**Better Together**

**Data Harmonization and Cross-Study Analysis of Abdominal MRI Data From UK Biobank and the German National Cohort**

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**Objectives:** The UK Biobank (UKBB) and German National Cohort (NAKO) are among the largest cohort studies, capturing a wide range of health-related data from the general population, including comprehensive magnetic resonance imaging (MRI) examinations. The purpose of this study was to demonstrate how MRI data from these large-scale studies can be jointly analyzed and to derive comprehensive quantitative image-based phenotypes across the general adult population.

**Materials and Methods:** Image-derived features of abdominal organs (volumes of liver, spleen, kidneys, and pancreas; volumes of kidney hilum adipose tissue; and fat fractions of liver and pancreas) were extracted from T1-weighted Dixon MRI data of 17,996 participants of UKBB and NAKO based on quality-controlled deep learning generated organ segmentations. To enable valid cross-study analysis, we first analyzed the data generating process using methods of causal discovery. We subsequently harmonized data from UKBB and NAKO using the ComBat approach for batch effect correction. We finally performed quantile regression on harmonized data across studies providing quantitative models for the variation of image-derived features stratified for sex and dependent on age, height, and weight.

**Results:** Data from 8791 UKBB participants (49.9% female; age, 63 ± 7.5 years) and 9205 NAKO participants (49.1% female; age, 51.8 ± 11.4 years) were analyzed. Analysis of the data generating process revealed direct effects of age, sex, height, weight, and the data source (UKBB vs NAKO) on image-derived features stratified for sex and dependent on age, height, and weight. Cross-study analysis of MRI data from UKBB and NAKO as proposed in this work can be helpful for future joint data analyses across cohorts linking genetic, environmental, and behavioral risk factors to MRI-derived phenotypes and provide reference values for clinical diagnostics.

**Key Words:** cohort study, MRI, deep learning, causality, cross-study analysis, abdomen, segmentation, age, NAKO, UK Biobank

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The UK Biobank (UKBB)† conducted in the United Kingdom and the German National Cohort (NAKO)‡ conducted in Germany are 2 of the largest ongoing population-scale cohort studies. Collecting a wide array of health-related information, including MR imaging data, these studies provide a unique level of individual phenotypic characterization of participants.³ UKBB enrols adults between ages 50 and 80 years, whereas NAKO enrols participants between ages 20 and 70 years.¹² This restriction naturally limits the generalizability of study results for each of these single studies.

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Merging study data performing cross-study analyses may potentially overcome such limitations and in addition yield higher statistical power, the opportunity to independently replicate results and improve resource efficiency. Data compatibility among different studies however poses challenges for proper merging. Recorded parameters and data structures might be substantially different with little overlap. From a statistical point of view, the presence of distribution shifts, or biases, in the observed data due to differences in the data-generating processes can result in data misinterpretation when data from different sources are merged. Cross-study analyses of imaging data are particularly challenging due to additional sources of variation regarding the image acquisition process such as different scanner types, varying imaging protocols, and study-specific image processing algorithms. These factors can influence image-derived biomarkers, especially when magnetic resonance imaging (MRI) is used—a modality that is inherently difficult to standardize. The practical relevance of such biases has previously been reported on different medical image data sets.7,8

In the case of UKBB and NAKO, image acquisition protocols are partially aligned with the strategic intention to potentially enable cross-study analyses. Similarities cover an overall agreement on anatomic coverage and partial agreement on MRI sequences. Still, central aspects of MR acquisition protocols vary significantly including scanner models, magnetic field strengths, sequence parameters, or the occurrence of artifacts. Thus, it is unclear whether image-derived features from UKBB and NAKO can be pooled in a meaningful way for subsequent combined analyses.

Aiming to overcome such challenges, several techniques for data harmonization across studies have been proposed including model-based approaches (eg, batch effect correction using ComBat11 [“Combining Batches”] and its modifications3,12–14). The advantage of model-based data harmonization is the possibility to selectively correct for undesired bias while preserving informative factors of variation. This has recently been demonstrated also in a medical imaging context, mainly in a neuroimaging and oncological imaging context.7,12,13

The effective and valid application of such model-based data correction techniques requires detailed understanding of the data generating process. Usually, prior (common sense) knowledge about causal interactions among observed variables is used to harmonize data. As an extension, methods of causal discovery15 may provide complementary information about the data generating process and thus inform the application of data harmonization techniques. This can be of particular relevance in large-scale studies with complex data interactions.16

The purpose of this study is to demonstrate how imaging data from large-scale studies such as UKBB and NAKO can be jointly analyzed and to derive comprehensive quantitative image-based organ phenotypes across the general adult population.

MATERIALS AND METHODS

Population Characteristics and Imaging Data

Data were obtained from UKBB and NAKO, which obtained written informed consents from all subjects and approved our data analysis. Analysis of anonymized data from these studies was approved by the local institutional ethics committee.

This study reports findings from the first 20,000 data sets including MRI data available to us from the 2 study cohorts (10,000 data sets per study). After exclusion of data samples with MRI acquisition artifacts and erroneous automated organ segmentations (see below), image data and related demographic information (age, sex, body weight, and height) from 17,996 participants (8791 from UKBB and 9205 from NAKO) were used for further analysis. Summary statistics describing the study cohorts are provided in Table 1 and visualized in Figure 1. All image data analyzed in this work have been part of a previously reported technical work on deep learning-based abdominal organ segmentation,17 which was the technical foundation for this present work. There is no overlap in data analysis or reported results between these 2 studies.

Both UKBB and NAKO acquire whole-body MRI data on a subset of participants using clinical MR scanners (UKBB: 1.5 T Siemens Magneton Avanto; NAKO: 3 T Siemens Magneton Skyra, Siemens Healthineers, Erlangen, Germany). In this study, whole-body T1-weighted images obtained from dual-echo gradient echo imaging—which is available in UKBB and NAKO—were used. This includes 4 tissue contrasts per participant and image volume (fat, water, in-phase, and opposed-phase). Although these tissue images are comparable between the 2 studies, other acquisition parameters vary markedly. Notably, voxel size is higher in UKBB (2.23 × 2.23 × 3 mm3) to 2.23 × 2.23 × 4.5 mm3) compared with NAKO (1.2 × 1.2 × 3 mm3), which has a direct impact on spatial resolution, image signal, and image noise.2,9

TABLE 1. Demographic Characteristics of Study Populations Included in This Analysis

<table>
<thead>
<tr>
<th></th>
<th>UKBB</th>
<th>NAKO</th>
<th>Combined</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. participants</td>
<td>8791</td>
<td>9205</td>
<td>17,996</td>
</tr>
<tr>
<td>Sex (M/F) %</td>
<td>49.9% / 50.1%</td>
<td>48.4% / 51.6%</td>
<td>49.1% / 50.9%</td>
</tr>
<tr>
<td>Age (SD), y</td>
<td>63.0 (7.5)</td>
<td>51.8 (11.4)</td>
<td>57.3 (11.2)</td>
</tr>
<tr>
<td>Weight (SD), kg</td>
<td>77.1 (15.1)</td>
<td>79.5 (16.2)</td>
<td>78.3 (15.7)</td>
</tr>
<tr>
<td>Height (SD), cm</td>
<td>169.4 (9.4)</td>
<td>171.8 (9.5)</td>
<td>170.7 (9.5)</td>
</tr>
<tr>
<td>BMI (SD), kg/m2</td>
<td>26.7 (4.3)</td>
<td>26.8 (4.7)</td>
<td>26.8 (4.5)</td>
</tr>
</tbody>
</table>

UKBB, UK Biobank; NAKO, German National Cohort; F, female; M, male; SD, standard deviation.

Extraction of Image-Derived Features

This study focuses on the phenotypic characterization of abdominal organs (liver, spleen, left and right kidneys, and pancreas). These target organs were automatically segmented on MRI scans of 10,000 data samples per study using a pretrained and publicly available deep learning model based on a 3D full resolution convolutional architecture (nnUNet17,18). Resulting organ segmentation masks were visually inspected for the purpose of quality control, and data samples with severe MR image artifacts or substantial automated segmentation errors were excluded. This resulted in a total of 17,996 data sets (8791 from UKBB and 9205 from NAKO) that were used for further analysis in this study. This entire process of organ segmentation and quality control is described in detail in previous work17 and was the technical basis for this work.

In a subsequent postprocessing step, the segmentation masks of the kidneys were split into a parenchymal kidney mask and a kidney hilum adipose tissue (AT) mask by applying a threshold of 0.5 to the relative signal of the fat image (=fat/[fat + water]). Thus, 7 segmentation masks were obtained per data set (5 organs + right and left kidney hilum AT). The corresponding organ and tissue volumes were calculated from these segmentation masks by multiplying the respective voxel count with the voxel volume. In addition to volume features, proton density fat fractions (PDDFs) of liver and pancreas were estimated. To this end, mean fat-image and water-image voxel signal intensities were extracted from liver and the pancreas segmentation masks, and relative fat signal intensities (=fat/[fat + water]) were computed as a measure for the relative organ fat content.19 Thus, 9 image-derived features were extracted in total (organ volumes, kidney hilum AT volumes, and PDDFs of liver and pancreas).

Analysis of the Data Generating Process

To acquire a comprehensive understanding of the data-generating process—a prerequisite for subsequent data harmonization—we combined prior knowledge with methods of causal discovery. Specifically, we used the knowledge that age was causally dependent on the data

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source (UKBB vs NAKO) due to different inclusion criteria among these studies. Based on common medical knowledge, we assumed that age and sex have a direct effect on height and weight, and that height has a direct effect on weight.\(^\text{29}\) Finally, based on scientific literature, it is well-established that age impacts at least a subset of the observed image features, for example, organ sizes of individuals decrease with age.\(^\text{21–23}\)

Beyond these causal relations established by prior knowledge, we aimed to investigate further potential causal relations among image-derived features, observed demographic features, and the data source. To this end, we used conditional independence testing as a method of causal discovery combined with the knowledge about the direction of potential causal relation. Specifically, we assumed that observed image features are purely children of a parent-child connection in the causal sense, whereas the data source has only a parent role in the causal sense.

To identify the causal graph, we performed nonparametric nonlinear conditional independence testing by Invariant Environment Prediction previously described by Heinze-Deml et al.\(^\text{24}\) Concretely, we implemented Invariant Environment Prediction using random forest classifiers/regressors (depending on the type of target variable) that were trained with 100 trees and 5-fold cross-validation. The predictive accuracies on the respective validation sets were statistically compared using nonparametric Wilcoxon testing with a significance value of 0.01 with Holm-Bonferroni correction as additional validation sets were statistically compared using nonparametric Wilcoxon testing with a significance value of 0.01 with Holm-Bonferroni correction as previously suggested for Invariant Environment Prediction.\(^\text{24}\) The null hypothesis of statistical independence was rejected below this threshold.

### Data Harmonization

Before cross-study analysis, we aimed to reduce undesired bias caused by differences in imaging protocols while preserving informative variation due to, for example, age-dependent biological effects. To this end, we used the ComBat technique initially described by Johnson et al.\(^\text{11}\) In summary, ComBat achieves batch effect correction by fitting a model to the observed data predicting the features that are to be corrected from the data source (in this case UKBB vs NAKO) and from observed covariates. Subsequently, the contribution of the data source is eliminated obtaining corrected features.

Formally, the value \(Y_{ijf}\) of a feature \(f\) of a participant \(j\) at site \(i\) is modeled as:

\[
Y_{ijf} = \alpha_f + \gamma_f + \beta_f k_j + \delta_f \varepsilon_{ijf}
\]

with \(\alpha_f\) being the feature mean, \(\gamma_f\) the site-specific deviation from the mean, \(\beta_f\) and \(k_j\) regression coefficients and input variables of which the (linear) effect should be preserved, and \(\delta_f\) a site- and feature-dependent scaling factor for the residue \(\varepsilon_{ijf}\) accounting for scaling effects. Harmonized feature values are then computed as:

\[
Y_{ijf}^{\text{corr}} = \frac{Y_{ijf} - \hat{\gamma}_f - \hat{\beta}_f k_j}{\hat{\delta}_f} + \hat{\alpha}_f + \gamma_f^T k_j
\]

preserving the influence of the input variables \(k_j\). As suggested in previous studies,\(^\text{7,25}\) we used a quadratic age-term to also account for nonlinear age-dependent feature variation.

We applied ComBat for harmonization of image features using the data source (UKBB vs NAKO) as the batch variable (of which the effect should be corrected) and based on the previous analysis of the data generating process using age, sex, height, and weight as covariates (of which the effects should be preserved). For ComBat harmonization, we chose UKBB as the reference data set in this study (ie, \(\gamma_f = 0\) and \(\delta_f = 1\) for all image features from UKBB).

### Cross-Study Analyses

Finally, we merged harmonized data from UKBB and NAKO for subsequent large-scale cross-study analyses. Specifically, we investigated age-dependent changes in extracted imaging features and performed multilinear quantile regression (with an additional quadratic age term accounting for nonlinear effects of age) describing the impact of available demographic parameters on image-derived abdominal phenotypes.

### Software

All analyses were performed in Python 3 using the packages Scikit-learn (for random forest implementation, quantile regression, and statistical testing) and neuroCombat (ComBat implementation,

![FIGURE 1](https://www.investigativeradiology.com)
RESULTS

Demographic Data

Image data and related demographic information from a total of 17,996 participants (8791 from UKBB and 9205 from NAKO) were included. Notably, due to different prospective inclusion criteria, participants of UKBB were on average significantly older than NAKO participants with peaks between ages 60 and 70 years in UKBB and around the age of 50 years in NAKO (Table 1, Fig. 1). Participant sex was largely balanced in both studies—a result of a balanced participant recruiting process. We observed similarly shaped empirical joint densities of body height and weight in participants from UKBB and NAKO stratified for sex (Fig. 1). Across data sets, a slight age-dependent decrease in height was observable resulting in slightly lower average height of UKBB participants (Fig. 1).

Image-Derived Features

Overall, the observed marginal densities of image-derived features showed varying degrees of deviation between UKBB and NAKO (Fig. 2A). Organ volumes of liver, spleen, and the kidney showed a slightly higher density in the UKBB cohort compared to NAKO. However, these differences were not statistically significant. The pancreas volume distribution did not show a significant difference between the two cohorts. The liver and spleen PDFF (Percent Fat Free Mass) distributions were similar, while the pancreas PDFF distribution was slightly lower in the UKBB cohort. The r kidney AT volume distribution showed a higher density in the UKBB cohort, while the r kidney volume distribution was similar in both cohorts.

FIGURE 2. Histograms of observed marginal distributions of image-derived features from UKBB (red) and NAKO (blue) before (A) and after (B) ComBat feature harmonization.
FIGURE 4. Exemplary visualization of the effect of data harmonization on cross-study analyses. A, Kernel density plot showing the change of liver proton density fat fraction (PDFF) with age on the original data (UKBB, red; NAKO, blue). B, Kernel density plot showing the change of liver PDFF with age on the harmonized data (UKBB, red; NAKO, blue). C, Kernel density plot showing the change of pancreas PDFF with age on the original data (UKBB, red; NAKO, blue). D, Kernel density plot showing the change of pancreas PDFF with age on the harmonized data (UKBB, red; NAKO, blue). Dashed red and blue lines represent 50% quantile regression separately for UKBB and NAKO data, respectively. Regression lines showed markedly better consistency between studies after feature harmonization.

FIGURE 5. Kernel density plots showing age-dependent changes of image-derived features based on harmonized data for the liver, spleen, pancreas, left (l) and right (r) kidney volume, kidney hilum adipose tissue (AT) volume, as well as liver and pancreas proton density fat fraction (PDFF). Red, UKBB; blue, NAKO. Dashed line, 50% quantile regression line; dotted lines, 25% and 75% quantile regression lines.
FIGURE 6. Representative examples for phenotypic organ alterations with increasing age (A) and with increasing body mass index (BMI) (B). A, Top row: water contrast images; bottom row: relative fat signal images depicting decrease in left kidney volume, increase in left kidney hilum adipose tissue (AT) (top), and increase in pancreatic fat content (bottom) in 6 representative participants (closest to the respective group median values) of increasing age (from left to right: 20–30, 30–40, 40–50, 50–60, 60–70, and 70–80 years). Orange lines mark pancreatic organ borders and the left kidney hilum. B, Top and bottom rows: relative fat signal images depicting increase in liver volume, liver fat content (top), and pancreas fat content (bottom) in 6 representative participants (closest to the respective group median values) of increasing BMI (from left to right: BMI of 15–20, 20–25, 25–30, 30–35, and 35–40 kg/m²). Orange lines mark pancreatic organ borders; asterisks mark the liver parenchyma.
tendency toward higher values in NAKO, whereas measured volumes of kidney hilar AT were slightly higher in UKBB.

Analysis of the Data Generating Process

To further understand these observed feature distribution shifts, we analyzed the data generating process using methods of causal discovery. We were able to use prior knowledge about the causal relation among subsets of observed variables to formulate a partial causal model of the data generating process as a starting point (Fig. 3A).

Further, using nonparametric nonlinear conditional independence testing, we were able to uncover direct causal effects of sex (P < 0.0001), height (P < 0.0001), and weight (P < 0.0001) on observed image features and, importantly, of the image source itself (UKBB vs NAKO, P < 0.0001) on image features. In contrast, no causal effect of the data source could be observed on weight (P = 0.95) or height (P = 0.99) beyond the effect mediated by age (Fig. 3B). These results confirm a direct effect (bias) of the data source (NAKO vs UKBB) on observed image features.

Data Harmonization

Image feature harmonization across studies resulted in a better alignment of empirical marginal feature densities between UKBB and NAKO in a subset of features, particularly for pancreas volume and liver PDFF (Fig. 2B). Interestingly, the above-described distribution shifts between unharmonized features from UKBB and NAKO (Fig. 2A) were slightly even further increased through harmonization in a subset of image features, most pronounced for pancreas PDFF and right kidney AT volume (Fig. 2B). Clearly, this was a result of preserving and enhancing age-related effects through feature harmonization. As shown for liver and pancreas PDFF in Figure 4, feature harmonization resulted in a markedly improved alignment of age-dependent empirical feature densities between UKBB and NAKO and thus enhanced conspicuity of age-related changes in liver and pancreas PDFF.

In a supplemental analysis (Supplemental Material 1, http://links.lww.com/RLI/A787), we assessed the success of data harmonization by predicting the data source (UKBB vs NAKO) based on image-derived features. The underlying rationale is that, after optimal data harmonization, identification of the data source should not be possible better than by random chance. We found that before data harmonization identification of the data source based on image features was possible to a high degree, whereas after data harmonization this classification accuracy was markedly decreased, pointing to successful harmonization of image-derived features (Supplemental Material 1, http://links.lww.com/RLI/A787).

Cross-Study Analyses

Using merged harmonized data from UKBB and NAKO, we assessed age-related changes of image-derived features over a wider age range (20–80 years) than would have been possible for UKBB (50–80 years) or NAKO (20–70 years) alone.

Overall, we observed a marked, nonlinear decrease in organ volumes with age with the steepest volume decline between ages 40 and 80 years. In contrast, volumes of left and right kidney AT compartments increased substantially with age with the steepest increase between ages 40 and 80 years (Figs. 5, 6).

Liver PDFF and pancreas PDFF both increased nonlinearly with age. This age-dependent increase in organ fat content was more pronounced for the pancreas. Regarding hepatic fat content, a slight age-dependent increase was observed, whereas a subpopulation of individuals with markedly increased hepatic fat content appeared after the age of approximately 40 years (Fig. 5).

Finally, joint analysis of harmonized data from UKBB and NAKO allowed us to generate quantitative models of interactions between epidemiological variables and image-derived features. Using quantile regression, we derived median feature values as well as 25% and 75% quantile feature values as a function of age (including a quadratic age term), weight, and height separately for male and female subpopulations. Interestingly, only the quadratic age term and body weight had nonzero coefficients in the final models (Table 2, Supplemental Material 2, http://links.lww.com/RLI/A788). These models provide a unique characterization of the expected phenotypic range of abdominal organ volumes and AT distributions in the investigated populations across a large age range. Beyond age-related changes described previously, these quantitative models revealed a positive effect of body weight on organ volumes and liver and pancreas PDFF of varying degree. Representative examples of abdominal organ phenotypes are shown in Figure 6.

DISCUSSION

In this study, we demonstrated joint, cross-study analysis of imaging data from UKBB and NAKO. We investigated the data generating process and corrected for undesired bias related to the data source. After data harmonization, we performed cross-study analyses characterizing abdominal organ phenotypes in the normal population across a wide age range.

### Table 2. Coefficients of Median (50% Quantile) Regression of Image-Derived Features Separated by Sex

<table>
<thead>
<tr>
<th>Feature</th>
<th>Female</th>
<th></th>
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<tbody>
<tr>
<td></td>
<td>Interc. (%)</td>
<td>Age² (%/y²)</td>
<td>Weight (%/kg)</td>
<td>Interc. (%)</td>
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<tr>
<td>Li PDFF (%)</td>
<td>−5.80E-03</td>
<td>2.80E-06</td>
<td>2.40E-04</td>
<td>Li PDFF (%)</td>
</tr>
<tr>
<td>Pc PDFF (%)</td>
<td>−7.40E-03</td>
<td>1.20E-05</td>
<td>8.60E-04</td>
<td>Pc PDFF (%)</td>
</tr>
<tr>
<td>Inter. (mL)</td>
<td>Li vol (mL)</td>
<td>6.60E+02</td>
<td>−2.10E-02</td>
<td>1.10E+01</td>
</tr>
<tr>
<td></td>
<td>Sp vol (mL)</td>
<td>9.20E+01</td>
<td>−1.10E-02</td>
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</tr>
<tr>
<td></td>
<td>Kd l vol (mL)</td>
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<td>−5.30E-03</td>
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<td></td>
<td>Kd r vol (mL)</td>
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<td></td>
<td>Pc vol (mL)</td>
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</table>

The coefficients for the linear age term and the height term were 0 in all regression models and were thus omitted in this table.

Li, liver; Pc, pancreas; Sp, spleen; Kd l, left kidney; Kd r, right kidney; Kd l AT, left kidney hilum adipose tissue; Kd r AT, right kidney hilum adipose tissue; PDFF, proton density fat fraction; Interc., intercept.
To understand data biases, we investigated the data generating process using a combination of prior knowledge and methods of causal discovery. We found that the data source (UKBB vs NAKO) had a direct effect on image-derived features beyond the effects of age, sex, height, and weight. This source-related bias is most likely the result of differences in the image acquisition process between the studies resulting in acquisition shift.26 Beyond the effects of different imaging protocols, however, it cannot be excluded that unobserved confounders (e.g., differences in ethnicity, lifestyle, or nutrition between UKBB and NAKO participants) mediate additional effects of the data source on image features. Overall, we expect these unobserved effects to be far less significant compared with the direct effects of different imaging protocols on image features.

Cross-study analysis of image features revealed how joint analysis of data from different sources enables a more comprehensive understanding of phenotypic variation. We were able to characterize age-related changes of abdominal organ phenotypes in a way that reflects the majority of the adult population in the United Kingdom and Germany. What has been previously reported for small cohorts with a focus on single organs was possible in this study on a large and representative data set thanks to a combination of a unique large-scale data, automated feature extraction using deep learning and cross-study analysis of harmonized data, grounded in causal analysis of the data generating process. We were thus able to provide quantitative models for abdominal organ volumes as well as abdominal AT distribution (liver PDFF, pancreas PDFF, kidney hilum AT volume). This information can potentially be used for defining normative and reference values also in clinical settings with diagnostic utility. To this end, however, the analysis of all data to be acquired in UKBB and NAKO as well as their joint interpretation with outcome data will be required.

The observed ranges of organ volumes in this study are in accordance with existing literature reports.26–31 Similarly, our findings on AT distribution are comparable to previous reports on liver PDFF,32 pancreas PDFF,33 and kidney hilum AT.34 In contrast to these previous studies, the size of the underlying data combined with the wide age range of participants in our study provide a much more comprehensive and general description of parameter distribution.

This study has limitations. Most importantly, feature extraction can be further improved for a subset of features by using dedicated image sequences available in UKBB and NAKO. For example, the analysis of dedicated multiecho sequences for estimation of liver and pancreas PDFF may increase accuracy for these parameters. Furthermore, the addition of further nonimaging data will allow for a more detailed understanding of the data generating process by considering information about, for example, lifestyle, patient history, or genetic predispositions. We will have to leave these analyses to future studies that can be performed once data collection in UKBB and NAKO are completed.

ComBat normalization (and comparable methods), by design, is performed relative to a reference, which can be one of the included data sets or their weighted combination. Without external calibration, the choice of this reference is not well-defined. In this study, we chose UKBB as the reference data set. The rationale for this choice was the assumption that particularly signal intensity measurements are more robust and less prone to artifacts on a 1.5 T scanner with larger voxel size due to higher field homogeneity and less noise or ghosting artifacts. To resolve the question of the choice of reference more definitely, additional external calibration measurements (e.g., multiecho acquisitions available in UKBB for precise PDFF estimation) will be required in future studies.

In this study, we provided a blueprint of how cross-study analyses can be performed in the context of epidemiological cohort imaging studies and demonstrated the remarkable potential of such analyses.

In conclusion, cross-study analysis of image-derived features from UKBB and NAKO is feasible and can provide unique, population-wide insights into imaging phenotypes and their relation to epidemiological data. Data from UKBB and NAKO harmonized as proposed in this work can be helpful for future joint data analyses across cohorts linking genetic, environmental, and behavioral risk factors to MRI-derived phenotypes and provide reference values for clinical diagnostics.

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