

Original Research Article

Optimising inter-patient image registration for image-based data mining in breast radiotherapy



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ABSTRACT

Background and purpose: Image-based data mining (IBDM) requires spatial normalisation to reference anatomy, which is challenging in breast radiotherapy due to variations in the treatment position, breast shape and volume. We aim to optimise spatial normalisation for breast IBDM.

Materials and methods: Data from 996 patients treated with radiotherapy for early-stage breast cancer, recruited in the REQUITE study, were included. Patients were treated supine ($n = 811$), with either bilateral or ipsilateral arm(s) raised (551/260, respectively) or in prone position ($n = 185$). Four deformable image registration (DIR) configurations for extrathoracic spatial normalisation were tested. We selected the best-performing DIR configuration and further investigated two pathways: i) registering prone/supine cohorts independently and ii) registering all patients to a supine reference. The impact of arm positioning in the supine cohort was quantified.

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DIR accuracy was estimated using Normalised Cross Correlation (NCC), Dice Similarity Coefficient (DSC), mean Distance to Agreement (MDA), 95 % Hausdorff Distance (95 %HD), and inter-patient landmark registration uncertainty (ILRU).

Results: DIR using B-spline and normalised mutual information (NMI) performed the best across all evaluation metrics. Supine-supine registrations yielded highest accuracy (0.98 ± 0.01 , 0.91 ± 0.04 , 0.23 ± 0.19 cm, 1.17 ± 1.18 cm, 0.51 ± 0.26 cm for NCC, DSC, MDA, 95 %HD, and ILRU), followed by prone-prone and supine-prone registrations. Arm positioning had no significant impact on registration performance. For the best DIR strategy, uncertainty of 0.44 and 0.81 cm in the breast and shoulder regions was found.

Conclusions: B-spline algorithm using NMI and registered supine and prone cohorts independently provides the most optimal spatial normalisation strategy for breast IBDM.

1. Introduction

Radiotherapy is an important treatment modality for breast cancer treatments. As the survival rate improves [1], long-term side effects become increasingly important to address. Adverse effects from radiotherapy, such as skin toxicity, breast fibrosis, lymphoedema, and shoulder stiffness, can cause treatment delays and negatively impact patients' quality of life [2,3]. Finding dose-outcome relationships can help clinicians prevent unwanted toxicity following breast radiotherapy.

Dose-outcome models in radiotherapy traditionally use simplified representations of dose distributions derived from dose-volume histograms (DVHs) or dose surface histograms (DSHs). However, this approach has some limitations, e.g., it requires a predefined hypothesis about structures or organs at risk (OARs) related to the investigated toxicity, where contour delineation uncertainties can bias prediction outcomes [4]. Generating a toxicity model from DVHs or DSHs [5–7] does not account for the heterogeneous dose distribution within OARs, as it summarises the complete 3D dose distribution to a single value, i.e., mean dose. Image-based data mining (IBDM), a voxel-based analysis technique, can help overcome these limitations [8]. IBDM involves analysing all voxels of the dose distribution and generating a spatial map to describe the dose–response relationship, including other confounding variables directly or through post-hoc analysis [9]. This is particularly relevant in breast cancer, where multimodal treatment such as surgery, chemotherapy, and hormonal therapy, as well as patient factors, e.g., comorbidities and body mass index (BMI), play a role in toxicity [10,11].

IBDM relies on spatial normalisation of the dose distribution to a common coordinate system (CCS), typically a reference template using Deformable Image Registration (DIR) before voxel-wise statistical analysis is performed [12]. While IBDM has been successfully applied to various tumour sites, including the lung, oesophagus, head and neck, and prostate cancer [13–17], its application in breast cancer remains largely unexplored. The dose–response relationships for many side effects of breast radiotherapy, such as lymphoedema and shoulder mobility, are poorly understood and may involve multiple structures or sub-structures. Therefore, implementing IBDM has the potential to advance our knowledge in this field.

However, spatial normalisation for breast radiotherapy presents significant challenges. There are considerable variations in breast size and shape between patients. Additionally, patients are treated in different positions: either prone or supine, the latter with either both arms raised or only the ipsilateral arm raised. These variations could affect the accuracy of spatial normalisation and the identification of sensitive sub-regions. This study aims to evaluate the accuracy of the spatial normalisation process and establish an optimal DIR strategy.

2. Methods

2.1. Patient dataset

Data from 996 early-stage breast cancer patients treated with radiotherapy after breast-conserving surgery, with/without systemic therapy [18], were included in this study, Table 1. These patients were

recruited prospectively in the REQUITE project (www.requite.eu) from 9 institutes in 5 countries. REQUITE established a resource for multi-national validation of models and biomarkers that predict the risk of late toxicity following radiotherapy [18]. REQUITE was approved by local ethics committees in participating countries (UK NRES Approval 14/NW/0035) and registered at <https://www.controlled-trials.com> (ISRCTN98496463) [19].

Due to the lack of anatomical landmarks and consistent contours in the full cohort, a smaller “representative cohort” (160 patients) was identified (see section *Identifying the Optimal Image Registration Pathway*). The purpose of the representative cohort is two-fold: 1) it allowed an in-depth exploration of the image registration performance, and 2) it

Table 1
The characteristics of 996 breast cancer patients recruited from the REQUITE dataset.

Characteristics	Supine	Prone
Number of patients	811	185
Age (years)	59 (23–90)	59 (35–79)
Treatment site (%)	Left = 52.6 %; Right = 47.4 %	Left = 48.6 %; Right = 51.4 %
Breast volume (cm ³)	670 (40–5450)	664 (107–1869)
Breast cup size (%) [*]	1 = 6.0 %; 2 = 8.6 %; 3 = 38.3 %; 4 = 27.2 %; 5 = 18.7 %; 6 = 5.4 %; 7 = 1.3 %; 8 = 0.0 %; 9 = 0.0 %; ≥ 10 = 0.0 %	1 = 0.6 %; 2 = 5.0 %; 3 = 29.3 %; 4 = 30.4 %; 5 = 26.5 %; 6 = 6.6 %; 7 = 0.6 %; 8 = 0.6 %; 9 = 0.6 %; ≥ 10 = 0.0 %
Breast band (%) ^{**}	1 = 0.0 %; 2 = 0.1 %; 3 = 8.2 %; 4 = 19.9 %; 5 = 26.0 %; 6 = 18.1 %; 7 = 9.8 %; 8 = 9.9 %; 9 = 3.5 %; 10 = 4.4 %	1 = 0.6 %; 2 = 0.0 %; 3 = 1.7 %; 4 = 12.2 %; 5 = 10.5 %; 6 = 17.1 %; 7 = 11.6 %; 8 = 23.8 %; 9 = 15.5 %; 10 = 7.2 %
BMI (kg/m ²)	25.0 (16.9–38.1)	25.2 (17.4–36.5)
Surgery (%)		
Breast	Yes = 98.4 % NA=1.6 %	Yes = 100.0 % NA=0.0 %
Axillary	Yes = 89.5 % No = 8.9 % NA=1.6 %	Yes = 94.5 % No = 5.5 % NA=0.0 %

^{*} Breast cup size was defined as 1 = AA; 2 = A; 3 = B; 4 = C; 5 = D; 6 = E/DD; 7 = F(E in Italy); 8 = G(F in Italy); 9 = H(FF in Italy); ≥ 10 = J(G in Italy); NA = not available.

^{**} Breast band was defined as 1 = 28 (UK); 2 = 30 (UK); 3 = 32 (UK), 70 (EU), 85 (FR), 1 (IT); 4 = 34 (UK), 75 (EU), 90 (FR), 2 (IT); 5 = 36 (UK), 80 (EU), 95 (FR), 3 (IT); 6 = 38 (UK), 85 (EU), 100 (FR), 4 (IT); 7 = 40 (UK), 90 (EU), 105 (FR), 5 (IT); 8 = 42 (UK), 95 (EU), 110 (FR), 6 (IT); 9 = 44 (UK), 100 (EU), 115 (FR), 7 (IT); 10 > above where EU = European Union; FR = France; IT = Italy; UK = United Kingdom.

facilitated the estimation of the residual uncertainty covering different patient characteristics in the full cohort. Multiple manual landmarks and breast contours were created on each patient's planning computed tomography (CT) scan in this cohort.

2.2. Selection of common coordinate system

To provide a suitable CCS for our patient subgroups, one patient was selected as a “reference patient” for 1) supine, 2) left-prone, and 3) right-prone positions. Note the prone patients were divided into left and right due to significant anatomical differences in breast shape and volume. To select the ‘most-average’ patient, we used variables collected in REQUITE: breast cup representing breast size, breast band representing chest diameter, and body mass index (BMI). Note that breast cup and breast band were mostly reported by the patient. For each subgroup, these patient characteristics were plotted as a 3D point cloud, and the patient at or nearest to the centroid was selected as CCSs.

2.3. Optimisation of image registration pathway

The performance of multiple DIR algorithms was investigated for each registration approach, Fig. 1. We compared the B-spline algorithm from NiftyReg [20] using either NMI (B-spline_NiftyReg-NMI) or sum of square difference (B-spline_NiftyReg-SSD) as a cost function, as well as B-spline and Demon algorithms in simple ITK [21] (B-spline_sITK-NMI and Demon_sITK-HistogramMatch, respectively). These algorithms were chosen to represent various solutions accessible to the research community. To reduce the bias related to DIR initialisation, all deformable registrations were performed after the same affine registration, which

was performed using Aladin (Affine_Aladin, part of NiftyReg).

2.3.1. Identifying the optimal image registration pathway

To identify the representative cohorts, patients in the full cohort were clustered according to breast cup size, breast band, and BMI using K-means: 100 clusters for the supine subgroup and 30 for each left-prone and right-prone subgroup. The selected representative patients were those at or nearest to the centroid of each cluster, see [supplementary Figure E.1](#).

1) Defining registration pathway for supine and prone set-up: We investigated two pathways of DIR for spatial normalisation for breast IBDM, Fig. 2. *Pathway 1* was a two-step process: In the first step, all patients in a similar set-up position were registered to their corresponding CCS, i.e., all supine patients were registered to the supine CCS, all left-prone patients to the left-prone CCS and all right-prone patients to the right-prone CCS. In the second step, the left- and right-prone CCS were registered to the supine CCS. In contrast, *Pathway 2* consisted in registering all patients in the representative supine, left- and right-prone subgroup directly to the supine CCS.

2) Evaluating the impact of arm position in the supine sub-group: We compared the registration performance between patients treated with bilateral arms raised (70/100) and ipsilateral arms raised (30/100) to the supine CCS patient (treated with bilateral arms raised) in the representative supine subgroup using Mann-Whitney test in SPSS version 29.

2.3.2. Evaluation of image registration performance

1) Evaluation of Image Registration for the Representative Cohort: Image registration performance was evaluated qualitatively using visual

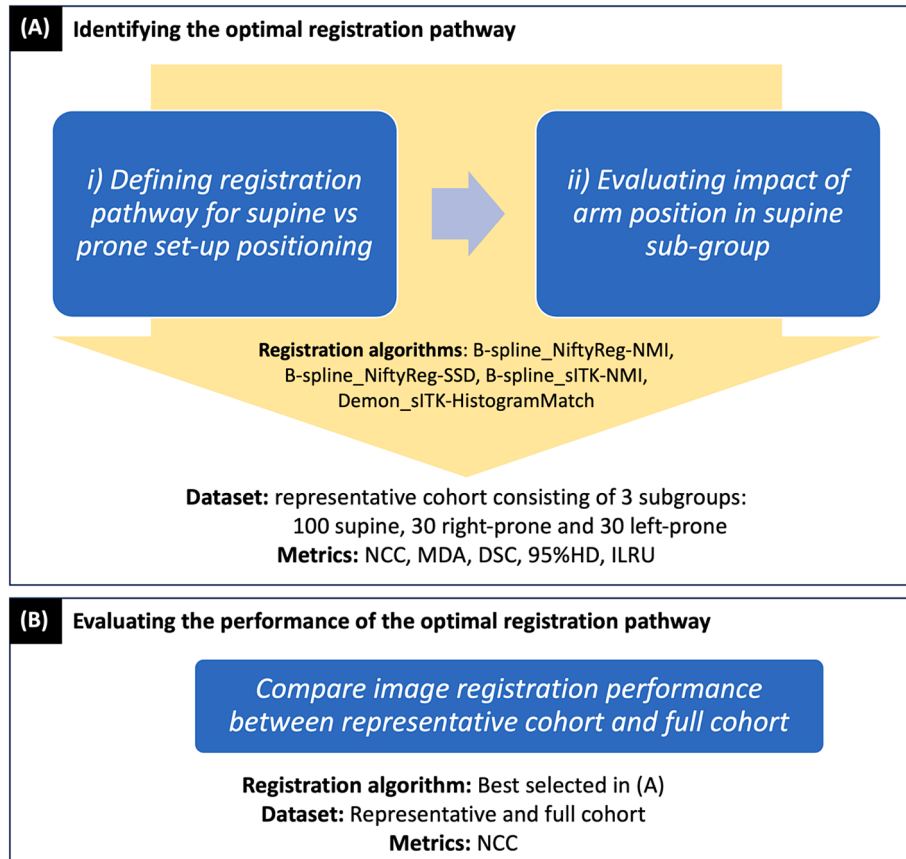


Fig. 1. Overview workflow for selecting the optimal spatial normalisation pathway for breast IBDM by considering (A) the representative cohort using difference image registration algorithm and evaluation metrics and (B) the full cohort. *NMI stands for normalised mutual information, SSD stands for sum of square difference, NCC stands for normalised correlation coefficient, MDA stands for mean distance to agreement, DSC stands for Dice similarity coefficient, HD stands for Hausdorff distance, and ILRU stands for interpatient landmark registration uncertainty.

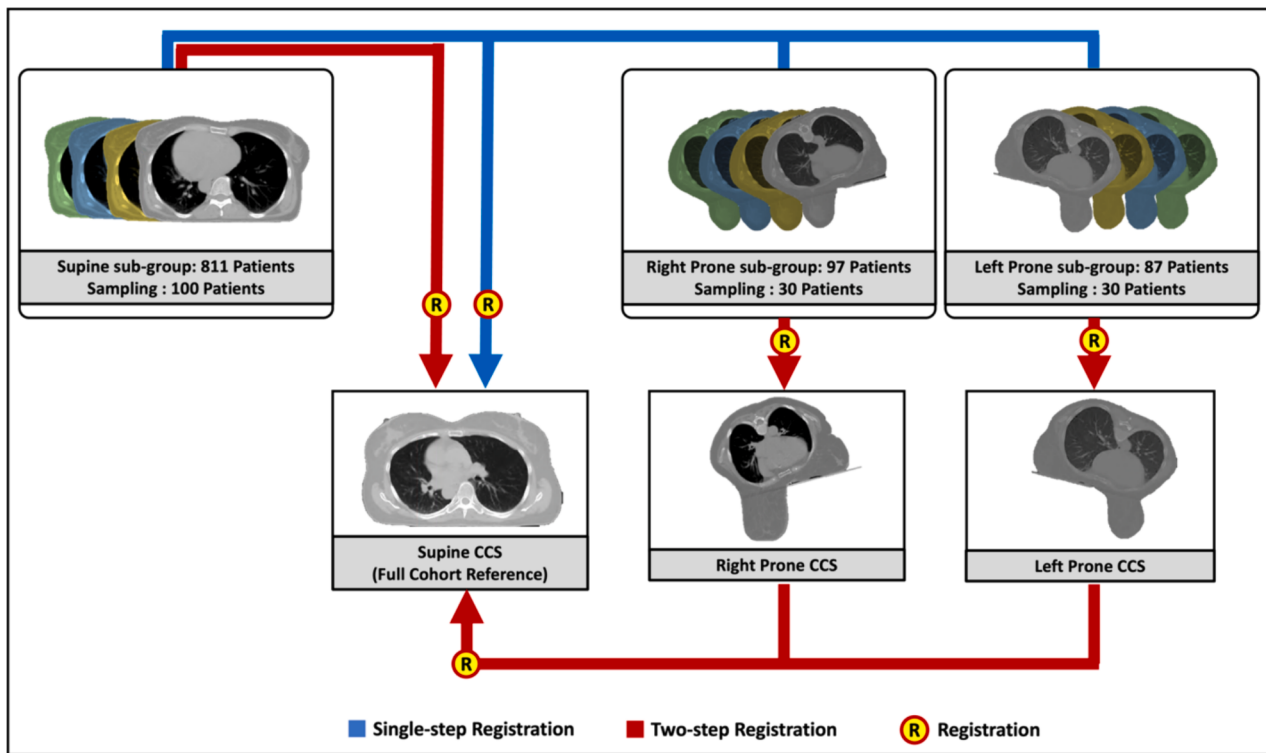


Fig. 2. Schematic of the two image registration pathways in our study. The red pathway represents the two-step registration (*Pathway 1*). First, the registration is performed within subgroups (using the CCS in the same set-up positioning), then the prone CCSs are registered to the supine CCSs and propagated this registration to the subgroup. The blue pathway represents a single-step registration (*Pathway 2*) where all patients were directly registered to the supine CCS, regardless of set-up positioning. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

inspection as well as quantitatively using five different metrics: 1) Normalised Cross Correlation (NCC), 2) Dice similarity coefficient (DSC), 3) mean distance to agreement (MDA), 4) 95 % Hausdorff distance (95 %HD) and 5) interpatient landmark registration uncertainty (ILRU).

NCC is an image-based metric used to estimate registration accuracy. We used a region of interest for NCC assessment on the CCSs covering 5 cm beyond the superior and inferior border of the breast (Fig. 3A). For

DSC, MDA and 95 % HD, we used ipsilateral and contralateral breast contours generated by atlas-based auto segmentation (Fig. 3B). For ILRU, bony and soft tissue landmarks were identified on the 3D CT images of the representative cohort by an expert clinician to estimate interpatient registration uncertainty, point-based evaluation similar to the transformation registration error (TRE) [22], (Fig. 3C). Definition and further information on these landmarks are detailed in [Supplementary Table E.1](#). Note that some landmarks were only identifiable for

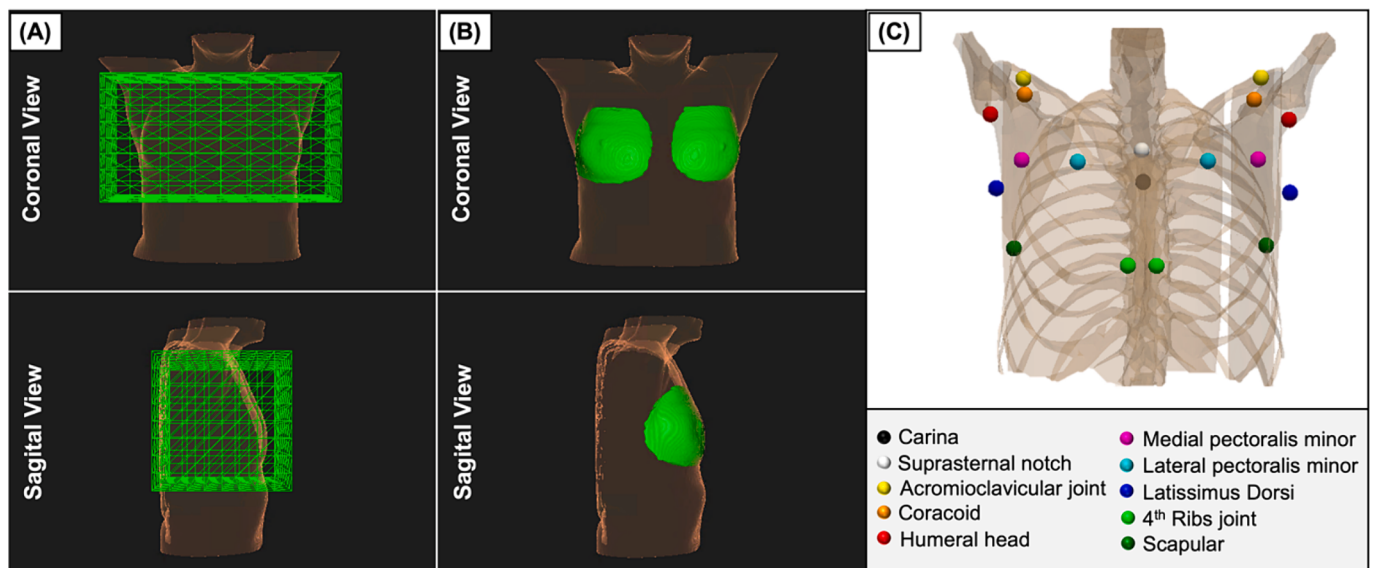


Fig. 3. The regions of interest for evaluating inter-patient image registration performance: (A) box over the chest region for NCC assessment in the representative and full cohort, (B) breast contour for DSC, MDA, and 95 %HD analysis, and (C) anatomical landmarks chosen to be relevant for breast radiotherapy for interpatient landmark registration uncertainty in the representative cohort.

raised arms due to large anatomical differences. The average uncertainty of image registration in each pathway was defined by the average standard deviation from ILRU as it provides the quantitative vector unit.

2) Evaluation of Image Registration for the Full Cohort: we evaluated the registration performance of the full cohort by using the image-based metric NCC. Differences in NCC between the representative cohort and full cohort were analysed using the Mann-Whitney test in SPSS version 29.

3) Consistency of Image Registration for Additional CCS: To investigate the reproducibility of image registration in multiple CCS, we randomly selected two patients around the centroid of the supine subgroup to be the new CCS (2nd and 3rd CCSs). DIR was performed for the supine subgroup to the first full cohort reference and two new CCSs. The performance of DIR among each CCS was reported in terms of NCC to ensure the consistency and repeatability of the spatial normalisation.

3. Results

3.1. Definition of image registration approach

Registration performance for all registration pathways and algorithms is shown in Fig. 4. The highest registration accuracy was obtained with B-spline_NiftyReg-NMI, achieving the lowest MDA (0.23 ± 0.19 cm, 0.38 ± 0.26 cm, and 1.19 ± 1.08 cm), 95 %HD (1.17 ± 1.18 cm, 1.55 ± 0.90 cm, and 4.18 ± 2.45 cm), and ILRU (0.51 cm, 0.53 cm, 0.90 cm) in supine-supine, prone-prone, and supine-prone registration, respectively. Likewise, it achieved a high NCC (0.98 ± 0.01 , 0.96 ± 0.03 , 0.95 ± 0.06) and high DSC for each sub-cohort (0.91 ± 0.04 , 0.86 ± 0.09 , and 0.61 ± 0.21). Registration pathway 1 (two-step) performed better than pathway 2. The best registration performance was observed for the supine-supine registrations ($\overline{sd} = 0.06$, 0.11, 0.36 cm, and 1.49 cm for NCC, DSC, MDA, and 95 %HD), common to both pathway 1 and pathway 2. Performance for the prone registration was better for pathway 1 (each prone patient registered to each prone specific CCS before registering to the supine CCS, $\overline{sd} = 0.07$, 0.16, 0.55 cm, and 1.57 cm for NCC, DSC, MDA, and 95 %HD), than for pathway 2 (all prone patients registered to the supine CCS, $\overline{sd} = 0.09$, 0.19, 1.19 cm, and 2.45 cm for NCC, DSC, MDA, and 95 %HD). Note that further dividing subgroups and using multiple CCSs did not improve registration performance, see [supplementary Figure E.2 and Table E.2](#).

Similar findings were observed when examining ILRU values presented in Table 2. ILRU for all landmarks were lower for supine-supine and prone-prone registrations compared to prone-supine CCS. We observed high variation in the shoulder region around 0.94 ± 0.22 cm and a low variation in the breast and chest region (0.52 ± 0.27 cm) when considering the ILRU of the B-spline_NiftyReg-NMI (Table 2). The average ILRU for Antero-posterior (AP), Superior-inferior (SI), and Right-left (RL) directions in the shoulder region were 0.16 cm, 0.62 cm, and 0.86 cm, and 0.21 cm, 0.43 cm, and 0.49 cm for chest and breast regions. The lowest mean variation was observed for the carina landmark (0.11 cm for similar position registration and 0.51 cm for difference position registration when considering at B-spline_NiftyReg-NMI), while the largest mean variation was found for the humeral head landmark, around 1 cm. For the best and worst registrations using B-spline_NiftyReg-NMI, see [supplementary Figure E.3](#).

In the representative supine subgroup, no difference in DIR accuracy was observed between ipsilateral and bilateral arm(s) raised (see [supplementary Figure E.4](#)). The NCC, DSC, MDA, 95 %HD, and ILRU for B-spline_NiftyReg-NMI registration from patients treated with bilateral arms raised were 0.98 ± 0.02 , 0.92 ± 0.03 , 0.21 ± 0.16 cm, 1.12 ± 1.11 cm, and 0.34 ± 0.15 cm, while these values were 0.98 ± 0.01 , 0.90 ± 0.04 , 0.27 ± 0.23 cm, 1.28 ± 1.33 cm, and 0.42 ± 0.18 cm for ipsilateral arm raised registration.

3.2. Evaluation of image registration for the full cohort

The median NCC for the full cohorts was 0.98 (IQR 0.01) when applying pathway 1, showing no significant difference from the representative cohort, $p = 0.534$ ([supplementary Figure E.5A](#)). By considering the performance of image registration in additional CCS, the median NCC for the first, second, and third CCS was 0.98 (IQR 0.01), 0.98 (IQR 0.01), and 0.98 (IQR 0.03), respectively ([supplementary Figure E.5B](#)).

4. Discussion

The feasibility of a spatial normalisation pathway for breast IBDM was demonstrated in our in-depth analysis. We identified a two-step registration approach using Nifty registration with NMI that is suitable for breast spatial normalisation. This approach optimised registration metrics in both the breast and the shoulder regions. To quantify the spatial normalisation uncertainty for both these regions, we propose and describe a set of landmarks other researchers can adopt. Quantifying the accuracy of spatial normalisation is key, as it affects dose mapping to the CCS and identifying any dose-sensitive regions [23,24].

We identified many challenges in applying spatial normalisation to breast patients. Our study shows the feasibility of using the patient at the centroid or nearby the centroid as a reference for entire sub-cohorts, divided by set-up position (supine, left-prone, right-prone). In previous IBDM work considered in other anatomical sites, the reference patient for IBDM was randomly selected from the entire cohort [16] or was represented by an anatomical phantom [25]. Our registration result from the selected approach for breast spatial normalisation complies with the criteria suggested by AAPM task group 132 when considering DSC (above 0.8) and MDA (0.2–0.3 cm) for the representative cohort and was in line with the recommendations in the “roadmap to clinical translation” proposed by McWilliam et al. [22,24]. Although the right-prone cohort registration showed a lower agreement of registration performance, it is still in the acceptable range of the recommendation.

We found that increasing the number of reference patients did not improve the registration accuracy, see [supplementary Figure E.2 and Table E.2](#); therefore, the use of a single reference patient is proposed for each set-up position. This result was also observed in the study by Vasquez Osorio et al. (2018) on the robustness of reference patient selection in lung cancer, which showed that the choice of reference patient did not change the identified region of IBDM [26]. However, a recent IBDM study on head and neck showed that in a small cohort of about 100 patients, different sub-regions could be identified when applying alternative reference patients [17]. Therefore, it will be important to use multiple CCS to assess the consistency of identified sub-regions in breast IBDM, even if the impact on registration accuracy is limited.

This work was performed with a cohort of about 1000 patients from 9 centres in 5 countries. As such, they represent a range of clinical practices, e.g. regarding CT image resolution, positioning approaches (prone vs supine, ranges of arm positions in the supine position), etc. as well as a range of BMI and breast size. However, 94 % of the cohort were European ancestry, and this lack of diversity constitutes a limitation. We investigated four different registration algorithms and showed that the choice of image registration algorithm did impact registration accuracy in the breast and shoulder area, depending on breast shape and volume. This is in agreement with published studies in lung cancer [27]. Therefore, consideration should be given to the selection of the best registration algorithm and optimisation parameters for breast IBDM. Because a gross misregistration will result in patients being excluded from the subsequent voxel-based analysis. Eliminating some patients could introduce biases and limit the applicability of the results to patients outside the cohort studied, as well as decrease the size or toxicity event of the cohort.

Evaluating image registration accuracy in breast cancer also requires assessment of the axilla and shoulder area in order to enable analysis of toxicity endpoints such as lymphoedema and shoulder morbidities.

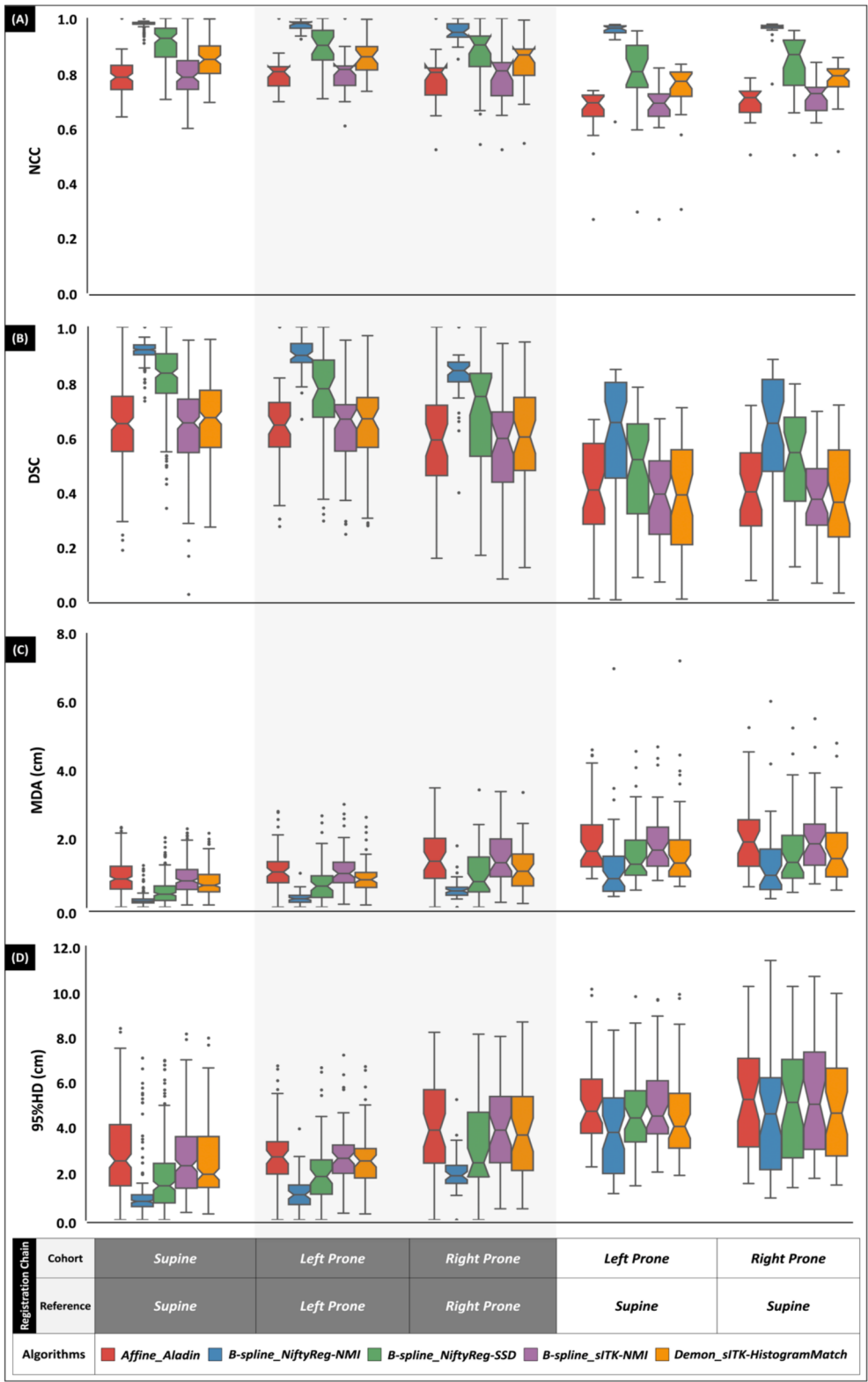


Fig. 4. The image registration accuracy for all registrations considering (A) NCC, (B) DSC, (C) MDA, and (D) 95% HD. For NCC and DSC, the ideal value is one, while for MDA and 95% HD, zero is ideal. Note that supine-supine CCS registrations are common for pathway 1 and pathway 2. Registrations for prone-prone CCS were part of pathway 1 (highlighted in grey), while registrations for prone-supine CCS were part of pathway 2. The performance of Affine_Aladin is presented as it is the preliminary step for all DIR pathways.

Table 2

Average standard deviation vector for evaluation of the interpatient landmark registration uncertainty (ILRU). Results are presented as combined for ipsi- and contralateral sides, and for ipsilateral side only. The landmarks representing the breast region include the 4thRibs Joint, Scapular, Suprasternal notch, Carina, Medial and Lateral pectoralis minor, and Latissimus dorsi, while the Acromioclavicular joint, Coracoid process, and Humeral head landmarks represent the shoulder region.

The average vector of standard deviation, $\overline{SD}$ (cm)							
Landmark		Approach 1 and 2:		Approach 1		Approach 2	
		Supine – Supine Registration		Prone – Prone Registration		Supine – Prone Registration	
		Affine_Aladi n	B-spline_NiftyReg- NMI	Affine_Aladi n	B-spline_NiftyReg- NMI	Affine_Aladin	B-spline_NiftyReg- NMI
Bony landmarks	Acromioclavicular joint	0.97	0.85	0.89	0.79	1.06	1.08
	Ipsilateral treatment side	0.93	0.63	0.97	0.94	0.99	0.74
	Coracoid process	1.35	0.73	0.74	0.67	0.95	0.95
	Ipsilateral treatment side	1.03	0.54	0.86	0.85	0.86	0.72
	Humeral head	0.94	0.86	1.24	1.15	1.59	1.36
	Ipsilateral treatment side	0.88	0.72	1.79	1.64	1.95	1.23
	4thRibs Joint	0.43	0.28	0.60	0.59	1.08	0.99
	Ipsilateral treatment side	0.32	0.23	0.76	0.80	0.94	0.51
	Scapular	0.55	0.56	0.42	0.44	0.86	0.82
	Ipsilateral treatment side	0.45	0.39	0.42	0.47	0.71	0.74
	Suprasternal notch	0.23	0.25	0.31	0.19	0.76	0.64
	Soft tissue	Carina	0.24	0.14	0.23	0.08	0.59
Lateral pectoralis minor		0.64	0.52	0.75	0.68	1.00	0.96
Ipsilateral treatment side		0.62	0.48	0.74	0.76	0.76	0.60
Latissimus dorsi		0.59	0.59	0.51	0.45	0.86	0.82
Ipsilateral treatment side		0.47	0.43	0.59	0.56	0.62	0.58
Medial pectoralis minor		0.31	0.30	0.34	0.28	0.98	0.90
Ipsilateral treatment side	0.28	0.24	0.36	0.26	0.96	0.87	

*Highlight is the best registration result among each pathway of the landmark investigated.

Image-based metric such as NCC may not be suitable for analysing the shoulder region due to the differences in the set-up of arm position. The shoulder and axillary anatomy are complexly linked bony, soft tissue and vascular structures which are not routinely delineated in breast radiotherapy. Our study focuses on breast registration. However, registration of OARs would also be of interest, for example, to investigate late cardiac effects. Though this is out of scope for this work, the pipeline could easily be modified to allow this investigation in the future.

In this study, we only evaluated the performance of image registration based on the breast contour and anatomical landmarks within the external thorax region. The inter-observer variation of both contour (e.g., breast and heart) and landmarks can reduce the validity of the geometrical evaluation of DIR [28,29]. Hence, delineating more relevant contours for breast radiotherapy, such as the humeral head, major- and minor-pectoralis muscle, and individual lymph node level, could increase the robustness of registration analysis. However, there is currently a lack of a universally accepted benchmark for evaluating image registration in the breast, shoulder, and axilla region, whether using contour-based or point-based methods. It is the fact that it is difficult to define soft tissue landmarks directly in the axillary region. Therefore, we propose a set of anatomical points, [Supplementary Table E.1](#), based on the feasibility of consistent, reproducible identification on CT scan and coverage of the relevant anatomy for the expected radiotherapy dose distribution (on and off target) in patients treated with whole breast and/or regional lymph node irradiation.

Even though our study found spatial normalisation for IBDM in breast RT was feasible, we still faced some limitations when using the best-performing algorithm of our study (B-spline NiftyReg-NMI) in the very large and pendulous breast where the breast could fall by the side of the body. Thus, investigating skin toxicity for breast radiotherapy may not be possible when applying our registration algorithm. Perhaps an additional landmark for point-point registration [30], surface-based registration, or using artificial intelligence for breast registration may further improve the performance of DIR [31]. However, those methods cannot yet achieve the performance observed in our study [32]. In addition, AI-based DIRs will require a large training model dataset to achieve robust results.

Our study includes only free-breathing patients. However, many breast cancer treatments are now delivered in deep inspiration breath-hold patients. As DIBH images are clearer and contain fewer artifacts, we expect that the performance of our pipeline would only improve, but this hypothesis will require confirmation.

Only one experienced clinician generated landmarks in our study; therefore, it might generate a subjective error, and multiple observers may be required. In future research, using our proposed landmarks in a point-based registration may be beneficial. In addition, dose metric evaluation is not performed in our spatial normalisation study. Still, it would benefit the dose mapping variation assessment, the next step of the voxel-based analysis [24]. Finally, though we have demonstrated the feasibility of DIR performance in breast RT, it remains important to test DIR performance in each cohort where IBDM is applied, especially when gathering large cohort data from multiple centres.

In conclusion, spatial normalisation for breast IBDM is feasible. Our findings will be useful for enabling IBDM investigations and integrating the registration uncertainty in the spatial normalisation processes for breast IBDM.

CRediT authorship contribution statement

Tanwihat Jaikuna: Conceptualization, Methodology, Software, Formal analysis, Investigation, Writing – original draft, Writing – review & editing, Visualization, Project administration. **Fiona Wilson:** Conceptualization, Methodology, Writing – original draft, Writing – review & editing. **David Azria:** Resources, Data curation, Writing – review & editing. **Jenny Chang-Claude:** Resources, Data curation,

Writing – review & editing. **Maria Carmen De Santis:** Resources, Data curation, Writing – review & editing. **Sara Gutiérrez-Enríquez:** Resources, Data curation, Writing – review & editing. **Marcel van Herk:** Writing – review & editing, Supervision. **Peter Hoskin:** Writing – review & editing, Supervision. **Lea Kotzki:** Resources, Data curation, Writing – review & editing. **Maarten Lambrecht:** Resources, Data curation, Writing – review & editing. **Zoe Lingard:** Resources, Data curation, Writing – review & editing. **Petra Seibold:** Resources, Data curation, Writing – review & editing. **Alejandro Seoane:** Resources, Data curation, Writing – review & editing. **Elena Sperk:** Resources, Data curation, Writing – review & editing. **R Paul Symonds:** Resources, Data curation, Writing – review & editing. **Christopher J. Talbot:** Resources, Data curation, Writing – review & editing. **Tiziana Rancati:** Resources, Data curation, Writing – review & editing. **Tim Rattay:** . **Victoria Reyes:** Resources, Data curation, Writing – review & editing. **Barry S. Rosenstein:** Resources, Data curation, Writing – review & editing. **Dirk de Ruyscher:** Resources, Data curation, Writing – review & editing. **Ana Vega:** Resources, Data curation, Writing – review & editing. **Liv Veldeman:** Resources, Data curation, Writing – review & editing. **Adam Webb:** Resources, Data curation, Writing – review & editing. **Catharine M L West:** Resources, Data curation, Writing – review & editing, Supervision. **Marianne C Aznar:** Conceptualization, Methodology, Writing – review & editing, Supervision. **Eliana Vasquez Osorio:** Conceptualization, Methodology, Writing – review & editing, Supervision.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.phro.2024.100635>.

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