

Conditional safety margins for less conservative peak local SAR assessment: A probabilistic approach

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Purpose: The introduction of a linear safety factor to address peak local specific absorption rate ($pSAR_{10g}$) uncertainties (eg, intersubject variation, modeling inaccuracies) bears one considerable drawback: It often results in over-conservative scanning constraints. We present a more efficient approach to define a variable safety margin based on the conditional probability density function of the effectively obtained $pSAR_{10g}$ value, given the estimated $pSAR_{10g}$ value.

Methods: The conditional probability density function can be estimated from previously simulated data. A representative set of true and estimated $pSAR_{10g}$ samples was generated by means of our database of 23 subject-specific models with an 8-fractionated dipole array for prostate imaging at 7 T. The conditional probability density function was calculated for each possible estimated $pSAR_{10g}$ value and used to determine the corresponding safety margin with an arbitrary low probability of underestimation. This approach was applied to five state-of-the-art local SAR estimation methods, namely: (1) using just the generic body model “Duke”; (2) using our model library to assess the maximum $pSAR_{10g}$ value over all models; (3) using the most representative “local SAR model”; (4) using the five most representative local SAR models; and (5) using a recently developed deep learning-based method.

Results: Compared with the more conventional safety factor, the conditional safety-margin approach results in lower (up to 30%) mean overestimation for all investigated local SAR estimation methods.

Conclusion: The proposed probabilistic approach for $pSAR_{10g}$ correction allows more accurate local SAR assessment with much lower overestimation, while a pre-defined level of underestimation is accepted (eg, 0.1%).

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KEYWORDS

parallel transmit, safety factor, SAR model library, SAR model selection, specific absorption rate, subject-specific local SAR assessment

1 | INTRODUCTION

Ultrahigh-field MRI (UHF-MRI) provides strong potential to achieve superior image quality compared with the current clinical systems at lower field strengths.¹⁻³ To address problems with B_1 homogeneity, often local multitransmit coil arrays are used.⁴⁻⁸ However, their application is restricted by the limits of the current methods for local specific absorption rate (SAR) assessment.^{9,10}

Multitransmit arrays produce a great variability of the electric field, and thereby of the absorbed power by the tissues,¹¹⁻¹⁷ making the local SAR difficult to predict, which contributes to making the local SAR limits more restrictive than the global SAR limits (as described in IEC 60601-2-33).¹⁸

The local SAR cannot be measured during an MRI examination and is usually evaluated by numerical simulations.⁹⁻¹⁷ New software tools to perform on-line simulations using patient-specific body models are being developed but are still quite time-consuming.¹⁹ Therefore, typically the electric-field distribution of each array element is simulated off-line on one or several generic models. After domain reduction by virtual observation points,²⁰⁻²² the peak 10g average SAR ($pSAR_{10g}$) for a given drive setting of the array is calculated on-line at the scanner.

Despite the previously mentioned progress, the resulting predicted SAR can still deviate from the true peak local SAR value in the patient being scanned. This is due to a variety of uncertainties in the actual examination setup. Indeed, even assuming that the reflected/lost power is properly monitored, and that there are no calibration errors, the used body model and its position within the MRI system could be very different compared with the patient under examination. To address this, a library of models can be used to cover a large patient population.^{10-12,17,23-26} Nevertheless, the residual uncertainties may still result in peak local SAR overestimation or underestimation error.

Although the overestimation error results in unnecessarily long scan times and/or suboptimal image quality, only peak local SAR underestimation error poses a safety risk. For this reason, to diminish the probability of underestimation, a linear safety factor is usually applied to increase the estimated peak local SAR value to such an extent that underestimation will not occur.^{10,12,23,24} This increased estimated peak local SAR level will be referred to as the “corrected” peak local SAR value.

Recently, an alternative deep learning-based method for subject-specific SAR estimation was presented.²⁷ This data-driven approach consists of training a convolutional neural

network to map the relation between subject-specific complex B_1^+ maps and the corresponding local SAR distribution. However, like the aforementioned methods, this method also suffers from residual underestimation errors that need to be addressed by applying a suitable safety factor.

As this study will show, the conventionally applied linear safety factor often results in unnecessary very conservative estimation, in particular when high peak local SAR value is estimated. Therefore, we propose an alternative approach to diminish the probability of peak local SAR underestimation based on the conditional probability density function of the true peak local SAR value, given the estimated peak local SAR value. This approach allows us to define a variable safety margin for each possible estimated peak local SAR value.

This approach is applied to five state-of-the-art local SAR estimation methods: local SAR prediction based on (1) one generic model,²⁵ (2) the largest value in a large database of models,^{11,12,17} (3) the model that fits best the subject in the scanner,²⁸ (4) a combination of methods 2 and 3, and (5) a deep learning method.²⁷

In the present work, these methods are applied to assess the peak local SAR with a multitransmit array of eight fractionated dipole antennas for prostate imaging at 7 T.^{29,30} For each method, the mean peak local SAR overestimation is evaluated for the conventional approach using one linear safety factor and the proposed approach, based on the probability density function. The results show that the proposed approach is better able to deal with the remaining uncertainty and reduces peak local SAR overestimation with respect to all other investigated local SAR estimation methods. In addition to the introduction of the conditional safety margin (CSM) approach based on the probability density function, this study presents a comparative evaluation of five potential local SAR assessment techniques that were described previously.

2 | THEORY

The introduction of a linear safety factor to avoid peak local SAR ($pSAR_{10g}$) underestimation bears one considerable drawback: If the estimated $pSAR_{10g}$ level is high, then the resulting corrected $pSAR_{10g}$ level shows severe overestimation of the true $pSAR_{10g}$ level. This is depicted in Figure 1. This figure shows for a certain transmit array configuration how estimated $pSAR_{10g}$ ($pSAR^E$) and truly obtained $pSAR_{10g}$ ($pSAR^T$) are related. In particular, each point in the scatter plot represents one out of 5750 $pSAR_{10g}$ estimations by the deep learning method

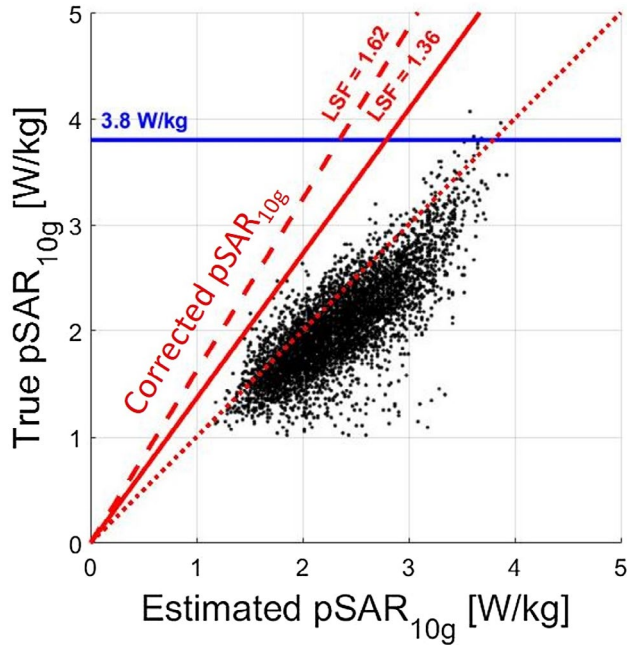


FIGURE 1 Scatter plot of the true peak local specific absorption rate ($pSAR_{10g}$) versus the estimated $pSAR_{10g}$ for 7T prostate imaging using a wide variety of random phase settings and 23 human models. The estimated $pSAR_{10g}$ is the $pSAR_{10g}$ value as determined by the deep learning-based method²⁷ using simulated B_1^+ distributions as input. The true $pSAR_{10g}$ is given by the simulated $pSAR_{10g}$ value. The diagonal red dotted line indicates where the estimated $pSAR_{10g}$ and true $pSAR_{10g}$ values are equal. The dashed red line denotes the corrected $pSAR_{10g}$ using the linear safety factor (LSF), defined to avoid underestimation errors in all cases evaluated (LSF = 1.62). The solid red line is the corrected $pSAR_{10g}$ by the LSF defined by removing the outliers (LSF = 1.36). The blue line is the 99.9% certain $pSAR_{10g}$ upper bound (3.8 W/kg) for random phase settings

with $8 \times 1W$ input power and random phase shimming. The diagonal red dotted line indicates where the estimated $pSAR_{10g}$ levels and true $pSAR_{10g}$ levels are equal. Points above the diagonal represent true $pSAR_{10g}$ levels larger than the corresponding estimate. The scatter plot clearly shows that there is a good correlation between estimated $pSAR_{10g}$ level and true $pSAR_{10g}$ level. However, the estimated $pSAR_{10g}$ level needs to be corrected to avoid underestimation, as there are many points above the dotted line. A linear safety factor of 1.62 is therefore determined to correct (increase) the estimated $pSAR_{10g}$ level, such that all of the potential true $pSAR_{10g}$ levels are lower than the corrected $pSAR_{10g}$ level.

The red dashed line shows how each estimated $pSAR_{10g}$ level on the horizontal axis results in a corrected $pSAR_{10g}$ ($pSAR^{E,C}$) level. If the correction factor is chosen appropriately, all points will be below the dashed red line. As Figure 1 shows, this condition is fulfilled.

However, the figure also clearly illustrates that for large estimated $pSAR_{10g}$ levels, the application of the linear safety factor results in corrected $pSAR_{10g}$ levels that have far too high overestimation. The overestimation could be reduced by

determining the safety factor using the upper outer fence.²⁷ This will make the safety factor less sensitive to outliers and/or small deviations in estimated or true $pSAR_{10g}$ of the one point that determines its slope. Then, using the upper outer fence definition,²⁷ the linear safety factor will be 1.36, as indicated by the solid red line in Figure 1. The figure shows that more than 99.9% of the points are below the line.

Unfortunately, for large estimated $pSAR_{10g}$ levels, the corrected $pSAR_{10g}$ level is still much larger than what realistically could be expected. In fact, a previous study¹² has shown that the $pSAR_{10g}$ level that is not exceeded for 99.9% of the cases (over all 23 models and any potential phase setting) is 3.8 W/kg. Therefore, a solution could be to combine these limits that use the corrected $pSAR_{10g}$ value if the value is lower than 3.8 W/kg, and set it equal to 3.8 W/kg otherwise (blue line in Figure 1).

However, from a mathematical perspective, this approach is arguably not the most appropriate solution to address this problem. Instead, a probabilistic approach should be followed. In a probabilistic setting, the scatter plot in Figure 1 represents samples from the joint probability distribution of estimated and true $pSAR_{10g}$ values, which is described by the probability density function $f_{E,T}(pSAR^E, pSAR^T)$. When a $pSAR^E$ value is estimated, we would like to know the probability of $pSAR_{10g}$ underestimation: $P(pSAR^T > pSAR^E)$. Thus, one needs to know the conditional probability density function $f_{T|E}(pSAR^T | pSAR^E)$, which describes the probability distribution of $pSAR^T$ for a given $pSAR^E$ value. This function is calculated in Eq. 1, where $f_E(pSAR^E)$ is the marginal probability density function that describes the probability distribution of estimating the $pSAR^E$ value regardless of $pSAR^T$ (Figure 2).

$$f_{T|E}(pSAR^T | pSAR^E) = \frac{f_{E,T}(pSAR^E, pSAR^T)}{f_E(pSAR^E)} \quad (1)$$

Now the probability of $pSAR_{10g}$ underestimation ($P_{underest}$) (ie, the probability that the true value $pSAR^T$ is actually larger than some estimated value $pSAR^E = E_1$) is given as

$$P_{underest} = P(pSAR^T > E_1 | pSAR^E = E_1) = \int_{E_1}^{\infty} f_{T|E}(pSAR^T | pSAR^E = E_1) d(pSAR^T). \quad (2)$$

Therefore, the conditional probability density function can be calculated for the estimated $pSAR^E = E_1$ value, and the integral can be numerically evaluated to determinate a corrected $pSAR^{E,C}$ value with probability of underestimation equal to an arbitrary (small) ϵ :

Find $pSAR^{E,C}$ such that:

$$P(pSAR^T > pSAR^{E,C} | pSAR^E = E_1) = \int_{pSAR^{E,C}}^{\infty} f_{T|E}(pSAR^T | pSAR^E = E_1) d(pSAR^T) = \epsilon. \quad (3)$$

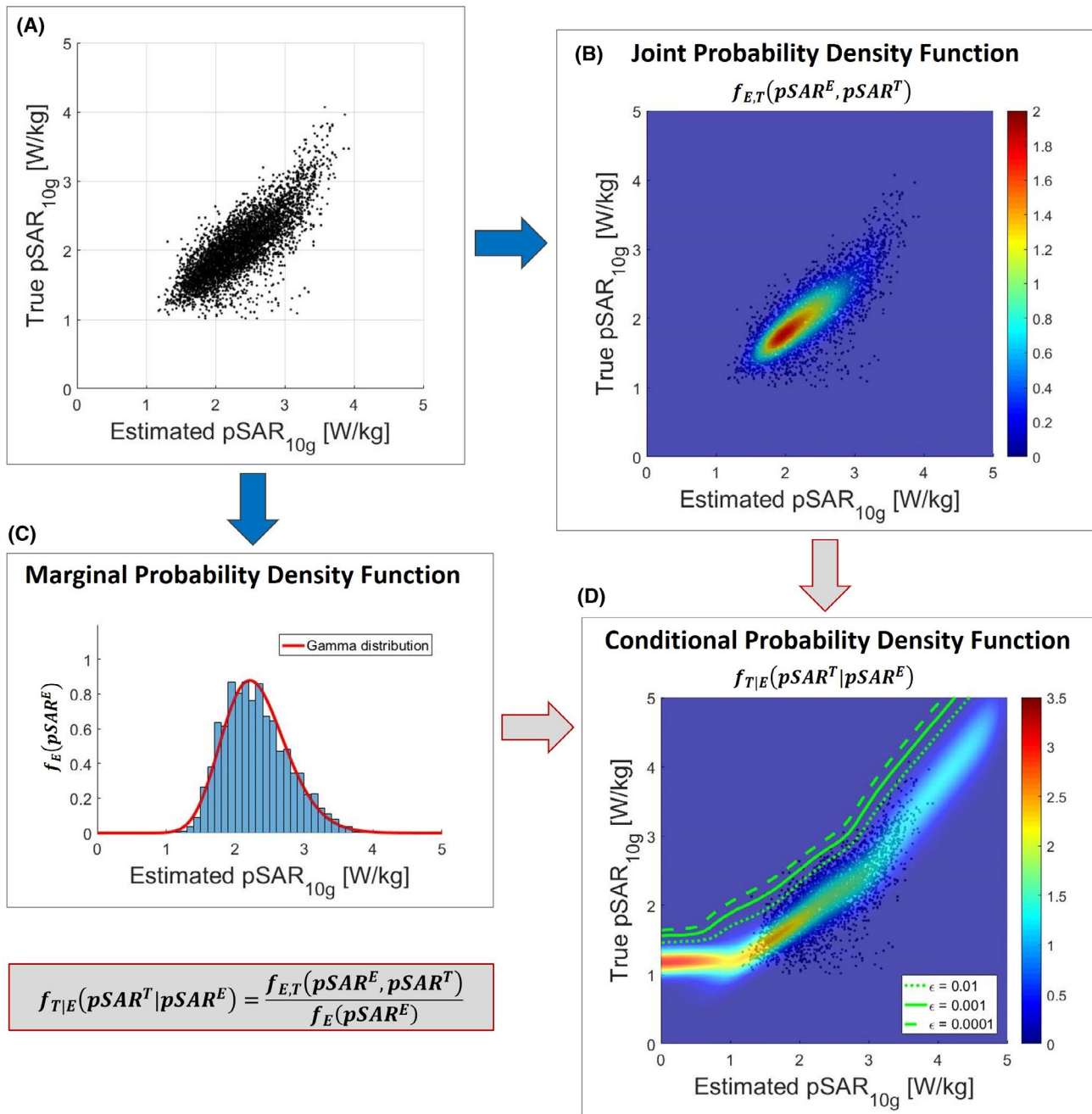


FIGURE 2 A, Scatter plot of the true pSAR^T versus the estimated pSAR^E using the deep learning method. B, Joint probability density function $f_{E,T}(pSAR^E, pSAR^T)$ of the estimated and true pSAR values. C, Marginal probability density function $f_E(pSAR^E)$ of the estimated pSAR values. D, Two-dimensional conditional probability density function $f_{T|E}(pSAR^T|pSAR^E)$ obtained by combining the conditional probability density functions for each possible pSAR^E value

This corrected pSAR^{E,C} value effectively results in a CSM that depends on the estimated pSAR_{10g} level and that can be determined for each possible estimated pSAR^E value.

Note that, from the observed data, a variety of approaches can be used to estimate the underlying probability density function,³¹ including kernel density estimator,³² histogram,³³ and mixture models.³⁴

3 | METHODS

The exact extent of reduction of overestimation using the proposed approach depends on the method of pSAR_{10g} estimation. In this study, the pSAR_{10g} values were predicted using five state-of-the-art local SAR estimation methods. Then, for each estimation method, the corrected pSAR_{10g}

values were obtained by applying the linear safety factor and the proposed approach based on probability density function.

To build the required set of ($pSAR^T$, $pSAR^E$) samples for the purpose of this study, our database of 23 subject-specific models¹² with an 8-channel transmit body array for 7T prostate imaging^{29,30} was used. For each model and every array channel, finite-difference time-domain simulations were performed (Sim4Life; ZMT, Zürich, Switzerland), and the results were processed to obtain the true $pSAR_{10g}$ and the estimated $pSAR_{10g}$ by each method, when the transmit array is driven with different drive vectors.

Three driving modes were considered:

- Random phase settings: Drive vectors with uniform amplitude ($8 \times 1W$ input power) and random relative phase settings with respect to the first channel (uniform distribution: $\mathcal{U}(-180^\circ, 180^\circ)$). This driving mode allows one to assess the performances during an arbitrary imaging examination in the lower abdomen region.
- Prostate shimmed phase settings: Drive vectors with uniform amplitude ($8 \times 1W$ input power) and relative phase normally distributed around the average prostate phase shimming set¹² $\mathcal{N}(\mu=0, \sigma=33^\circ)$. This driving mode allows one to assess the likely performances during a prostate imaging examination.
- Random amplitudes and phases settings: Drive vectors with random amplitude ($\mathcal{U}(0,1)$) and random relative phase settings ($\mathcal{U}(-180^\circ, 180^\circ)$). This driving mode allows one to assess the performances even in the case of sophisticated RF pulse design strategies (eg, SPINS.⁵ k_T -points⁶). Two scenarios were investigated: one in which the drive vectors were normalized to 1W maximum input power per channel, and one in which the drive vectors were normalized to 8W total input power.

3.1 | Peak local SAR correction

3.1.1 | Linear safety factor

For each model, 250 different drive vectors were used to calculate the true and the estimated $pSAR_{10g}$ values. Then, for each $pSAR_{10g}$ estimation method, 5750 (23×250) validation sets are used to determine the required linear safety factor using the upper outer fence method²⁷ as described in the section 2.

The safety factor application can produce very highly corrected $pSAR_{10g}$ values, particularly for high estimated $pSAR_{10g}$ levels. In our previous study,¹² based on the probability distribution of the true $pSAR_{10g}$, we defined a 99.9% certain $pSAR_{10g}$ upper bounds of 3.8 W/kg and 3.2 W/kg for

random phases and prostate shimmed phases, respectively. In the same way, we can define a 99.9% certain $pSAR_{10g}$ upper bound of 2.6 W/kg and 6.8 W/kg for random amplitudes and phase settings normalized to 1W maximum input power per channel and 8W total input power, respectively. Therefore, as explained in section 2, we can further reduce overestimation of the linear safety factor approach by combining it with these $pSAR_{10g}$ upper bounds, when higher corrected $pSAR_{10g}$ values are obtained (Figure 1, blue line).

3.1.2 | Conditional safety margin

We assumed that the 23 models are representative of the entire patient population. Then, for each $pSAR_{10g}$ estimation method, the pairs-estimated and true $pSAR_{10g}$ values ($pSAR^E$, $pSAR^T$) are samples from the joint probability density function $f_{E,T}(pSAR^E, pSAR^T)$ (Figure 2A) that can be modeled as a Gaussian mixture (Figure 2B). The estimated $pSAR_{10g}$ values are samples from the marginal probability density function $f_E(pSAR^E)$ and follow what appears to be a gamma distribution (Figure 2C).

Therefore, the conditional probability density function can be calculated, and the CSM with an arbitrary small probability of underestimation ϵ ($\epsilon=0.001$ in our study) can be determined using Equation 3 (Figure 2D).

For the practical implementation of the proposed approach, the domain $f_{E,T}(pSAR^E, pSAR^T)$ is discretized (eg, $\Delta(pSAR^E)$, $\Delta(pSAR^T)=0.01$ W/kg) and the estimated statistical models are used to obtain the 2D array of $f_{E,T}(pSAR^E, pSAR^T)$ values and the 1-dimensional array of $f_{E,T}(pSAR^E)$ values. Then, from the ratio between each column of the first array and the corresponding value of the second array, the 2D array of $f_{TE}(pSAR^T|pSAR^E)$ values are determined. Each column of the obtained array represents the conditional probability density function for the corresponding $pSAR^E$ value, and the CSM is determined by integrating numerically along the column until the required probability of underestimation is reached (the integral along the entire column will be equal to 1). Finally, the obtained CSMs for the discretized $pSAR^E$ values are interpolated on-line to determine the CSM for any possible estimated $pSAR^E$ value.

3.2 | True peak local SAR

To obtain the true $pSAR_{10g}$ value, for each model m and each drive setting s , the 10g-averaged Q -matrices (Q_{10g})^{20,21} are calculated as follows:

$$pSAR_{m,s}^T = \max_r (s^H Q_{10g}(r)_m s), \quad (4)$$

where r is the spatial location of each Q_{10g} matrix.

3.3 | Peak local SAR estimation methods

For each model m and each drive setting s , the corresponding $pSAR_{10g}$ values are estimated using five different methods.

3.3.1 | Generic body model

A very common approach consists of performing off-line electromagnetic simulations using a generic body model. Then, assuming that the investigated model is representative for the current subject, on-line $pSAR_{10g}$ estimation based on the actual drive scheme can be performed.

To assess the performance of this method, the generic model “Duke” of the Virtual Family²⁵ with 77 tissues (version 3.0; voxel resolution $0.5 \times 0.5 \times 0.5 \text{ mm}^3$) is used as follows:

$$pSAR_{m,s}^{E,GM} = \max_{\mathbf{r}} (s^H \mathbf{Q}_{10g}(\mathbf{r})_{Duke} \mathbf{s}). \quad (5)$$

3.3.2 | Model library

To cover an entire population of patients, a model library can be used to predict the maximum $pSAR_{10g}$ over all models.^{11,12,17} Therefore, for each model m , and each drive setting s , the maximum $pSAR_{10g}$ over the other models is determined as reported in Equation 6:

$$pSAR_{m,s}^{E,ML} = \max_{k \neq m} (pSAR_{k,s}^T). \quad (6)$$

3.3.3 | Model selection

The estimation error might be reduced using a large database of models and selecting the most representative local SAR model for the patient under examination.¹⁷ However, in our previous work,¹² no relationship was found between measurable body features and $pSAR_{10g}$. Therefore, as presented in our preliminary study,²⁸ for each model m we consider the model n , which shows the most similar worst-case local SAR distribution with uniform amplitude SAR_{10g}^{WoC} (ie, for each voxel, the maximum SAR level that could possibly be achieved with $8 \times 1W$ input power and worst-case phase settings),³⁵ as its most representative local SAR model. Indeed, the SAR_{10g}^{WoC} distribution contains both patient anatomy information and electric-field information (transmit-array information). In particular, we select the model n , which presents the SAR_{10g}^{WoC} distribution that minimized the RMS error with respect to the registered SAR_{10g}^{WoC} distribution of the model m (Supporting Information Figures S1-S3 report the SAR_{10g}^{WoC} distributions, the RMS error matrix, and the most representative local SAR models). Then, for each drive setting s , the $pSAR_{10g}$ value is estimated using model n as follows:

$$pSAR_{m,s}^{E,MS} = pSAR_{n,s}^T. \quad (7)$$

3.3.4 | Multiple models selection

As results will show, the model library method results in large mean overestimation, while the model selection method has large underestimation errors. For these reasons, a good compromise could be the selection of multiple models.

Using the same model selection approach of the previous method, for each model m , the group of five most representative local SAR models A_m is identified (Supporting Information Table S1), and the maximum $pSAR_{10g}$ of these five models is used to determine the performance of the multiple models selection method as follows:

$$pSAR_{m,s}^{E,MMS} = \max_{k \in A_m} (pSAR_{k,s}^T). \quad (8)$$

3.3.5 | Deep learning-based method

The deep learning-based method is a new image-based method.²⁷ In this data-driven approach, a convolutional neural network is trained to learn a “surrogate SAR model” to map the relation between subject-specific complex B_1^+ maps and the corresponding local SAR distribution.

Therefore, for each model m and each drive setting s , the simulated B_1^+ maps of each channel i are processed to produce the shimmed B_1^+ map as follows:

$$B_1^+(\mathbf{r})_{m,s} = \sum_{i=1}^{Nch.} B_1^+(\mathbf{r}, i)_m s(i), \quad (9)$$

where $Nch. = 8$ is the number of transmit channels. Then, with the obtained complex B_1^+ maps, realistic synthetic MR images are generated (magnitude and relative transmit phase maps with noise)²⁷ and used to produce the input data for the trained convolutional neural network to infer the corresponding SAR_{10g} distribution and determine the peak value, as follows:

$$SAR_{10g}^{DL}(\mathbf{r})_{m,s} = CNN(B_1^+(\mathbf{r})_{m,s}) \quad (10)$$

$$pSAR_{m,s}^{E,DL} = \max_{\mathbf{r}} (SAR_{10g}^{DL}(\mathbf{r})_{m,s}) \quad (11)$$

Note that the performance of the deep learning-based method is evaluated by performing a leave-one-out cross-validation. This means that 23 separate times, the convolutional neural network is trained on all of the data samples from all models except for one model ($22 \times 250 = 5500$ training samples with random phase shimming).

3.4 | Performance evaluation

To assess the performance of all methods covered by this study, for each body model, 1000 test sets were generated for each driving mode. Then, using the obtained 23 000 (23×1000) test sets, the mean pSAR_{10g} overestimation error of all pSAR_{10g} estimation methods with the linear safety factor and with the proposed CSM were evaluated and compared with each other for each driving mode.

4 | RESULTS

For each pSAR_{10g} estimation method and for each driving mode, 5750 (23×250) validation sets were used to define the required linear safety factor and the CSM. For random phase and prostate shimmed-phase drive modes, the required linear safety factors were similar and ranged from 1.36 for the deep learning method to 1.96 for the model selection method. For the amplitude and phase shimming driving mode, the obtained linear safety factors range was a bit wider, ranging from 1.22 to 2.09 for deep learning and model selection methods, respectively.

To determine the joint and marginal probability density functions of each pSAR_{10g} estimation method, each scatter plot of true versus estimated pSAR_{10g} was fitted with a Gaussian mixture distribution (the number of Gaussian terms was determined with the Bayesian information criterion³⁶ and ranged between 3 and 5), and the estimated pSAR_{10g} histogram was fitted with a gamma distribution (*MATLAB and Statistics and Machine Learning Toolbox*, The MathWorks, Natick, MA). Subsequently, the conditional probability density function was determined and the CSM was evaluated to have a probability of underestimation of 0.1% ($\epsilon = 0.001$). All probability density functions are reported in Supporting Information Figures S4-S8).

The performances of the defined safety factors were assessed for each pSAR_{10g} estimation method and each driving mode with 23 000 (23×1000) test sets.

Figure 3 shows the scatter plot of true versus estimated pSAR_{10g} considering the use of a single generic body model to evaluate the pSAR_{10g} with random phases (Figure 3A), prostate shimmed phases (Figure 3B) and random amplitudes and phases normalized to 1W maximum input power per channel (Figure 3C), and 8W total input power (Figure 3D). The orange line denotes the corrected pSAR_{10g} after the application of the linear safety factor. The red line denotes the corrected pSAR_{10g} after application of the linear safety factor limited by the 99.9% certain pSAR_{10g} upper bound (dashed blue line). The green line denotes the corrected pSAR_{10g} using the CSM. Figure 3E shows the mean overestimation for each pSAR_{10g} correction method.

Figure 4 presents the scatter plot of true versus estimated pSAR_{10g} and the mean overestimation considering the model library pSAR_{10g} estimation method. Figures 5 and 6 present the same results for the model selection and multiple models selection methods, and Figure 7 presents the same plot for the deep learning method. The histogram of the pSAR_{10g} estimation error of each pSAR_{10g} estimation method, each driving mode, and each pSAR_{10g} correction method is presented in Supporting information Figures S9-S13.

The bar diagram in Figure 8 makes it easy to compare the pSAR_{10g} estimation methods covered by this study. It shows the mean overestimation after the correction for all considered pSAR_{10g} estimation methods for each driving mode. The deep learning-based pSAR_{10g} estimation method outperforms conventional methods. Compared with the pSAR_{10g} estimation methods based on the selection of the most similar models from a database, after the application of the CSM, it achieves a mean overestimation reduction of 44%-28% for phase shimming and 13%-9% for amplitude and phase shimming. The multiple models selection and the model selection methods are the second and third best pSAR_{10g} estimation methods, respectively.

5 | DISCUSSION

The linear safety factor is a common approach to avoid potential underestimation due to errors in predicted peak local SAR levels.^{10-12,23,24} This work has shown that the linear safety factor results in unnecessarily severe scanning constraints. The correction of estimated peak local SAR levels (pSAR_{10g}) using a linear safety factor produces an overconservative pSAR_{10g} prediction, even when the safety factor is defined with outlier rejection and a low underestimation probability is allowed.²⁷ This is because for high estimated pSAR_{10g} values, the corrected pSAR_{10g} values are unreasonably high. If we consider the mean estimated pSAR_{10g} value with almost all methods, it already produces corrected pSAR_{10g} values higher than the 99.9% certain pSAR_{10g} upper bound.

Therefore, it could be reasonable to limit the corrected pSAR_{10g} values to this upper bound. This produces a slightly higher probability of underestimation ($< 0.2\%$) but severely reduces the mean overestimation. Such a combination of corrected pSAR_{10g} values is in principle a first attempt at defining variable margin factors depending on the estimated SAR level.

However, a more appropriate way of tackling this problem is by making use of probability theory, to take into account the statistical distribution of realistically attainable pSAR_{10g} values. Therefore, a new approach to define a variable safety margin based on the conditional probability distribution was presented. This approach allows one to define a CSM to obtain, for each estimated pSAR_{10g} value, a corrected pSAR_{10g} value that will

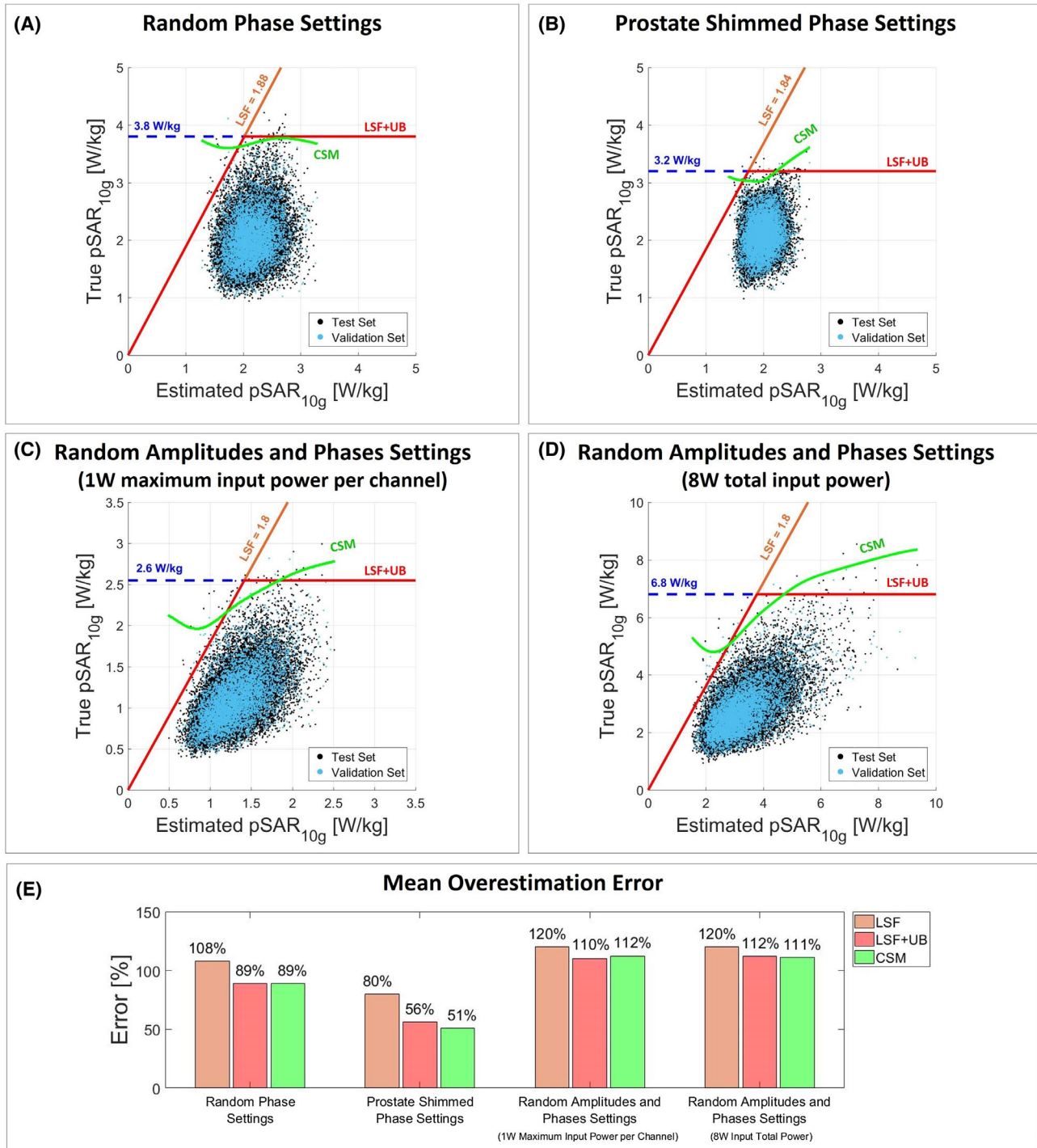


FIGURE 3 Generic body model. Scatter plot of true versus estimated pSAR_{10g} values with random phases (A), prostate shimmed phase settings (B), random amplitudes and phases normalized to 1W maximum input power per channel (C), and random amplitudes and phases normalized to 8W input total power (D). The orange line denotes the corrected pSAR_{10g} by the LSF; the red line denotes the corrected pSAR_{10g} by the LSF limited by the 99.9% certain pSAR_{10g} upper bound (UB) (dashed blue line); and the green line denotes the corrected pSAR_{10g} by the conditional safety margin (CSM). The LSF and the CSM are determined using the validation set (validation set: cyan dots; test set: black dots). E, Bar plot of the mean pSAR_{10g} overestimation error for each pSAR_{10g} correction method and each driving mode

allow for only a very small predefined probability of underestimation (eg, 0.1%). The 0.1% probability pSAR_{10g} threshold violations will be modest with rapidly decreasing likelihood

for higher pSAR_{10g} levels. Nevertheless, some users may prefer using the proposed approach with lower underestimation probabilities, such as 0.001% (Supporting information Figure S14).

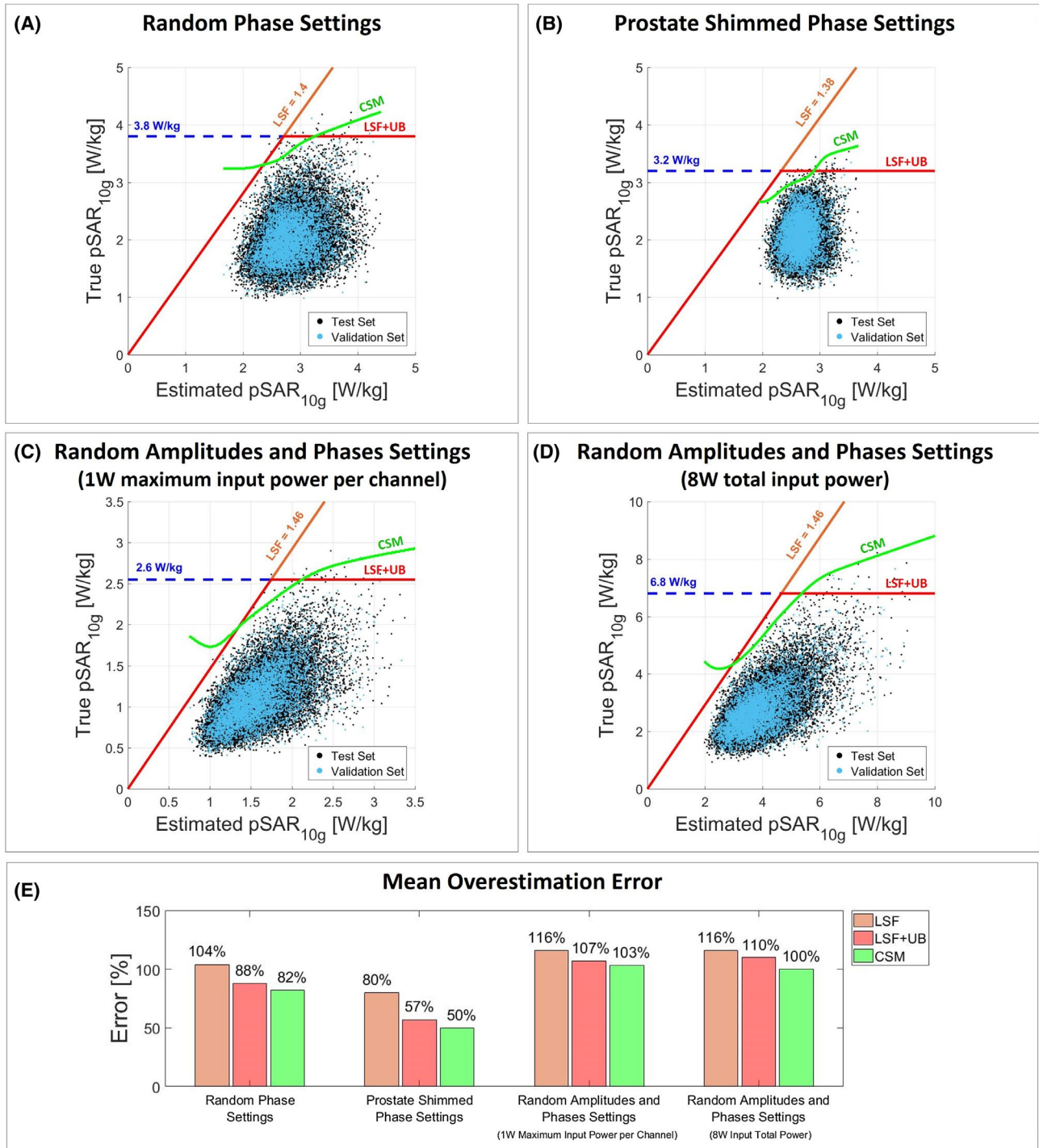


FIGURE 4 Model library. Scatter plot of true versus estimated $pSAR_{10g}$ with random phases (A), prostate shimmed phase settings (B), random amplitudes and phases normalized to 1W maximum input power per channel (C), and random amplitudes and phases normalized to 8W input total power (D). The orange line denotes the corrected $pSAR_{10g}$ by the LSF; the red line denotes the corrected $pSAR_{10g}$ by the LSF limited by the 99.9% certain $pSAR_{10g}$ upper bound (dashed blue line); and the green line denotes the corrected $pSAR_{10g}$ by the CSM. The LSF and the CSM are determined using the validation set (validation set: cyan dots; test set: black dots). E, Bar plot of the mean $pSAR_{10g}$ overestimation error for each $pSAR_{10g}$ correction method and each driving mode

This approach can be applied to any $pSAR_{10g}$ estimation method. It was applied to five state-of-the-art $pSAR_{10g}$ estimation methods for a multitransmit array of eight fractionated dipole antennas for prostate imaging at 7 T^{29,30} with

three different drive modes: random phase settings, prostate shimmed phase settings, and random amplitude and phase settings. For each examined case, 23 × 250 validation sets were used to determine the linear safety factor and the CSM. A larger

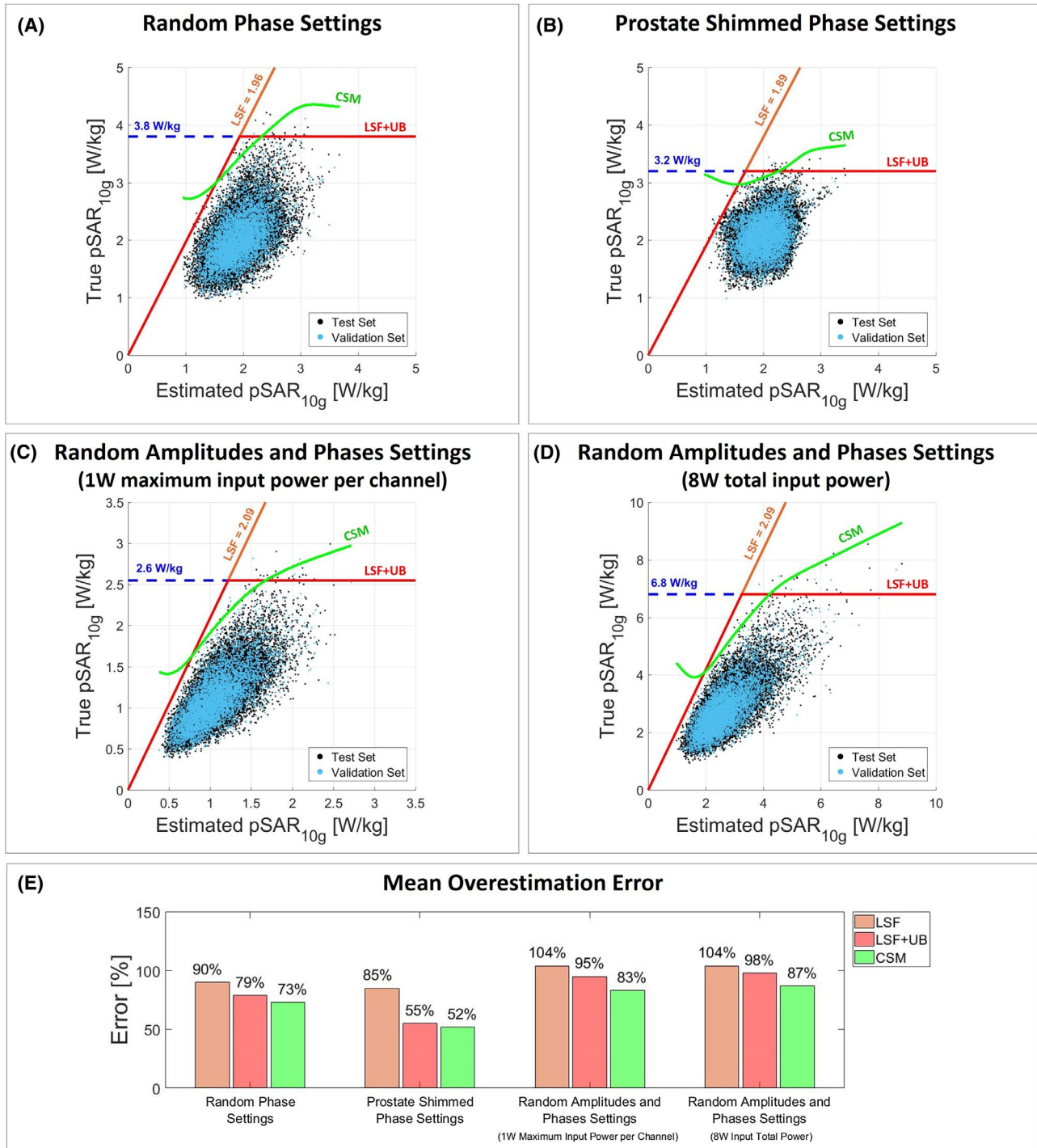


FIGURE 5 Model selection. Scatter plot of true versus estimated pSAR_{10g} with random phases (A), prostate shimmed phase settings (B), random amplitudes and phases normalized to 1W maximum input power per channel (C), and random amplitudes and phases normalized to 8W input total power (D). The orange line denotes the corrected pSAR_{10g} by the LSF; the red line denotes the corrected pSAR_{10g} by LSF limited by the 99.9% certain pSAR_{10g} upper bound (dashed blue line); and the green line denotes the corrected pSAR_{10g} by the CSM. The LSF and the CSM are determined using the validation set (validation set: cyan dots; test set: black dots). E, Bar plot of the mean pSAR_{10g} overestimation error for each pSAR_{10g} correction method and each driving mode

number of validation sets did not produce a significant change in the determined safety factor nor in performance (Supporting Information Figure S15).

Compared with the linear safety factor limited by the 99.9% certain pSAR_{10g} upper bound, the performance increase achievable with this method depends on the accuracy

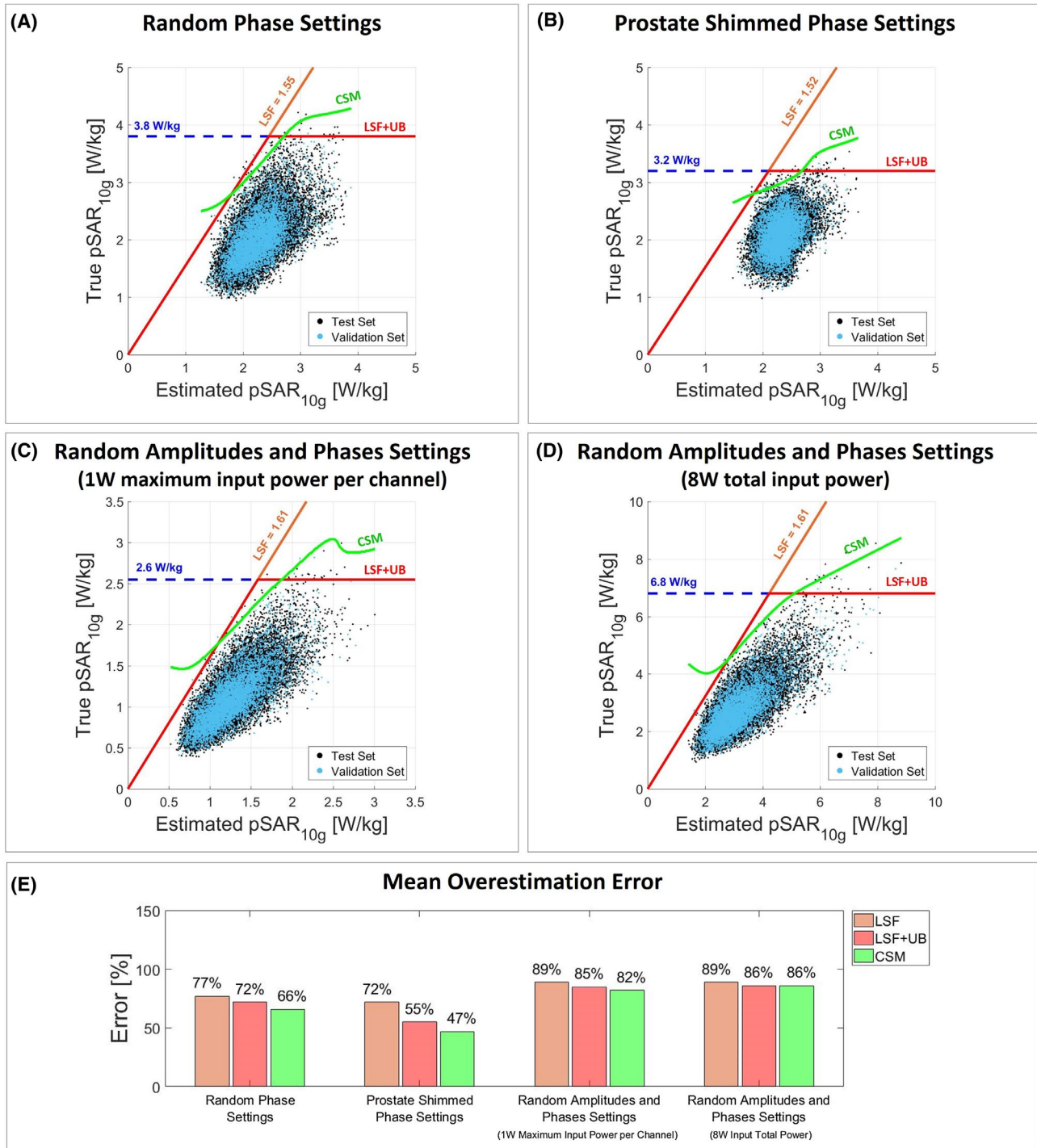


FIGURE 6 Multiple models selection. Scatter plot of true versus estimated pSAR_{10g} with random phases (A), prostate shimmed phase settings (B), random amplitudes and phases normalized to 1W maximum input power per channel (C), and random amplitudes and phases normalized to 8W input total power (D). The orange line denotes the corrected pSAR_{10g} by the LSF; the red line denotes the corrected pSAR_{10g} by the LSF limited by the 99.9% certain pSAR_{10g} upper bound (dashed blue line); and the green line denotes the corrected pSAR_{10g} by the CSM. The LSF and the CSM are determined using the validation set (validation set: cyan dots; test set: black dots). E, Bar plot of the mean pSAR_{10g} overestimation error for each pSAR_{10g} correction method and each driving mode

of the used pSAR_{10g} estimation method. With pSAR_{10g} estimation methods that show a good correlation between estimated and true pSAR_{10g} values (eg, deep learning method), the proposed approach allows a significant reduction of the

mean pSAR_{10g} overestimation (16%-28%). For methods that have a poor performance in predicting the true peak local SAR level (one generic model or worst-case of all models), the benefit of the proposed approach is much less pronounced.

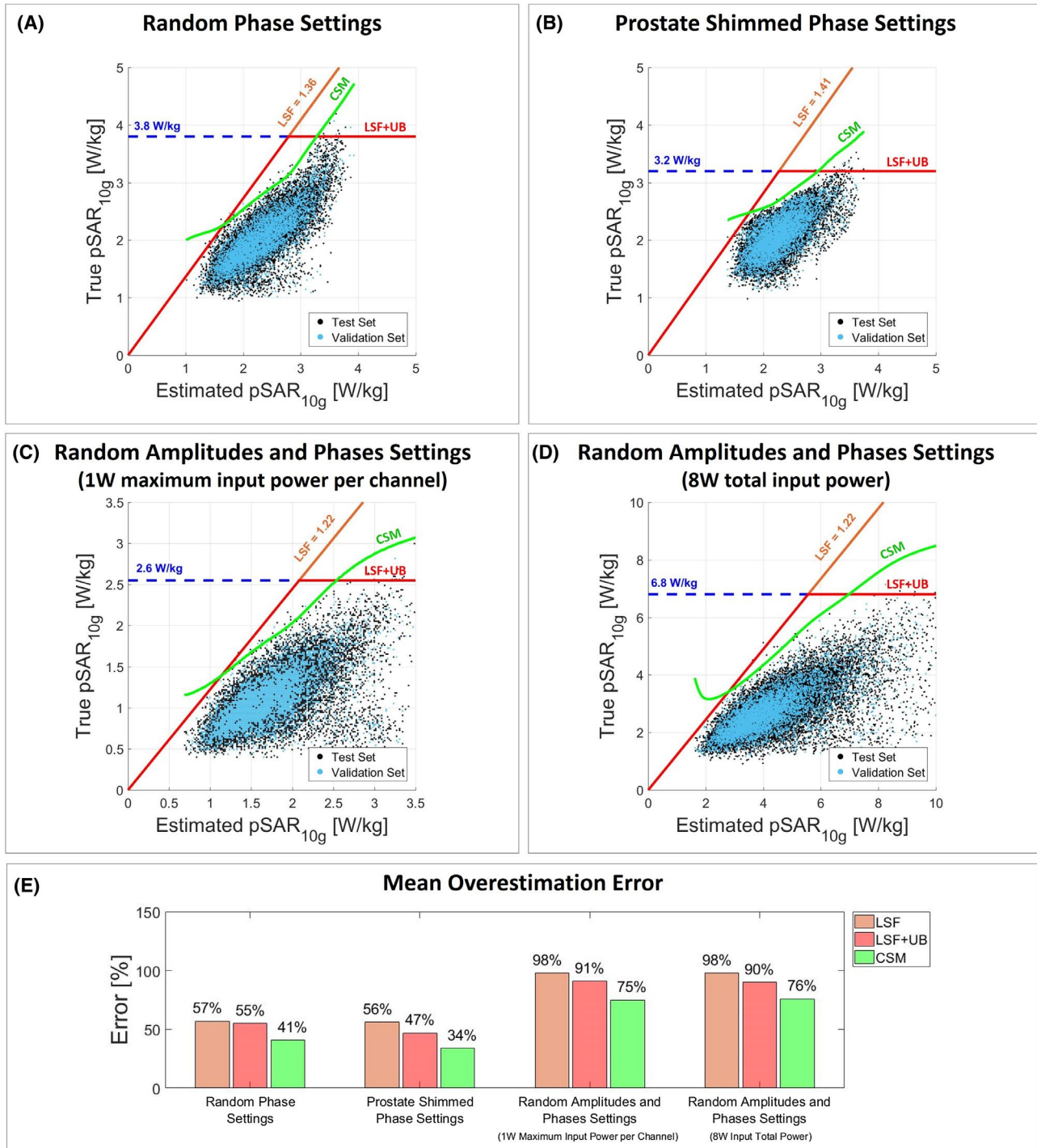


FIGURE 7 Deep learning. Scatter plot of true versus estimated pSAR_{10g} with random phases (A), prostate shimmed phase settings (B), random amplitudes and phases normalized to 1W maximum input power per channel (C), and random amplitudes and phases normalized to 8W input total power (D). The orange line denotes the corrected pSAR_{10g} by the LSF; the red line denotes the corrected pSAR_{10g} by the LSF limited by the 99.9% certain pSAR_{10g} upper bound (dashed blue line); and the green line denotes the corrected pSAR_{10g} by the CSM. The LSF and the CSM are determined using the validation set (validation set: cyan dots; test set: black dots). E, Bar plot of the mean pSAR_{10g} overestimation error for each pSAR_{10g} correction method and each driving mode

For these methods, the reduction in mean pSAR_{10g} overestimation is less than 12%.

Note that a special case exists for extremely poor estimation methods. In that case, the estimated and true

pSAR_{10g} values exhibit low correlation, or, in other words, are essentially independent; in this case, the joint probability is the product of their marginal probabilities $P(pSAR^E, pSAR^T) = P(pSAR^E)P(pSAR^T)$. Then, the conditional

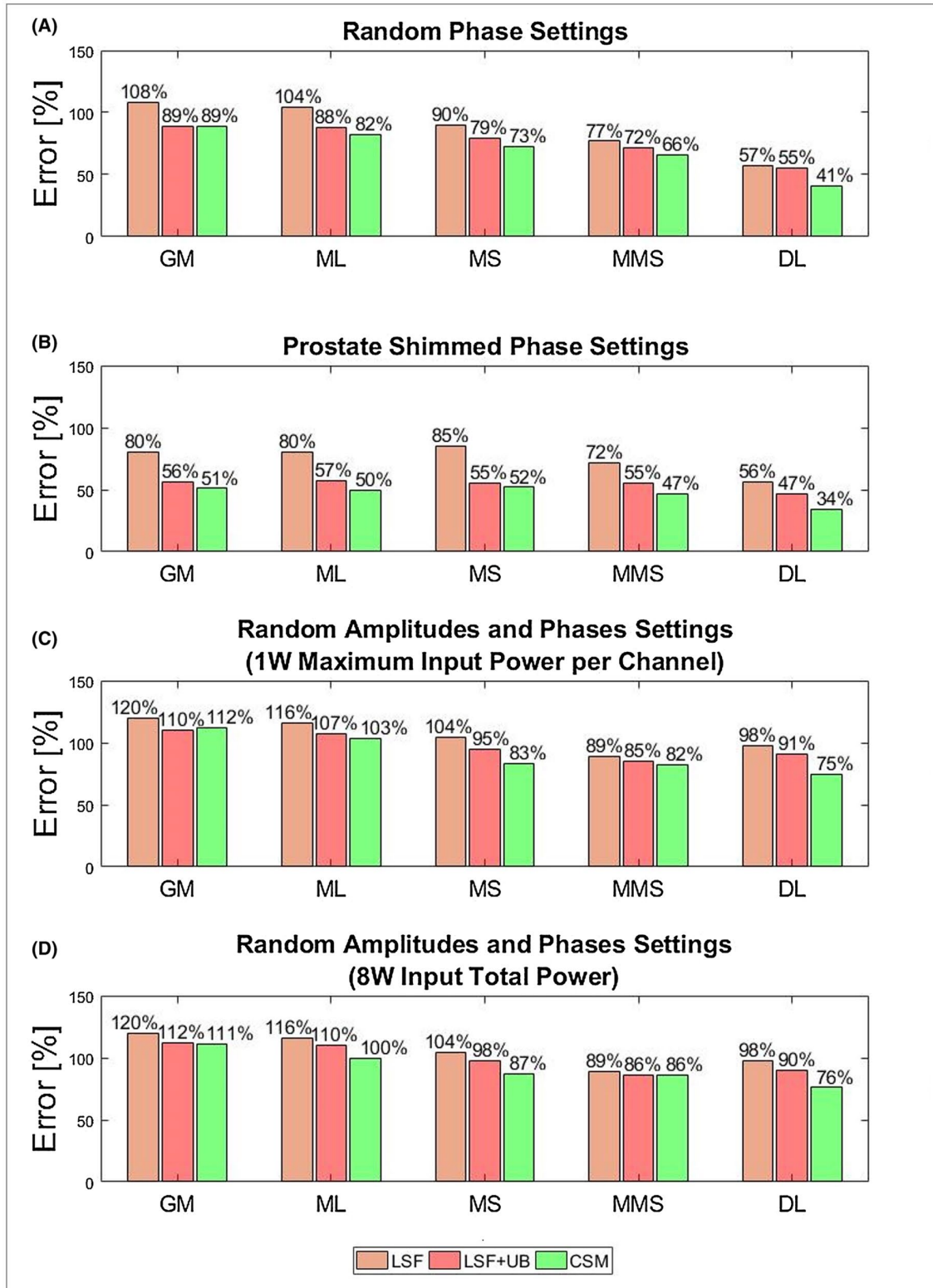


FIGURE 8 Bar plot of the mean overestimation error for each pSAR_{10g} estimation method with random phases (A), prostate shimmed phase settings (B), random amplitudes and phases normalized to 1W maximum input power per channel (C), and random amplitudes and phases normalized to 8W input total power (D). The orange bar denotes the mean overestimation of the corrected pSAR_{10g} by the LSF; the red bar denotes the mean overestimation of the corrected pSAR_{10g} by the LSF limited by the 99.9% certain pSAR_{10g} upper bound; and the green bar denotes the mean overestimation of the corrected pSAR_{10g} by the CSM. Abbreviations: DL, deep learning model; GM, general body model; ML, model library; MMS, multiple models selection; MS, model selection

probability coincides with the marginal probability of true value $P(pSAR^T | pSAR^E) = P(pSAR^E, pSAR^T) / P(pSAR^E) = P(pSAR^T)$. This means that from the knowledge of the estimated value, nothing can be inferred, and if the CSM is defined considering $\epsilon = 0.001$, for any estimated $pSAR_{10g}$ the CSM will determine a corrected $pSAR_{10g}$ value that will approximately coincide with the 99.9% certain $pSAR_{10g}$ upper bound. This is more or less the case for $pSAR_{10g}$ prediction using one generic model with random phase settings. As indicated in Figure 3A, the scatter cloud shows very poor correlation of true $pSAR_{10g}$ versus estimated $pSAR_{10g}$. Therefore, the CSM results in an almost horizontal line following the 3.8 W/kg 99.9% confident upper limit.

Note that, for low estimated $pSAR_{10g}$ values, the linear safety factor produces lower corrected $pSAR_{10g}$ values than the conditional safety factor. Indeed, with the linear safety factor, the probability of underestimation is larger than 0.1% when a low $pSAR_{10g}$ value is estimated, whereas with the CSM it is 0.1% for any estimated $pSAR_{10g}$ value. Likewise, with the 99.9% certain $pSAR_{10g}$ upper bound, the probability of underestimation is larger than 0.1% when a high $pSAR_{10g}$ value is estimated. Moreover, because the histogram of estimated $pSAR_{10g}$ values follows a gamma distribution, close to the range boundaries, the number of estimated $pSAR_{10g}$ values is less and less. With a very small number of samples, the estimated conditional probability density function is larger (greater uncertainty) and less accurate (because it is defined as the ratio between very small numbers, the relative error is larger). These are probably the causes of the observed large overestimation for low estimated $pSAR_{10g}$ values and the anomalies in the conditional probability density functions outside of the feasibly estimated $pSAR_{10g}$ ranges (Supporting Information Figures S4-S8). Note that the relatively large overestimation of these low expected $pSAR_{10g}$ values is not at all problematic, because their occurrence is quite rare.

It is worth noting that, for the random amplitudes and phases settings, we assumed that each drive vector is equally likely to occur. This assumption is probably far from the truth for many RF pulse design strategies. Therefore, for the considered RF pulse type, the probability of occurrence of the drive vectors should be assessed and taken into account for the validation set generation (similarly to what has been done for prostate shimmed phase settings).

Furthermore, it must be considered that the time-dependent drive vector of these custom RF pulses produces a time-dependent SAR distribution with the peak value in a different location for each time step. The CSM, as well as the linear safety factor, is a global factor that does not take into account the spatial distribution of the SAR. Nevertheless, it can be used to define a correction factor (CSM/ $pSAR^E$) to apply to the whole SAR distribution (or virtual observation

points) for each time step (in the same way that the linear safety factor is usually applied). Subsequently, the SAR distributions are integrated over time, and then the global peak SAR value can be assessed.

Results on phase-amplitude shimming have been presented for two normalization methods: normalization on maximum input power per channel and normalization on total input power. Both normalizations resulted in different scatter clouds and different CSM curves. However, the performance in terms of mean overestimation was basically equivalent (Figure 8).

In addition, a comparison was made among the state-of-the-art $pSAR_{10g}$ estimation methods. We showed that when using a single generic body model for $pSAR_{10g}$ estimation, a poor relation between estimated and true $pSAR_{10g}$ is observed and the mean overestimation error after the correction with the CSM is 89% for random phase settings, 51% for prostate shimmed phase settings, 112% for random amplitude and phase settings normalized to 1W maximum input power per channel, and 111% for random amplitude and phase settings normalized to 8W input total power (Figure 8). Figure 3 clearly shows that the point cloud does not follow the diagonal and the corrected $pSAR_{10g}$ values, almost coincident with the 99.9% certain $pSAR_{10g}$ upper bound. Thus, the proposed correction approach allows a little mean overestimation reduction (less than 10%).

Similar results were found using a model library to predict $pSAR_{10g}$ over all models. This $pSAR_{10g}$ estimation is obviously more conservative and requires smaller linear safety factors (Figure 4). However, after the correction the mean overestimation was comparable with the generic body model method (82%, 50%, 103%, and 100%). In this case, the CSM allows a mean overestimation reduction from 4% to 12%, because if a low $pSAR_{10g}$ value is estimated over all models, it is less likely that a high true $pSAR_{10g}$ value is observed, and this weak relationship is exploited by the proposed correction method.

The $pSAR_{10g}$ estimation methods based on the selection of the most similar model(s) from a database show better performance than the conventional estimation methods. If the most “similar” model is used, the $pSAR_{10g}$ estimation is more accurate but not very precise (Figure 5). To manage the low precision due to the intersubject $pSAR_{10g}$ variability, a considerable linear safety factor is required. Nevertheless, the model selection method achieves a lower mean overestimation after the correction with the proposed approach (73%, 52%, 83%, and 87%).

The use of multiple models reduces the probability of underestimation due to the intersubject $pSAR_{10g}$ variability, and allows us to reduce the required safety factors. This results in an even lower mean overestimation error (66%, 47%, 82%, and 86%). With the CSM, an overestimation reduction up to 15% is reached with these estimation methods.

The proposed $pSAR_{10g}$ correction method exploits the correlation between estimated and true $pSAR_{10g}$ levels. Therefore, with more accurate and precise $pSAR_{10g}$ estimation methods, the deep learning method provides the best performance (Figure 7). It reduces the mean $pSAR_{10g}$ overestimation by almost 30%, achieving a mean overestimation error of 41% for random phase settings and 34% for prostate shimmed phase settings.

Considerable mean overestimation is still observed for random amplitude and phase shimming (75%-76%). However, it should be noted that the neural network was trained with only phase-shimmed training samples.²⁷ This means that better performance might be obtained by training the network with amplitude and phase-shimmed training samples. Nonetheless, even in this case the CSM allows one to reduce the mean overestimation of almost 20%.

It should also be noted that the deep learning-based method makes use of a network that was trained by penalizing the underestimation error more.²⁷ This was done in an attempt to avoid underestimation, yet some underestimation error remained. Therefore, the method was included in this study as one of the investigated SAR assessment methods. If the network is trained by equally penalizing the underestimation and overestimation error, the deep learning-based method would have shown a larger degree of underestimation. The points in the scatter clouds in Figure 7 would shift to the left, the linear safety factor required to avoid underestimation would increase (red lines in Figure 7 become steeper), and the linear safety factor approach would result in a larger overestimation. Therefore, the benefit of the CSM approach would appear larger. We chose to use the deep learning-based method as it was published, which results in a conservative estimate of what benefit the CSM could provide for such methods.

6 | CONCLUSIONS

To avoid underestimation of predicted peak local SAR ($pSAR_{10g}$) levels for multitransmit arrays, a safety factor is often applied to correct (increase) the predicted $pSAR_{10g}$ levels. This work has shown that this approach results in drastic and unnecessary overestimation of $pSAR_{10g}$ levels, particularly if the estimated $pSAR_{10g}$ level is relatively high.

In this work, an alternative approach for safety margin definition is presented using probability theory. This CSM approach allows us to define a variable safety margin for each possible estimated $pSAR_{10g}$ value.

For prostate imaging at 7 T, the proposed CSM approach results in lower mean overestimation for all investigated local SAR estimation methods. Compared with the linear safety

factor in combination with the 99.9% upper bound, the reduction of overestimation up to 30% is reached for the more accurate local SAR assessment methods.

CONFLICT OF INTEREST

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REFERENCES

- Collins CM, Smith MB. Calculations of B(1) distribution, SNR, and SAR for a surface coil adjacent to an anatomically-accurate human body model. *J Magn Reson*. 2001;45:692-699.
- Vaughan JT, Garwood M, Collins CM, et al. 7T vs. 4T: RF power, homogeneity, and signal-to-noise comparison in head images. *Magn Reson Med*. 2001;46:24-30.
- Uğurbil K. Imaging at ultrahigh magnetic fields: history, challenges, and solutions. *NeuroImage*. 2018;168:7-32.
- Padormo F, Beqiri A, Hajnal JV, Malik SJ. Parallel transmission for ultrahigh-field imaging. *NMR Biomed*. 2016;29:1145-1161.
- Malik SJ, Keihaninejad S, Hammers A, Hajnal JV. Tailored excitation in 3D with spiral nonselective (SPINS) RF pulses. *Magn Reson Med*. 2012;67:1303-1315.
- Cloos MA, Boulant N, Luong M, et al. kT-Points: short three-dimensional tailored RF pulses for flip-angle homogenization over an extended volume. *Magn Reson Med*. 2012;67:72-80.
- Sbrizzi A, Hoogduin H, Lagendijk JJ, Luijten PR, Sleijpen GL, Van Den Berg CAT. Fast design of local N-gram-specific absorption rate-optimized radiofrequency pulses for parallel transmit systems. *Magn Reson Med*. 2012;67:824-834.
- Sbrizzi A, Hoogduin H, Hajnal JV, van den Berg CAT, Luijten PR, Malik SJ. Optimal control design of turbo spin-echo sequences with applications to parallel-transmit systems. *Magn Reson Med*. 2017;77:361-373.
- Fiedler TM, Ladd ME, Bitz AK. SAR simulations & safety. *NeuroImage*. 2018;168:33-58.
- Boulant N, Gras V, Amadon A, Luong M, Ferrand G, Vignaud A. Workflow proposal for defining SAR safety margins in parallel transmission. In: Proceedings of the 26th Annual ISMRM Meeting, Paris, France, 2018. p 295.
- Ipek Ö, Raaijmakers AJE, Lagendijk JJ, Luijten PR, Van Den Berg CAT. Intersubject local SAR variation for 7T prostate MR imaging with an eight-channel single-side adapted dipole antenna array. *Magn Reson Med*. 2014;71:1559-1567.
- Meliadó EF, van den Berg CAT, Luijten PR, Raaijmakers AJE. Intersubject specific absorption rate variability analysis through construction of 23 realistic body models for prostate imaging at 7T. *Magn Reson Med*. 2019;81:2106-2119.
- Graesslin I, Vernickel P, Bornert P, et al. Comprehensive RF safety concept for parallel transmission MR. *Magn Reson Med*. 2014;74:589-598.
- Zhu Y, Alon L, Deniz CM, Brown R, Sodickson DK. System and SAR characterization in parallel RF transmission. *Magn Reson Med*. 2012;67:1367-1378.

15. Orzada S, Ladd ME, Bitz AK. A method to approximate maximum local SAR in multichannel transmit MR systems without transmit phase information. *Magn Reson Med.* 2017;78:805-811.
16. Van Den Bergen B, Van Den Berg CAT, Klomp DWJ, Lagendijk JJW. SAR and power implications of different RF shimming strategies in the pelvis for 7T MRI. *J Magn Reson Imaging.* 2009;30:194-202.
17. Homann H, Börner P, Eggers H, Nehrke K, Dössel O, Graesslin I. Toward individualized SAR models and in vivo validation. *Magn Reson Med.* 2011;66:1767-1776.
18. IEC 60601-2-33. *Medical Electrical Equipment Part 2-33: Particular Requirements for the Basic Safety and Essential Performance of Magnetic Resonance Equipment for Medical Diagnosis.* Edition 3.0. International Standard. Geneva, Switzerland: IEC; 2010.
19. Villena JF, Polimeridis AG, Eryaman Y, et al. Fast electromagnetic analysis of MRI transmit RF coils based on accelerated integral equation methods. *IEEE Trans Biomed Eng.* 2016;63:2250-2261.
20. Graesslin I, Homann H, Biederer S, et al. A specific absorption rate prediction concept for parallel transmission MR. *Magn Reson Med.* 2012;68:1664-1674.
21. Caputa K, Okoniewski M, Stuchly MA. An algorithm for computations of the power deposition in human tissue. *IEEE Antennas Propag Mag.* 1999;41:102-107.
22. Eichfelder G, Gebhardt M. Local specific absorption rate control for parallel transmission by virtual observation points. *Magn Reson Med.* 2011;66:1468-1476.
23. De Greef M, Ipek O, Raaijmakers AJE, Crezee J, Van Den Berg CAT. Specific absorption rate intersubject variability in 7T parallel transmit MRI of the head. *Magn Reson Med.* 2013;69:1476-1485.
24. Le Garrec M, Gras V, Hang MF, Ferrand G, Luong M, Boulant N. Probabilistic analysis of the specific absorption rate intersubject variability safety factor in parallel transmission MRI. *Magn Reson Med.* 2017;78:1217-1223.
25. Christ A, Kainz W, Hahn EG, et al. The virtual family—development of surface-based anatomical models of two adults and two children for dosimetric simulations. *Phys Med Biol.* 2010;55:N23-N38.
26. Gosselin M-C, Neufeld E, Moser H, et al. Development of a new generation of high-resolution anatomical models for medical device evaluation: the Virtual Population 3.0. *Phys Med Biol.* 2014;59:5287-5303.
27. Meliadó EF, Raaijmakers AJE, Sbrizzi A, et al. A deep learning method for image-based subject-specific local SAR assessment. *Magn Reson Med.* 2020;83:695-711.
28. Meliadó EF, Raaijmakers AJE, Restivo MC, Maspero M, Luijten PR, van den Berg CAT. Database construction for local SAR prediction: preliminary assessment of the intra and inter subject SAR variability in pelvic region. In: Proceedings of the 24th Annual Meeting of ISMRM, Singapore, 2016, p 3660.
29. Steensma BR, Luttje M, Voogt IJ, et al. Comparing signal-to-noise ratio for prostate imaging at 7T and 3T. *J Magn Reson Imaging.* 2019;49:1446-1455.
30. Raaijmakers AJE, Italiaander M, Voogt IJ, et al. The fractionated dipole antenna: a new antenna for body imaging at 7 Tesla. *Magn Reson Med.* 2016;75:1366-1374.
31. Gut A. *Probability: A Graduate Course.* 2nd Edition. New York, NY: Springer; 2013.
32. Parzen E. On estimation of a probability density function and mode. *Ann Math Stat.* 1962;33:1065-1076.
33. Lopez-Rubio E. A histogram transform for probability density function estimation. *IEEE Trans Pattern Anal Mach Intell.* 2014;36:644-656.
34. Scott D, Sain S. Data mining and computational statistics. In: *Handbook of Statistics.* Vol. 24. Amsterdam: Elsevier; 2005;229-261.
35. Meliadó EF, Raaijmakers AJE, Luijten PR, van den Berg CAT. Fast method to get an upper bound of the maximum SAR_{10g} for body coil arrays. In: Proceedings of the 25th Annual Meeting of ISMRM, Honolulu, 2017. p 4312.
36. Schwarz GE. Estimating the dimension of a model. *Ann Stat.* 1978;6:461-464.

SUPPORTING INFORMATION

Additional Supporting Information may be found online in the Supporting Information section.

FIGURE S1 Transverse maximum intensity projection of the worst-case peak 10g average specific absorption rate (pSAR_{10g}) distributions with uniform amplitude ($8 \times 1W$ input power)

FIGURE S2 Root-mean-square error (RMSE) matrix. Each entry RMSE $[n,m]$ represents the RMSE between the worst-case pSAR_{10g} distribution of the model n and the registered worst-case pSAR_{10g} distribution of the model m

FIGURE S3 Worst-case pSAR_{10g} distribution of each model and registered worst-case pSAR_{10g} distribution of the most representative “local SAR model”

FIGURE S4 Generic body model (validation set): scatter plot of the true pSAR^T versus the estimated pSAR^E (first column); marginal probability density function $f_E(pSAR^E)$ of the estimated pSAR values (second column); joint probability density function $f_{E,T}(pSAR^E, pSAR^T)$ of the estimated and true pSAR values (third column); and 2D conditional probability density function $f_{T|E}(pSAR^T|pSAR^E)$ obtained by combining the conditional probability density functions for each possible pSAR^E value (fourth column)

FIGURE S5 Model library (validation set): scatter plot of the true pSAR^T versus the estimated pSAR^E (first column); marginal probability density function $f_E(pSAR^E)$ of the estimated pSAR values (second column); joint probability density function $f_{E,T}(pSAR^E, pSAR^T)$ of the estimated and true pSAR values (third column); and 2D conditional probability density function $f_{T|E}(pSAR^T|pSAR^E)$ obtained by combining the conditional probability density functions for each possible pSAR^E value (fourth column)

FIGURE S6 Model selection (validation set): scatter plot of the true pSAR^T versus the estimated pSAR^E (first column); marginal probability density function $f_E(pSAR^E)$ of the estimated pSAR values (second column); joint probability density function $f_{E,T}(pSAR^E, pSAR^T)$ of the estimated and true pSAR values (third column); and 2D conditional probability density function $f_{T|E}(pSAR^T|pSAR^E)$ obtained by combining the conditional probability density functions for each possible pSAR^E value (fourth column)

FIGURE S7 Multiple models selection (validation set): scatter plot of the true pSAR^T versus the estimated pSAR^E (first column); marginal probability density function $f_E(pSAR^E)$ of

the estimated pSAR values (second column); joint probability density function $f_{E,T}(pSAR^E, pSAR^T)$ of the estimated and true pSAR values (third column); and 2D conditional probability density function $f_{T|E}(pSAR^T|pSAR^E)$ obtained by combining the conditional probability density functions for each possible pSAR^E value (fourth column)

FIGURE S8 Deep learning (validation set): scatter plot of the true pSAR^T versus the estimated pSAR^E (first column); marginal probability density $f_E(pSAR^E)$ of the estimated pSAR values (second column); joint probability density function $f_{E,T}(pSAR^E, pSAR^T)$ of the estimated and true pSAR values (third column); and 2D conditional probability density function $f_{T|E}(pSAR^T|pSAR^E)$ obtained by combining the conditional probability density functions for each possible pSAR^E value (fourth column)

FIGURE S9 Generic body model (test set). Scatter plot of true versus estimated pSAR_{10g} and histogram of the pSAR_{10g} estimation error for each driving mode and each pSAR_{10g} correction method. The linear safety factor (LSF) and the conditional safety margin (CSM) were determined using the validation set (cyan dots)

FIGURE S10 Model library (test set). Scatter plot of true versus estimated pSAR_{10g} and histogram of the pSAR_{10g} estimation error for each driving mode and each pSAR_{10g} correction method. The LSF and the CSM were determined using the validation set (cyan dots)

FIGURE S11 Model selection (test set). Scatter plot of true versus estimated pSAR_{10g} and histogram of the pSAR_{10g} estimation error for each driving mode and each pSAR_{10g} correction method. The LSF and the CSM were determined using the validation set (cyan dots)

FIGURE S12 Multiple models selection (test set). Scatter plot of true versus estimated pSAR_{10g} and histogram of the pSAR_{10g} estimation error for each driving mode and each pSAR_{10g} correction method. The LSF and the CSM were determined using the validation set (cyan dots)

FIGURE S13 Deep learning (test set). Scatter plot of true versus estimated pSAR_{10g} and histogram of the pSAR_{10g} estimation error for each driving mode and each pSAR_{10g}

correction method. The LSF and the CSM were determined using the validation set (cyan dots)

FIGURE S14 A, Scatter plot of true versus estimated pSAR_{10g} using the deep learning method with random phase settings (test set: black dots; validation set: cyan dots). The orange line denotes the corrected pSAR_{10g} by the LSF based on the worst-case ratio of true and estimated pSAR_{10g}. The green line denotes the corrected pSAR_{10g} by the CSM with probability of underestimation of 0.001% ($\epsilon = 0.00001$). B, Histogram of the pSAR_{10g} estimation error for the corrected pSAR_{10g} by the LSF. C, Histogram of the pSAR_{10g} estimation error for the corrected pSAR_{10g} by the CSM. The LSF and the CSM were determined using the validation set

FIGURE S15 A, Scatter plot of true versus estimated pSAR_{10g} using the deep learning method with random phase settings (23 × 1000 validation sets: black dots; 23 × 250 validation sets: cyan dots). The orange line denotes the corrected pSAR_{10g} by the LSF determined with 23 × 1000 validation sets (LSF = 1.37). It practically coincides with the corrected pSAR_{10g} by the LSF determined with 23 × 250 validation sets (LSF = 1.36). The solid and dotted green lines denote the corrected pSAR_{10g} by the CSM determined with 23 × 1000 validation sets and with 23 × 250 validation sets, respectively ($\epsilon = 0.001$). B, Histogram of the pSAR_{10g} estimation error for the corrected pSAR_{10g} by the LSF determined with 23 × 1000 validation sets. C, Histogram of the pSAR_{10g} estimation error for the corrected pSAR_{10g} by the CSM determined with 23 × 1000 validation sets

TABLE S1 The five most representative local SAR models for each model

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