Inter- and Intra-software Reproducibility of Computed Tomography Lung Density
 Measurements

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38 ABSTRACT (376/500 words)

Purpose: Multiple commercial, open-source, and academic software tools exist for objective
quantification of lung density in computed tomography (CT) images. The purpose of this study
was evaluate the inter-software reproducibility of CT lung density measurements.

Methods: CT images from 50 participants from the COPDGeneTM cohort study were randomly 42 selected for analysis; n=10 participants across each Global Initiative for Chronic Obstructive 43 Lung Disease (GOLD) grade (GOLD 0-IV). Academic-based groups (n=4) and commercial 44 vendors (n=4) participated anonymously to generate CT lung density measurements using their 45 software tools. CT total lung volume (TLV), percentage of the low attenuation areas in the lung 46 with Hounsfield unit (HU) values below -950HU (LAA₉₅₀), and the HU value corresponding to 47 the 15th percentile on the parenchymal density histogram (Perc15) were included in the analysis. 48 49 The inter-software bias and reproducibility coefficient (RDC) was generated with and without quality assurance (QA) for manual correction of the lung segmentation; intra-software bias and 50 RDC was also generated by repeated measurements on the same images. 51

Results: Inter-software mean bias was within ± 0.22 mL, $\pm 0.46\%$, and ± 0.97 HU for TLV, LAA₉₅₀ 52 and Perc15, respectively. The reproducibility coefficient (RDC) was 0.35L, 1.2% and 1.8HU for 53 TLV, LAA₉₅₀ and Perc15, respectively. Inter-software RDC remained unchanged following QA: 54 0.35L, 1.2% and 1.8HU for TLV, LAA₉₅₀ and Perc15, respectively. All software investigated had 55 56 an intra-software RDC of 0. The RDC was comparable for TLV, LAA₉₅₀ and Perc15 measurements, respectively, for academic-based groups/commercial vendor-based software 57 58 tools: 0.39L/0.32L, 1.2%/1.2%, and 1.7HU/1.6 HU. Multivariable regression analysis showed that academic-based software tools had greater within-subject standard deviation of TLV than 59

commercial vendors, but no significant differences between academic and commercial groups
were found for LAA₉₅₀ or Perc15 measurements.

62 Conclusions: CT total lung volume and lung density measurement bias and reproducibility was reported across eight different software tools. Bias was negligible across vendors, reproducibility 63 was comparable for software tools generated by academic-based groups and commercial 64 vendors, and segmentation QA had negligible impact on measurement variability between 65 software tools. In summary, results from this study report the amount of additional measurement 66 variability that should be accounted for when using different software tools to measure lung 67 density longitudinally with well-standardized image acquisition protocols. However, intra-68 software reproducibility was deterministic for all cases so use of the same software tool to reduce 69 variability for serial studies is highly recommended. 70

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Keywords (up to 5): computed tomography, imaging biomarker, emphysema, COPD, lung
density

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75 INTRODUCTION

Computed tomography (CT) lung density is an imaging biomarker used to objectively and non-76 invasively quantify the extent of emphysema in the lung. Over the last three decades, numerous 77 studies in patients with chronic obstructive pulmonary disease (COPD) have demonstrated that 78 79 CT lung density measurements are correlated with emphysema measured in excised lungs by histology (2,3), are associated with mortality (5) and exacerbations (6), and can identify 80 81 subgroups of patients with better responses following lung-volume-reduction surgery (7) and endobronchial valve implantation (8). Further, in patients with alpha 1-antitrysin deficiency, a 82 83 significant response to augmentation therapy was shown using CT lung density as a surrogate of emphysema, but not with conventional spirometry measurements (9). These findings all 84 highlight the potential role of quantitative CT for COPD patient management, such as 85 longitudinal monitoring of disease progression and assessing treatment response. 86

Maintaining standardized image acquisition parameters, however, is critically important for serial assessments that aim to quantify CT lung density. It is well-established that there are technical challenges for generating reproducible CT measurements. Submaximal inspiration breath-hold volume (10), dose (11,12) as well as image reconstruction parameters, including 91 slice thickness (13,14) and reconstruction kernel (15–17), have all been shown to impact CT 92 measurements. However, several large, multicenter, longitudinal cohort studies, such as 93 COPDGeneTM (18), have utilized breath-hold coaching and dedicated lung phantoms to 94 standardize image acquisition and reconstruction parameters across all sites to minimize 95 variability introduced by image acquisition related parameters.

Another factor that has the potential to impact the reproducibility of CT measurements is the 96 specific software used to generate the measurements. Lung density measurements are derived 97 from the parenchymal density histogram of CT Hounsfield unit (HU) values and thus are 98 deterministic computations and are directly computed given an accurate lung segmentation mask 99 100 (1-4). However, measurement variability may be introduced by differences in the thoracic cavity segmentation, as well as segmentation of the large airways and pulmonary vessels, even 101 102 when consistent image acquisition and reconstruction settings are utilized. Previous studies investigating the influence of different software tools have shown conflicting results, and in 103 104 some studies high inter-software variation for CT lung density measurements have been reported (19-21).105

In an effort to standardize methodology, the Lung Density Committee of the Quantitative 106 Imaging Biomarker Alliance (QIBA) has released for public comment a profile regarding the CT 107 lung density measurement (22). Given the multitude of software tools used by different 108 commercial, open-source, and academic research laboratories, an evaluation of the inter-software 109 variability of CT lung density measurements is warranted to support this profile, particularly in 110 the context of serial investigations. Further, quantifying inter-software CT measurement 111 reproducibility requires a cohort with minimal variability introduced by image acquisition 112 parameters. Therefore, here our objective was to investigate and report CT lung volume and 113 lung density measurement inter-software bias and reproducibility using CT images from the 114 COPDGeneTM cohort study, with various academic groups and commercial vendors participating 115 in the reproducibility study. 116

117

118 **METHODS**

119 Details of the Software Comparison

CT images from 50 participants from the COPDGeneTM cohort study (18) were selected for 120 analysis; n=10 participants across each COPD GOLD grade (GOLD 0-IV) were randomly 121 122 selected. Participation was solicited from academic groups and commercial vendors, and the solicitation letter indicated that the results would be anonymized (i.e. the software packages were 123 provided on the condition they would not be individually identified). The anonymization was 124 performed by The Radiological Society of North America (RSNA) that acted as a neutral broker 125 between all participating groups and the QIBA Lung Density committee, to ensure the committee 126 was blinded to the participants' identity. The CT datasets used in this study are accessible in the 127 Quantitative Imaging Data Warehouse (QIDW): https://qidw.rsna.org/. 128

All vendors indicated if their software tool was for academic use only or commercial. Vendors were instructed to generate measurements: 1) without segmentation quality assurance (QA) or manual correction to evaluate inter-software reproducibility; 2) a repeated set of measurements on the same images, to evaluate intra-software reproducibility; and, 3) a third set of measurements repeated on the same images following segmentation QA and manual correction.

134 CT Image Acquisition

135 CT images were acquired using CT systems of various makes and models, including GE, 136 Siemens and Philips models, with the participant supine at suspended full-inspiration from apex 137 to base of the lung as previously described (18). In general, CT images were reconstructed with 138 smooth convolution kernels (Siemens B31f, GE STANDARD, or Philips B) and slice 139 thicknesses and intervals between 0.625 and 0.75 mm. The full-dose protocol used an effective 140 dose of 200 mAs without dose modulation. A more detailed description of the CT image 141 acquisition protocol is described elsewhere (18).

142 CT Image Analysis

143 CT images were processed using academic and commercial CT lung density software. All groups 144 were instructed to generate CT measurements for each image dataset using none or a minimal 145 amount of manual software interaction. We also requested no image auto-calibration or pre-146 processing (e.g. noise reduction filtering). All vendors were asked to perform the following 147 steps for lung segmentation:

- 148 1. Segmentation of the lung parenchyma from the rest of the thoracic cavity;
- Removal of airways from the segmentation (no strict definition of which airways were
 required to be removed was provided, but the software was required to at least remove

the trachea and major bronchi from the air-space prior to computing the CT lung densitymetrics);

153 3. Blood vessel removal (no instruction was provided on the amount of acceptable blood154 vessel exclusion from the lung volume).

Next, groups were instructed to repeat each of these steps on the same image dataset in order to assess the intra-software repeatability. Finally, the vendors were asked to perform quality assurance (QA) by reviewing and manually correcting any lung segmentation errors to generate a third set of CT measurements using the corrected segmented lung volume.

The measurements generated include: the total lung volume (TLV), percentage of the low attenuation areas in the lung with HU values below -950 (LAA₉₅₀) (1–3), and the HU unit value corresponding to the 15th percentile on the parenchymal density histogram (Perc15) (4).

162 Statistical Analysis

All statistical analysis was performed using SAS 9.4 software (Cary, NC, USA) and MATLAB 163 R2018a (Natick, MA, USA). A one-way analysis of variance (ANOVA) with a Tukey test for 164 multiple comparison correction was performed for statistical comparison between GOLD groups 165 166 for age; for sex and race, a Fisher's Exact test was used. MATLAB was used for Bland-Altman analysis to compare measurements generated by each possible pair of software tools; 167 measurements included TLV, LAA₉₅₀ and Perc15 without QA. The reproducibility coefficient 168 (RDC) (23) was calculated for each software tool, as described below, to compare between the 169 170 different software tools for each lung measurement with and without QA, and by group type (academic-based, commercial). The RDC is the value under which the difference between 171 repeated measurements on the same participant acquired under different conditions (ie. different 172 software tools) should fall within 95% probability. To estimate the RDC for any given software 173 174 tool, we must estimate the variance relative to the other K-1 software tools in the comparison 175 (K=8 in our study). Therefore, for a specific software tool, l, we calculated the mean variance, σ_l^2 , for the measurements, subscript *i*, across the 50 image sets, where $M_{i,l}$ represents 176 measurement *i* of software *l* and $\sigma_{i,k,l}^2$ represents the variance between software *l* and software *k* 177 178 for measurement *i*:

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$$\sigma_{i,k,l}^2 = \frac{1}{2} (M_{i,k} - M_{i,l})^2$$

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180 Next, $\sigma_{k,l}^2$ represents the variance between software k and software l averaged over all 181 measurements N:

$$\sigma_{k,l}^2 = \frac{1}{N} \sum_{i=1}^{N} \sigma_{i,k,l}^2 = \frac{1}{N} \sum_{i=1}^{N} \frac{1}{2} \left(M_{i,k} - M_{i,l} \right)^2 = \frac{1}{2N} \sum_{i=1}^{N} \left(M_{i,k} - M_{i,l} \right)^2$$

183 Then, the average variance over the other K - 1 software tools is calculated to generate the 184 average variance for software *l*:

$$\sigma_l^2 = \frac{1}{K - 1} \sum_{k=1}^{K - 1} \sigma_{k,l}^2$$

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186 The average RDC for software l is then given by:

ada

$$RDC_l = 1.96 * \sqrt{2\sigma_l^2}$$

Low RDC values indicate high reproducibility between software tools. The 95% confidenceintervals for the RDC were constructed using bootstrapping with 5000 resamples.

Multivariable linear regression models were built to assess whether group type (academic-based, commercial) was a predictor of the within-subject standard deviation of TLV, LAA₉₅₀ and Perc15 measurements. If group type was found to be a significant predictor, it would indicate that the standard deviation between software tool measurements is different for commercial vendors and academic groups; in other words, it would indicate that CT measurements are more similar between commercial vendors or academic groups. Generalized estimating equations (GEEs) were used to account for the clustered nature of the data.

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198 **RESULTS**

A total of 50 participants were investigated: n=10 in each GOLD grade. As shown in Table 1, 199 200 there were no differences between the groups for age, sex or race. A total of 9 software tools participated in the study; software tools 1-4 were from academic-based groups and software tools 201 202 5-9 were from commercial vendors. A single commercial vendor withdrew from the study and therefore a total of 8 software tools, n=4 research-based and n=4 commercial, were included in 203 204 the analysis. All eight software tools were able to generate measurements for all images provided. A total of 3 of 8 software tools reported some manual editing of the segmentation 205 206 masks for some of the CT images as part of the QA step.

Figure 1 shows an example of the CT lung volume (in blue) and LAA₉₅₀ segmentation masks (in red) for two different software tools. The differences observed for exclusion of airways and vessels from the lung volume segmentation mask between the two software tools are subtle and representative of the type of differences that would be expected given acceptable segmentation quality for both images (i.e. no major segmentation errors).

212 Bland-Altman Analysis

Bland-Altman analysis was performed for TLV, LAA₉₅₀, and Perc15 measurements for each software tool compared with all other software tools. Table 2 provides the summary of the Bland-Altman analysis for measurements generated by each software tool with the average of all the other software tools for TLV, LAA₉₅₀, and Perc15 measurements. There was negligible bias for all software tools to within $\pm 0.22L$, $\pm 0.46\%$, and $\pm 0.97HU$, for TLV, LAA₉₅₀, and Perc15 respectively.

219 Reproducibility Coefficients

Table 3 shows the RDC for TLV, LAA₉₅₀ and Perc15 measurements for eight different software 220 tools with and without quality assurance (QA) using manual correction of the lung volume 221 segmentation. Overall, inter-software RDC was 0.35L, 1.2% and 1.8HU for TLV, LAA₉₅₀ and 222 Perc15, respectively. Inter-software RDC remained unchanged following QA: 0.35L, 1.2% and 223 1.8HU for TLV, LAA₉₅₀ and Perc15, respectively. Intra-software RDC was generated by 224 performing repeated measurements using the same software tool without QA; all software had an 225 intra-software RDC of 0, indicating that image processing workflows were deterministic for all 226 software tools. 227

Table 4 shows the RDC for TLV, LAA₉₅₀ and Perc15 measurements for software tools by group type (academic or commercial) with and without QA. Academic groups and commercial vendor's software tools generated comparable RDC measurements for TLV, LAA₉₅₀ and Perc15: 0.39L / 0.32L, 1.2% / 1.2%, and 1.7HU / 1.6 HU, respectively. As shown in Table 4, QA had negligible impact on measurement reproducibility between software.

233 Multivariable Linear Regression Models

Table 5 shows multivariable linear regression models for within-subject standard deviation of TLV, LAA₉₅₀ and Perc15 measurements generated by the different software tools with group type (academic, commercial) as a predictor. In the multivariable linear regression model for within-subject standard deviation of TLV, group type (academic=1, commercial=2) was a significant predictor (p<0.0001); this indicates that academic vendors had greater within-subject standard deviation of TLV measurements than commercial vendors. However, group type was not a significant predictor for within-subject standard deviation in the multivariable linear regression model for LAA₉₅₀ (p=0.46) or Perc15 measurements (p=0.24).

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243 **DISCUSSION**

There have been numerous clinical and research studies demonstrating that quantitative CT lung 244 density measurements are related to important outcomes in COPD patients (5-8) and in patients 245 with alpha 1-antitrysin deficiency (9). Potential clinical applications include patient selection for 246 treatment (e.g. by lung volume reduction surgery or endobronchial valves), or for evaluating 247 treatment response over time. However, in order for CT lung density measurements to be used as 248 a surrogate of emphysema in clinical applications, the variability of the CT measurements must 249 be carefully controlled. Several large, multicenter, longitudinal cohort studies, including 250 COPDGene (18), SPIROMICS (24), ECLIPSE (25), MESA (26) and CanCOLD (27), have 251 implemented standardized image acquisition protocols to carefully control for known factors that 252 253 impact CT measurements. However, the number of software tools developed by academic groups and commercial vendors to generate CT lung density measurements is increasing, with several 254 255 well-established commercial and prototype software packages now available, and each has their own proprietary segmentation algorithms. For serial assessments or longitudinal evaluations 256 257 where there is potential to change software tools at different time-points the reproducibility of CT measurements generated for various software tools must be evaluated. 258

259 In this study we evaluated reproducibility for eight different software tools, including wellestablished software from both academic groups and commercial vendors. We investigated 260 never-smokers and participants with a range of COPD severities. Our results indicate relatively 261 high reproducibility across the different software tools for TLV, LAA₉₅₀ and Perc15 262 measurements. Although the Bland-Altman analysis and Figure 1 indicate there are clear 263 differences for total lung volume segmentation between some of the vendors, which may result 264 in the slight deviations observed in the Bland-Altman analysis for LAA₉₅₀, the bias overall was 265 quite low and for LAA₉₅₀ the bias was less than 1% between all vendors. This bias is much less 266 than reported previously by Wielputz and colleagues (20) who investigated five software tools 267 268 (two academic and three commercial) for lung density measurements in COPD. The more

reproducible findings reported here may be related to several factors: the wider range of severity of the patients investigated (the patients evaluated by Wielputz and colleagues (20) were mainly end-stage COPD); the fact that a more standardized image acquisition protocol was used for COPDGene; or potentially improvements in image processing techniques over the last several years leading to more reproducible measurements between software tools.

274 In addition to assessing inter-software agreement for CT measurements, we also generated reproducibility coefficients (RDC) to determine how much variability may be introduced by 275 using different software tools when repeated measurements are made on the same patient. 276 Again, although the measurements generated by some software tools agreed slightly better than 277 others, the RDC values were low, and overall the RDC between all software tools was only 1.2% 278 for LAA₉₅₀. For example, this indicates that if the software tool was changed during a 279 longitudinal study, whereby there were repeated measurements on the same patient but 280 measurements were made using different software, the variability attributed to the software 281 would be 1.2% for LAA₉₅₀. In other words, to detect real emphysema progression, the 282 variability due to inter-software reproducibility measured in this study is 1.2% for LAA₉₅₀. 283 284 However, to determine the true overall RDC, the inter-software reproducibility would need to be combined with expected test/re-test measurement repeatability arising from differences in patient 285 positioning, scanner model, scanner calibration, breath hold volumes, etc., and a detection of 286 progression would need to be greater than the combined variability to be considered significant. 287 288 Obuchowski et. al. (28) has described the RDC calculations required to compute measurement reproducibility and repeatability. In general, however, we recommend that the same software be 289 290 used for sequential measures during a longitudinal study, especially given that all methods showed deterministic intra-software reproducibility. 291

Intra-software reproducibility was evaluated by having all groups run their software tool on the same CT images a second time. The RDC for the intra-software comparison was zero. We also requested each vendor run their software a third time and perform more rigorous QA. Although 3 of 8 vendors reported that manual edits were required in some of the participants evaluated (eg. lung volume edits or airway and vessel removal), the RDC did not change between the first run when there was no QA and the third run when QA was performed. This finding suggests that the results generated between the software tools were similar regardless of whether QA was performed. This may indicate that lung segmentation and airway and vessel removal algorithmsgenerate similar results between vendors, before manual editing.

301 Finally, we investigated the RDC for CT measurements stratified by whether the software was developed by academic-based groups or commercial vendors. Although based on the RDC we 302 found that the lung volume segmentation results tended to agree slightly better within 303 304 commercial vendors than academic groups, the difference was very small and the RDC for LAA₉₅₀ was 1.2% for both commercial and research vendors. This observation was consistent 305 with the results of the multivariable linear regression analysis in which we investigated group 306 type as a predictor of the standard deviation between the CT measurements generated by the 307 different software tools. We found commercial vendors had lower within-subject standard 308 deviation of TLV than academic groups, but no difference was found for LAA₉₅₀ or Perc15 309 These findings indicate that for CT lung density measurements, the 310 measurements. reproducibility within academic-based and commercial vendors is similar. 311

312 Although efforts must be made to standardize CT measurements, including image acquisition protocols and image analysis software, there are other sources of variability that may impact CT 313 314 measurements that were not considered in our study that must be acknowledged. For studies that acquire multiple CT image series over a short period of time, there is the potential for variability 315 316 to be introduced due to physiological or patient-related factors, but not disease related factors, such as the patient orientation in the bore, slightly different lung inflation volumes at breath-317 318 hold, etc. Previous studies have investigated the short-term repeatability of CT lung density measurements within the same-day (29), over two-weeks (30) and over a 1-year period (31) in 319 320 healthy volunteers and COPD patients. Although all studies report high short-term repeatability for CT measurements, these patient related factors may also impact how the software performs, 321 322 and may add additional variability between groups. Therefore, an important limitation in our 323 study is that we did not investigate both the reproducibility and short-term repeatability of the CT measurements between software tools. Our study is also limited by the fact that assessment 324 of CT lung segmentation accuracy is ultimately subjective, and therefore we were only able to 325 compare measurement reproducibility between the various software tools rather than accuracy, 326 327 as ground truth segmentation is not available. Another factor that should be considered is the potential for individual commercial or academic groups to upgrade their software over time. For 328 serial and longitudinal studies, even when the same software tool is used for CT analysis, CT 329

measurement reproducibility may need to be reassessed. Further, we note that we did not acquire 330 CT measurements by lung lobe from software tools and therefore we did not investigate CT 331 332 measurement reproducibility at the lobar level. Lobar segmentation algorithms between software tools may be more variable than whole lung segmentation. Reporting CT lung volume and 333 density measurements by lobe is relevant for lung volume reduction applications, and therefore 334 should be investigated in future studies. We also acknowledge that instruction was provided to 335 the academic-based groups and commercial vendors using their software tools for performing the 336 analysis, including how much manual intervention was permitted and that there should be no 337 pre-processing of the images. This may or may not mimic how these vendors generate CT 338 measurements routinely. However, the goal of our study was to assess the reproducibility of 339 their software for generating CT lung density measurements under standardized conditions. 340 341 Finally, as a result of the well-standardized CT image acquisition parameters used in this study, these findings may only be applicable to other well-standardized studies, or to clinical trials. 342 Further investigation is required to determine CT measurement reproducibility between software 343 tools for studies involving a wider range of CT acquisition parameters, such as those used in 344 clinical practice. 345

In conclusion, we evaluated CT lung volume and lung density measurement reproducibility 346 347 between eight different software tools using CT images acquired with standardized image acquisition protocols. The bias was negligible and measurement reproducibility was high 348 349 between software tools, and was comparable for software developed by academic-based groups and commercial vendors. While using the same software tool for serial studies is highly 350 351 recommended, these findings report how much added measurement variability will be introduced should it be necessary to include different software tools in serial studies with standardized 352 353 image acquisition parameters, and provides guidance on how to incorporate such information into longitudinal studies. 354

355 FIGURE LEGENDS

356

Figure 1. CT Lung and Emphysema Segmentation Generated by Two Different Software Tools. Shown above are two examples of CT lung segmentation images from two different software tools. Areas of the lung greater than or equal to -950 HU are colored in blue, areas less than -950 HU are colored in red. Differences in the inclusion of blood vessels (yellow arrows) and airways 361 (white arrows) can impact lung volume and low-attenuation area calculations. Note that the CT
 362 slice in this figure was the slice with the largest disagreement in segmentation volume over the
 363 entire image series.

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381 DISCLOSURE OF CONFILICTS OF INTEREST

382

JS was an employee and shareholder of VIDA Diagnostics Inc.; MK is a consultant at VIDA
Diagnostics Inc.; CH is an employee of Imbio.

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Parameter*	GOLD 0	GOLD I	GOLD II	GOLD III	GOLD IV
	(n=10)	(n=10)	(n=10)	(n=10)	(n=10)
Age, yrs	68 (8)	69 (9)	63 (10)	68 (9)	62 (6)
Female Sex, n (%) Race, n (%)	4 (40)	4 (40)	4 (40)	3 (30)	5 (50)
Non-Hispanic White	10 (100)	9 (90)	8 (80)	8 (80)	6 (60)
African American	0 (0)	1 (10)	2 (20)	2 (20)	4 (40)

Table 1. Subject Demographics

* All parameter values are mean (+/- SD) unless otherwise noted.

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 Table 2. Bland-Altman Analysis for Each Software Compared to the Average of All Other

 Software Tools

	Mean Bias	Median Bias	SD of Bias	Upper 95%	Lower 95%
				CI	CI
TLV (L)					
Software 1	-0.15	-0.14	0.05	-0.06	-0.25
Software 2	0.04	0.02	0.04	0.12	-0.04
Software 3	0.22	0.23	0.05	0.32	0.13
Software 4	0.05	0.04	0.02	0.09	0.01
Software 5	-0.04	-0.04	0.03	0.01	-0.09
Software 6	-0.01	-0.01	0.02	0.03	-0.06
Software 7	-0.21	-0.19	0.05	-0.10	-0.31
Software 8	0.10	0.09	0.04	0.18	0.02
LAA ₉₅₀ (%)					
Software 1	0.33	0.22	0.37	1.05	-0.40
Software 2	-0.24	-0.18	0.28	0.31	-0.80
Software 3	-0.29	-0.14	0.34	0.37	-0.95

Software 4	-0.42	-0.39	0.29	0.15	-0.98
Software 5	-0.34	-0.34	0.19	0.03	-0.71
Software 6	0.42	0.39	0.20	0.82	0.02
Software 7	0.46	0.26	0.49	1.42	-0.50
Software 8	0.09	0.10	0.12	0.32	-0.15
Perc15 (HU)					
Software 1	-0.33	-0.39	0.47	0.58	-1.24
Software 2	0.20	0.14	0.36	0.90	-0.51
Software 3	0.97	0.95	0.46	1.87	0.07
Software 4	0.49	0.54	0.39	1.25	-0.27
Software 5	0.24	0.17	0.36	0.95	-0.47
Software 6	-0.88	-0.80	0.27	-0.35	-1.40
Software 7	-0.58	-0.57	0.57	0.54	-1.70
Software 8	-0.11	-0.18	0.32	0.52	-0.74

Table 3. The RDC for TLV, LAA₉₅₀ and Perc15 for All Software Tools with and without QA

	Inter-so	oftware RDC	Inter-software RDC	
	Without QA		Wi	th QA
Parameter	RDC	95% CI	RDC	95% CI
TLV (L)				
Total	0.35	0.32 - 0.37	0.35	0.32 - 0.37
Software 1	0.38	0.35 - 0.41	0.38	0.35 - 0.42
Software 2	0.26	0.24 - 0.27	0.26	0.24 - 0.28
Software 3	0.26	0.24 - 0.29	0.26	0.24 - 0.29
Software 4	0.48	0.46 - 0.51	0.48	0.45 - 0.51
Software 5	0.25	0.23 - 0.27	0.25	0.23 - 0.27
Software 6	0.46	0.43 - 0.49	0.46	0.43 - 0.49
Software 7	0.31	0.28 - 0.34	0.31	0.28 - 0.34
Software 8	-	-	-	-
Software 9	0.26	0.24 - 0.28	0.26	0.24 - 0.28
LAA ₉₅₀ (%)				

T (1	1.0	10 14	1.0	10 14
Total	1.2	1.0 - 1.4	1.2	1.0 - 1.4
Software 1	1.2	1.0 - 1.5	1.2	1.0 - 1.5
Software 2	1.1	0.9 - 1.2	1.1	0.9 - 1.2
Software 3	1.1	0.9 - 1.2	1.1	0.9 - 1.2
Software 4	1.2	0.9 - 1.4	1.2	0.9 - 1.4
Software 5	1.2	1.0 - 1.3	1.2	1.0 - 1.3
Software 6	1.5	1.2 - 1.8	1.5	1.2 - 1.8
Software 7	0.9	0.7 - 1.0	0.9	0.7 - 1.0
Software 8	-	-	-	-
Software 9	1.2	1.0 - 1.4	1.2	1.0 - 1.4
Perc15 (HU)				
Total	1.8	1.6 - 2.0	1.8	1.6 - 2.1
Software 1	1.6	1.4 - 1.9	1.7	1.4 - 1.9
Software 2	1.5	1.3 – 1.7	1.6	1.3 – 1.8
Software 3	1.5	1.3 – 1.6	1.5	1.3 – 1.6
Software 4	2.3	2.1 - 2.6	2.3	2.1 - 2.6
Software 5	2.1	1.9 - 2.3	2.1	1.9 - 2.3
Software 6	2.0	1.6 - 2.3	2.0	1.6 - 2.4
Software 7	1.4	1.2 - 1.7	1.4	1.2 - 1.6
Software 8	-	-	-	-
Software 9	1.7	1.5 - 1.9	1.7	1.5 – 1.9

 Table 4. The RDC for TLV, LAA₉₅₀ and Perc15 for Academic-based and Commercial Software

 Tools

Parameter	Inter-software RDC	95% CI	Inter-software RDC	95% CI
	without QA		with QA	
TLV (L)				
Academic	0.39	0.36 - 0.41	0.39	0.36 - 0.41
Commercial	0.32	0.29 - 0.34	0.32	0.29 - 0.35

$LAA_{950}(\%)$				
Academic	1.2	0.9 - 1.4	1.2	0.9 - 1.4
Commercial	1.2	1.0 - 1.3	1.1	1.0 - 1.3
Perc15 (HU)				
Academic	1.7	1.5 - 1.9	1.7	1.5 - 1.9
Commercial	1.6	1.3 – 1.9	1.6	1.3 - 2.0
0				

Table 5. Multivariable Linear Regression Analysis for Software Tool Type with StandardDeviation of TLV, LAA950 and Perc15

	Estimate	Standard Error	Significance of
			Difference (p)
TLV [SD]	-0.03	0.004	< 0.0001
LAA ₉₅₀ [SD]	-0.009	0.01	0.46
Perc15 [SD]	-0.04	0.03	0.24

*Software type (academic=1, commercial=2)

Author

