

Radiomics feature reliability assessed by intraclass correlation coefficient: a systematic review

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Abstract: Radiomics research is rapidly growing in recent years, but more concerns on radiomics reliability are also raised. This review attempts to update and overview the current status of radiomics reliability research in the ever expanding medical literature from the perspective of a single reliability metric of intraclass correlation coefficient (ICC). To conduct this systematic review, Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines were followed. After literature search and selection, a total of 481 radiomics studies using CT, PET, or MRI, covering a wide range of subject and disease types, were included for review. In these highly heterogeneous studies, feature reliability to image segmentation was much more investigated than reliability to other factors, such as image acquisition, reconstruction, postprocessing, and feature quantification. The reported ICCs also suggested high radiomics feature reliability to image segmentation. Image acquisition was found to introduce much more feature variability than image segmentation, in particular for MRI, based on the reported ICC values. Image post-processing and feature quantification yielded different levels of radiomics reliability and might be used to mitigate image acquisitioninduced variability. Some common flaws and pitfalls in ICC use were identified, and suggestions on better ICC use were given. Due to the extremely high study heterogeneities and possible risks of bias, the degree of radiomics feature reliability that has been achieved could not yet be safely synthesized or derived in this review. More future researches on radiomics reliability are warranted.

Keywords: Radiomics; reliability; intraclass correlation coefficient (ICC); quantitative imaging; oncology

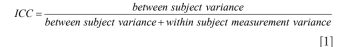
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Introduction

Radiomics has become one of the most popular research areas in medical imaging, in particular for clinical oncology, since its first introduction by Lambin et al. in 2012 (1). According to Gillies et al., radiomics is defined as "the conversion of images to higher dimensional data and the subsequent mining of these data for improved decision support" (2). These higher dimension data are normally understood as the information contained in a large number of quantitative radiomics features derived from the original or transformed medical images, which are usually artificially engineered with mathematical definition and have continuous values. By utilizing the radiomics models built on the selected radiomics features, radiomics promises to increase diagnosis accuracy and precision, assessment of prognosis, and therapy response prediction for different clinical applications, bridging between medical imaging and personalized medicine (3,4). A tremendous number of papers on radiomics have been published in recent years (5). However, despite the promising results reported, the broad validity, and generality of radiomics are still much hindered by the concerns on its reliability (6-11). Variability and uncertainty of radiomics can be introduced in many procedures of its complicated workflow. These procedures include but not limited to imaging hardware configuration, patient setup, image acquisition, reconstruction, image post-processing (filtering, segmentation and registration, etc.), radiomics feature quantification (such as feature definition, calculation setting like image discretization, software implementation, calculation result harmonization, etc.), and radiomics modeling.

The term reliability is commonly used with other terms like agreement, repeatability, stability, reproducibility, accuracy/precision, and robustness in varying degrees of consistency in the medical literature. In this study, a general and mathematically expressible definition of reliability (R) is adopted to be the extent to which measurements can be replicated. It is expressed as the ratio of true (error-free) variance (σT^2) over true variance plus error variance (σ_E^2), i.e., $R = \sigma_T^2/(\sigma_T^2 + \sigma_E^2)$. This definition of reliability is compliant with the classical definition of intraclass correlation coefficient (ICC), using the between-subject variance in the trait of interest to represent the true variance since it cannot be directly measured in reality.



ICC is one of the most widely adopted reliability indexes based on the analysis of variance (ANOVA) in medical literature (12). ICC is applicable for all radiomics features that have continuous values. In addition, ICC is a ratio index ranged from 0 to 1, so it is useful in the cross-study reliability comparison. For these reasons, ICC is chosen in this study as the single statistical metric for radiomics feature reliability assessment. We adopted the ICC forms in McGraw and Wong Convention, including three components of model (one-way or two-way, random-effects or mixed-effects), type (single or multiple measurements/ raters), and definition (absolute agreement or consistency), following the guideline proposed by Koo *et al.* (13).

Reliability of radiomics has to be carefully and rigorously measured and assessed prior to its real clinical deployment, but generic radiomics reliability is still not yet fully explored, so not well known. An excellent systematic review on repeatability and reproducibility of radiomics features was published in 2018 (14), in which the qualitative synthesis on 41 studies revealed the status of radiomics reliability research until April 2017. Since then, the status of radiomics reliability research has not been timely updated in the pace of an ever fast increasing number of radiomics publications. Few strongly evident consensus has been reached and well-acknowledged so far.

Thus, this review attempts to serve multi-fold purposes: (I) to have a timely updated overview on the current status of radiomics reliability research, mainly from the perspective of ICC use in the medical literature; (II) to survey what ICC was used for, and what were the major findings of radiomics reliability as revealed by the reported ICCs; (III) to critically review how ICC was used, reported and interpreted; (IV) to give some suggestions on ICC use to mitigate the flaws and pitfalls, if applicable, so as to improve radiomics reliability assessment for future studies.

Methods

Systematic search strategy

The major research question for the literature search was described as: "What are the known radiomics studies that used ICC as a radiomics feature reliability index, and reported the quantitative ICC results (as either major or secondary outcome)". Thus, a comprehensive literature search was conducted by two authors (JY and CX) to identify the relevant published studies in the database of MEDLINE/PubMed (National Center for Biotechnology Information, NCBI), from 1 January 2012 to 8 December 2020 (ePub date).

A combination of the following terms and their common variations: "CT/PET/MRI", "radiomics/radiomic/texture analysis/quantitative (heterogeneity) feature", "ICC/intraclass correlation" were comprehensively used for literature search. Imaging modalities other than CT, PET, and MRI, such as ultrasound, X-ray, cone-beam CT, and Megavoltage CT, were not included in this search due to their relative minority and immaturity in the radiomics research. Image analysis based solely on the gray level histogram, i.e., histogram analysis, does not provide any voxel positional/distributive information on the images, so it was not included.

Study selection

Only full-text journal or conference articles written in English were eligible and included. Conference abstracts, case reports, (systematic) reviews, editorials/commentaries, expert opinion papers, and non-English papers were excluded from selection.

After article type exclusion, all publications that involved the use of ICC for feature reliability assessment were identified through full text (and Supplementary materials if needed) examination in the searching results. If a study mentioned the ICC use in the method but reported no ICC results, it was also excluded.

Three authors (JY, CX, and OW) worked jointly on the study selection procedures as described above. Disagreements were resolved by consensus. Reasons for exclusion were documented.

Data extraction

Four authors (JY, CX, OW, and YZ) jointly performed record extraction. The study information on publication date, imaging modality, study design, study subject (phantom, animal or human), organ, disease, radiomics feature type/number was extracted. In terms of ICC use and reporting, the purpose of using ICC, the sample size for ICC calculation, ICC form, ICC reporting format, and major ICC results were extracted. Despite the high heterogeneity of ICC result reporting in different studies, we attempted to extract, synthesize, or harmonize the ICC results in the form of satisfactory feature rate (SFR), i.e., the percent of features showing satisfactory (determined by excellent, good, or other ICC criteria in each individual publication) ICC in the total investigated features, as much as possible. In this way, cross-study ICC comparison might become feasible to some extent. Radiomics quality score (RQS) of the extracted studies were not individually appraised since RQS might not be applicable for many of these studies because they were not completely clinical application studies (4). The quality of ICC use and reporting was not scored either. QUADAS-2 was not applied for study appraisal either since the diagnostic accuracy was not the common purpose of the included radiomics studies (15).

Outcomes and prioritizations

The primary outcome of interest in this review was radiomics feature reliability in different aspects as assessed by ICC. Quality of ICC use and reporting was the secondary outcome. Other statistic metrics used in combination with ICC were only noted but not further analyzed. The outcome was not prioritized on specific imaging modality or disease type.

Risk of bias analysis

Two authors (CX and JY) jointly assessed the possible risk of bias in the included studies from the extracted study information with consensus in the following perspectives. (I) study characteristics such as the study design (retrospective or prospective), cohort, sample size and feature number; (II) appropriateness of methodology, and sufficiency of method description and disclosure, such as the details of imaging acquisition, post-processing, segmentation, as well as feature definition (standardization) and quantification; (III) the quality of ICC use and reporting, such as the ICC form selection, confidence interval reporting, threshold values, and interpretation.

The TRIPOD (Transparent Reporting of a multivariable prediction model for Individual Prognosis Or Diagnosis) Statement (16) was not applied for two major reasons. First, it was not applicable for many studies because the investigation of radiomics feature reliability did not necessarily lead to the final prognosis or diagnosis performance report. Second, the clinical purposes were beyond the scope of this review, and they were also highly heterogeneous, including but not limited to prognosis or diagnosis.

Results

Literature search and selection

The PubMed search yielded 2,596 records. Records were

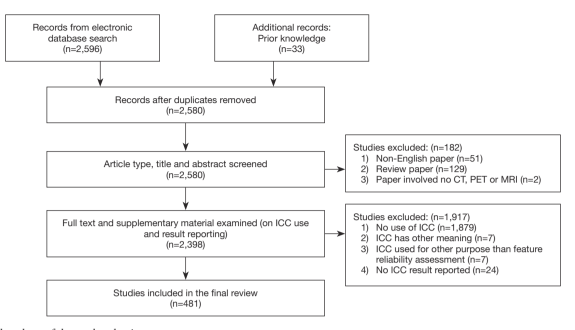


Figure 1 Flowchart of the study selection process.

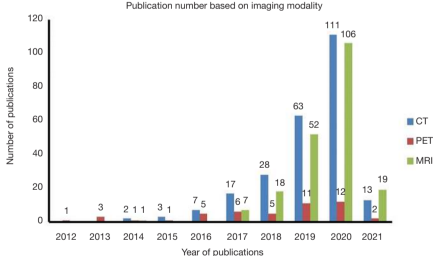


Figure 2 Publication number based on imaging modality in recent years.

reduced to 2,580 after duplicates were removed. The subsequent article type, title, and abstract screening excluded 182 records. Then the remaining 2,398 records underwent full text (and Supplementary materials) examination, and 1,917 records were further excluded. Finally, 481 studies were included in this systematic review. The selection process is illustrated in Figure 1.

Statistics of the included publications

Figure 2 shows the increasing number of published radiomics studies in which ICC was used for radiomics reliability assessment from 2012 to 2021. In recent years since 2017, MRI articles show a much faster increase than CT, and PET.

In terms of publication number, CT, PET and MRI

Publication distribution based on subject type and anatomical region

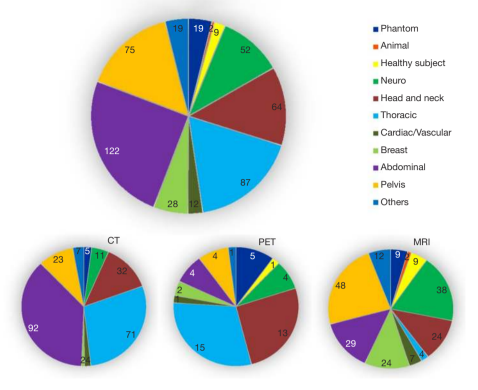


Figure 3 Study number based on subject type and anatomical region. The study number is counted repeatedly if multiple subject types or anatomical regions were involved in a study.

account for 50.73% (n=244), 9.77% (n=47), and 42.20% (n=203) of the total publications (N=481), respectively. Note that some publications involved multi-modality radiomics, so the sum was slightly over 100%.

In the included articles, 18.09% (87/481) and 81.91% (394/481) of them assessed reliability using ICC as the major study outcome and secondary (partial) outcome, respectively. 5.82% (n=28) and 94.18% (n=453) studies were prospective and retrospective (including those studies using prospectively acquired imaging data for clinical purposes other than radiomics) in nature.

The numbers of studies based on subject type and anatomical region are illustrated in *Figure 3*.

Clinical oncology is the most common clinical application in these studies, accounting for 86.69% (417/481) of the total publications. Lung cancer is the most common type, followed by head and neck cancer and neuro-oncology. Imaging modality reflects variation in predominant type of cancer, as PET and CT studies showed a higher proportion of lung and head and neck patients, while MRI studies comprised more neuro-oncology, breast

cancer, cervical cancer, and prostate cancer. The publication distribution based on cancer types, and imaging modalities is illustrated in *Figure 4*.

The characteristics of human radiomics studies

The use of ICC in the included human radiomics studies was categorized by the ICC purpose. The characteristics of the studies were summarized in the following tables. If a study involved more than one ICC purpose, each purpose was separately listed in the corresponding table. SFR was directly extracted or synthesized from the original data from each study as much as possible. However, in many cases, SFR was not available or could not be clearly and reliably extracted, which was thus labeled NA in the tables. One common reason was that ICC was reported in other forms like mean \pm SD. Another reason was that ICC was applied for some but not all features, e.g., for those features after correlation assessment or clustering. Meanwhile, for some other studies, very comprehensive ICC results were reported. The SFRs could not be simply extracted and

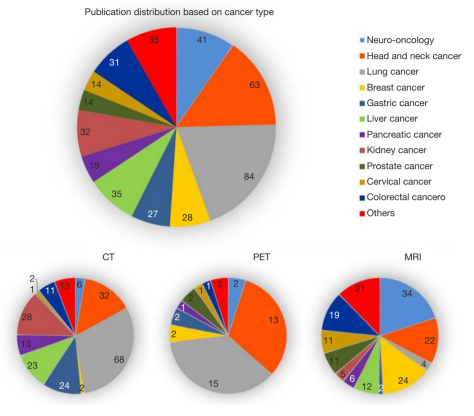


Figure 4 Distribution of oncological patient studies. The study number is counted repeatedly for each type of cancer if a study investigated more than one cancer type.

listed. They were labeled as NE.

Radiomics feature reliability due to image acquisition

The use of ICC in the included human radiomics studies to assess the feature reliability due to image acquisition was summarized in *Table 1*.

Radiomics feature reliability due to image reconstruction

The use of ICC in the included human radiomics studies to assess the feature reliability due to image reconstruction was summarized in *Table 2*.

Radiomics feature reliability due to image segmentation

There were 416 out of 481 studies reported the ICC results regarding the feature reliability influenced by image segmentation. Only those studies that substantially investigated the radiomics feature reliability to image segmentation as the primary study endpoint were listed in *Table 3*. For other studies, most simply mentioned the

ICC use for intra-/inter-observer agreement, and reported a very short ICC (usually optimistic) result, in which SFR was extracted from 206 studies. The SFR distribution was provided by the histogram in Figure S1, showing no apparent differences between imaging modalities (Figure S2).

Radiomics feature reliability due to image processing

The use of ICC in the included human radiomics studies to assess the feature reliability due to image processing was summarized in *Table 4*.

Radiomics feature reliability due to feature quantification

The use of ICC in the included human radiomics studies to assess the feature reliability due to feature quantification was summarized in *Table 5*.

The characteristics of phantom and animal radiomics studies

The study characteristics in the included phantom and

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First author, year	Disease	Patient number/feature number/ICC sample size	Acquisition factors	Satisfactory feature rate	ICC threshold
СТ					
Aerts et al., 2016 (17)	Lung cancer	47/183/31	Test-retest	NA	NA
Hu <i>et al</i> ., 2016 (18)	Rectal cancer	40/775/40	Test-retest	64.00%	0.8
Hyunh <i>et al</i> ., 2016 (19)	Lung cancer	113/1605/31	Test-retest	53.27%	0.8
Hyunh <i>et al</i> ., 2017 (20)	Lung cancer	112/644/31	Test-retest	44.41%	0.8
Hosny <i>et al</i> ., 2018 (21)	Lung cancer	1194/NA/32	Test-retest	NA	0.8
Soufi <i>et al</i> ., 2018 (22)	Lung cancer	162+143/432/31	Test-retest	NE	0.75
Dou <i>et al.</i> , 2018 (23)	Lung cancer	200/2175/31	Test-retest	33.15–51.17%	0.85
Huang et al., 2019 (24)	Lung cancer	54/203/31	Test-retest	65.02%	0.9
Khorrami <i>et al</i> ., 2019 (25)	Lung cancer	125/1542/31	Test-retest	76.00%	0.8
Khorrami <i>et al</i> ., 2019 (26)	Lung cancer	90/1542/31	Test-retest	67%	0.85
Osman <i>et al.</i> , 2019 (27)	Prostate cancer	342/1618/20	Test-retest	32.26%	0.8
Zwanenburg e <i>t al</i> ., 2019 (28)	Lung cancer, HNC	31+19/4032/31+19	Test-retest	57.3% (lung cancer), 14.4% (HNC)	0.9
Kadoya <i>et al</i> ., 2020 (29)	Lung cancer	287/107/31	Test-retest	79.44%	0.8
Khorrami <i>et al.</i> , 2020 (30)	Lung cancer	350/133/31	Test-retest	NA	0.75
Ligero et al., 2021 (31)	Colorectal/kidney cancer	43 (97 liver mets)/93/43 (97 liver mets)	Acquisition voltage	65.59-81.72%	0.8
Prayer et al., 2021 (32)	Lung disease	60/86/30	Test-retest, inter-scanner	75.6% (test-retest), NE (inter-scanner)	0.9
Vuong <i>et al.</i> , 2020 (33)	Lung cancer	124/1404/10+11	Motion, contrast	41.30% (motion), 29% (contrast)	0.9
Yamashita et al., 2020 (34)	Pancreatic cancer	37/266/37	Inter-scanner, contrast agent	NE	0.8
MRI					
Fiset <i>et al.</i> , 2019 (35)	Cervical cancer	62/1761/8+20	Test-retest, inter-scanner	22.6% (test-retest), 6.2% (inter-scanner)	0.9
Li <i>et al</i> ., 2019 (36)	Hippocampus	60/55/60	Test-retest	NE	0.4
Zinn <i>et al</i> ., 2018 (37)	GBM	79/300/79	Inter-scanner	NA	NA
Bologna <i>et al</i> ., 2020 (38)	NPC	142/1072/142	Imaging parameters, geometrical transformation	24.72%	0.75
Jang <i>et al.</i> , 2020 (39)	Healthy, cardiac referral	10+51/1023/61	Test-retest	2.6-28.8% (healthy), 8.9–34.8% (patient)	0.8
			Test-retest (repositioning)	0.7-3.1%	
Merisaari <i>et al</i> ., 2020 (40)	Prostate cancer	112/1694/34	Test-retest	NA	0.75
Pandey et al., 2021 (41)	Healthy	87+4+8/93/4+8	Test-retest	~71.04%	0.5
			Inter-scanner	18% (GM), 21.5% (WM)	

Table 1 Summary of human radiomics studies using ICC for image acquisition

Table 1 (continued)

Xue et al. Systematic review of radiomics feature reliability assessed by ICC

First author, year	Disease	Patient number/feature number/ICC sample size	Acquisition factors	Satisfactory feature rate	ICC threshold
Gutmann <i>et al.</i> , 2020 (42)	Diabetes	310/684/310	Test-retest	26.5%	0.85
Scalco et al., 2020 (43)	Prostate cancer	14/91/14	Test-retest repositioning	7–11%	0.9
Shiri <i>et al</i> ., 2020 (44)	GBM	17/26295192/17	Test-retest	NE	0.95
Ta et al., 2020 (45)	Healthy	6/22/6	Test-retest, inter-scanner	NE	0.75
Zhang et al., 2020 (46)	Brain cancer	1728/1595/50	Inter-scanner	100%	0.75
Han <i>et al</i> ., 2021 (47)	GBM	57/45/57	Test-retest	NA	0.75
PET					
Leijenaar et al., 2013 (48)	Lung cancer	34/106/11	Test-retest	71%	0.8
Willaime <i>et al</i> ., 2013 (49)	Breast cancer	15/28/9	Test-retest, tissue	~46.43 (test-retest), ~64.29% (across tissue)	0.7
van Velden <i>et al.</i> , 2014 (50)	Colorectal carcino- ma	- 29/18 /29	Test-retest	83.33%	0.7
Cheng et al., 2016 (51)	Lung cancer	56/12/56	Different tracer	NA	0.95
van Rossum <i>et al</i> ., 2016 (52)	HNC	217/80/7	Test-retest	NE	0.9
Carvalho <i>et al</i> ., 2018 (53)	Lung cancer	215/118/215	Test-retest	65.25%	0.85
Jiang <i>et al</i> ., 2018 (54)	Gastric cancer	214/80/30	Inter-scanner	NA	0.8
Lin <i>et al</i> ., 2019 (55)	Prostate cancer	18/47/18	Test-retest	83%	0.75
Manabe et al., 2019 (56)	Cardiac disease	89/36/33	Inter-scanner	47.22%	0.8
Vuong et al., 2019 (57)	Lung cancer	10+9/1,355/10	Test-retest (inhale, exhale phase)	55.6% (shape), 39.3 (wavelet)	0.9
			Inter-scanner (PET/CT <i>vs</i> . PET/MRI)	61% (shape and intensity), 28% (wavelet)	0.9
PET, CT					
Desseroit <i>et al.</i> , 2017 (58)	Lung cancer	73/40/73	Test-retest	NE	NA
PET, MRI					
Jiang et al., 2017 (59)	Healthy	86/NA/66	Test-retest	NA	0.75

Table 1 (continued)

NA, not available or not clear; NE, could not be simply extracted due to comprehensive results, see the reference for detail; HNC, head and neck cancer; liver mets, liver metastases; GBM, glioblastoma multiforme. NPC, nasopharyngeal carcinoma. GM, gray matter. WM, white matter.

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First author, year	Disease	Patient number/feature number/ICC sample size	Reconstruction factors	Satisfactory feature rate	ICC threshold
СТ					
Ahn <i>et al</i> ., 2019 (60)	Liver lesion and renal cyst	1,462/11/1,462	Inter-method	NE	0.75
Kolossváry <i>et al.</i> , 2019 (61)	Vascular disease	60/171/60	Inter-method	~97.04%	0.9
Koo <i>et al.</i> , 2017 (62)	Lung cancer	194/10/194	Reconstruction intervals	~70%	0.8
Lee et al., 2019 (63)	Lung nodule	194 patients (260 scans) /252/114 patients (180 scans)	Voxel size	9.13%	0.7
Ligero <i>et al.</i> , 2021 (31)	Colorectal/kidney	43 (97 liver mets)/93/43 (97	Voxel size	48.40-78.49%	0.8
	disease	liver mets)	Slice spacing	43.01-86.02%	0.8
			Slice thickness	75.25-88.17%	0.8
			Convolution kernel	55.92-97.85%	0.8
Prayer <i>et al.</i> , 2021 (32)	Lung disease	60/86/60	Reconstruction kernel and slice thickness,	NE	NE
Vuong et al., 2020 (33)	Lung cancer	124/1,404/23	Convolution kernel	17.20%	0.9
Yamashita et al., 2020 (34)	Pancreatic cancer	37/266/37	Voxel size	NE	0.8
MRI					
Suter et al., 2020 (64)	GBM	63+76/8,327/19	K-space subsampling	NE	0.85
PET					
Altazi <i>et al.</i> , 2017 (65)	Cervical cancer	88/79/8	Inter-method	NE	NA
van Velden <i>et al.</i> , 2016 (66)	Lung cancer	11/105/11	Inter-method	63% (segmentation or reconstruction)	0.9

Table 2 Summary of human radiomics studies using ICC for image reconstruction

NA, not available or not clear; NE, could not be simply extracted due to comprehensive results, see the reference for detail; liver mets, liver metastases; GBM, glioblastoma multiforme.

Table 3 Summary of human radiomics studies using ICC for image segmentation

First author, year	Disease	Patient number/ feature number/ICC sample size	Segmentation factors	Satisfactory feature rate	ICC threshold
СТ					
Parmar <i>et al</i> ., 2014 (67)	Lung cancer	20/56/20	Intra-/inter-observer	51.8% (manual),	0.8
			Software	73.2% (software)	
Echegaray <i>et al</i> ., 2015 (68)	Liver cancer	29/745/29	Inter-observer	78.39%	0.8
Echegaray <i>et al</i> ., 2016 (69)	Lung cancer	100/94/100	Semi-automatic	89%	0.7
Qiu <i>et al</i> ., 2017 (70)	Liver cancer	15/71/15	Intra-/inter-observer, semi-automatic	81% (GrowCut), 77% (GraphCut), 73% (manual)	0.75
Owens <i>et al.</i> , 2018 (71)	Lung cancer	10/83/10	Intra-/inter-observer, software	NE	0.75

Table 3 (continued)

Xue et al. Systematic review of radiomics feature reliability assessed by ICC

First author, year	Disease	Patient number/ feature number/ICC sample size	Segmentation factors	Satisfactory feature rate	ICC threshold
Pavic et al., 2018 (72)	Lung cancer, HNC	11+11+11/1, 404/11+11+11	Inter-observer	90% (NSCLC); 59% (HNC); 36% (MPM)	0.8
Kocak <i>et al.</i> , 2019 (73)	Kidney cancer	47/828/25	Inter-method	86.2% (contour-focused); 93.2% (margin shrinkage)	0.9
Kocak <i>et al</i> ., 2019 (74)	Kidney cancer	30/744/30	Intra- observer	84.4–92.2% (CT); 85.5–93.1% (CECT)	0.75
			Inter-observer	76.7% (CT); 84.9% (CECT)	
Moltz et al., 2019 (75)	Liver cancer	13/110/13	Inter-observer	13.64%	0.9
Mori <i>et al</i> ., 2019 (76)	Pancreatic cancer	31/69/31	Inter-observer	94.20%	0.8
Qiu <i>et al</i> ., 2019 (77)	Liver cancer	106/71/26	Semi-automatic	79% (GrowCut); 73% (GraphCut); 69% (manual)	0.75
Sung et al., 2019 (78)	Liver disease	58/5/58	Intra- inter- observer	NE	0.75
Uthoff et al., 2019 (79)	Lung cancer	71/101/71	Intra- inter- observer	NA	NA
Caballo <i>et al.</i> , 2020 (80)	Breast cancer	69/672/35	Automatic	90%	0.75
Haarburger <i>et al</i> ., 2020 (81)	Lung, kidney, liver tumors	1,536/92/1,536	Automatic	84% (PHiSeg); 88% (expert)	0.8
Kakino <i>et al.</i> , 2020 (82)	Lung cancer	256/93/31	Inter-observer	64.52%	0.5
Kulkarni <i>et al</i> ., 2021 (83)	Pancreatic cancer	128/14/128	Inter-method	NA	0.9
Liu <i>et al.</i> , 2020 (84)	HNC	436/109/436	Inter-observer, inter-method	NE	0.9
Nguyen <i>et al</i> ., 2020 (85)	Kidney cancer	165/25/165	Inter-observer	NE	0.8
Ren <i>et al</i> ., 2020 (86)	HNC	47/1032/47	Intra-observer	21.6% (2D) 30.4% (3D)	0.9
			Inter-observer	14.1% (2D) 31.4% (3D)	
MRI					
Adduru <i>et al</i> ., 2017 (87)	Healthy	38/NA/38	Software	NE	0.56
Lee et al., 2017 (88)	GBM	45/180/45	Inter-observer, software	~91.1% (tumor Prism 3D), 90% (3D slicer)	0.8
Bologna <i>et al</i> ., 2018 (89)	Sarcoma, HNC	18+18/69/18+18	Inter-method	~78.26% (HNC), 85.5% (sarcoma)	0.78
Saha et al., 2018 (90)	Breast cancer	50/529/50	Inter-observer	NE	0.9
Duron <i>et al</i> ., 2019 (91)	Lachrymal gland tumors, breast cancer	74+30/69/74+30	Intra-/inter- observer	NE	0.8
Fiset <i>et al</i> ., 2019 (35)	Cervical cancer	62/1,761/62	Inter-observer	95.2%	0.9
Koçak, 2019 (92)	GBM	25/1,116/25	Inter-method	~25%	0.9
Lecler et al., 2019 (93)	Breast cancer	37/510/37	Intra-/inter- observer	53%	0.8

Table 3 (continued)

Table 3 (continued)

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First author, year	Disease	Patient number/ feature number/ICC sample size	Segmentation factors	Satisfactory feature rate	ICC threshold
Tixier et al., 2019 (94)	GBM	90/108/90	Semi-automatic	NE	0.9
Traverso <i>et al</i> ., 2019 (95)	Cervical cancer	81/552/81	Inter-observer	NE	0.9
Alis <i>et al.</i> , 2021 (96)	Healthy cardiac	59/352/59	Inter-observer	92%	0.8
Chen <i>et al.</i> , 2021 (97)	Cervical cancer	20/105/20	Intra- inter- observer	NE	0.8
Granzier <i>et al.</i> , 2020 (98)	Breast cancer	129 (tumors)/1,328 (Radiomix), 833 (pyradiomics)/129	Inter-observer, software	41.6% (RadiomiX); 32.8% (Pyradiomics); 41.1% (unfiltered RadiomiX); 16.2% (unfiltered Pyradiomics)	0.9
Jang <i>et al.</i> , 2020 (39)	Healthy, cardiac	10+51/1,023/15	Intra-observer	61–73%	0.8
	referral		Inter-observer	32-47%	
Lin <i>et al.</i> , 2020 (99)	Cervical cancer	169/51/169	Automatic	NA	NA
Gutmann <i>et al.</i> , 2020 (42)	Diabetes	310/684/310	Inter-observer	82.9%	0.85
Pati <i>et al.</i> , 2020 (100)	GBM	31/11,700/31	Inter-observer	NE	0.8
Scalco <i>et al.</i> , 2020 (43)	Prostate cancer	14/91/14	Inter-observer	NE	0.9
Suter <i>et al.</i> , 2020 (64)	GBM	63+76/8,327/19	Inter-observer	NE	0.85
PET					
Leijenaar <i>et al.</i> , 2013 (48)	Lung cancer	34/106/23	Inter-observer	91%	0.8
Lu <i>et al</i> ., 2016 (101)	NPC	40/88/40	Inter-observer, inter-method	50% (¹⁸ F-FDG), 62% (11C-choline)	0.8
van Velden <i>et al</i> ., 2016 (66)	Lung cancer	11/105/11	Inter-method	63% (segmentation or reconstruction)	0.9
Bashir e <i>t al</i> ., 2017 (102)	Lung cancer	53/83/53	Inter-observer, inter-method	NE	0.85
Belli <i>et al</i> ., 2018 (103)	HNC, pancreatic cancer	25+25/72/25+25	(Semi)-automatic	19% (HN-N); 19% (HN-T); 47% (pancreas)	0.8
Manabe et al., 2019 (56)	Cardiac disease	89/36/33	Inter-observer	77.78%	0.8
PET, MRI					
Yang et al., 2020 (104)	NPC	21/540/21	Inter-method	85.74% (PET), 84.81% (T2), 89.07% (DWI)	0.95

Table 3 (continued)

NA, not available or not clear; NE, could not be simply extracted due to comprehensive results, see the reference for detail; HNC, head and neck cancer; NSCLC, non-small-cell lung carcinoma; MPM, malignant pleural mesothelioma; CECT, contrast-enhanced CT; GBM, glioblastoma multiforme; HN-N, head and neck cancer-positive lymph node; HN-T, head and neck cancer-tumor; NPC, nasopharyngeal carcinoma.

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Table 4 Summary of human radiomics studies using ICC for image processing

First author, year	Disease	Patient number/feature number/ICC sample size	Image processing factors	Satisfactory feature rate	ICC threshold
СТ					
Bogowicz et al., 2016 (105)) Lung cancer, HNC	; 11+11/315/11+11	HU threshold	~35%	0.9
			Voxel size resampling	~5%	
			Temporal resolution	~30%	
			Artery contouring	~45%	
			Noise threshold	~52.5%	
Ger et al., 2018 (106)	Lung cancer HNC	20+30/49/20+30	Voxel size resampling	71–79.6%	0.9
Shafiq-UI-Hassan <i>et al.,</i> 2018 (107)	Lung cancer	18/24/18	Gray level normalization	100.00%	0.8
Lee et al., 2019 (63)	Lung nodule	194 patients (260 scans)/252/114 patients (180 scans)	Voxel size resampling	15%	0.7
Zwanenburg et al., 2019	Lung cancer HNC	31+19/4,032/31+19	Noise	95.0–97.4%	0.9
(28)			Rotation	~75–80%	
			Translation, volume adaptation, contrast	16.6–32.9%	
			Rotation, volume adaptation, contrast	16.8–33.3%	
			Noise, translation, volume adaptation, contrast	16.7–33.7%	
			Rotation, noise, volume adaptation, contrast	16.7–34.2%	
			Other	NE	
Defeudis <i>et al.</i> , 2020 (108)	Colorectal cancer	14/35/14	Standardization	~36–60%	0.9
Park <i>et al.</i> , 2020 (109)	Bladder cancer	83/55/83	SNR, and outlier inclusion	NE	0.75
MRI					
Kim <i>et al.</i> , 2019 (110)	GBM	167/356/167	Pertubation	77.20%	0.75
Li <i>et al.</i> , 2019 (36)	Hippocampus	60/55/60	Normalization	~36.36– 56.36%	0.4
Schwier <i>et al.</i> , 2019 (111)	Prostate cancer	15/1,120/15	Normalization	NE	0.85
Traverso <i>et al.</i> , 2020 (95)	Cervical cancer	81/552/81	Normalization	NE	0.9
Fan <i>et al.</i> , 2020 (112)	Breast cancer	322/107/322	Voxel size resampling	NE	0.7
Moradmand et al., 2020	GBM	65/107/65	Noise	21.4%	0.9
(113)			Noise + bias field	20.4%	
			Bias field	23.2%	
			Bias field + noise	22.5%	
Scalco <i>et al.</i> , 2020 (43)	Prostate cancer	14/91/14	Normalization	~12–14%	0.9
Shiri e <i>t al.,</i> 2020 (44)	GBM	17/26,295,192/17	Transformation, bias field removal	NE	0.95

Table 4 (continued)

First author, year	Disease	Patient number/feature number/ICC sample size	Image processing factors	Satisfactory feature rate	ICC threshold
Suter <i>et al.</i> , 2020 (64)	GBM	63+76/8,327/19	Perturbation	42.5%	0.85
PET					
Branchini <i>et al.</i> , 2019 (114)	Pediatric	21/106/21	Activity reduction simulation	NE	0.9
Whybra et al., 2019 (115)	HNC	441/141/441	Voxel size resampling	66%	0.9
PET, MRI					
Yang <i>et al.</i> , 2020 (104)	NPC	21/540/21	Pixel size resampling	55.74% (T2), 60.37% (DWI), 58.33% (PET)	0.95
			Slice thickness	24.07% (T2 and DWI), 23.89% (PET)	

Table 4 (continued)

NA, not available or not clear; NE, could not be simply extracted due to comprehensive results, see the reference for detail; HNC, head and neck cancer; HU, Hounsfield unit; SNR, signal-to-noise ratio; GBM, glioblastoma multiforme; NPC, nasopharyngeal carcinoma; DWI, diffusion-weighted imaging.

animal radiomics studies were summarized in Table 6.

Quality of ICC use and reporting

Generally speaking, the quality of ICC use and reporting was found unsatisfactory in many publications, associating with various flaws and pitfalls. Only 63 studies (13.10%) explicitly and precisely reported the selected ICC form, in which the ICC definition of absolute agreement rather than consistency on feature values predominated. The rationale of ICC form selection was seldom explained. In the remaining 418 articles, the ICC form was either unavailable, implicit (e.g., giving the general ICC formula not specific to a certain ICC form), or incomplete. The available ICC forms described in the studies, either completely or not, were summarized in the Table S1. Very few studies tested the normal distribution of data prior to ICC use (as ICC was based on ANOVA). The adopted reliability criteria/level and the corresponding threshold ICC values could be found in most studies but were heterogeneous. The reliability levels could be binary (low/ high, acceptable/unacceptable, stable/unstable, repeatable/ unrepeatable, etc.), three (e.g., poor/moderate/good), four (e.g., poor/moderate/good/excellent), five (e.g., poor/fair/ moderate/good/excellent) and even six. The thresholds of >0.7, >0.75, >0.8, and >0.9 were frequently used to determine the highest reliability level. The thresholds of <0.2, <0.4, and <0.5 were frequently adopted to determine the lowest reliability level. The reported ICC values were normally

presented in the mean (± SD), median, range, or interquartile range (IQR). Confidence interval (usually 95% CI) was reported along with ICC only in 64 studies. Many studies seemed to interpret reliability levels based on the estimated ICC values without giving or referring to the reported ICC confidence interval.

Notable findings of radiomics feature reliability as revealed by ICC

The reported ICC results were highly heterogeneous, varying by imaging modality, ICC purpose, disease, lesion type, sample size, as well as feature types. Meanwhile, they were also frequently reported with pitfalls. Therefore, it was impractical to conduct quantitative data synthesis and metaanalysis based on the reported ICCs to reliably estimate the achievable absolute radiomics reliability levels for different modalities, purposes, or diseases. But there were still a few notable consistent findings observed on the report ICCs even in the presence of high study heterogeneities.

High satisfactory feature rates were reported for most intra/inter-observer segmentation studies, indicating the high robustness of many radiomics features to intra/inter-observer segmentation variability

In the hundreds of articles using ICC for assessing intra/ inter-observer segmentation reliability of radiomics features, only a small number reported relatively negative

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Table 5 Summary of human radiomics studies using ICC	for feature quantification
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First author, year	Disease	Patient number/feature number/ICC sample size	Quantification factor	Satisfactory feature rate	ICC threshold
СТ					
Bogowicz <i>et al.</i> , 2016 (105)	Lung cancer, HNC	11+11/315/11+11	Discretization	~40%	0.9
Foy <i>et al</i> ., 2018 (116)	HNC	39/12/39	Radiomics calculation	33%	0.9
Soufi <i>et al.</i> , 2018 (22)	Lung cancer	162+143/432/305	Discretization	NE	0.75
Lee <i>et al</i> ., 2019 (63)	Lung nodule	194 patients (260 scans) /252/114 patients (180 scans)	Discretization	~9.5–11.5%	0.7
Zwanenburg <i>et al</i> ., 2019 (28)	Lung cancer, HNC	31+19/4,032/31+19	Discretization	~10–65% (FBW); 10–40%(FBN)	0.9
Park et al., 2020 (109)	Bladder cancer	83/55/83	Discretization	NE	0.75
MRI					
Duron <i>et al</i> ., 2019 (91)	Lachrymal gland tumors, breast cancer	74+30/69/74+30	Discretization	NE	0.8
Schwier et al., 2019 (111)	Prostate cancer	15/1,120/15	Discretization	NE	0.85
Traverso <i>et al.</i> , 2020 (95)	Cervical cancer	81/552/81	Discretization	NE	0.9
Pandey <i>et al.</i> , 2021 (41)	Healthy	87+4+8/93/4+8	Harmonization	60.33% (GM); 62% (WM)	0.5
Shiri <i>et al</i> ., 2020 (44)	GBM	17/26,295,192/17	Discretization	NE	0.95
Suter <i>et al.</i> , 2020 (64)	GBM	63+76/8,327/19	Discretization	NE	0.85
PET					
Tixier et al., 2012 (117)	HNC	16/25/16	Discretization	NE	NA
Leijenaar <i>et al.</i> , 2015 (118)	Lung cancer	35/44/35	Discretization	NE	NA
Lu et al., 2016 (101)	NPC	40/88/40	Discretization	23% (18F-FDG), 21% (11C-choline)	0.8
Altazi et al., 2017 (65)	Cervical cancer	88/79/80	Discretization	18% (GLCM and GLRLM)	0.9
Bogowicz et al., 2017 (119)	HNC	128+50/649/178	Feature implementation	12%	0.8
Lv <i>et al.</i> , 2018 (120)	NPC	106/57/106	Averaging, symmetry, distance	NE	0.8
Branchini <i>et al</i> ., 2019 (114)	Pediatric	21/106/21	Discretization	NE	0.9
PET, MRI					
Yang <i>et al.</i> , 2020 (104)	NPC	21/540/21	Discretization	12.96% (PET and T2), 11.30% (DWI)	0.95

NA, not available or not clear; NE, could not be simply extracted due to comprehensive results, see the reference for detail; HNC, head and neck cancer; FBW, fixed bin width; FBN, fixed bin number; GM, gray matter; WM, white matter; NPC, nasopharyngeal carcinoma; GBM, glioblastoma multiforme; GLCM, gray level co-occurrence matrix; GLRLM, gray level run length matrix; DWI, diffusion-weighted imaging.

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First author, year	Phantom or animal	ICC purpose	Factors	Satisfactory feature rate	ICC threshold
СТ					
Panth e <i>t al</i> ., 2015 (121)	Animal (mice)	Acquisition, segmentation	Test-retest, inter-observer	NE	NA
-	Anthropomorphic pelvic	Acquisition	Test-retest	93.80%	0.9
2018 (122)	phantom, multi-material phantom		Imaging parameters	69.89%	0.9
Ger <i>et al.</i> , 2018 (106)	Credence Cartridge Radiomics phantom	Processing	Resampling	71.43%	0.9
Mannil <i>et al</i> ., 2018 (123)	Human urinary stones	Segmentation	Intra-/inter-observer	29.87%	0.8
Li <i>et al</i> ., 2020 (124)	Anthropomorphic	Acquisition,	Effective dose and pitch	86.90%	0.8
	thoracic phantom	reconstruction, segmentation,	Slice thickness and filter	89.75%	
		quantification	Intra-observer	89.80%	
			Inter-observer	92.00%	
			Gray-level range (HU)	86.40%	
			Bin size (HU)	58.00%	
Nardone <i>et al</i> ., 2020	Commercial phantom	Acquisition	Inter-scanner	0%	0.75
(125)	(CIRS model 467)		Inter-scanner (delta)	45.90%	
MRI					
Song <i>et al</i> ., 2014 (126)) Animal (rats)	Acquisition, segmentation	Test-retest, Intra-observer	NE	0.75
Baeßler et al., 2019	Fruits	Segmentation	Intra-observer	100.00%	0.75
(127)			Inter-observer	62.22%	
-	Phantom (virtual) (brain-	Acquisition,	Varying TE and TR	76.00%	0.75
(128)	web)	processing	Normalization	81.00%	0.75
Bologna <i>et al</i> ., 2019 (129)	Virtual phantom (brainweb)	Acquisition, processing	Varying TE and TR, denoising, voxel size resampling, bias field correction (sequence dependent T1 & T2)	59.80%	0.75
Cattell <i>et al</i> ., 2019	Fruits	Segmentation,	Erosion and dilation (normalized mean ± 3 SD)	75.89%	0.9
(130)		processing	Erosion and dilation (normalized 0-max)	90.18%	
			Denoising (normalized mean \pm 3SD)	34.78%	
			Denoising (normalized 0-max)	47.82%	
			Voxel size resampling (normalized mean \pm 3SD)	64.28%	
			Voxel size resampling (normalized 0-max)	83.92%	
Bianchini <i>et al</i> ., 2020	Customised female	Acquisition	Test-retest (T1)	74.76%	0.9
(131)	pelvis phantom		Test-retest (T2)	95.26%	

Table 6 Summary of phantom and animal studies

Table 6 (continued)

First author, year	Phantom or animal	ICC purpose	Factors	Satisfactory feature rate	ICC threshold
Dreher C <i>et al</i> ., 2020	DWI phantom and fruits	Acquisition,	Test-retest	84.78%	0.9
(132)		segmentation	Intra-observer	97.80%	
			Inter-observer	95.65%	
Eresen <i>et al</i> ., 2020 (133)	Animal (mice)	Segmentation		~58.33% (T2), ~61.74% (T1)	0.75
Jang et al., 2020 (39)	Fruits and vegetables	Acquisition	Test-retest	~43.44%	0.8
			Test-retest re-positioning	~9.82%	
Rai <i>et al</i> ., 2020 (134)	Customized phantom	Acquisition	Inter-scanner	53.62%	0.8
	printed by a Connex 260 polyjet printer		Phantom dependence	55.07%	
Bianchini <i>et al</i> ., 2021	Pelvic-shaped	Acquisition	Test-retest	82.90%	0.9
(135)	container (NEMA IEC Body Phantom Set:		Test-retest re-positioning	58.33%	
Spectrum Corporation)			Inter-scanner	3.30%	
PET					
Gallivanone <i>et al.</i> , 2018 (136)	Anthropomorphic Alderson Thorax phantom	Acquisition	Test-retest	53.45%	0.6
Ger <i>et al</i> ., 2019 (137)	3-dimensional	Processing,	Pixel size resampling	~97.00%	0.9
	Hoffman brain phantom	reconstruction	Filter cutoff	~82.30%	
	I		Effective iteration	~86.16%	
			Time per bed position alteration	~92.00%	
			Q.clear	92.00%	
Pfaehler et al., 2019	NEMA NU 2-2012 IQ	Acquisition, reconstruction, quantification	Scan duration (noise)	NE	0.8
(138)	phantom		High uptake	32.00%	
		1	Low uptake	21.00%	
			Point-spread-function (PSF)	53.00%	
			Osem	30.00%	
			Time-of-flight	32.00%	
			Discretization (FBW)	89.00%	
			Discretization (FBN)	35.00%	
Pfaehler et al., 2020	3D printed phantom	Acquisition,	Test-retest (statistically equal replicates) (FBN)	90.70%	0.9
(139)	inserts in NEMA NU 2-2012 IQ phantom	quantification	Test-retest (statistically equal replicates) (FBW)	97.30%	
			Test-retest (FBN)	79.13%	0.6
			Test-retest (FBW)	89.96%	
			Multicenter (FBN)	45.60%	
			Multicenter (FBW)	62.97%	

Table 6 (continued)

Table 6 (continued)

Table 6 (continued)					
First author, year	Phantom or animal	ICC purpose	Factors	Satisfactory feature rate	ICC threshold
Yang et al., 2020 (140)	Digital phantom	Segmentation,	Inter-observer (bin number 32 or 64)	84.00%	0.9
	(Zubal anthropomorphic phantom)	quantification	Inter-observer (bin number 128 or 256)	88.00%	

NA, not available or not clear; NE, could not be simply extracted due to comprehensive results, see the reference for detail; HU, Hounsfield unit; TE, echo time; TR, repetition time; DWI, diffusion-weighted imaging; OSEM, ordered subset expectation maximization; FBW, fixed bin width; FBN, fixed bin number.

reliability results. For instance, Jang et al. (39) showed that in the inter-observer segmentation reproducibility study in cardiac patients, only 32.1%, 46.7%, and 35.5% of MRI radiomics features were reproducible with the cine bSSFP, T1 mapping, and T2 mapping, worse than the corresponding 73.1%, 66.8%, and 61.1% reproducible features in the intra-observer segmentation. Liu et al. (84) investigated 109 radiomics features on 436 contrastenhanced CT images of oropharyngeal cancer patients and found that "most radiomic features in this study varied a lot when the ROIs were not well segmented. For both the representation agreement and predictive agreement, the ICC and CCC were below 0.5 for all the features." Uthoff et al. (79) reported that "observers had perfect intra-repeatability (ICC =1.0)" but "demonstrated fair inter-reader variability (ICC =0.52)" for 4 observers (2 radiologists, 2 pulmonologists) in 100 cases of non-small cell lung cancer CT scans. Many other studies generally reported high SFRs, implying excellent robustness to intra/ inter-observer segmentation disagreement, independent of modalities and diseases, although different ICC thresholds were applied. Meanwhile, radiomics feature robustness to inter-observer segmentation seemed not notably inferior to intra-observer segmentation.

Comparable or better radiomics feature reliability was reported for (semi-)automated segmentation than manual segmentation with much shorter segmentation time

Manual image segmentation in radiomics analysis involved intensive labor work of clinicians and was timeconsuming, and also suffered from intra-/inter-observer segmentation disagreement, leading to the low costeffectiveness/efficiency of radiomics so greatly hampering its wide application in clinical practice. Thus, lots of efforts were taken to develop (semi-)automated segmentation as a potential alternative in radiomics research. Moreover, (semi-)automatic segmentation was frequently reported useful to further reduce the intra-/inter-observer radiomics feature variability induced by manual segmentation in the included studies in addition to its advantage in segmentation time, suggesting the future role of (semi-)automated segmentation in more reliable and cost-effective/efficient radiomics analysis (67,69,70,77,80,99,103,141,142).

Acquisition had substantial impacts on radiomics feature values, and their impact on feature reliability was larger than the impact by intra-/inter-observer segmentation Based on the reported ICC results, image acquisition had substantial impacts on the radiomics feature values for all imaging modalities and acquisition protocols. Regarding modality dependence, the reported SFRs seemed to be highest in PET and lowest in MRI [excluding the outlier of 100% inter-scanner SFR reported by Zhang J et al. (46)], which might be partially explained by the relatively smooth low-resolution PET image and the multi-contrast highresolution MRI images with considerable anatomical details acquired by different sequences. The simple intrascanner test-retest could introduce considerable feature value variations(17,18,24,28,32,36,40,44,139). The inter-scanner (or inter-center) acquisition (with similar imaging protocols or imaging parameter changes) induced even more radiomics feature variability than the intrascanner test-retest (32,35,45,135,137,139). In the studies investigating both acquisition and segmentation, acquisition was consistently reported to have much larger impacts on feature variability (always smaller ICCs) than segmentation (35,39,42,44,48,56,126,132) for all modalities.

Feature reliability and ICCs were heterogeneous for post-processing and feature quantification; optimized post-processing and feature quantification could be used to mitigate acquisition-induced radiomics variability

Image post-processing and feature quantification were

usually used to explore the robustness of radiomics features and improve feature reliability by optimized or standardized approaches. Among these approaches, image intensity discretization and normalization were most frequently investigated. Actually, there could be tremendous types of image post-processing and feature quantification methods, algorithms, and tools that were applicable to the acquired original images and thus had remarkable influences on feature values. In many studies, various post-processing and feature quantification approaches were conducted and optimized to mitigate the possible radiomics feature variability introduced in the acquisition procedure (28,31,36,41,44,111,114,129). The results suggested that comprehensive image perturbation and quantification might be helpful to improve radiomics reliability, in particular for those retrospective radiomics studies in which existing imaging data were used without control on imaging acquisition protocol. For example, in a study by Zwanenburg et al. (28), image perturbation chains were proposed to be used as an alternative to test-retest imaging to assess feature robustness. Most robust features in acquisition test-retest were successfully identified by comprehensive image perturbations. In another study by Suter et al. (64), single-center MRI data was perturbed to simulate unseen multi-center MRI data with greater variabilities, which generated and conducted over 16 million tests of typical perturbations and to identify robust radiomics features for multi-center radiomics study. In contrast, post-processing and feature quantification were seldom proposed for mitigation or compensation for the radiomics feature variability affected by segmentation.

Shape and first-order (FO) radiomics features were frequently reported to be more robust to various variability factors than texture features in the original image domain

Different types of radiomics features could subject to different levels of variability influenced by different factors. Among the heterogeneous reported results, it was noticed that shape or first-order (FO, or named histogram) features in the original image domain were often reported to be more robust than texture (also named second-order or higher-order) features to different variability factors of acquisition (18,36,44,57,134), post-processing and quantification (36,115,116,128,130,132), and segmentation (94,99,104,142-145). In different types of texture features, GLCM (gray-level co-occurrence matrix) features were observed to be more robust than other texture features in a few studies (18,44,94,116,128,143,145). On the other hand, opposite or deviant results on the low reliability of shape features were also occasionally reported. For instance, Rai et al. reported that none of the shape features exhibited high inter-(MRI) scanner stability (ICC >0.8), the lowest among all feature types (134). Tixier et al. showed that shape features in MRI (ICC =0.74) were among the most impacted feature types by the choice of segmentation method, with poorer reliability than first-order and GLCM features (ICC >0.96) (94). Beyond the radiomics features in the original image domain, radiomics features in the transformed domains, most frequently in the Laplacian of Gaussian (LoG) filtered domain and wavelet domain, were also investigated in many studies. No uniform robustness of these transformed features compared to those original features could be derived from the included studies.

Other statistical metrics in conjunction with ICC

A variety of statistical metrics were used in conjunction with ICC for different purposes. For segmentation purposes, dice similarity coefficient (DSC) was often reported. Bland-Altman analysis was conducted in some studies involving paired comparison of two observers/acquisitions/ measurements. Other types of statistical metrics such as concordance correlation coefficient (CCC), coefficient of variation (CV), Pearson/Spearman correlation coefficient, false discovery rate (FDR), (normalized) dynamic range, Krippendorff's alpha, percentage difference, and betweenclass distance (BD) were also used in combination with ICC.

Risk of bias

There were different levels and aspects of potential risks of bias in the included studies for many reasons. Many clinical studies were limited in their retrospective study nature and were usually conducted without phantom validation and control on acquisition protocol. Many technical studies utilized public imaging data, and the heterogeneity in these data might not be well understood or compensated. Very few studies described the imaging protocol sufficiently to the desired level of detail, as suggested in (4). Similarly, details in the intra-/inter-observer segmentation process were normally insufficiently described. For many clinical studies aiming for radiomics diagnosis or prognosis performance, the possible publication bias on the very high feature reliability to intra/inter-observer segmentation might not be neglected in that much lower ICC and SFRs were reported in the studies that substantially assessed feature robustness to segmentation as the primary study endpoint. The statistical power of the calculated ICC might not be strong enough due to the limited sample size and observer/acquisition/measurement number. The investigated radiomics features in many studies might have different definitions even with the same or similar name. They might not have been well standardized due to the different implementations in a variety of software and inhouse built programs, in particular in the studies before the proposal of feature standardization by the image biomarker standardization initiative (IBSI) (146). Besides, risks of bias could also be induced by the flaws and pitfalls of using and reporting ICC as identified in the articles.

Discussion

Radiomics research is experiencing an increasing explosive rate both in publication volume and diversity in recent years (5). However, along with the soaring publication numbers of radiomics, more concerns, questions, and/or criticisms on radiomics reliability are also increasingly raised in the last few years (11,147,148). Some recent systematic review papers (149-153) also showed that many radiomics publications had suboptimal or poor study quality, as revealed by the low RQS.

Reliability is highly correlated to RQS criteria. For instance, image protocol quality (+1 point), phantom study on all scanners (+1 point) and imaging at multiple time points (+1 point) are all related to acquisition reproducibility/replicability/reliability; Multiple segmentations (+1 point) is related to intra-/inter-observer agreement analysis; Feature reduction or adjustment for multiple testing (+3 point if implemented or -3 if not implemented) is related to feature correlation and redundancy analysis. ICC could be used to fulfill these criteria on RQS to improve radiomics study quality.

The increasing publication number with time reflects the fact that much more efforts have been taken to investigate radiomics reliability in recent years, in particular for MRI. Clinical oncology is still the major arena of radiomics research as revealed in this review, consistent with a recent bibliometric review (5), while different imaging modalities have substantially different roles in different cancers, as reflected in the proportions of the publications.

Regarding the ICC purpose, it is within the expectation that ICC was most frequently used for segmentation reliability assessment, particularly for intra-/inter-observer agreement. This could be mainly explained by the large fractions of retrospective studies. It also reflects the common interests and concerns on radiomics reliability from clinicians. It is interesting to notice that many intra-/ inter-observer agreement studies reported high ICC values or SFRs, which might suggest that many radiomics features are quite robust to (manual) lesion segmentation. In other words, radiomics reliability to lesion segmentation might not be much concerned.

Although segmentation dominated the ICC use, the importance of radiomics reliability in other aspects could not be overlooked or underestimated. Image acquisition and reconstruction are at the very front-end of the complex radiomics workflow, and greatly impact the quality of the original imaging data for radiomics reliability assessment. Indeed, image acquisition and reconstruction were strongly suggested to impact much more on radiomics reliability than segmentation. However, the influence of image acquisition and reconstruction on radiomics feature reliability was still much underexplored relative to segmentation studies. There are still many unknowns about how image acquisition and reconstruction affect radiomics reliability. Much research work is warranted in the future, in particular for MRI, due to its semi-quantitative image intensity nature, various image contrasts, and much greater variability in image acquisition and reconstruction compared to CT and PET.

ICC was also frequently used for reliability assessment attributed to post-acquisition image processing and feature quantification. The reported ICCs were heterogeneous in these studies, much dependent on the various processing types and different implementations. In theory, there could be infinite types of post-processing methods applicable to original images, so potentially lead to even bigger radiomics feature variability than acquisition and reconstruction. But, in practice, post-processing and feature quantification are investigated and utilized to mitigate acquisition and reconstruction-related feature variability by taking advantage of comprehensive and powerful computation capability for robust feature selection, without the need for prior knowledge on image acquisition variability. However, the evidence is so far not strong enough. More rigorous validation and evaluation are definitely warranted.

Some common pitfalls of using and reporting ICC were frequently identified in the articles. First, the information on ICC form and its selection was missing, ambiguous or incomplete in a large number of articles. Meanwhile, relevant information like scanner/observer/measurement

numbers sometimes was not clarified to facilitate ICC form selection. When ICC is used as a reliability metric, it is important for researchers to carefully select the most appropriate ICC form. The inappropriate selection of ICC form might mathematically yield similar ICC values but could lead to substantially different and even misleading interpretations. No article conducted sample size estimation for ICC calculation, which could be helpful, although might not be necessary, for ICC precision estimation. In terms of ICC reporting, ICC values were often reported without the confidence interval. Without reporting the confidence interval, the precision of ICC could not be known. For instance, a very high ICC value but associating with a very wide width of confidence interval (large uncertainty and low precision) could not guarantee the high reliability. Heterogeneous ICC thresholds were used for reliability assessment, also hampering rigorous data synthesis for crossstudy comparison. Occasionally, ICC threshold values were only implicitly indicated or unavailable. The reliability levels of poor (ICC <0.5), moderate (ICC: 0.50-0.75), good (ICC: 0.75-0.90), and excellent (ICC >0.90) as suggested in (13) were frequently adopted. But, on the other hand, the conditions under which the criteria were suggested were usually neglected, i.e., "As a rule of thumb, researchers should try to obtain at least 30 heterogeneous samples and involve at least 3 raters whenever possible when conducting a reliability study." (13). The reliability levels were also seemed to be inappropriately determined on the basis of the ICC value itself rather than its confidence interval. ICC was normally interpreted without further quantifying the underlying true variance $(\sigma_{\rm T}^2)$ and error variance $(\sigma_{\rm E}^2)$. Actually, a high ICC value might mainly reflect the high between-subject heterogeneity (such as malignant tumors) in the sampled population but does not guarantee the accuracy or precision of radiomics feature quantification. Vice versa, a low ICC might probably be resulted from the high homogeneity (such as normal tissues) in the subjects, even with high measurement accuracy and precision.

Some suggestions could be given to mitigate the identified pitfalls for future radiomics studies. Overall, if radiomics reliability itself is the major purpose of a radiomics study, guidelines for reporting reliability and agreement studies (GRRAS) should be helpful in the study planning (154). In order to facilitate ICC form selection, the model, type, and definition of the ICC form should be justified or explained. Relevant information like scanner/ observer/measurement numbers need to be sufficiently disclosed. The guideline proposed by Koo *et al.* is an

excellent reference and is easy to follow (13). It would be very helpful to conduct sample size estimation for ICC calculation in order to assure that the study could have an adequate chance of achieving the desired ICC precision (155,156). After ICC form selection, the tool used for ICC calculation should be reported with software name, version, and setting. The ICC calculation results should be reported along with the confidence interval. Meanwhile, the criteria for ICC appraisal should be clearly described. It should also be kept in mind that it is the estimated CI forms the basis to evaluate the reliability level, but not the ICC value itself. Along with ICC, the joint use of other statistical metrics could strengthen the study quality and statistical power. For instance, if paired observers/acquisitions/measurements were involved, Bland-Altman analysis is anticipated and beneficial. For segmentation reliability assessment, dice similarity coefficient (DSC) is desirable. Last but not least, the acceptability of ICC should be determined on the requirements by each specific study and clinical application, rather than simply on the calculated values from the specific sample populations and pre-defined thresholds.

There are some limitations to this study. First, the literature search in a single database was one limitation, although partially compensated by the prior knowledge on additional papers. Meanwhile, even in a single database, there are tremendous numbers of publications relevant to radiomics, but it is not uncommon that a variety of terms are used instead, which makes the precise localization of these publications even more difficult. So, there might still be potentially eligible studies missed for analysis. The ICC use and its result reporting had to be recognized and extracted through full text (and even Supplementary materials) examination rather than title and abstract screening. This procedure involved tremendous work and might slightly affect the inclusion and exclusion of papers. Nonetheless, the large sample size of 481 studies should not considerably weaken or bias the statistics in this review. Second, this review concentrated on a single metric of reliability, i.e., ICC, which tackles only a very narrow topic on general radiomics reliability. ICC is only applicable to continuous variables, so the radiomics reliability revealed by ICC is usually on the level of radiomics feature values. The role of ICC is relatively minor in the reliability aspects of radiomics feature reduction and modeling, as well as model outcome/performance assessment. It is acknowledged that many other statistical metrics could be applicable or more suitable in radiomics reliability assessment in various scenarios, providing complementary or additional information on radiomics reliability beyond ICC. Thus, the current status of radiomics reliability could only be partially reflected in the included papers. This study by no means formed a systematic review and meta-analysis on the diagnostic accuracy of radiomics, so study quality was not individually assessed in each article by following QUADAS-2 (15), TRIPOD (16) or RQS (4), but PRISMA statement was followed (157). Third, there were great difficulties in study quality normalization, data synthesis, and harmonization on the highly heterogeneous study characteristics along with the pitfalls in ICC use and reporting. It was of great difficulty to conduct quantitative analysis on cross-study ICC assessment. The use of SFR slightly mitigated this issue, but SFR itself also had pitfalls such as different ICC thresholds. Therefore, the consensus on the degree of radiomics reliability that has been achieved, or could be achievable in radiomics research could not be safely derived. Fourth, radiomics feature reliability has been suggested to be dependent on imaging modality, organ, disease, and other factors, which was also noticed in some included individual studies (72,89,91,103,104,158). But these dependencies could not be further generalized in this review. Our study collected, analyzed and presented data in a modality-neutral and disease-neutral way. Moreover, we also recognized that it was still an extremely difficult task for this dependency investigation in the presence of high heterogeneities of study characteristics even though hundreds of studies had been included. But, on the other hand, it should be cautioned that there might be a potential risk of bias by trying to present modality-neutral or disease-neutral common findings in the study. The validity of these findings might be violated if applied to some fewer common diseases or other modalities. Therefore, future research efforts on disease-specific and modality-specific feature reliability are desirable. Fifth, some flaws and pitfalls in selecting, reporting, and interpreting ICC were identified in many radiomics studies, so some suggestions were given. But we did not intend to specifically propose a standardized form of ICC use for future radiomics studies. The standardization of QIB metrology (159), the IBSI radiomics feature standardization (146,160), the guidelines for reporting reliability and agreement studies (GRRAS) (154), the general guideline of selecting and reporting ICCs (13), and statistical methods for clinical reliability in different aspects (121,161-164), have been well established in the medical literature. They could act as excellent guidelines or references for radiomics study planning. But, consensus toward the standardized radiomics reliability assessment and reporting is yet to be reached by the whole community.

Conclusions

This study attempted to have an updated overview on the current status of radiomics reliability research from the perspective of using and reporting ICC in the ever-fastexpanding radiomics literature. The 481 eligible CT, PET, and MRI radiomics studies yielded from the literature search partially revealed the fact that much more efforts have been taken to rigorously assess radiomics reliability for clinical use, in particular in the recent two years. ICC was used for assessing different aspects of radiomics feature reliability in these studies, but feature reliability with respect to image segmentation was much more reported than reliability to other factors such as image acquisition, reconstruction, post-processing, and feature quantification. As indicated by the reported satisfactory ICCs in intra/inter-observer segmentation agreement, manual segmentation seems to be the least influential factor on radiomics reliability, but the risk of bias might be cautioned. The (semi-)automated segmentation may further increase segmentation agreement to further increase radiomics feature reliability with better cost-effectiveness/ efficiency in the future. Image acquisition could introduce much more feature variability than image segmentation. More research on radiomics reliability with respect to image acquisition and reconstruction is desired. Comprehensive image post-processing and feature quantification techniques could be applied for radiomics analysis and yield different levels of radiomics reliability. Optimized comprehensive image post-processing and feature quantification could be used to mitigate image acquisition-induced variability and thus improve reliability. There were some common flaws and pitfalls in ICC use, as identified in many studies. Thus, some suggestions were given in order to mitigate them and to improve radiomics reliability research quality for future studies. Unfortunately, it was also recognized that the included studies were highly heterogeneous in characteristics and quality, greatly hampering the reliable data synthesis for further meta-analysis. Therefore, no consensus on the degree of radiomics reliability that has been achieved or could be achievable in radiomics research could be safely derived and reached by this review. More research works are warranted in the future.

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Footnote

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Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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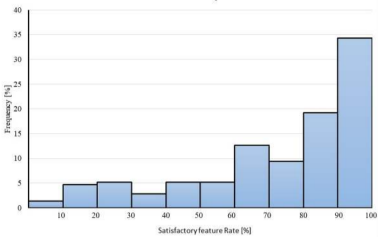
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Distribution of Satisfactory Feature Rate

Figure S1 Satisfactory feature rate distribution in the studies reported ICC results regarding the feature reliability due to image segmentation for intra-/inter-observer agreement.

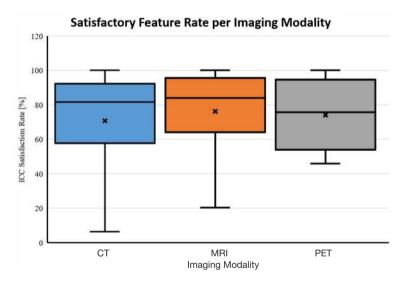


Figure S2 The difference of satisfactory feature rate in different imaging modality in the studies reported ICC results regarding the feature reliability due to image segmentation for intra-/inter-observer agreement.

PMID/doi	First author	Publication year	Disease or organ	Patient number	ICC purpose	ICC form described in the article	ICC model	ICC type	ICC definition
23807457	van Velden FHP et al.	2014	Colorectal cancer	29	Acquisition	2-way random single measure model	Yes	Yes	No
24047337	Leijenaar RTH et al.	2013	Lung cancer	34	Acquisition	ICC(1,1) for test -retest reliability and ICC(3,k) for inter-rater simulation	Yes	Yes	Yes
25025374	Parmar C et al.	2014	Lung cancer	20	Segmentation	Inter-observer segmentation (two-way mixed effect model, case 3A(McGraw), absolute agreement), intra-observer segmentation (one-way model, case 1(McGraw), form-specific formula given	Yes	Yes	Yes
26163091	Panth KM et al.	2015	Phantom	23	Segmentation	ICC(1,1), ICC(2,1)	Yes	Yes	Yes
26242464	Leijenaar RTH et al.	2015	Lung cancer	35	Quantification	Formula given, but cannot be specified to a certain form	No	No	No
26587549	Echegaray S et al.	2015	Liver cancer	29	Segmentation	A1 method (McGraw)	Yes	Yes	Yes
26795288	Rossum PSNR et al.	2016	HNC	217	Acquisition	An absolute agreement definition in a 2-way mixed-effects model	Yes	No	Yes
26920355	van Velden FHP et al.	2016	Lung cancer	11	Reconstruction	One-way random single-measure	Yes	Yes	No
27669756	Hu P et al.	2016	Colorectal cancer	40	Acquisition	Two-way mixed effect model, with formula given	Yes	Yes	Yes
27893446	Bogowicz M et al.	2016	HNC and lung cancer	22	Processing	Two-way mixed model, consistency	Yes	No	Yes
28612050	Echegaray S et al.	2016	Lung cancer	100	Segmentation	A1 method (McGraw)	Yes	Yes	Yes
29122358	Bogowicz M et al.	2017	HNC	178	Quantification	Two-way mixed single measures	Yes	Yes	Yes
29494598	Carvalho S et al.	2018	Lung cancer	215	Acquisition	ICC(1,k)	Yes	Yes	Yes
29513054	Pavic M et al.	2018	HNC and Lung Cancer	11	Segmentation	ICC(3,1), consistency	Yes	Yes	Yes
29520429	Lv W et al.	2018	NPC	106	Quantification	Formula given, but cannot be specified to a certain form	No	No	No
29633002	Shi Z et al.	2018	Vascular disease	96	Segmentation	Two-way random model with absolute measurements	Yes	Yes	Yes
29663411	Saha A et al.	2018	Breast cancer	50	Segmentation	ICC(3,1), consistency	Yes	Yes	Yes
29725965	Bologna M et al.	2018	HNC	36	Segmentation	Two-way mixed effect model	Yes	No	No
30002441	Hassan M et al.	2018	Lung cancer	18	Processing	Form-specific formula given	Yes	Yes	Yes
30135549	Bibault J et al.	2018	Colorectal cancer	95	Segmentation	Two-way mixed effect model, formula given	Yes	Yes	Yes
30158540	Ger RB et al.	2018	HNC and lung cancer	50	Processing	ICC (2, 1) (two-way random effects, absolute agreement, single ratter/measurement) and ICC (3, 1) (two-way random effects, consistency, single rater/measurement)	Yes	Yes	Yes
30230556	Soufi M et al.	2018	Lung cancer	305	Acquisition	Two-ray random effect model, absolute agreement condition	Yes	No	Yes
30286184	Owens CA et al.	2018	Lung cancer	10	Segmentation	Two-way mixed-effects model	Yes	No	No
30463529	Zheng B et al.	2018	Liver cancer	319	Segmentation	A two-way random, single measure (absolute agreement)	Yes	Yes	Yes

Table S1 The available ICC forms described in the studies

Table S1 (continued)

Table S1	(continued)
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PMID/doi	First author	Publication year	Disease or organ	Patient number	ICC purpose	ICC form described in the article	ICC model	ICC type	ICC definition
30506687	Pfaehler E et al.	2019	Phantom	1	Reconstruction, segmentation	Two-way single measure model for consistency	Yes	Yes	Yes
30679599	Alex Z et al.	2019	HNC and Lung Cancer	50	Acquisition	ICC(1,1)	Yes	Yes	Yes
30714158	Vuong D et al.	2019	Lung cancer	19	Acquisition	ICC(1,1) for 4DPETMR, ICC(3,1) for PETCT-PETMR	Yes	Yes	Yes
30738530	Mori M et al.	2019	Pancreatic cancer	31	Segmentation	One-way random single-measure model	Yes	Yes	No
30765732	Lecler A et al.	2019	Breast cancer	207	Segmentation	ICC(2,1), absolute agreement	Yes	Yes	Yes
30840747	Foy JJ et al.	2018	Healthy	39	Quantification	ICC(a,1)	Yes	Yes	Yes
30845221	Duron L et al.	2019	Breast cancer	104	Segmentation	2-way random intraclass correlation coefficient (ICC) (absolute agreement, average type)	Yes	Yes	Yes
30928060	Branchini M et al.	2019	Pediatric	21	Processing	One way random single measure model	Yes	Yes	No
30961895	Peeken JC et al.	2019	Soft tissue sarcoma	221	Segmentation	ICC(3,1)	Yes	Yes	Yes
31015155	Fiset S et al.	2019	Cervical cancer	62	Acquisition	ICC(1,1) for test -retest reliability and diagnostic-simulation, ICC(2,1) for inter-observer	Yes	Yes	Yes
31032192	Qiu Q et al.	2019	Liver cancer	106	Segmentation	Form-specific formula given	Yes	Yes	Yes
31063427	Kocak B et al.	2019	Kidney cancer	30	Segmentation	Two-way model, single-rating, and absolute agreement	Yes	Yes	Yes
31131906	Tixier F et al.	2019	GBM	90	Segmentation	Two-way random effects model	Yes	No	No
31273242	Whybra P et al.	2019	HNC	441	Processing	2-way mixed-effects model, single rater, absolute agreement	Yes	Yes	Yes
31375452	Huang L et al.	2019	Lung cancer	31	Acquisition	2-way fixed effect, absolute agreement, and single measurement model	Yes	Yes	Yes
31375883	Ugga L et al.	2019	HNC	89	Segmentation	Absolute agreement ICC value	No	No	Yes
31392481	Yamashita R et al.	2020	Pancreatic cancer	55	Acquisition	Mixed effects linear model	Yes	No	No
31420497	Pfaehler E et al.	2020	Phantom	3	Acquisition, quantification	2-way single-measure model for consistency of features	Yes	Yes	Yes
31423714	Ta D et al.	2020	Healthy	6	Acquisition	Two-way mixed-effects model	Yes	No	No
31477335	Traverso A et al.	2020	Cervical cancer	81	Processing	ICC(2,1)	Yes	Yes	Yes
31539450	Bologna M et al.	2019	Phantom	1	Acquisition, processing	Measure agreement in mixed-effect models, equivalent to the (A,1) model described in McGraw et al	Yes	Yes	Yes
31703155	Merisaari H et al.	2020	Prostate cancer	112	Acquisition	ICC(3,1)	Yes	Yes	Yes
31743889	Cong M et al.	2020	Lung cancer	50	Segmentation	Absolute agreement method	No	No	Yes
31761666	Wang Y et al.	2019	Gastric cancer	30	Segmentation	Two-way mixed effects, absolute agreement, single rater ICC test	Yes	Yes	Yes

Table S1 (continued)

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PMID/doi	First author	Publication year	Disease or organ	Patient number	ICC purpose	ICC form described in the article	ICC model	ICC type	ICC definition
31824843	Wu J et al.	2019	Colorectal cancer	102	Segmentation	Single rater; definition: absolute agreement; model: inter-user ICC: two-way random effects; intra-user ICC: two-way mixed effects	Yes	Yes	Yes
31824847	Chen W et al.	2019	Gastric cancer	30	Segmentation	Intraclass (single-rating, consistency, 2-way mixed effects model), interclass (multiple-rating, consistency, 2-way random-effects model)	Yes	Yes	Yes
31903240	Xu X et al.	2019	Lung cancer	132	Segmentation	Two-way fixed effect, absolute agreement, and single measurement	Yes	Yes	Yes
31918370	Kakino R et al.	2020	Lung cancer	256	Segmentation	Two-way random effects model under an absolute agreement condition	Yes	No	Yes
31941949	Yang F et al.	2020	Phantom	1	Segmentation	2-way random effect model to estimate the absolute agreement of multiple raters per measurement	Yes	Yes	Yes
31971614	Scalco E et al.	2020	Prostate cancer	14	Acquisition	Two-way mixed effect model, single rater type, both consistency and absolute agreement were used	Yes	Yes	Yes
32123281	Park BW et al.	2020	Bladder cancer	83	Processing	ICC(c,1)	Yes	Yes	Yes
32174316	Caballo M et al.	2020	Breast cancer	93	Segmentation	ICC(3,1)	Yes	Yes	Yes
32229081	Palle S et al.	2020	Endometrial cancer	30	Segmentation	Two-way random effects, single rater, agreement	Yes	Yes	Yes
32236847	Nardone V et al.	2020	Phantom	1	Acquisition	Mean rating, absolute agreement, and two-way mixed effects model	Yes	Yes	Yes
32240418	Cattell R et al.	2019	Phantom	5	Quantification	2-way mixed-effects model, single rater, absolute agreement	Yes	Yes	Yes
32277703	Rai R et al.	2020	Phantom	11	Acquisition	Two-way random effect with absolute agreement model	Yes	No	Yes
32395833	Vuong D et al.	2020	Lung cancer	124	Acquisition	ICC(3,1)	Yes	Yes	Yes
32399621	Haider SP et al.	2020	HNC	50	Segmentation	Two-way mixed effects, absolute agreement, single rater	Yes	Yes	Yes
32557189	Yang P et al.	2020	NPC	21	Processing	ICC(3,10)	Yes	Yes	Yes
32593138	Ramon R et al.	2020	GBM	100	Segmentation	Two-way random effects, absolute agreement, single rater/measurement	Yes	Yes	Yes
32705290	Cuocolo R et al.	2020	HNC	89	Segmentation	Two-way, absolute agreement and single rater ICC	Yes	Yes	Yes
32723501	Dreher C et al.	2020	Phantom	17	Segmentation	Two-way mixed model for absolute agreement and single measures	Yes	Yes	Yes
32728098	Haarburger C et al.	2020	Kidney cancer, liver cancer, lung cancer	1216	Segmentation	ICC(1,1)	Yes	Yes	Yes
32737518	Cysouw MCF et al.	2021	Prostate cancer	76	Segmentation	Two-way mixed effect model, absolute agreement	Yes	No	Yes

Table S1 (continued)

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PMID/doi	First author	Publication year	Disease or organ	Patient number	ICC purpose	ICC form described in the article	ICC model	ICC type	ICC definition
32758279	Suter Y et al.	2020	GBM	63	Reconstruction	ICC(2,1), absolute agreement	Yes	Yes	Yes
32767049	Song X et al.	2021	Ovarian cancer	104	Segmentation	Single-rater, absolute- agreement, 2-way mixed-effects model	Yes	Yes	Yes
32800693	Gutmann DAP et al.	2020	Liver cancer	310	Processing	ICC(1,1) for test reliability and ICC(3,k) for inter-rater agreement	Yes	Yes	Yes
32822054	Kim D et al.	2020	Lung cancer	35	Segmentation	Inter-rater agreement (absolute agreement, 2-way random effect model, $k = 2$), intra-rater agreement (absolute agreement, 2-way mixed effect model, $k = 2$)	Yes	Yes	Yes
32827069	Park YW et al.	2021	Lung cancer	51	Segmentation	One-way random effects model	Yes	No	No
32843663	Granzier RWY et al.	2020	Breast cancer	138	Segmentation	Two-way random single measure ICC(2,1)	Yes	Yes	Yes
32939634	Kulkarni A et al.	2020	Pancreatic cancer	128	Segmentation	Two-way model with single rating and absolute agreement	Yes	Yes	Yes
32968131	Crobe A et al.	2020	Soft tissue sarcoma	70	Segmentation	2-way random model, agreement between raters and 6 raters	Yes	Yes	Yes
32970859	Bianchini L et al.	2021	Phantom	1	Acquisition	Two-way random effects for absolute agreement and single rater/measurement	Yes	Yes	Yes
33118182	Pati S et al.	2020	GBM	31	Segmentation	ICC(3,1)	Yes	Yes	Yes
33128598	Park JE et al.	2020	GBM	422	Segmentation	Two-way mixed-effects model	Yes	No	No
33137621	Tsarochi M et al.	2020	Breast cancer	73	Segmentation	Two-way mixed effect model, single measurement for absolute agreement	Yes	Yes	Yes
33228815	Wang X et al.	2020	Gastric cancer	539	Segmentation	Multiple-rating, consistency, 2-way random-effects model	Yes	Yes	No
10.1109/ACCESS. 2019.2923755	Li Z et al.	2019	Brain	15	Acquisition, pro- cessing	ICC(1,1) for intra-observer repeatability, ICC(3,1) for inter-observer repeatability	Yes	Yes	Yes
10.1117/12.2512406	Hendrik MJ et al.	2019	Liver cancer	13	Segmentation	ICC(2,1)	Yes	Yes	Yes
10.21037/ trc.2017.09.47	Qingtao Q et al.	2017	Liver cancer	15	Segmentation	ICC(A,1) (inter-observer segmentation), ICC(C,1) (intra-observer segmentation), formulas given	Yes	Yes	Yes
10.4274/imj. galenos.2019.09582	Burak K et al.	2019	GBM	70	Segmentation	Two-way model, single-rating, and absolute agreement	Yes	Yes	Yes