Adaptive statistical iterative reconstruction (ASIR) affects CT radiomics quantification in primary colorectal cancer

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Abstract

Objectives To investigate whether adaptive statistical iterative reconstruction (ASIR), a hybrid iterative CT image reconstruction algorithm, affects radiomics feature quantification in primary colorectal cancer compared to filtered back projection. Additionally, to establish whether radiomics from single-slice analysis undergo greater change than those from multi-slice analysis.

Methods Following review board approval, contrast-enhanced CT studies from 32 prospective primary colorectal cancer patients were reconstructed with 20\% ASIR level increments, from 0 to 100\%. Radiomics analysis was applied to single-slice and multi-slice regions of interest outlining the tumour: 70 features, including statistical (first-, second- and high-order) and fractal radiomics, were generated per dataset. The effect of ASIR was calculated by means of multilevel linear regression.

Results Twenty-eight CT datasets were suitable for analysis. Incremental ASIR levels determined a significant change ($p < 0.001$) in most statistical radiomics features, best described by a simple linear relationship. First-order statistical features, including mean, standard deviation, skewness, kurtosis, energy and entropy, underwent a relatively small change in both single-slice and multi-slice analysis (median standardised effect size $B = 0.08$). Second-order statistical features, including grey-level co-occurrence and difference matrices, underwent a greater change in single-slice analysis (median $B = 0.36$) than in multi-slice analysis (median $B = 0.13$). Fractal features underwent a significant change only in single-slice analysis (median $B = 0.49$).

Conclusions Incremental levels of ASIR affect significantly CT radiomics quantification in primary colorectal cancer. Second-order statistical and fractal features derived from single-slice analysis undergo greater change than those from multi-slice analysis.

Key Points

\begin{itemize}
  \item Incremental levels of ASIR determine a significant change in most statistical (first-, second- and high-order) CT radiomics features measured in primary colorectal cancer, best described by a linear relationship.
  \item First-order statistical features undergo a small change, both from single-slice and multi-slice radiomics analyses.
  \item Most second-order statistical features undergo a greater change in single-slice analysis than in multi-slice analysis. Fractal features are only affected in single-slice analysis.
\end{itemize}

Keywords Multidetector computed tomography \& Image processing \& Computer-assisted \& Colorectal neoplasms \& Fractals
Abbreviations
2D 2-dimensional
3D 3-dimensional
ASIR Adaptive statistical iterative reconstruction
CT Computed tomography
FBP Filtered back projection
GLCM Grey-level co-occurrence matrix
GLDM Grey-level difference matrix
GLRL Grey-level run length
GLZSM Grey-level zone-size matrix
MBIR Model-based iterative reconstruction
NGTDM Neighborhood grey-tone difference matrix
ROI Region of interest
SD Standard deviation

Introduction

There has been a growing interest in radiomics approaches that extract quantitative image features to improve cancer phenotyping [1]. Radiomics have shown promise for tumour characterisation, prognostication, therapy planning and therapy assessment in a number of cancers including non-small cell lung cancer [2–8], breast [9–12], prostate [13, 14] and colorectal cancer [15–21].

Advances in computed tomography (CT) technology, linked to dose reduction, have led to the implementation of iterative image reconstruction algorithms in order to compensate for the increase in image noise with low-dose CT acquisitions [22–26]. Adaptive statistical iterative reconstruction (ASIR) is a hybrid algorithm that uses the image information obtained from filtered back projection (FBP) as the basis for iterative reconstruction to optimise image quality. It has enabled dose reductions between 32 and 65%, without substantially affecting image quality in phantom studies [27, 28].

To date, the majority of published retrospective CT radiomics studies have been based on images reconstructed with FBP alone. Given that iterative reconstruction has been adopted by all manufacturers in current CT scanners, the impact of iterative reconstruction algorithms on quantitative image features is highly relevant to the field of radiomics. We hypothesised that ASIR would alter feature values significantly compared with FBP, and alter features progressively with incremental ASIR weightings (percentage increments). Thus, the primary aim of our study was to investigate whether hybrid iterative reconstruction, specifically ASIR applied with incremental weightings, affects the quantification of radiomics features, including first-, second- and high-order statistical as well as fractal parameters, using primary colorectal cancer at peak contrast enhancement as an exemplar in light of promising data [29]. Our secondary aim was to establish whether features calculated from single-slice analysis (2-dimensional [2D] radiomics) are influenced to a greater degree than those from multi-slice analysis (3-dimensional [3D] radiomics).

Materials and methods

Participants

Following institutional review board approval and informed consent, 32 consecutive patients with primary colorectal cancer underwent contrast-enhanced CT from January 2012 to July 2014 from a single institution (prospective trial, ISCTRNR 95037515). Exclusion criteria were tumour diameter <2 cm (to assure a sufficient number of CT voxels for analysis), impaired renal function (estimated glomerular filtration rate <50 mL/min) and previous iodinated contrast media allergic reaction precluding administration of an iodinated contrast agent.

CT acquisition and reconstruction

CT was performed on a single Discovery 750 HD multidetector CT scanner (GE Healthcare). As part of a prospective research protocol, a dynamic contrast-enhanced CT centred on the primary cancer was acquired using the following protocol: 100 kV; 75 mAs; z-axis coverage, 4 cm; scan field of view, 50 cm; matrix, 512 × 512 mm; B30 soft reconstruction kernel; 5 mm reconstructed slice thickness; axial mode with 35 acquisition time points at a 1.5-s temporal resolution for 45 s and a 5-s temporal resolution thereafter for 120 s. To minimise bowel peristaltic movement, 20 mg of the spasmyloytic agent hyoscine butylbromide (Buscopan; Boehringer Ingelheim) was administered intravenously prior to data acquisition unless contraindicated. The contrast agent was administered as follows: 50 mL of 370 mg/mL iodinated contrast agent (Niopam, Bracco) via a pump injector (Medrad Stellant dual syringe, Bayer Healthcare) at a rate of 5 mL/s, followed by a 50 mL saline chaser at the same rate. Mean CTDIvol and DLP were 137.8 ± 15.3 mGy and 551.0 ± 61.2 mGy cm, respectively. The dynamic acquisition was reconstructed at the scanner with six different ASIR percentages: 0%, (equivalent to FBP) 20%, 40%, 60%, 80% and 100%, resulting in six separate datasets per patient (Fig. 1).

Image analysis

Image analysis was carried out by two readers in consensus (a clinical oncologist and a radiologist with 5 and 10 years CT experience, respectively) using the CT acquisition corresponding to peak tumour enhancement, in order to maximise the tumour contrast-to-noise ratio. A free-hand region of interest (ROI) was drawn around the tumour on 0% ASIR reconstructions to generate two datasets per patient: (1) a single axial
image corresponding to the largest tumour area; (2) five contiguous axial images including tumour. ROIs were then copied onto 20–100% ASIR reconstructions to ensure these were identical.

Radiomics features were extracted using an in-house software based on Matlab (Mathworks), previously validated on a digital phantom developed as part of the Image Biomarker Standardisation Initiative [30]. A medium smoothing filter and 32 bin width were applied to the native DICOM images.

First-order (histogram), second-order (grey-level co-occurrence matrix, GLCM; grey-level difference matrix, GLDM) and high-order (neighbourhood grey-tone difference matrix, NGTDM; grey-level run length, GLRL; grey-level zone-size matrix, GLZSM) statistical, as well as model-based fractal features were extracted. The extracted features are summarised in Table 1.

**Statistical analysis**

All radiomics features were regarded as continuous in nature. To allow for the fact that measurements from the same subject were likely to be more similar than from different subjects, the analysis was performed using multilevel linear regression. Two-level models were used, with individual measurements nested within patients. The shape of the relationship between ASIR levels and radiomics parameter values was examined. Initially, cubic, squared and linear terms for ASIR were included in the analysis. If the higher order terms (i.e. cubic and squared terms) were not found to be statistically significant, they were omitted and a simple linear relationship between variables was assumed. The distributions of the majority of radiomics parameters were such that the assumptions of the statistical methods were met. However, one parameter (fractal dimension lacunarity) had a particularly positively skewed distribution, and was thus analysed on the log scale. Regression coefficients along with corresponding confidence intervals were reported, representing the change in the CT parameter value for a 20% increase in ASIR. Standardised effect sizes ($B$) were calculated by the absolute regression coefficient divided by the between-subject standard deviation for a 20-unit increase in ASIR. Conventionally, $B$ values $\leq 0.3$ are considered representative of a small effect; $B$ values $> 0.3$ and $< 0.8$ are considered a moderate effect; $\geq 0.8$, a large effect. All analyses were undertaken by a statistician using Stata, version 13.1 (StataCorp LP). A $p$ value $< 0.05$ was taken to represent statistical significance.
Results

Participants

No tumour was identified on CT in 1 participant (confirmed histological diagnosis of colorectal adenocarcinoma based on a resected polyp). Voxel spatial mismatch between 0% ASIR and subsequent ASIR reconstructed series precluded the analysis in 3 further participants. Therefore, 28 CT datasets were suitable for single-slice (2D) radiomics analysis. Multi-slice (3D) analysis was only possible in a subset of 23 participants, as 5 tumours were not sufficiently large or appropriately oriented to allow segmentation on at least 5 consecutive axial slices. Tumours had a mean diameter of 5.7 ± 1.74 cm and were located as follows: caecum, n = 4; ascending colon, n = 2; sigmoid colon, n = 7; rectum, n = 15. Radiological tumour (T) and nodal (N) staging, evaluated on CT images and based on the AJCC/UICC TNM classification of malignant tumours (8th edition), was as follows: T2, n = 9; T3, n = 17; T4, n = 2; N0, n = 16; N1, n = 10; N2, n = 2.

Radiomics analysis

Absolute regression coefficients, standardised effect size $B$ values and corresponding $p$ values for first-order, second-order and fractal radiomics features from single-slice and multi-slice datasets are summarised in Table 2; values for high-order features from multi-slice datasets are summarised in Table 3.

Single-slice analysis

All first-order, second-order and fractal features varied significantly and according to a linear relationship with increasing ASIR values, with the exception of GLCM sum entropy (Table 2). Some features significantly increased; others decreased.

The relative effect was small on all first-order features (median standardised effect size $B = 0.08$; range, 0.02–0.24). Most second-order features, including all grey-level difference matrix (GLDM) and the majority of grey-level co-occurrence matrix (GLCM) features, were moderately affected (median $B = 0.36$;
## Table 2
Summary of results for first-order, second-order and fractal features from single-slice (2D) and multi-slice (3D) analysis. For each radiomics feature, the regression coefficients (95% confidence intervals) and standardised effect size (B) for a 20% increase in ASIR, and the corresponding p values, are presented. Significant p values are marked in italics.

| CT radiomics                  | 2D regression coefficient (95% CI) | 2D p value | 3D regression coefficient (95% CI) | 3D p value |
|-------------------------------|------------------------------------|------------|-----------------------------------|------------|
| **First-order features**      |                                    |            |                                   |            |
| Mean                          | $-0.05 (-0.06, -0.03)$             | 0.01       | $-0.18 (-0.30, -0.05)$            | 0.01       |
| Maximum                       | $-5.9 (-6.3, -5.2)$                | 0.24       | $-6.1 (-6.8, -5.3)$              | 0.27       |
| Minimum                       | 3.9 (3.0, 4.8)                     | 0.04       | $-8.0 (-13.6, -2.5)$             | 0.04       |
| Range                         | $-9.8 (-11.2, -8.3)$               | 0.09       | $2.0 (-3.6, 7.6)$                | 0.01       |
| Standard deviation            | $-1.27 (-1.4, -1.1)$               | 0.18       | $-0.78 (-1.04, -0.52)$           | 0.06       |
| Coefficient of variation      | $-0.022 (-0.025, -0.018)$          | 0.10       | $-0.010 (-0.019, -0.002)$        | 0.02       |
| Kurtosis                      | 0.33 (0.21, 0.45)                  | 0.05       | $2.1 (1.4, 2.8)$                 | 0.10       |
| Skewness                      | $-0.064 (-0.078, -0.052)$          | 0.08       | $-0.18 (-0.23, -0.13)$           | 0.12       |
| Energy                        | 0.0010 (0.0007, 0.0015)            | 0.04       | $0.005 (0.004, 0.007)$           | 0.11       |
| Entropy                       | $-0.008 (-0.013, -0.003)$          | 0.02       | $-0.047 (-0.060, -0.035)$        | 0.10       |
| **Second-order features**     |                                    |            |                                   |            |
| GLCM Autocorrelation          | 8.5 (7.2, 9.9)                     | 0.19       | $68 (54, 82)$                    | 0.11       |
| GLCM Cluster prominence       | 787 (555, 1018)                    | 0.07       | $70 (-92, -48)$                  | 0.10       |
| GLCM cluster shade            | $-28 (-35, -20)$                   | 0.12       | $4147 (2692, 5602)$              | 0.09       |
| GLCM contrast                 | $-2.9 (-3.2, -2.5)$                | 0.35       | $-2.4 (-3.0, -1.8)$              | 0.10       |
| GLCM correlation              | 0.038 (0.034, 0.042)               | 0.50       | $0.017 (0.014, 0.020)$           | 0.31       |
| GLCM difference entropy       | $-0.053 (-0.060, -0.048)$          | 0.44       | $-0.05 (-0.06, -0.04)$           | 0.16       |
| GLCM difference variance      | $-2.9 (-3.2, -2.5)$                | 0.35       | $-2.4 (-3.0, -1.8)$              | 0.10       |
| GLCM dissimilarity            | $-0.23 (-0.26, -0.20)$             | 0.38       | $-0.20 (-0.24, -0.16)$           | 0.14       |
| GLCM entropy                  | $-0.056 (-0.063, -0.050)$          | 0.37       | $-0.09 (-0.11, -0.07)$           | 0.13       |
| GLCM energy                   | 0.0003 (0.0003, 0.0004)            | 0.50       | $0.0010 (0.0008, 0.0012)$        | 0.16       |
| GLCM homogeneity              | 0.012 (0.011, 0.014)               | 0.43       | $0.013 (0.011, 0.015)$           | 0.19       |
| GLCM Information measure correlation 1 | $-0.010 (-0.012, -0.009)$        | 0.24       | $-0.008 (-0.010, -0.007)$        | 0.35       |
| GLCM Information measure correlation 2 | 0.017 (0.014, 0.019)               | 0.16       | $0.013 (0.010, 0.015)$           | 0.15       |
| GLCM Inverse difference moment normalised | 0.0024 (0.0021, 0.0027)          | 0.36       | $0.0006 (0.0004, 0.0007)$        | 0.11       |
| GLCM inverse difference moment normalised | 0.0052 (0.0046, 0.0057)         | 0.39       | $0.0027 (0.0022, 0.0032)$        | 0.14       |
| GLCM maximum probability      | 0.0009 (0.0007, 0.0010)            | 0.40       | $0.0024 (0.0019, 0.0029)$        | 0.17       |
| GLCM sum average              | 0.40 (0.33, 0.47)                  | 0.17       | $1.40 (1.11, 1.68)$              | 0.11       |
| GLCM sum entropy              | 0.0000 (0.0001, 0.0002)            | 0.01       | $0.026 (-0.035, -0.018)$         | 0.07       |
| GLCM Sum of squares variance  | 7.1 (5.9, 8.3)                     | 0.17       | $67 (53, 81)$                    | 0.11       |
| GLDM mean                     | $-0.23 (-0.26, -0.20)$             | 0.38       | $-0.20 (-0.24, -0.16)$           | 0.14       |
| GLDM entropy                  | $-0.054 (-0.060, -0.048)$          | 0.44       | $-0.05 (-0.06, -0.04)$           | 0.16       |
| GLDM variance                 | $-0.96 (-1.08, -0.84)$             | 0.36       | $-0.64 (-0.83, -0.44)$           | 0.08       |
| GLDM contrast                 | $-2.86 (-3.21, -2.50)$             | 0.35       | $-2.38 (-2.96, -1.81)$           | 0.10       |
| **Fractal features**          |                                    |            |                                   |            |
| Fractal lacunarity            | 0.09 (0.08, 0.10)                  | 0.55       | $-0.001$                         |            |
| Fractal dimension mean        | $-0.029 (-0.033, -0.026)$          | 0.82       | $-0.001$                         |            |
| Fractal dimension SD          | 0.011 (0.010, 0.013)               | 0.43       | $-0.001$                         |            |
| Hurst exponent                | 0.029 (0.026, 0.033)               | 0.82       | $-0.001$                         |            |
| Blanket mean                  | $-0.056 (-0.062, -0.050)$          | 0.25       | $-0.001$                         |            |
| Blanket max                   | $-0.056 (-0.062, -0.050)$          | 0.25       | $-0.001$                         |            |

*Calculated as absolute regression coefficient divided by between-subject standard deviation; * Variable analysed on the log scale
range, 0.01–0.44). The relative effect on most fractal features was moderate to large (median $B = 0.49$; range, 0.25–0.82).

**Multi-slice analysis**

Nearly all first-order, second-order and high-order features varied significantly with increasing ASIR values: some increasing, others decreasing. Exceptions were first-order range, neighbourhood grey-tone difference matrix (NGTDM) complexity and 3 out of 13 grey-level run length (GLRL) features (Tables 2 and 3).

A small effect was confirmed on first-order statistical features (median $B = 0.08$; range, 0.01–0.27). Second-order features were affected to a lesser degree than from single-slice analysis (median $B = 0.13$; range, 0.07–0.35): 21 out of 23 features were affected in small measure, compared to 9 out of 23 from single-slice analysis. ASIR effect was small on high-order features (median $B = 0.09$; range, 0.01–0.31). In contrast to single-slice analysis and to all other features, multi-slice fractal features did not change significantly with increasing ASIR levels.

**Discussion**

Despite the rising number of studies investigating the clinical potential of radiomics in cancer imaging, relatively little is known

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**Table 3** Summary of results for high-order features obtained from multi-slice (3D) analysis. For each texture feature, the regression coefficients (95% confidence intervals [CI]) and standardised effect size ($B$) for a 20% increase in ASIR, and the corresponding $p$ values, are presented. Significant $p$ values are marked in italics.

| CT radiomics | Regression coefficient (95% CI) | $B$ value* | $p$ value |
|--------------|---------------------------------|------------|-----------|
| High-order features | | | |
| NGTDM coarseness | $-0.07 (-0.09, -0.05)$ | 0.05 | $< 0.001$ |
| NGTDM contrast | $-0.004 (-0.006, -0.002)$ | 0.04 | $< 0.001$ |
| NGTDM busyness | $0.025 (0.016, 0.034)$ | 0.04 | $< 0.001$ |
| NGTDM complexity | | | |
| NGTDM texture strength | $0.039 (0.024, 0.054)$ | 0.08 | $< 0.001$ |
| GLRL run percentage | $-0.010 (-0.012, -0.009)$ | 0.22 | $< 0.001$ |
| GLRL high grey-level run emphasis | $0.25 (0.17, 0.32)$ | 0.01 | $< 0.001$ |
| GLRL short-run low grey-level emphasis | | | |
| GLRL short-run high grey-level emphasis | $0.16 (0.10, 0.22)$ | 0.006 | $< 0.001$ |
| GLRL short-run emphasis | $0.0004 (0.0003, 0.0005)$ | 0.02 | $< 0.001$ |
| GLRL long-run emphasis | $48 (37, 58)$ | 0.11 | |
| GLRL grey-level non-uniformity | $-209 (-273, -146)$ | 0.06 | $< 0.001$ |
| GLRL run length non-uniformity | | | |
| GLRL low grey-level run emphasis | $-0.009 (-0.010, -0.007)$ | 0.22 | $< 0.001$ |
| GLRL long-run low grey-level emphasis | $18.9 (11.1, 26.7)$ | 0.06 | $< 0.001$ |
| GLRL long-run high grey-level emphasis | $344 (279, 410)$ | 0.24 | $< 0.001$ |
| GLRL intensity variability | $-37,623 (-56,176, -19,070)$ | 0.05 | $< 0.001$ |
| GLRL run length variability | | | |
| GLZLSM short-zone emphasis | $0.05 (0.04, 0.06)$ | 0.17 | $< 0.001$ |
| GLZLSM short-zone low-intensity emphasis | $5.3 (2.9, 7.7)$ | 0.13 | $< 0.001$ |
| GLZLSM short-zone high-intensity emphasis | $0.048 (0.033, 0.063)$ | 0.17 | $< 0.001$ |
| GLZLSM long-zone low-intensity emphasis | $-9.8 (-16.4, -3.2)$ | 0.04 | $0.004$ |
| GLZLSM long-zone high-intensity emphasis | $544,100 (368,325; 719,875)$ | 0.18 | $< 0.001$ |
| GLZLSM long-zone emphasis | $47 (35, 59)$ | 0.09 | $< 0.001$ |
| GLZLSM intensity non-uniformity | $-145 (-190, -102)$ | 0.14 | $< 0.001$ |
| GLZLSM zone length non-uniformity | $-11.5 (-15.3, -7.7)$ | 0.10 | $< 0.001$ |
| GLZLSM zone percentage | $-0.25 (-0.028, -0.021)$ | 0.20 | $< 0.001$ |
| GLZLSM low-intensity zone emphasis | $0.015 (-0.017, -0.013)$ | 0.31 | $< 0.001$ |
| GLZLSM high-intensity zone emphasis | $161 (110, 212)$ | 0.17 | $< 0.001$ |
| GLZLSM intensity variability | $-3503 (-5140, -1866)$ | 0.14 | $< 0.001$ |
| GLZLSM size zone variability | $-339 (-511, -167)$ | 0.13 | $< 0.001$ |

* Calculated as absolute regression coefficient divided by between-subject standard deviation; " variable analysed on the log scale. ^ Figures reported in units of $10^{-5}$
on how the shift of CT image reconstruction from filtered back projection to hybrid iterative algorithms might affect radiomics features quantitation. In our prospective study, we found that the application of incremental ASIR levels altered most statistical (first-, second- and high-order) features according to a linear relationship, both in single-slice and multi-slice analysis. Fractal features changed significantly only in single-slice datasets. While first-order features underwent small relative effects across all datasets, second-order features underwent a greater change in single-slice than in multi-slice analysis.

A fundamental aspect of CT is the assignment of an attenuation value to a voxel. Analytical reconstruction algorithms, namely filtered back projection, have long been the backbone of CT reconstruction. However, as image noise from the Poisson statistical variation across an image is not accounted for by FBP; this has been a limiting factor for low-dose CT imaging [31]. Iterative reconstruction algorithms have thus risen to the fore with advances in scanner hardware and increasing computing power [31]. Hybrid algorithms such as ASIR combine both analytical and iterative methods, optimising image characteristics by decreasing image noise. However, there is a perceptive alteration in image characteristics with high ASIR increments often described as a ‘waxy’ appearance. Given this, we hypothesised that radiomics features may be altered by using ASIR compared to FBP and the ASIR weighting.

To date, only one other clinical study has compared the effect of different reconstruction algorithms on quantitative CT data including a limited number of histogram and second-order GLCM features. The effect of FBP, 50% ASIR and model-based iterative reconstruction (MBIR) was compared for volumes of interest in 20 patients with liver lesions (n = 13), lung nodules (n = 9) or renal calculi (n = 25) who underwent either non-enhanced CT or contrast-enhanced CT at a fixed 120 kVp but 2 different dose levels (full and half-dose) [32]. MBIR had the highest impact on features. Fifty percent ASIR had a significant effect on standard deviation (SD) but not on other first-order (entropy, kurtosis or skewness) or second-order GLCM features studied. These differences in part reflect a difference in study design, i.e. assessment of an incremental effect of ASIR as opposed to a comparison of a single ASIR percentage to FBP and MBIR.

We acknowledge that the reconstruction algorithm is only one of several factors that potentially affect CT radiomics features. Other factors include acquisition factors, e.g. kVp, mAs, reconstruction kernel, voxel size, grey-level discretisation and contrast administration [32–39]. For example, Zhao et al investigated the effect of reconstruction kernel on 89 unenhanced CT radiomics features including shape, first-order, second-order statistical, wavelet and fractal features for 32 lung cancers using 2D and 3D images. The reconstruction kernel had a significant effect on extracted features, with smooth images having a smaller effect than noisier images. 3D images were also more reproducible than 2D images [34]. This was also echoed in our study where 3D features were more stable.

He et al assessed the effects of reconstruction slice thickness, reconstruction kernel and contrast-enhancement on the diagnostic performance of 150 radiomics features in 240 patients with solitary pulmonary nodules (malignant, n = 180; benign, n = 60). This study demonstrated better discrimination and classification for malignant versus benign nodules when based on unenhanced versus contrast-enhanced CT, thin (1.25 mm) versus thick-slice CT (5 mm) and standard versus lung reconstruction kernel [35]. In a further study, Shafiq-Ul-Hassan et al found that voxel size and discretisation were also important factors affecting radiomics features including shape, intensity, GLCM, GLZSM, GLRL, NGTDM, fractal and wavelet features in a digital phantom [39].

From our study and published data, minimisation of variation in reconstruction kernel, reconstruction algorithm, voxel size and grey-level discretisation would improve the quantification and stability of CT radiomics features.

We acknowledge some limitations to our study. Firstly, the study cohort was small (n = 28) but with the advantage of fixed acquisition parameters, including kVp and mAs, reconstruction kernel, and voxel size, allowing us to focus on the effect of the reconstruction algorithm alone. Secondly, we did not assess the effect of other iterative reconstruction algorithms as these were not available to us at the time of acquisition; we acknowledge that this would be of value going forward in future studies.

In conclusion, we confirmed that the application of a hybrid reconstruction algorithm versus traditional FBP affects CT radiomics quantification. The use of multi-slice (3D) rather than single-slice (2D) data will minimise this effect, particularly for second-order statistical and model-based fractal features. The reconstruction algorithm should be taken into account and standardised when acquiring data for future multicentre CT radiomics studies.

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Compliance with ethical standards

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Informed consent Written informed consent was obtained from all subjects (patients) in this study.

Ethical approval Institutional Review Board approval was obtained.

Methodology
- prospective
- observational
- performed at one institution

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References

1. Gillies RJ, Kinahan PE, Hricak H (2016) Radiomics: images are more than pictures, they are data. Radiology 278:563–577
2. Aerts HJ, Grossmann P, Tan Y et al (2016) Defining a radiomic response phenotype: a pilot study using targeted therapy in NSCLC. Sci Rep 6:33860
3. Aerts HJ, Velazquez ER, Leijenaar RT et al (2014) Decoding tumour phenotype by noninvasive imaging using a quantitative radiomics approach. Nat Commun 5:4006
4. Coroller TP, Grossmann P, Hou Y et al (2015) CT-based radiomic signature predicts distant metastasis in lung adenocarcinoma. Radiother Oncol 114:345–350
5. Grossmann P, Stringfield O, El-Hachem N et al (2017) Defining the biological basis of radiomic phenotypes in lung cancer. Elife 6
6. Song SH, Park H, Lee G et al (2017) Imaging phenotyping using Radiomics to predict microcircular pattern within lung adenocarcinoma. J Thorac Oncol 12:624–632
7. Yip C, Tacelli N, Remy-Jardin M et al (2015) Imaging tumor response and tumoral heterogeneity in non-small cell lung cancer treated with antiangiogenic therapy: comparison of the prognostic ability of RECIST 1.1, an alternate method (Crabb), and image heterogeneity analysis. J Thorac Imaging 30:300–307
8. Huang Y, Liu Z, He L et al (2016) Radiomics signature: a potential biomarker for the prediction of disease-free survival in early-stage (I or II) non-small cell lung cancer. Radiology 281:947–957
9. Ha S, Park S, Bang JI, Kim EK, Lee HY (2017) Metabolic radiomics for pretreatment 18F-FDG PET/CT to characterize locally advanced breast cancer: histopathologic characteristics, response to neoadjuvant chemotherapy, and prognosis. Sci Rep 7:1556
10. Kim JH, Ko ES, Lim Y et al (2017) Breast cancer heterogeneity: MR imaging texture analysis and survival outcomes. Radiology 282:665–675
11. Li H, Zhu Y, Burnside ES et al (2016) MR imaging Radiomics signatures for predicting the risk of breast cancer recurrence as given by research versions of MammaPrint, Oncotype DX, and PAM50 gene assays. Radiology 281:382–391
12. Parikh J, Selmi M, Charles-Edwards G et al (2014) Changes in primary breast cancer heterogeneity may augment midtreatment MR imaging assessment of response to neoadjuvant chemotherapy. Radiology 272:100–112
13. Cameron A, Khalvati F, Haider MA, Wong A (2016) MAPS: a quantitative radiomics approach for prostate cancer detection. IEEE Trans Biomed Eng 63:1145–1156
14. Nektiah G, Elshoh M, Kim E et al (2017) T2-weighted MRI-derived textural features reflect prostate cancer aggressiveness: preliminary results. Eur Radiol 27:3050–3059
15. Huang YQ, Liang CH, He L et al (2016) Development and validation of a radiomics nomogram for preoperative prediction of lymph node metastasis in colorectal cancer. J Clin Oncol 34:2157–2164
16. Liang HY, Huang YQ, Yang ZX, Ying D, Zeng MS, Rao SX (2016) Potential of MR histogram analyses for prediction of response to chemotherapy in patients with colorectal hepatic metastases. Eur Radiol 26:2009–2018
17. Ng F, Ganeshan B, Kozarski R, Miles KA, Goh V (2013) Assessment of primary colorectal cancer heterogeneity by using whole-tumor texture analysis: contrast-enhanced CT texture as a biomarker of 5-year survival. Radiology 266:177–184
18. De Cecco CN, Ganeshan B, Ciolina M et al (2015) Texture analysis as imaging biomarker of tumoral response to neoadjuvant chemoradiotherapy in rectal cancer patients studied with 3-T magnetic resonance. Investig Radiol 50:239–245
19. Ahn SJ, Kim JH, Park SJ, Han JK (2016) Prediction of the therapeutic response after FOLFOX and FOLFIRI treatment for patients with liver metastasis from colorectal cancer using computerized CT texture analysis. Eur J Radiol 85:1867–1874
20. Julit O, Afaq A, Ganeshan B et al (2017) Magnetic resonance based texture parameters as potential imaging biomarkers for predicting long-term survival in locally advanced rectal cancer treated by chemoradiotherapy. Color Dis 19:349–362
21. Liu M, Lv H, Liu LH, Yang ZH, Jin EH, Wang ZC (2017) Locally advanced rectal cancer: predicting non-responders to neoadjuvant chemoradiotherapy using apparent diffusion coefficient textures. Int J Color Dis 32:1009–1012
22. Leipisc J, Labounty TM, Heilbron B et al (2010) Estimated radiation dose reduction using adaptive statistical iterative reconstruction in coronary CT angiography: the ERASIR study. AJR Am J Roentgenol 195:655–660
23. Marin D, Nelson RC, Schindera ST et al (2010) Low-tube-voltage, high-tube-current multidetector abdominal CT: improved image quality and decreased radiation dose with adaptive statistical iterative reconstruction algorithm–initial clinical experience. Radiology 254:145–153
24. Baker ME, Dong F, Primak A et al (2012) Contrast-to-noise ratio and low-contrast object resolution on full- and low-dose MDCT: SAFIRE versus filtered back projection in a low-contrast object phantom and in the liver. AJR Am J Roentgenol 199:8–18
25. Singh S, Kalra MK, Do S et al (2012) Comparison of hybrid and pure iterative reconstruction techniques with conventional filtered back projection: dose reduction potential in the abdomen. J Comput Assist Tomogr 36:347–353
26. Deak Z, Grimm JM, Treitl M et al (2013) Filtered back projection, adaptive statistical iterative reconstruction, and a model-based iterative reconstruction algorithm–initial clinical experience. Radiology 266:197–206
27. Hara AK, Paden RG, Silva AC, Kujak JL, Lawder HJ, Pavlicek W (2009) Iterative reconstruction technique for reducing body radiation dose at CT: feasibility study. AJR Am J Roentgenol 193:764–771
28. Silva AC, Lawder HJ, Hara A, Kujak J, Pavlicek W (2010) Innovations in CT dose reduction strategy: application of the adaptive statistical iterative reconstruction algorithm. AJR Am J Roentgenol 194:191–199
29. Ng F, Kozarski R, Ganeshan B, Goh V (2013) Assessment of tumor heterogeneity by CT texture analysis: can the largest cross-sectional area be used as an alternative to whole tumor analysis? Eur J Radiol 82:342–348

30. Zwanenburg A, Leger S, Vallières M, Löck S for the image biomarker standardisation initiative (2018) Image biomarker standardisation initiative. Available via https://arxiv.org/abs/1612.07003. Accessed 02 Jan 2018

31. Geyer LL, Schoepf UJ, Meinel FG et al (2015) State of the art: iterative CT reconstruction techniques. Radiology 276:339–357

32. Solomon J, Mileto A, Nelson RC, Roy Choudhury K, Samei E (2016) Quantitative features of liver lesions, lung nodules, and renal stones at multi-detector row CT examinations: dependency on radiation dose and reconstruction algorithm. Radiology 279:185–194

33. Zhao B, Tan Y, Tsai WY, Schwartz LH, Lu L (2014) Exploring variability in CT characterization of tumors: a preliminary phantom study. Transl Oncol 7:88–93

34. Zhao B, Tan Y, Tsai WY et al (2016) Reproducibility of radiomics for deciphering tumor phenotype with imaging. Sci Rep 6:23428

35. He L, Huang Y, Ma Z, Liang C, Liang C, Liu Z (2016) Effects of contrast-enhancement, reconstruction slice thickness and convolution kernel on the diagnostic performance of radiomics signature in solitary pulmonary nodule. Sci Rep 6:34921

36. Buch K, Li B, Qureshi MM, Kuno H, Anderson SW, Sakai O (2017) Quantitative assessment of variation in CT parameters on texture features: pilot study using a nonanatomic phantom. AJNR Am J Neuroradiol 38:981–985

37. Lo P, Young S, Kim HJ, Brown MS, McNitt-Gray MF (2016) Variability in CT lung-nodule quantification: effects of dose reduction and reconstruction methods on density and texture based features. Med Phys 43:4854

38. Lu L, Ehmke RC, Schwartz LH, Zhao B (2016) Assessing agreement between radiomic features computed for multiple CT imaging settings. PLoS One 11:e0166550

39. Shafiq-Ul-Hassan M, Zhang GG, Latifi K et al (2017) Intrinsic dependencies of CT radiomic features on voxel size and number of gray levels. Med Phys 44:1050–1062

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