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Non invasive measurement of liver iron concentration using 3 Tesla magnetic resonance imaging: validation against biopsy

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ABSTRACT

OBJECTIVES: To evaluate the performance and limitations of the R2* and signal intensity ratio (SIR) methods for quantifying liver iron concentration (LIC) at 3T.

METHODS: One hundred and five patients who underwent a liver biopsy with biochemical LIC (LIC_b) were included prospectively. A 3T MRI scan with a breath-hold multiple-echo gradient-echo sequence (mGRE) was undertaken for all patients. LIC calculated by 3T SIR algorithm (LIC_{SIR}) and by R2* (LIC_{R2*}) were correlated to LIC_b. Sensitivity and specificity were calculated. The comparison of methods was analyzed for successive classes.

RESULTS: LIC_b was strongly correlated to R2* (r = 0.95 p < 0.001) and LIC_{SIR} (r = 0.92 p < 0.001). In comparison to LIC_b, LIC_{R2*} and LIC_{SIR} detect liver iron overload with a sensitivity/specificity of 0.96 / 0.93 and 0.92 / 0.95, respectively and a bias \pm SD of 7.6 \pm 73.4 and 14.8 \pm 37.6 µmol/g, respectively. LIC_{R2*} presented the lowest differences for patients with LIC_b values under 130 µmol/g. Above this value, LIC_{SIR} has the lowest differences.

CONCLUSIONS: At 3T, R2* provides precise LIC quantification for lower overload but the SIR method is recommended to overcome R2* limitations in higher overload. Our software, available on mrquantif.org, uses jointly both methods and selects the best one.

KEYWORDS

Iron, Liver, Magnetic resonance imaging, Hemosiderosis, Non-alcoholic fatty liver disease, Hemochromatosis.

KEY POINTS

- Liver iron can be accurately quantified by MRI at 3T
- At 3T, R2* provides precise quantification of slight liver iron overload
- At 3T, SIR method is recommended in case of high iron overload
- Slight liver iron overload present in metabolic syndrome can be depicted.
- Treatment can be monitored with great confidence.

ABBREVIATIONS

SIR: Signal intensity ratio
LIC: liver iron concentration
LIC_b: LIC assessed by biopsy using biochemical analysis
LIC_{sIR}: LIC calculated by SIR method
LIC_{R2*}: LIC calculated by T2* conversion
mGRE: Multiple-echo gradient-echo sequence
NASH: Non-alcoholic steatohepatitis
DIOS: Dysmetabolic iron overload syndrome
MRI: Magnetic resonance imaging
BMI: Body mass index
AUC: Area under the curve

MAIN TEXT

Introduction

Liver iron content (LIC) is a surrogate marker of whole-body iron load. In overload diseases such as primary or secondary hemochromatosis, LIC measurement is mandatory for guiding therapeutic decisions. Liver iron overload may also be present in non-alcoholic steatohepatitis (NASH) and dysmetabolic iron overload syndrome (DIOS), which are both highly prevalent in the Western population (1). The main complications are cirrhosis and hepatocellular carcinoma. Many studies (2,3) have suggested a close correlation between iron deposition and carcinogenesis.

The gold-standard method for detecting and quantifying liver iron overload is histopathological analysis of a liver sample collected by biopsy with biochemical analysis of the core fragment. The biopsy procedure is both invasive and painful and carries some risk of complications (4). In addition, the very small liver sample may not be representative of the whole liver in cases of heterogeneous iron distribution (5).

Non-invasive, quantitative assessment of LIC by 1.5T magnetic resonance imaging (MRI) has been extensively validated against histology by calculating the relaxation rates R2 and R2* (6–11) and/or the signal intensity ratio (SIR) between the liver and paraspinal muscles (12–14). MRI is thus now used in routine clinical practice to diagnose, quantify and monitor iron overload (15).

In recent years, 3T MRI has become more widespread. In view of the shift in magnetic field strength, acquisition parameters need to be adapted and new reference values proposed.

Better sensitivity and accuracy can be expected at 3T, improving diagnosis of DIOS with low iron burden. Conversely, quantification of high overload cases may prove more difficult (16).

Recently, the SIR method, based on several single-echo GRE sequences, has been validated against histology at 3T (17).

The purpose of our study was to evaluate the ability of the R2* method to detect and quantify liver iron at 3T using biochemical quantification as the reference method. Our secondary goal was to compare, at the 3T field strength, two major LIC quantification methods: R2* and SIR.

Materials and Methods

Patient population

Between January 2007 and January 2013, all patients referred for liver biopsy and in whom liver iron overload was suspected according to their disease were prospectively recruited. All patients provided written informed consent to participate in this prospective single-center clinical trial. In addition to usual care, an MRI scan was scheduled to assess hepatic iron stores. Age, sex and body mass index were recorded.

Biochemical liver iron concentration

Liver biopsy was indicated as per the guidelines of the American Association for the Study of Liver Diseases (18,19). A biopsy sample was taken from the right lobe of the liver using a 16 gauge needle (Hepafix 16G, Braun, Melsingen, Germany) under ultrasound guidance. Biochemical liver iron concentration (LIC_b) was measured using Barry and Sherlock's method for biopsy samples taken from paraffin-embedded blocks (20). Liver iron overload was defined as an LIC_b greater than 35 µmol/g (dry liver). Biochemical analysis was blinded to MRI results.

Magnetic resonance imaging protocol

The study was performed with two 3T MRI scanners: first with Achieva (Philips, Best, Netherlands) and then with Magnetom Verio (Siemens Healthcare, Erlangen, Germany). The body coil was used as the receive coil to achieve homogeneous signal intensity in the imaged section and avoid signal depth fall-off. Only the Siemens scanner had a compensation method for better B1 homogeneity. There was a slight difference in resonance frequency (127.79 vs. 123.24 MHz) between the two scanners. Using the body coil, one multi-echo gradient echo (mGRE) sequence was performed, with 11 echoes. The selected TEs were slightly different depending on the scanner: a multiple of 1.15 ms for the Philips group and 1.23 ms for the Siemens group. Pixel bandwidth was 1161 Hz for the Philips group and

1048 Hz for the Siemens group. The remaining parameters were identical for both machines: 400x400mm² field of view; 128*121 acquisition matrix; 256x256 reconstruction matrix with a pixel size of 1.56x1.56 mm²; 120 ms repetition time; 20° flip angle; 7 mm slice thickness; 1 excitation. The breath-hold acquisition lasted 15s.

MRI data analysis

Measurements were conducted using an in-house Java program integrating ImageJ functions (NIH, Bethesda, USA). All data were analyzed by a radiologist (with 10 years' experience in abdominal radiology) who was blind to clinical information and to the biopsy result.

On the selected slice, 3 ROIs with a diameter of $2.5 \text{ cm} (4.9 \text{ cm}^2)$ were placed in the right liver area, taking care to avoid large vessels, biliary tracts, parenchymatous lesions and artifacts, 2 ROIs with a diameter of $2 \text{ cm} (3.1 \text{ cm}^2)$ in the right and left paraspinal muscles and 1 ROI with a diameter of $3 \text{ cm} (7.1 \text{ cm}^2)$ in the air outside of the body for noise measurement. All ROIs were automatically copied to the same place on each echo of this selected location. The placement of the ROIs is illustrated in Figure 1.

Before performing fitting, we applied a noise subtraction algorithm to subtract the mean background noise from the liver signal. Then T2* values were automatically calculated using a simplex non-linear algorithm to fit the magnitude of the complex signal from all echoes or only from in-phase echoes when the signal of the first out-of-phase echo was lower than the signal of the first in-phase echo.

Thus T2* was calculated according to the formula: Liver signal – noise = $M0.e^{-\left(\frac{TE}{T2*}\right)}$

R2* was calculated as follows: R2* = $1/T2^*$, and we used the linear correlation with LIC_b to determine LIC_{R2*}.

The liver-to-muscle signal intensity ratio (SIR) method was used to calculate LIC_{SIR} with the algorithm derived from the same patient series using 5 single-echo GRE sequences (17). Only the first four echoes of this formula were used to calculate LIC_{SIR} since the longest fifth echo (14 ms) was not obtained in the mGRE acquisition.

Statistical analysis

Statistical analyses were performed using SAS 9.4 (SAS Institute, Cary, NC).

Qualitative variables were expressed as numbers and percentages. Quantitative data were expressed as means \pm standard deviations (SD) if normally distributed and medians (Q1-Q3) if not normally distributed.

Given that LIC quantification variables were not normally distributed, we calculated non-linear correlation coefficients (Spearman) to estimate the strength of the linear relationship between LIC_{R2*} or LIC_{SIR} and LIC_{b} .

Similarly, in order to compare measurements using the Philips or Siemens scanner, Generalized Poisson Mixed Models (GLIMMIX procedure) were used with or without adjustment for sex, BMI and age.

Agreement between LIC quantifications was assessed using the Bland and Altman method, calculating the mean difference (estimated bias, d), the standard deviation of the differences (precision, SD), and the limits of agreement ($d\pm1.96$ SD). Student's t-test was used to determine whether the bias between measurement methods was significant.

Optimal cut-off values for the threshold of LIC_b at 36 μ mol/g were obtained by optimization of the Youden index from AUROC curve analysis.

The area under the curve, sensitivity, specificity, positive and negative predictive values were calculated for both LIC_{R2*} and LIC_{SIR} .

In order to compare the two methods at different levels, the cohort was divided into equal successive classes according to the values of LIC_b . Then, $LIC_b - LIC_{R2*}$ and $LIC_b - LIC_{SIR}$ were calculated and compared at the different LIC_b levels. A similar comparison, corresponding more to the practical intent to diagnose, was also done by using LIC_{R2*} classes.

P values less than 0.05 were considered statistically significant.

RESULTS

Patient population

One hundred and five patients were prospectively included between January 2007 and January 2013 (Figure 2). Fifty-eight (55%) had hyperferritinemia, 52 (50%) had metabolic syndrome, 15 (14%) had

chronic alcohol intoxication, 15 (14%) had either hepatitis B or C, and 6 (6%) had other liver diseases (autoimmune hepatitis, primary biliary cirrhosis).

The first 36 patients (34%) included were scanned with the Philips scanner and the following 69 (66%) with the Siemens scanner. Mean (\pm SD) age was 55.8 \pm 12.7 and 50 \pm 12.8 years for women and men, respectively (p=0.03). One hundred and one patients underwent the MRI examination and the biopsy on the same day while 4 patients experienced an interval between the biopsy and MRI of less than 15 days. The LIC_b concentration ranged from 0 to 630 µmol/g, and 49 patients (47%) had normal LIC_b values < 36 µmol/g. Fifty six patients had a liver iron overload. It was due to genetic hemochromatosis in 31 patients (LIC_b mean=286 \pm 148 µmol/g, range=43-630 µmol/g), to dysmetabolic syndrom in 22 patients (LIC_b mean=62 \pm 27 µmol/g, range=36-123 µmol/g) and to other causes (alcoholic or viral hepatitis) in 3 patients (LIC_b mean= 42 \pm 5 µmol/g, range=36-46 µmol/g). Patient characteristics are provided in Table 1.

Our analysis without / with the adjustment for BMI, sex and age yielded no difference in the distribution of LIC_b (p=0.65 / p=0.19), LIC_{R2*} (p=0.49 / p=0.14) and LIC_{SIR} (p=0.50 / 0.27) results between the two groups using MRI machines from different manufacturers.

R2* and LIC_{R2*} measurements

Linear regression between LIC_b and R2* is shown in Figure 3a and yielded the following equation:

$$LIC_{R2*}$$
 (µmol/g) = 0.316 R2* +7.6

The Spearman correlation coefficient (r = 0.95 p < 0.001) indicates a strong positive correlation between LIC_b and R2^{*}.

Figure 3b shows the Bland-Altman plot of the difference vs. mean values of LIC_b and LIC_{R2*} measurements. The bias (SD) or average difference between the results of the two methods was 7.6 (73.4) µmol/g and the 95% limits of agreement were -136.4 µmol/g and 151.5 µmol/g. The bias was not statistically significantly different to zero (p= 0.74).

With the reference threshold established at $LIC_b = 36 \mu mol/g$, ROC curves obtained with LIC_{R2*} results

showed an area under the curve (AUC) of 0.987. The best threshold was given for LIC_{R2*} at 32 µmol/g, corresponding to an R2* of 77 s⁻¹, and a T2* of 13 ms, with 47 true positives, 4 false positives, 52 true negatives and 2 false negatives. The sensitivity was 0.96 (95% CI: 0.9; 1.01) and the specificity 0.93 (95% CI: 0.86; 1.0).

Taking in consideration only the 76 patients with LIC_b below 130 μ mol/g, linear regression yielded the following equation:

$$LIC_{R2*} (\mu mol/g) = 0.314 R2* - 0.96$$

The best threshold was then given for LIC_{R2*} at 27 µmol/g, corresponding to an R2* of 89 s⁻¹, and a T2* of 11 ms, with 44 true positives, 2 false positives, 25 true negatives and 5 false negatives. The sensitivity was 0.90 (95% CI: 0.81; 0.98) and the specificity 0.93 (95% CI: 0.83; 1.0).

LIC_{SIR} measurement

Linear regression between LIC_b and LIC_{SIR} is shown in Figure 3c. The Spearman correlation coefficient (r = 0.92 p < 0.001) indicates a strong positive correlation between LIC_{SIR} and LIC_b . Figure 3d shows the Bland-Altman plot of the difference vs. mean values of LIC_b and LIC_{SIR} measurements. The bias (SD) or average difference between the results of the two methods was 14.8 (37.6) and the 95% limits of agreement were -59.0 and 88.5 µmol/g. The bias was statistically significantly different to zero (p<0.0001).

With the reference threshold established at $LIC_b = 36 \,\mu mol/g$, the ROC curves obtained with LIC_{SIR} results showed an AUC of 0.965. The best threshold was given for $LIC_{SIR} = 20 \,\mu mol/g$ with 45 true positives, 3 false positives, 53 true negatives and 4 false negatives. The sensitivity was 0.92 (95% CI: 0.84; 0.99) and the specificity 0.95 (95% CI: 0.89; 1.0).

Comparison between LIC_{R2*} and LIC_{SIR} measurements

The Spearman correlation coefficient (r = 0.95 p < 0.001) indicates a strong positive correlation between LIC_{R2*} and LIC_{SIR} . Figure 4 shows the mean differences (absolute values) between LIC_{SIR} and LIC_b or between LIC_{R2*} and LIC_b according to LIC_b or LIC_{R2*} class. With LIC_b classes, LIC_{R2*} presented the lowest differences for patients with LIC_b values under 130 µmol/g and the highest differences for patients above 190 µmol/g (Figure 4a). Using LIC_{R2*} classes, the differences increased above 130 µmol/g (Figure 4b).

DISCUSSION

With a shortest TE of 1.2 ms, liver iron overload can be reliably quantified by MRI at 3T with the R2* for patients with biopsy-proven LIC under 130 μ mol/g but the SIR method appears more robust for higher iron overload.

The R2* calculation is well known and its clinical use is well established at 1.5T. In the literature, there are 5 main publications, validating R2* against LIC determined by biopsy (6,8–11). Conversion formulas have been proposed to estimate LIC from R2* (s⁻¹) with a slope of 0.025 to 0.032 to obtain the LIC value in mg/g. Pooling the data from the main publications, Henninger found a mean slope of 0.029 (11). Then, to obtain the LIC in µmol instead of mg/g, we multiplied this mean slope by 18 to obtain 0.52. So, at 1.5T, simply by dividing by 2 the value of R2* expressed in s⁻¹ we have a correct approximation of LIC expressed in µmol/g.

No such validation with biopsies has been done at 3T. Theoretical calculations suggest a doubling of R2* from 1.5 to 3T (21). Then the mean slope to obtain the LIC value in μ mol/g should be divided by 2 and should be approximately 0.26. Anwar's (22) results in 5 patients seem to confirm this hypothesis but with significant delay between MRI and biopsy. However, in our series we obtained a slope of 0.316, slightly higher than the slope expected by extrapolation of 1.5T polled data but close to half the higher slope proposed at 1.5T by Garbowski, who used the same laboratory reference (10). In our series, the background noise subtraction leads to a higher value of R2* and partly explains the residual difference with Garbowski's results. This emphasizes the need for a standardized protocol to obtain more comparable results.

The higher magnetic susceptibility observed at 3T introduced limitations in the R2* calculation. For high overloads, there is a strong decrease in liver signal intensity. It is then difficult to obtain a correct exponential curve fit.

The SIR method is also widely recognized and used for hepatic iron quantification at 1.5T. Our study evaluated this method at 3T using the algorithm defined from single-echo sequences (17). The results we obtained with an mGRE sequence showed good correlations but with a slight overall overestimation and a slight underestimation for low values because the longest TE, around 14 ms, was not included in the mGRE sequence. For slight to moderate overloads, below 130 μ mol/g, almost exclusively patients with DIOS, our study showed a better correlation of the R2* method than the SIR method to LIC_b. However, in patients with high LIC_b above 130 μ mol/g, corresponding exclusively, in our study, to patient with genetic hemochromatosis, the SIR method provides a better correlation to LIC_b. At 1.5T, quantification was possible by SIR up to 350 μ mol/g by using the shortest in-phase TE of 4 ms. Rose overcomes this limit by using a shorter first TE of 1.8 ms (23). At 3T, a first TE of 1.2 ms is short enough to give a liver signal over the signal noise and to allow a SIR estimation in high overload.

Our study is the largest series calibrating R2* versus LIC_b, for any magnetic field strength. It validates the use of 3T MRI for hepatic iron quantification. In comparison to the biopsy with biochemical determination of iron, we propose a formula to convert R2* at 3T to LIC_b. Despite variation in technical characteristics, there was no significant difference between the two machines used. Although the use of 3T MRI is becoming more widespread, some centers only have a 3T magnetic field for abdominal imaging. There is a strong need for reference values at 3T. Moreover, the use of a 3T magnetic field allows for more accurate quantification of slight to moderate overloads. Improving sensitivity is clinically relevant regarding the increasing incidence of DIOS with low iron overload.

Our study has certain limitations. First, the shortest TE was about 1.2 ms, a value which is also the first TE usually proposed by MR vendors in most built-in protocols dedicated to hepatic iron and fat quantification. Obviously, this TE is not short enough at 3T to correctly calculate R2* in the case of

high overload. It is technically difficult to use a first TE of 0.4 ms, which is half the shortest TE of 0.8 ms proposed by Wood at 1.5T (8). Very short TEs will be available using ultrashort echo time (UTE) imaging (24). In the meantime, the main risk is not being unable to quantify correctly a high overload, which has only a small impact on patient management, but miscalculating R2* and hence underestimating liver iron overload. For example a patient with an R2* of 512 s⁻¹, corresponding to an LIC_{R^*} of 130 µmol/g, actually had an LIC_b of 480 µmol/g. This type of error explains how the difference between LIC_{R2*} and LIC_b increases faster with LIC_{R2*} classes than with LIC_b classes. So, to overcome this limitation, we propose either greatly reducing the shortest TE or combining both SIR and T2* methods. Second, we used two different machines with a slight magnetic field difference (3%). Acquisition parameters were as close as possible. However, there were also slight TE differences (8%). This could have produced errors particularly for the SIR method which does not take into account TE differences between the two units. The absence of B1 heterogeneity correction with the first machine may also lead, in some cases, to an overestimation of LIC_{SIR} through reduction of the paraspinal muscle signal, as described with single-echo sequences (17). Third, we used the body coil for both methods. This coil is necessary for the SIR method. A surface coil allows a higher signal for R2* calculation but this is offset by larger voxels (17mm³) and T2* fitting to the entire ROI instead of producing a pixel-wise map. Fourth, we only use 4 of the 5 echoes used by the 3T SIR algorithm based on single-echo sequences. This explains the bias observed for the low values of LIC_b with a LIC_{SIR} cut-off of 20 µmol/g for determining overloaded patients. A new version of the algorithm taking into account the reduction in the number of echoes obtained has now been incorporated into our dedicated software. Nevertheless, this has no practical impact since at that level of overload R2* is the most precise method.

This study validates hepatic iron quantification by MRI at 3T, with a conversion formula to LIC_b obtained from biopsy material. With the selected TEs, the R2* method is more accurate for slight to moderate hepatic iron overload whereas the SIR method is more accurate for high overloads. Shorter TEs are needed to improve performance for quantifying massive iron overload by R2* [24]. In the meantime, both methods should be used simultaneously with a breath-hold mGRE sequence acquired

using the body coil. The sequence protocol we propose can be applied to the majority of MRI scanners without the need to purchase a specific option. Detailed sequence parameters and a dedicated DICOM software program, incorporating both calculations with cross-checks, are available at www.mrquantif.org

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TABLES

Table 1: Patient characteristics (n=105)	
Male/Female	68/37
Mean age (SD)	52.1 (+/-13)
MRI scanner manufacturer (Siemens/Philips)	69/36
Body Mass Index, mean (SD)	27.8 (4.2)
LIC _b µmol/g, median [interquartile range]	37.5 [18.2 - 130.1]
LIC _{SIR} µmol/g, median [interquartile range]	30 [0 - 120]
$LIC_{R2*} \mu mol/g$, median [interquartile range]	32.9 [18.5-117.7]

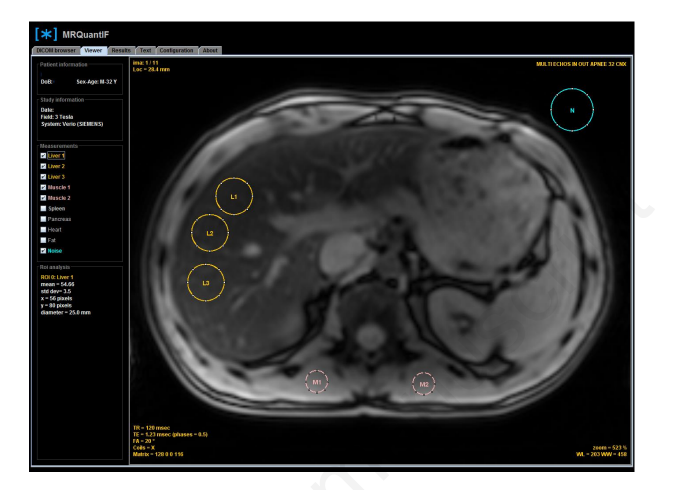
FIGURE CAPTIONS

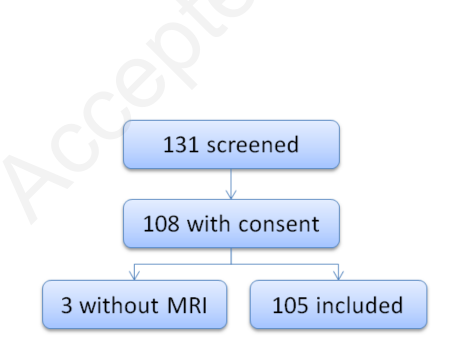
Figure 1: Screen-copy of the in-house software viewer with example of ROIs placement.

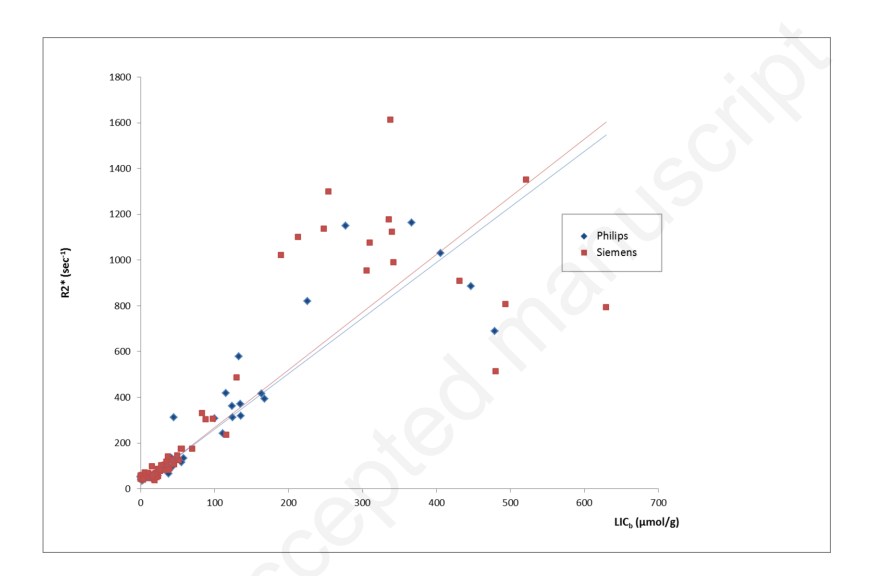
Figure 2: Flow chart.

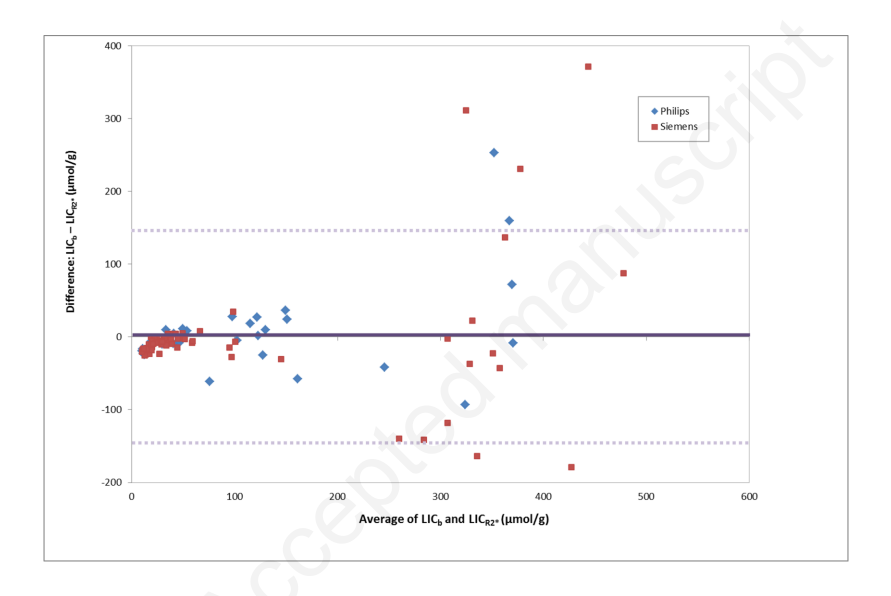
Figure 3: Comparison between LICb and R2* or LIC_{SIR} : a) Linear regression between R2* and LIC_b , b) Bland-Altman plot of the difference vs. average of LIC_{R2*} , c) Linear regression between LIC_{SIR} and LIC_b , and d) Bland-Altman plot of the difference vs. average of LIC_{SIR} in comparison to LIC_b

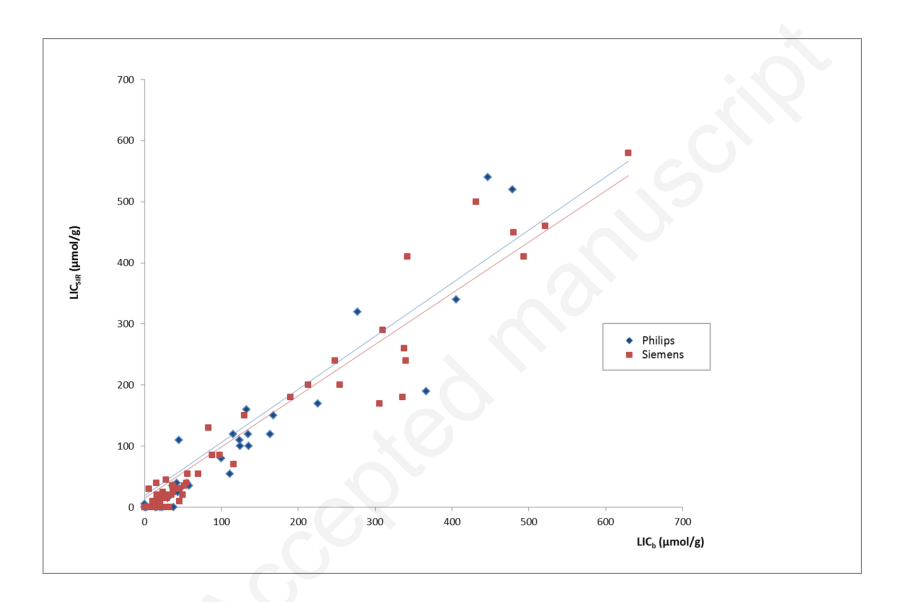
Figure 4: Mean of absolute differences between LIC_b and LIC_{R2*} or LIC_{SIR} according to a) (Figure 5a) LIC_b classes or b) (Figure 5b) LIC_{R2*} classes

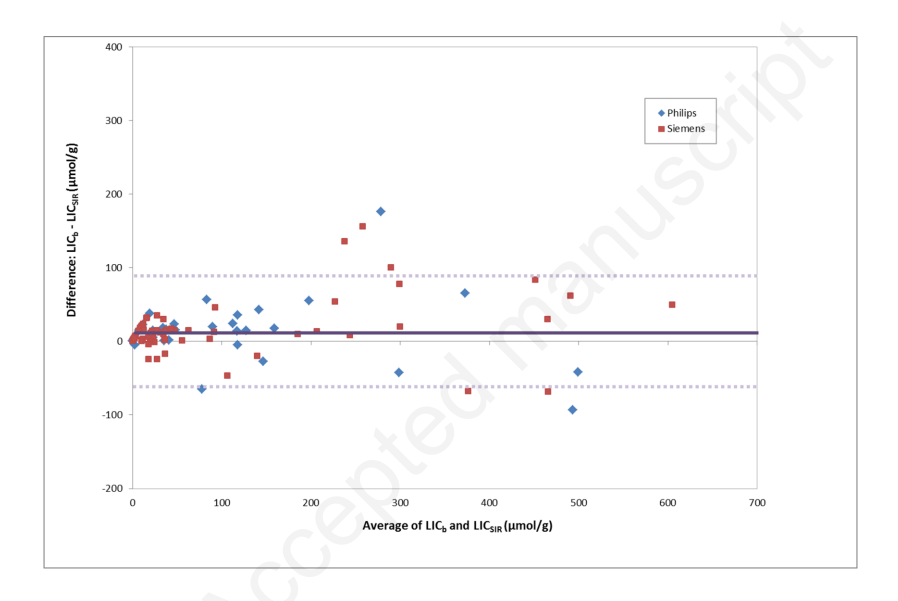


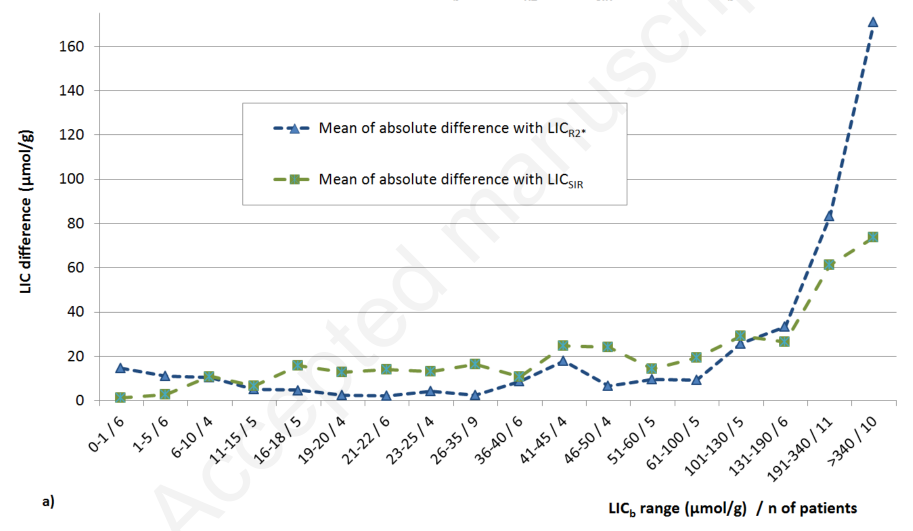




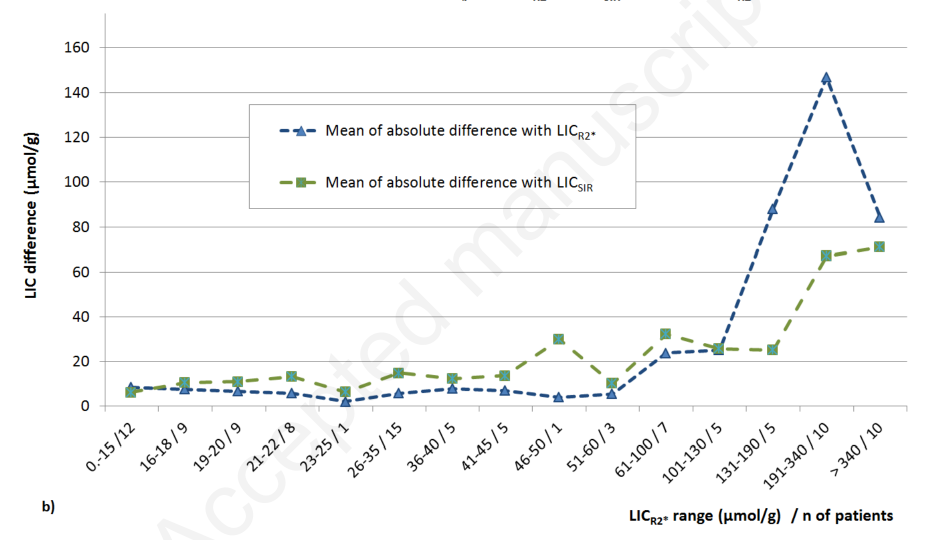








Absolute difference mean between LIC_{b} and LIC_{R2*} or LIC_{SIR} , for each LIC_{b} class



Absolute difference mean between LIC_{b} and LIC_{R2*} or LIC_{SIR} , for each LIC_{R2*} class

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