angle, dose measurement in points 1–3 resulted in 3.5 mGy, 3.2 mGy and 3.8 mGy respectively. The imaging dose of the Halcyon kV CBCT is comparable with the Synergy Agility CBCT. Dose values in points 1–3 was 1.0 mGy, 0.6 mGy and 1.0 mGy respectively. For treatment on the Halcyon unit setup imaging must be performed daily. The cumulated dose of all treatment and imaging fractions for the Halcyon plan would result in
lower doses with 102.3 mGy, 141.1 mGy and 261.9 mGy in points 1, 2 and 3 of the contralateral breast, respectively (Table 4). In this case, dose was accumulated for 15 fractions of CBCT imaging and 15 treatment fractions. Thus, our standard approach with weekly MV imaging at the Synergy linac resulted in a higher dose for the contralateral breast than the daily kV imaging at the Halcyon linac.

Discussion

The presented work shows that the Halcyon linac is suitable for whole-breast irradiation using a tangential beam technique. A practical and efficient planning method for delivering 3D conformal breast radiotherapy using the Halcyon was implemented in our department. The simplified workflow on the Halcyon system enables a fast delivery with a reduced probability of patient collisions due to the closed ring gantry design and collision warning during treatment planning.

The overall plan quality regarding target volumes and OARs was comparable for plans delivered on Halcyon and Synergy Agility machines. Homogeneity index was 0.13 and 0.12 for Synergy and Halcyon plans, respectively. Similar studies of treatment plan comparisons for Halcyon and C-arm linacs were published [14, 15]. Flores et al. used the irregular surface compensator technique and reported lower homogeneity indices of 0.09 and 0.08 for C-arm linacs and the Halcyon plans respectively [14]. The increased number of MUs for delivering the Halcyon plans was also observed by other researchers [14–16]. For the current investigation, the MUs were 50% higher than the field-in-field breast treatment plans using tangential beams with Synergy Agility. The reason is the sliding window technique in combination with the un-flattened beam for the Halcyon plans.

In the current work, lower mean doses in the OARs were observed for the Halcyon plans, while the maximum heart dose was on average 1.3 Gy higher for the Halcyon plans. In contrast, Flores et al. and Morris et al. observed an increase in the mean dose in the contralateral lung and the heart for the Halcyon plans compared to C-arm linacs [14, 15].

Dose to OARs needs to be considered when choosing the setup imaging technique (MV portals or CBCT). Comparable imaging dose for kV CBCT was observed for both systems in the current work. However, the Halcyon requires CBCT imaging to be performed at every fraction. Regarding the sum of imaging and treatment dose for our standard protocol, much lower doses to the contralateral breast were determined with the Halcyon unit. This is mainly due to the lower interleaf transmission for the Halcyon MLC and a contribution of scattered dose from the collimator system. The Halcyon MLC has 77 mm thick leaves in each bank, creating a combined thickness of 154 mm. The combined transmission through the dual layer MLC is less than 0.01% at 6 MV FFF [15, 17]. Whereas for the Agility MLC the transmission through the MLC is 0.44% at 6 MV and 0.52% at 10 MV [18]. Flores et al. suggested a dose reduction for the OARs by using angled MV portals or non-ionizing techniques like surface guided RT [14].

The plan specific QA indicate a good agreement between the measured and calculated dose. The point dose measurements in the ArcCHECK phantom showed an average and maximum deviation of 0.97% and 2.67% respectively. Similar findings were reported by Morris et al. with an average and maximum point dose deviation of 1.5% and 2.23% in a solid water phantom [15]. In the current work, the average ArcCHECK gamma passing rate was 99.5% for 3%/2 mm criterion. Morris et al. published high pass rates of 99.4% for plan specific QA with Delta4 [15]. Portal dosimetry yielded pass rates of 100% for the 3%/2 mm criterion which is consistent with the
results published by Morris et al. [15]. In order to get some variation the 2%/1 mm gamma criterion was used, still resulting in mean gamma pass rates above 96%.

A common technique to spare the heart for left-sided treatments is deep inspiration breath-hold (DIBH). Barsky et al. reported about initial experience in treating patients with breast cancer on a Halcyon system in either supine position, with or without DIBH, or prone position. DIBH was used in 29% of the patients with average treatment times of 5.8 min [19]. Kennedy stated the total treatment time can be drastically reduced using irregular surface compensator technique as opposed to the Dynamic Flattening Beam-enabled field-in-field technique [16].

A bolus is commonly applied to improve target coverage near the surface while also enhancing the risk of severe skin reactions and a possible negative impact on cosmesis. The effect on superficial dose for 6 MV FFF beams and the need for bolus was evaluated by O’Grady et al. [20]. An increased superficial dose was observed for the use of a Halcyon 6MV FFF beam compared to 6 MV beams on C-Arm linacs for whole breast irradiation. It was assumed by O’ Grady et al. that the increased dose is primarily caused by a higher proportion of low energy photons. The traditional application of a bolus should be carefully re-evaluated for 6 MV beams [20].

In the current work, we only investigated tangential field arrangements. Several modalities are available for whole breast irradiation, e.g., the Irregular Surface Compensation technique and dynamic flattening beam sequences. For target volumes including the mammaria interna lymph nodes, we currently use VMAT and multi field IMRTs. The possibilities of treating such cases with the Halcyon will be investigated in a subsequent examination.

The field size of the Halcyon system is limited to $28 \times 28$ cm². A multi isocenter technique allows an extension up to 36 cm field size in longitudinal direction. Patients with a PTV extension of more than 36 cm are currently not suited for treatment on a Halcyon unit (Version 2.0).

**Conclusions**

We implemented a treatment technique with a tangential beam arrangement for whole breast irradiation with a tangential beam arrangement. Plan quality was equally good on Halcyon and Synergy Agility machines. Monitor units increased on average by 50% for Halcyon plans due to the sliding window technique. A significantly lower dose to the contralateral breast was measured for the Halcyon plans due to less interleaf transmission of Halcyon’s dual layer MLC and some scattered dose from the collimator system.

**List Of Abbreviations**

| Abbreviation | Description                     |
|--------------|---------------------------------|
| 3D CRT       | Three-dimensional conformal radiotherapy |
| BEV          | Beams eye view                  |
| CBCT         | Cone beam computer tomography   |
| DIBH         | Deep inspiration breath-hold    |
| DVH          | Dose volume histogram           |
| FFF          | Flattening filter free          |
Declarations

Ethics approval and consent to participate

Ethical review and written informed consent was not required for participation in this retrospective analysis in accordance with the local legislation (BayKrG Art. 27 (4)) and institutional requirements. Written informed consent for treatment was provided by all patients.

Availability of data and materials

Research data are stored in an institutional repository and will be shared upon request to the corresponding author.

Competing interests

The authors declare that they have no competing interests.

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None.

Authors’ contributions

AR, SoW, BP, KIB, OS and MF designed the analysis. AR, KaB, GR, SoW, StW and FE participated in the data collection and performed the analysis. All authors performed critical review of the manuscript and finally approved the manuscript.

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Breast surrogates for the phantom measurements were made by Jonas Ringholz using a 3D printer approach.

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**Figures**
CIRS phantom with breast surrogates Measurement of imaging and treatment dose was performed using a CIRS phantom with breast surrogates. Point dose was measured for the contralateral (1-3) and ipsilateral side (4-6). A treatment plan was generated for the Synergy Agility in Pinnacle (a) and for the Halcyon in Eclipse (b).
Figure 2

Imaging and treatment dose Dose contribution by imaging with the standard protocol (a, b) and by one treatment fraction (c, d) is displayed for the Synergy Agility (grey) and Halcyon unit (striped area). Results for the contralateral breast (points 1-3) are plotted on the left (a, c). The results for the ipsilateral breast (points 4-6) are shown at the right (b, d).