



	Rethinking MRI as a measurement device through modular and portable pipelines
	Agah Karakuzu, Nadia Blostein, Alex Valcourt Caron, Arnaud Boré, François Rheault, Maxime Descoteaux, & Nikola Stikov
Date:	2025
Туре:	Article de revue / Article
	Karakuzu, A., Blostein, N., Caron, A. V., Boré, A., Rheault, F., Descoteaux, M., & Stikov, N. (2025). Rethinking MRI as a measurement device through modular and portable pipelines. Magnetic Resonance Materials in Physics, Biology and Medicine (MAGMA), 38, 423-439. https://doi.org/10.1007/s10334-025-01245-3

Document en libre accès dans PolyPublie Open Access document in PolyPublie

URL de PolyPublie: PolyPublie URL:	https://publications.polymtl.ca/64682/
Version:	Version officielle de l'éditeur / Published version Révisé par les pairs / Refereed
Conditions d'utilisation: Terms of Use:	Creative Commons Attribution 4.0 International (CC BY)

Document publié chez l'éditeur officiel Document issued by the official publisher

	Magnetic Resonance Materials in Physics, Biology and Medicine (MAGMA) (vol. 38)
Maison d'édition: Publisher:	
URL officiel: Official URL:	https://doi.org/10.1007/s10334-025-01245-3
Mention légale: Legal notice:	This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate Magnetic Resonance Materials in Physics, Biology and Medicine if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

REVIEW



Rethinking MRI as a measurement device through modular and portable pipelines

Agah Karakuzu^{1,2} · Nadia Blostein⁶ · Alex Valcourt Caron³ · Arnaud Boré³ · François Rheault³ · Maxime Descoteaux³ · Nikola Stikov^{1,2,4,5}

Received: 30 November 2024 / Revised: 27 February 2025 / Accepted: 11 March 2025 / Published online: 24 April 2025 © The Author(s) 2025, corrected publication 2025

Abstract

The premise of MRI as a reliable measurement device is limited by proprietary barriers and inconsistent implementations, which prevent the establishment of measurement uncertainties. As a result, biomedical studies that rely on these methods are plagued by systematic variance, undermining the perceived promise of quantitative imaging biomarkers (QIBs) and hindering their clinical translation. This review explores the added value of open-source measurement pipelines in minimizing variability sources that would otherwise remain unknown. First, we introduce a tiered benchmarking framework (from black-box to glass-box) that exposes how opacity at different workflow stages propagates measurement uncertainty. Second, we provide a concise glossary to promote consistent terminology for strategies that enhance reproducibility before acquisition or enable valid post-hoc pooling of QIBs. Building on this foundation, we present two illustrative measurement workflows that decouple workflow logic from the orchestration of computational processes in an MRI measurement pipeline, rooted in the core principles of modularity and portability. Designed as accessible entry points for implementation, these examples serve as practical guides, helping users adapt the frameworks to their specific needs and facilitating collaboration. Through critical evaluation of existing approaches, we discuss how standardized workflows can help identify outstanding challenges in translating glass-box frameworks into clinical scanner environments. Ultimately, achieving this goal will require coordinated efforts from QIB developers, regulators, industry partners, and clinicians alike.

 $\textbf{Keywords} \ \ MRI \ workflows \cdot Reproducibility \cdot Standardization \cdot Metrology \cdot Vendor-neutral \cdot Quantitative \ MRI$

Agah Karakuzu and Nadia Blostein contributed equally.

✓ Nadia Blostein nadia.blostein@gmail.comNikola Stikov nikola.stikov@polymtl.ca

- NeuroPoly Lab, Polytechnique Montreal, Montreal, Québec, Canada
- Montreal Heart Institute, University of Montreal, Montreal, Québec, Canada
- Sherbrooke Connectivity Imaging Laboratory (SCIL), Computer Science Department, Université de Sherbrooke, Sherbrooke, Québec, Canada
- Center for Advanced Interdisciplinary Research, Ss. Cyril and Methodius University, Skopje, North Macedonia
- NYUAD Research Institute, New York University Abu Dhabi, Abu Dhabi, UAE
- ⁶ School of Medicine, University Collage Cork, Cork, Ireland

Introduction

"The principal task of a measurement system is the sensing of the measured value of a physical quantity or of a measurement signal representing the wanted measured value."

Basics of measurement technology—Part 2: Terms for measuring instruments [1].

When evaluated against this standard, MRI scanners are not considered true measurement systems. Their intended use lacks the metrological rigor required for quantitative applications in clinical settings [2, 3]. As such, they are clinical tools designed for qualitative imaging, not for quantitative science. Nevertheless, measurements derived from MRI are integral to most research in the field.

Quantitative imaging biomarkers (QIBs) are numerical characteristics derived from quantitative imaging techniques [4]. However, specific definitions of MRI measurements



depend on context. For instance, quantitative MRI (qMRI) can measure the relaxation time constants T1 and T2 (in seconds). Such *physics-based QIBs* are obtained by producing systematic variations in voxel values, which can then be fitted to a signal representation. Translating complex tissue properties into measurable QIBs necessitates a more layered modeling approach [5], such as the quantification of white matter microstructure with diffusion MRI (dMRI) [6], or g-ratio by combining dMRI, magnetization transfer or relaxometry imaging [7].

On the other hand, the measurement concept goes beyond purely quantitative frameworks, incorporating quantifiable features extracted from morphological characteristics in imaging data. These *data-driven QIBs* are generated by computational image analysis methods, such as radiomic feature extraction [8, 9] using conventional processing or deep learning [10]. For example, tumor volume (mm³) qualifies as a data-driven QIB [4].

All of these measurement procedures ultimately boil down to three main methodological steps: pulse sequence implementations (*Acq*), image reconstruction algorithms (*Recon*), and downstream processing methods (*Post*). For the rest of this article, we use *workflow* for the overall process and *pipeline* for the software that automates it.

Setting aside the impact of inevitable variations in experimental conditions (e.g., hardware differences) and random factors, the generalizability of these three primary steps (Acq, Recon, and Post) is crucial to the success of a QIB. Nevertheless, most MRI software development and execution environments are segregated by vendor-specific boundaries, which makes it a challenge to control these three main methodological steps across various experimental settings. In addition to triggering the well-known "but it works on my machine!" syndrome, the isolated and proprietary nature of commercial MRI development environments often hinders reproducibility and collaboration between imaging sites. This fragmentation creates significant barriers to innovation and hampers the consistency of QIBs in research and clinical applications.

For example, a recent editorial on radiomics identified this issue as the elephant in the scanner room that must be addressed for data-driven QIBs [11]. Authors emphasized that trustworthy radiomic features should not be influenced by the texture variations of different scanners, to the point where one could infer the vendor based on the images. The inevitable capture of these patterns by deep learning applications clearly impacts the performance and generalizability of the developed models trained on multicenter data, primarily due to a phenomenon known as domain shift [12]. To compensate for such effects when multiple large-scale datasets are pooled in search of true biological variability, statistics or deep learning based data harmonization methods [13] for normative modeling [14] have become increasingly popular.

Nevertheless, this drawback is not unique to data-driven QIBs. For nearly 50 years, physics-based QIBs have been suggested as a powerful alternative to improve the consistency of qualitative pattern recognition (e.g., using T2 to eliminate inter-site contrast variations in T2-weighted images due to protocol differences [15]) and to bring about a whole new era of precision MRI (e.g., leveraging T2 values themselves to establish normative thresholds for routine diagnostics [16]). However, this aspirational vision for physics-based QIBs—that they might serve as universal scalars to ease radiological interpretations across centers and/or as reliable tools to convert such interpretations into an objective numbers game—remains out of reach.

The past two decades in MRI research have seen synergistic forces to lift QIBs from the swamp of variability: grassroots open-source development [17-19] and more concerted efforts like the Quantitative Imaging Biomarkers Alliance (QIBA), which united, clinicians, industry partners, and regulatory scientists with the mission of reducing variability from QIBs [20]. While independent groups create open-source solutions targeting specific workflow steps, QIBA established profile stages that evaluate a QIB's ability to support its stated claim across four levels of confidence [21]. As of today, only MR elastography of the liver [22] and the apparent diffusion coefficient [23] are listed as level three (clinically feasible), while cerebral blood volume [24], T1_p and T2 for cartilage [25], and dynamic contrast enhancement quantification [26] are designated as level two (consensus) profiles. These are promising developments that have emerged over the past five years. Undoubtedly, the next iterations of QIBA [27] will continue to build on its previous successes.

The momentum of open-source community initiatives is a powerful driver in advancing QIBs. To leverage this momentum effectively, we must implement overarching strategies throughout the development process. In doing so, respecting the boundaries of regulatory requirements and commercial interests is key to landing innovations in clinical practice. Playfully captured in the ISMRM 2024 Lauterbur Lecture by Andrew Webb [28], a chicken-egg dilemma arises from the conflict between the open-science hype and skepticism. This debate playfully highlights the need to balance enthusiasm for open science with the caution required to tackle its practical challenges. We believe the first step to achieving this balance is to clearly determine how open a QIB must be for its claim to hold. To this end, we introduce a tiered benchmarking framework for MRI measurement workflows by borrowing the concepts of software testing implementations [29]: black-box, gray-box, frosted-box, and glass-box (Fig. 1). The next crucial need is clearly communicating the methods implemented to reduce the variability from MRI measurements. To support this, we have created a mini glossary to foster consistent use of relevant terminology (Fig. 2).



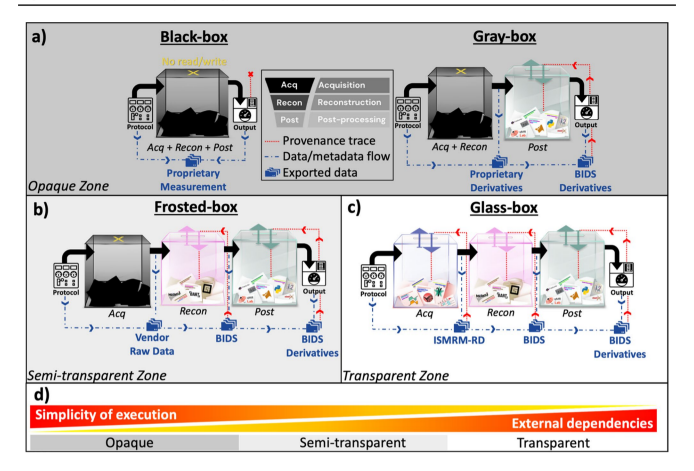


Fig. 1 Benchmarking of MRI measurement workflows based on the openness of their key steps: acquisition (*Acq*), reconstruction (*Recon*), and downstream post-processing (*Post*). The extent of traceability (red dashed lines) of the generated data flow (blue dashed lines) are indicated along with the preferred use of community data standards, wherever possible. **a** The opaque zone encompasses black- and gray-

box workflows, where Acq and Recon steps are hindered. **b** The semi-transparent zone corresponds to frosted-box workflows, which limit openness at the Acq step. **c** The transparent zone requires glass-box workflows, ensuring full openness across all MRI measurement steps. **d** Variations in execution simplicity and external dependencies across transparency zones

As an overarching strategy for creating interoperable, modular, and reproducible measurement pipelines that can propel QIBs through levels of confidence, we propose the use of NextFlow [30] in tandem with well-maintained containers and community data standards. We also provide two example pipelines to illustrate this approach and further discuss implementation challenges and the latest solutions for bringing QIBs closer to practical use.

Benchmarking the openness of MRI measurement workflows

MRI workflows can be categorized into four benchmarks distributed across three zones of transparency:

Opaque zone: When the internal implementations of MRI workflows (*Acq* and *Recon*) are unknown and inaccessible (i.e., no read/write access).

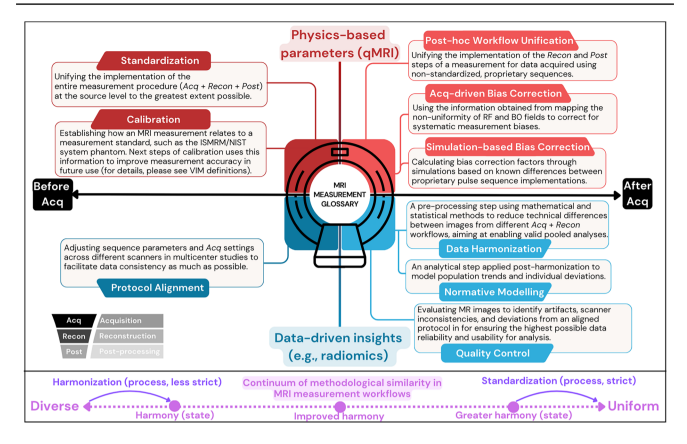
Black-box benchmark: No visibility into any key step.

Gray-box benchmark: Provides access to *Post* but remains closed at earlier stages (*Acq* and *Recon*).

Semi-Transparent zone (Fig. 1b): A vendor-native acquisition followed by open-source *Recon* and *Post*.

Frosted-box benchmark: This partial transparency allows limited modifications to on-site steps of an MRI measurement.





ment bias.

Fig. 2 Infographic defining MRI measurement terms used consistently throughout this review. The horizontal axis separates terms based on their relevance before or after acquisition (k-space sampling), while the vertical axis indicates whether their use is geared towards physics-based parameter quantification (i.e., qMRI) or data-driven insights (i.e., radiomics). The continuum of methodological

similarity in MRI measurement workflows, based on the definitions by [48]: standardization involves the unification of all steps (*Acq*, *Recon*, and *Post*) to converge towards uniformity, whereas harmonization aims at moving away from total diversity to achieve better harmony. Please refer to VIM and Kessler et al. [4] for a more comprehensive list of terms relevant to MRI measurements [4, 47]

Transparent zone: (Fig. 1c): When all implementations (*Acq*, *Recon*, and *Post*) are fully accessible for both viewing and editing.

Glass-box benchmark: Represents the ideal case of full visibility and control over MRI workflows.

Making sense of numbers within the opaque zone of MRI measurements

Freed from the complexity of setting up third-party execution environments and dependencies at the scanner site, black-box workflows are often the most user-friendly choice (Fig. 1d). They offer turnkey measurements whose complexities are isolated from the end users by proprietary barriers. An alternative to this monolithic architecture is offered by typical gray-box workflows that offload *Post* tasks to a remote server to extract measurements from vendor reconstructed images. When equipped with advanced

data management and off-site network solutions, gray-box measurement workflows can also remove the complexity away from the end user's experience. Nevertheless, these conveniences come at the expense of dialing into the implementation details to identify potential sources of measure-

Case-1: A notable black-box example is the comparison of cardiac T1 maps produced by the proprietary implementations of the MOLLI, shMOLLI, and SASHA sequences [31]. Reliable observation of the T1 changes would help quantify the severity of conditions such as myocardial iron overload (lower T1) and edema (higher T1) [32]. However, when the measurement bias between different methods (or between implementations of the same method across vendors) exceeds the differences observed between diagnostic groups, the clinical utility of the measurement becomes compromised.

Other common examples of commercially available quantification packages are fraction measurement methods, typically used for investigating musculoskeletal and hepatic anomalies [33, 34]. As noted by Cashmore et al. (2021), the



main issue with such proprietary packages is their infeasibility for modeling uncertainty propagation (through *Acq*, *Recon*, and *Post*), rendering it impractical to determine measurement uncertainty [2].

Case-2: Notably, the lack of inter-vendor agreement in MRI measurements is not specific to a single domain and black-box workflows. Similar challenges arise in other gray-box benchmarks, such as whole-brain T2 mapping using dual-echo FSE data from the ADNI dataset [35], even when protocol alignment and phantom-based quality control (Fig. 2) are carried out in a highly regimented manner [36].

This shortcoming remains evident across a spectrum of measurements, from the most fundamental T1 mapping methods [37, 38] to fractional anisotropy (FA) in diffusion tensor imaging (DTI) [39], and more sophisticated microstructure imaging approaches [40]. While these examples illustrate the issue, they are far from exhaustive, as many other cases follow a similar pattern [41].

Under more carefully controlled multi-site conditions managed by the same research group, the reproducibility of gray-box workflows can be improved. A notable example is proton-density fat fraction, where each major vendor's implementation of chemical-shift encoded spoiled gradient echo was used, followed by a standardized *Post* procedure [42]. A similar outcome was also apparent from a global reproducibility challenge for a gold-standard T1 mapping method [37] and MR fingerprinting [43].

Another common roadblock researchers encounter is the challenge posed by proprietary system upgrades. These upgrades may introduce variability that is difficult to account for, even within the same site on the same MRI system [44]. For example, it has been shown that the T1 parameter varies as much as 30% after a scanner upgrade, and a large part of this variation can be attributed to B1 effects. However, B1 calibration is often done at the prescan level, and changes in the RF transmit gain are rarely disclosed by the vendors. The combination of all these factors make the selection of a proper bias correction method a challenge of its own [45]. Note that there is also the impact of variations from the Acq, Recon, and Post steps. Even with full transparency, these steps may not protect the measurement from variations arising from deeper-level system modifications that the acquisition controllers depend on [46].

A glossary for navigating the zones of MRI measurements

Correcting for the non-biological variance commonly observed in MRI measurements is a non-trivial effort—one that has taken on a life of its own, giving rise to a nomenclature that is increasingly intricate and sometimes difficult

to parse. Therefore, we provide a glossary of terms (Fig. 2) and use them consistently throughout this review to clearly distinguish between different approaches striving for consistency in MRI measurements.

It is important to note that both QIBA and the International Vocabulary of Metrology (VIM) offer comprehensive terminology relevant to measurements [4, 47]. Rather than redefining these established terms, our glossary builds upon them, providing additional clarification within the specific context of this review.

The terms standardization and harmonization are used rather loosely by MRI and neuroimaging researchers to refer to the process of "removing scanner effects" either before or after data collection. Even though these two terms are closely related, they actually represent distinct concepts. In the field of metrology, harmonization is an overarching term used to describe ensuring consistency of values measured by different methods [49]. On the other hand, standardization is a more formal and rigorous approach to harmonization, where consistency is assured by making sure that different machines measure the same quantity in the same way, using reference measurement procedures and objects [50]. Even in fields unrelated to MRI and measurement, a similar distinction applies: harmonization is the process of moving away from total diversity of practice (i.e., achieving harmony by reducing variance), while standardization describes a movement toward uniformity (i.e., the removal of variance at its source) [48, 51].

The definitions outlined in Fig. 2 are closely linked to their intended applications—whether physics-based or data-driven measurements—as well as the timing of their use relative to data acquisition (i.e., pre- or post-acquisition). In the subsequent sections of this review, we illustrate each term from Fig. 2 with practical use cases organized according to these classifications.

Opening up the implementations, that's how the light gets in

In earlier sections, we presented two examples—Case-1 for cardiac T1 mapping and Case-2 for brain T2 mapping—that highlight the challenges of operating in the opaque zone of MRI measurements. In this section, we introduce two approaches taken by independent research groups to address these challenges and highlight additional efforts that tackle this issue at various levels.

Case-1: Open-MOLLI introduced a glass-box solution that enabled informed comparisons of this specific cardiac T1 measurement procedure across different scanners and vendors [52]. Through *standardization* (Fig. 2), Gaspar et al. (2024) demonstrated improved within-site repeatability, while providing a foundation for further



advancements in multi-site studies and enhancing the reliability of MOLLI. Furthermore, the open-source availability of Open-MOLLI nucleates the development of other cardiac T1 mapping methods—such as SASHA and shMOLLI—on any vendor's system, given that the sequence implementations share similar patterns for sampling the longitudinal relaxation recovery curve.

Standardizing MRI measurement workflows through glass-box implementations provides the state-of-the-art approach for systematically comparing measurement procedures across different systems. This standardization can even be leveraged to evaluate the performance of systems expected to operate identically. Keenan et al. (2025) demonstrated this by deploying a standardized relaxometry workflow on both a commercial low-field 0.55 T system and a prototype system ramped down from 1.5 T to 0.55 T [53]. Their findings showed no significant differences between the prototype and commercial systems for the tested relaxometry measurements, offering a powerful example of the confidence that standardized measurement workflows can provide.

In addition to enabling multi-center collaboration and providing generalizable templates that help MRI researchers build on each other's work [54], standardized workflows significantly enhance inter-vendor agreement in measurements. The first evidence in support of this claim is presented by Karakuzu et al. (2022), reporting up to 23% reduction of inter-vendor bias for the MTsat [55] multiparametric mapping method [56]. Even though this was achieved using a proprietary vendor-neutral operating system [57], following standardized workflows developed using Pulseq [58] demonstrated similar effectiveness. For a standardized single-shell diffusion MRI workflow, a nearly 2.5 fold reduction of standard error in diffusion indices was reported using Pulseq [59], building on the initial efforts by Nunes et al. (2020) that made the sequence implementation available [60]. Within the same reproducible framework, the application of time-division multiplexing led to more accurate relaxometry and diffusion measurements [61]. More recently, extending the application of Pulseq-based diffusion imaging to cardiac studies has led to significant SNR gains across scanners from different vendors [62].

Case-2: Even when the implementation cannot be opened up, obtaining some information on the precise timing and component attributes of a pulse sequence can be valuable. Fortunately, vendors may be willing to cooperate with researchers to disclose such information within the framework of their legal guidelines. With these priors, differences between the nominal and actual (physical) cross-vendor effects can be identified by devising mathematical relationships to derive respective correction factors. In addressing Case-2, a successful implementation of this strategy is

provided by Chehetri et al. (2021) using Bloch modeling for retrospective correction of T2 bias in the ADNI dataset [63]. In this particular case, the *simulation-based bias correction* (Fig. 2) was performed within the gray-box benchmark, given that the source dataset is reconstructed in a proprietary pipeline.

Similarly, Rowley et al. (2021) developed a simulation framework that models the impact of magnetization transfer pulses, which vary between vendors, for sequence-informed removal of B1 effects from the MTsat maps [64]. Another gray-box bias correction method aimed to control these saturation effects at the sequence implementation level, guided by simulation-based insights [65]. Other examples of simulation-based bias correction include [66–68]. A more common approach to post-hoc correction of systematic bias is acquiring field maps, adding yet another layer of MRI measurement [69]. Consequently, field maps used for Acqdriven bias correction (Fig. 2) are subject to the same limitations as QIBs. Lee et al. [70] demonstrated that, despite their corrective intent, different B1⁺ mapping implementations can actually exacerbate inter-vendor variability in T1 measurements [70]. Nevertheless, B1 and B0 mapping remain essential for the viable application of numerous QIBs, such as the variable flip angle framework and diffusion imaging, respectively.

Beyond the domain of MRI physics, data harmonization represents a post-hoc methodological approach aimed at improving intersite comparability of data-driven QIBs (Fig. 2). This strategy seeks to retrospectively mitigate variability arising from differences in acquisition protocols, scanner hardware, and reconstruction pipelines, thereby facilitating pooled analyses of large-scale multisite datasets [13]. Recently, Warrington et al. [13] recommended the combined application of (i) implicit harmonization, determining the optimal pipeline that is the most immune to the scanner effects for a given set of QIBs (referred to as imagederived phenotype by the authors) and (ii) explicit harmonization, which leverages intra-site measurements of QIBs under repeatability conditions to estimate global scaling factors that correct for the said scanner effects. In addition to providing a valuable comparison framework for harmonization methods, the authors reported comparable performance of ComBat [71] and CovBat [72] in reducing inter-scanner variability across cortical and subcortical volumes, as well as in physics-based QIBs of T2* and diffusion indices.

The ComBat has been successfully applied to enhance the statistical power of a mega-analysis using MRI data from 33 sites and 6000 subjects to distinguish healthy participants from schizophrenia patients [73]. In addition to structural and diffusion data, it was also applied to MR spectroscopy dataset, revealing so-called biological effects that would have otherwise remained hidden [74]. CovBat extends ComBat by accounting for covariance structures



among harmonized features [75]. A recent CT radiomics study demonstrated that CovBat outperforms ComBat in this regard [76]. Other data harmonization approaches exist, such as RELIEF, which has been reported to surpass both ComBat and CovBat [77]. Most applications of these methods in diffusion MRI have focused on derived indices. However, De Luca et al. [78] demonstrated the feasibility of harmonizing reconstructed diffusion images directly [78].

In addition to statistics-driven harmonization methods, recent literature has seen a growing number of deep learning-based approaches, such as DeepHarmony [79] and MIS-PEL [80]. More recently, PhyCHarm introduced a physics-informed constraint by integrating Bloch simulations into the model [81]. For a more comprehensive analysis of data harmonization methods, the reader is referred to [75, 82–85].

A practical guideline for streamlining portable MRI measurement pipelines

A key consideration in QIB development is balancing complexity and modularity. As summarized in Fig. 1d, modular design enables greater flexibility when each step of the measurement workflow is accessible. However, this openness comes at the cost of increased external dependencies, which can, in turn, reduce the simplicity of execution.

To address Webb's chicken-egg dilemma, a good starting point is to avoid putting all the eggs in one basket. Container-mediated workflow orchestration frameworks address this problem because they simplify the management of complex computational pipelines by breaking them down into smaller, manageable tasks. One such workflow engine is Nextflow [30]: a domain-specific framework for bioinformatics, genomics, and medical imaging, designed to develop scalable, reproducible, and portable data-intensive computational pipelines.

Basics of nextflow

Nextflow combines a declarative domain-specific language (DSL) with a reactive dataflow model. The reactive model enables Nextflow to dynamically manage data flow in response to changing data streams, eliminating the need for explicit callbacks to handle updates. For instance, a Nextflow pipeline can be linked to the raw data stream of an MRI scanner, automatically triggering reconstruction processes based on predefined patterns.

Its declarative nature enables users to define what their pipeline should achieve rather than how to execute it. Nextflow manages parallel execution, error recovery, and workflow dependencies, allowing users to focus on data transformations and file-naming conventions. This abstraction ensures seamless scalability across diverse platforms—including any operating system, cloud environments, or high-performance computing clusters—without requiring changes to the pipeline's logic.

Nextflow can be installed on any POSIX-compatible system with a single command. Pipelines are defined in scripts using Nextflow's DSL in scripts with the.nf extension. Comprehensive documentation on DSL is available at https://nextflow.io, and https://training.nextflow.io serves as an excellent resource for getting started. For the sake of completeness, below is a high-level overview of how Nextflow pipelines are structured within an MRI-specific context:

To build a Nextflow pipeline, users define tasks as *processes*—self-contained code blocks that specify inputs (e.g., structural MRI files like *sub-*_Tlw.nii.gz*), outputs (e.g., defaced images such as *sub-*_desc-defaced_Tlw.nii.gz*), and a script (e.g., *pydeface \$input*). Inputs are ingested from *channels*, which dynamically match regex patterns (similar to wildcards for BIDS-structured datasets) and stream data into processes. For instance, a channel matching *sub-*_Tlw.nii.gz* can automatically feed all participant scans into a defacing process, with outputs systematically organized into a BIDS-compliant derivatives directory.

By default, Nextflow delegates command execution to the underlying environment, meaning external packages like PyDeface [86] must be installed and accessible (e.g., available in the system PATH for local execution). Note that alternative approaches exist to bypass local dependency management, as detailed later. Connecting an array of *processes* through *channels* in a.nf script creates a pipeline description (e.g., *my_pipeline.nf*).

Finally, running Nextflow on a pipeline (e.g., *nextflow run my_pipeline.nf*) initiates execution of all the tasks based on the data channels that become available. Nextflow also provides interactive HTML reports, real-time tracing, directed acyclic graph (DAG) visualization, and execution logs for monitoring and debugging.

A simple update to the pipeline configuration file, *nextflow.config*, is all it takes to switch between supported executors. For instance, setting *process.executor='awsbatch'* enables job submission in the cloud without the need to manually provision or manage a virtual machine cluster.

DSL2: a modular framework

Nextflow's original DSL constrained pipelines to monolithic scripts, requiring error-prone code duplication or complex channel logic to reuse components. With the introduction of DSL2, Nextflow revolutionized workflow design by enabling native modularity: processes, subworkflows, and functions



can now be encapsulated in separate, reusable files. This paradigm shift is particularly transformative for MRI measurement pipelines, where evolving tools and analyses demand adaptable, compartmentalized workflows.

DSL2 allows critical pipeline components—such as preprocessing steps, quality control checks, or analysis modules—to be stored in dedicated files (e.g., modules/preprocess.nf, subworkflows/qc.nf) and imported via the include directive. This modular architecture enforces strict input/ output interfaces, clarifies data flow, and supports independent versioning of components (e.g., through GitHub or the nf-core module registry). Teams can now develop, test, and update pipeline stages in isolation before integrating them into larger workflows, streamlining collaboration.

These advancements are pivotal for MRI measurement pipelines, where evolving open-source reconstruction tools, quality control checks, and analysis steps demand adaptable, compartmentalized workflows. Unlike Snakemake [87], which relies on external templates for reuse, or Nipype and Airflow's rigid DAGs [88, 89], DSL2 natively supports dynamic, hierarchical workflows. For a more comprehensive comparison of pipeline management systems, the reader is referred to [90].

Key principles for optimizing nextflow in QIB development

Although most of Nextflow's advantages come out of the box, there is a need to establish a set of principles to fully leverage its potential in QIB development:

 Apply one process to one container mapping for modular dependency management at ease

Each process in a Nextflow pipeline can be linked to a Docker or Singularity container either by specifying the container directly within the process (Fig. 3) or by defining the corresponding image name in the nextflow.config file. This enables the pipeline to execute the process within an isolated, reproducible environment that includes all necessary dependencies, in contrast to the default approach, which searches for executables in the local system environment.

When combined with the modularity introduced by DSL2, the following example outlines the high-level steps of this principle from the user's perspective:

 Include motion_correction process from ANTs subworkflow



Fig. 3 Example Nextflow snippets demonstrating how different processes—FSL (red, *fsl_bet.nf*) and ANTs (green, *ants_brain_extraction.nf*)—performing equivalent tasks through standardized input/out-

put interfaces, enabling zero-maintenance swaps in the reconstruction pipeline (main.nf)



- 2. Pull qmrlab/antsfsl image from Docker Hub
- 3. Edit *nextflow.config* (or the process block) to link *motion_correction* process to the *qmrlab/antsfsl*
- 4. Repeat 1–3 for *wm_segment* process from FSL subworkflow using the same container
- Repeat 1–3 for mt_sat from qMRFlow using the qmrlab/ minimal container
- 6. Define the dataflow between these processes to define the pipeline
- 7. Execute the pipeline

Instead of consolidating all these tools into a single monolithic container, the modular approach offers several advantages:

Simplified dependency management: Users can seamlessly pull pre-configured images maintained by their developers, allowing Nextflow to orchestrate them effortlessly.

Conflict-free dependencies: Isolating processes within dedicated containers eliminates dependency conflicts. Developer autonomy: Each development team can focus exclusively on their specific runtime environment.

Avoidance of excessive wrappers: This methodology discourages the common but inefficient practice of layering academic software packages with too many wrappers.

2) Use data standards to enable declarative interlinking of the workflow processes

Nextflow's reactive dataflow model relies on file patterns to dynamically populate channels. Therefore, community standards like ISMRM-RD [91], BIDS [92], and BIDS derivatives [93] act as structural backbones by enabling:

Declarative routing: Naming conventions (e.g., BIDS sub-*_ses-*_Tlw.nii.gz) enable automatic file discovery and channel population without hardcoded paths. For example, a motion correction process can declare it consumes *_Tlw.nii.gz files, trusting they follow anatomical scan conventions.

Interface contracts: Standards define clear input/output relationship between workflow stages. For example, a BIDS derivatives rule that segmentation masks follow *_label-WM_probseg.nii.gz lets subsequent processes reliably consume these outputs regardless of which tool generated them.

3) Swap modules, do not reinvent them

Once data standards and containerization are established, pipeline modules become interchangeable "LEGO blocks". Figure 3 demonstrates, in DSL2, how one can easily replace one *Post* method with another due to adherence to these principles.

This approach can drastically reduce code complexity, particularly when the pipeline description includes all steps. Figure 4 expands on the previous DSL2 example with a schematic representation, demonstrating how numerous pipeline variations (3⁴ = 81 possible pipeline alternatives) can be generated by leveraging this easy swap feature. The ultimate vision is the creation of decentralized, portable workflows within a BIDS-apps like framework [94]. Nevertheless, unlike platforms like Brainlife [95], CBRAIN [96], or OpenNeuro [97], these workflows are designed for seamless execution across diverse infrastructures, making them essential for deployment at the point-of-care.

Glass-box examples with qMRFlow

qMRFlow is a suite of container-mediated, data-driven and transparent qMRI pipelines, ideally starting with a vendor-neutral *Acq* [57, 58, 98]. However, when preceded by a vendor-native acquisition, the pipeline transitions from a *glass-box* to a *frosted-box* benchmark (Fig. 1c), shifting the respective strategy from *standardization* to *post-hoc work-flow unification* (Fig. 2).

Powered by Nextflow, qMRFlow adapts the principles outlined in this review to enable reproducible and modular measurements. Its modules are developed to work primarily with qMRLab [30, 99], which implements a wide range of qMRI methods, including relaxometry, magnetization transfer and diffusion imaging, field mapping, and susceptibility mapping which are essential for microstructural characterization of in-vivo tissue, particularly in neurological applications.

While qMRLab offers core functionality for processing quantitative MRI data, qMRFlow enhances this by seamlessly integrating with nearly all *Recon* and *Post* software. Additionally, qMRFlow's glass-box compatibility is a significant benefit. This compatibility was first demonstrated in [56], where qMRFlow was integrated with RTHawk [57]. The source code is available at https://github.com/qmrlab/venus, with an overview provided below:

 The modules folder in this repository contains several Nextflow scripts, most of which implement methodspecific processes (e.g., mt_sat.nf) that utilize qMRLab. Additionally, bids_patterns.nf provides helper functions to manage BIDS-specific directory structures and rules.



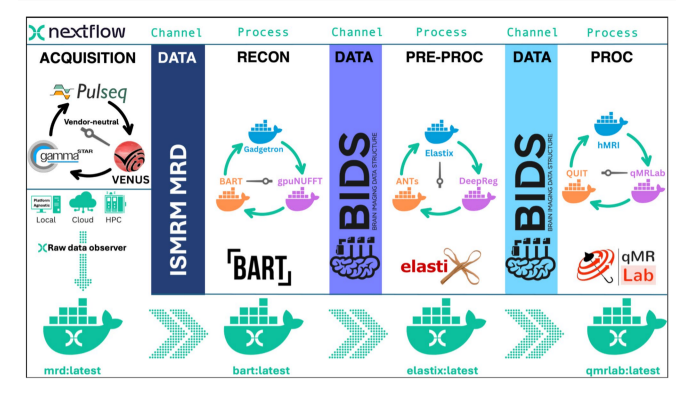


Fig. 4 This illustration captures one of many possible (3⁴=81 in total) process combinations—such as Pulseq, BART, Elastix, and qMRLab—executing within their dedicated containers in isolation from each other. Each Nextflow channel's data flow is guided by

established data standards and derivative conventions and the modules can be easily swapped thanks to DSL2. Platform agnostic Nextflow executors allow for running the pipeline on a local workstation, cloud computing platforms, or high-performance computing clusters

- There are two main pipeline descriptions that make use of these modules: *venus-process-invivo.nf* and *venus-process-phantom.nf*. As their file names suggest, these workflows share certain processes, such as the *fitMtsat* process from *mt_sat.nf*. However, only the in-vivo pipeline imports the *generateRegionMasks* process from *ants. nf*, as this step is not required for phantom data.
- The *nextflow.config* file starts with the specification of resource allocations, including the number of CPUs, memory limits, and the maximum number of parallel runs. It also defines container configurations, linking each process to the appropriate software environment. It follows with a code block that enables the use of docker, which is then followed by process-specific parameters such as *use_blcor* that can be changed by the users to alter the dataflow.
- Since the pipeline follows qMRI-BIDS, data channels can be structured using standardized patterns, simplifying the integration of processes and ensuring seamless workflow connectivity.

To repeat the *Recon* and *Post* steps offline, users simply provide a root folder containing multiple subject datasets, and qMRFlow automatically organizes and executes the

necessary preprocessing and analysis steps. Furthermore, qMRFlow's integration into the broader imaging neuroscience ecosystem, including connections to next-generation platforms like NeuroLibre [100], facilitates large-scale, multi-center neuroimaging studies with enhanced reproducibility and ease of use.

A more recent demonstration of qMRFlow's glass-box compatibility is shown through its integration with Pulseq [58]. This study leveraged the open-source MPRAGE implementation provided by Pulseq to develop an end-to-end MP2RAGE [101] pipeline, deployed it to five scanners from three different vendors [102].

Gray-box examples with tractoflow (vendor-native data)

The TractoFlow pipeline processes dMRI dataset from the raw data to the tractography as a modular entity. Tractoflow was quickly adopted by researchers outside of the immediate network of developers. Powered by Nextflow DSL1 and containerization, here are the five main highlights of Tractoflow (Fig. 5):



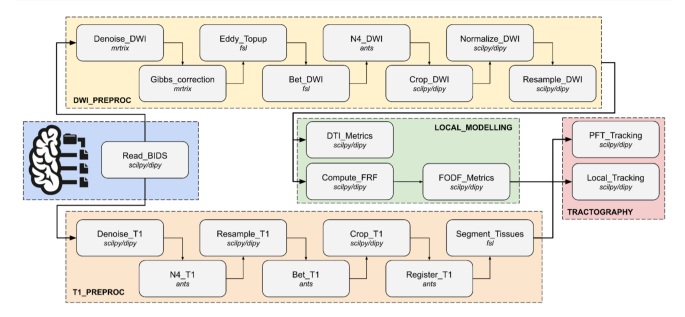


Fig. 5 This figure represents the TractoFlow pipeline, designed as a gray-box workflow: it uses BIDS-formatted inputs from scanner reconstructions. It outputs a variety of files, ensuring reproducibility when inputs are identical. Each box in the diagram denotes a modular step, capable of saving outputs and configured for durability using dedicated Docker containers. Parameters are fully accessible and cus-

tomizable through a comprehensive configuration file, allowing users to adapt the pipeline for diverse applications (e.g., healthy brain, tumor analysis, aging studies). Steps like tissue segmentation can be reconfigured to use tools like FreeSurfer, aligning with the gray-box principles

- Efficient diffusion MRI processing pipeline from raw (i.e., vendor-reconstructed) diffusion data to tractography, which has allowed processing of the HCP [103, 104], UKBiobank [105], ADNI [36], PPMI [106], ABCD [107], and most large open-access databases by non-developers and non-experts in dMRI.
- The stochastic nature of the Tractoflow tractography pipeline makes reproducible results particularly challenging. Efforts are made to reduce variability when relaunching Tractoflow to ensure reproducible and replicable results today, tomorrow, and over time [108].
- Tractoflow provides a tradeoff between customization and curation of configuration files, making it easy-to-use for non-technical and clinician users. Profiles are provided to adapt to different use cases (aging, development, white matter hyperintensities, amongst others [108]).
- Little to no installation steps and compatibility with High-Performance Computing thanks to containers and Nextflow executors (e.g. SLURM, AWS).
- Support for multiple vendors through BIDS.

Tractoflow includes more than 20 steps that are automatically distributed depending on their dependencies and requirements and grouped into two main blocks of processes (see Fig. 3). The first is diffusion MRI pre-processing, local reconstruction of DTI and constrained spherical deconvolution (CSD), and anatomically constrained deterministic or probabilistic tractography. The second involves the T1

pre-processing and T1 analysis to register to the final diffusion MRI space and extract white matter, gray matter and CSF partial volume maps. The Tractoflow container is based on scilpy [109], dipy [110], mrtrix3 [111], ANTS [112], FSL [113] and freesurfer [114].

From the Tractoflow output tree, there are a series of optional flows available to automatically extract the main white matter bundles (rbx_flow) [115], extract NODDI (noddi_flow) [6, 116] and FreeWater (FW)-corrected DTI (FW_flow) metrics [116, 117], perform tractometry (tractometry_flow) [118], and connectomics (connectoflow) [119].

However, Tractoflow does not address completely the *gray-box principles* as stated. While the curation of configuration makes it easy to execute its workflow seamlessly, it does hinder the transparency of each process executed. In addition, each process is hardly independent from the next, from design requirements imposed by the DSL1 version of Nextflow. Its modularity at the pipeline level forces chaining to be done manually outside of the controlled environments provided by containerization and Nextflow.

Nextflow DSL2 enabled the creation of *versaFlow* [120], a modular pipeline that reaches down to the level of each process. It provides fully transparent access to all inputs, outputs and the corresponding parameters, while still curating easy-to-use configurations using Nextflow's parameters inheritance (something that could not be done by Tractoflow because it uses a single centralized file to store this



information). versaFlow can also be applied to the processing of macaque subjects, by selecting parameters and steps that optimally adapt Tractoflow, carefully validated on the PRIME-DE database [121].

Discussion

In the section titled "Opening up the implementations, that's how the light gets in," we reviewed recent QIB studies that use glass-box workflows as an effective approach to tackle variability at its source, all while staying true to the first-principles foundation of MRI acquisitions. This title purposefully draws inspiration from Leonard Cohen's famous lyrics, "There's a crack in everything, that's how the light gets in". While glass-box workflows are designed for transparency, their increasing complexity introduces fractures where dependencies accumulate (Fig. 1d), particularly when interfacing with multiple vendor-specific scanner control systems.

Are numbers extracted from MRI clinically superior to mere images?

Nearly half a century of efforts have fallen short in giving a generalizable answer to this question. MRI research silos answer it by championing a particular QIB attached to a fitting clinical use case (and even some de-facto standards) to make it reproducible across scanners [122]. Nevertheless, when we restate the same question in more practical terms—"What is to MRI what the Hounsfield Unit is to CT?"—it is difficult to provide a confident answer.

The idea that the future of MRI is quantitative emerges, fades, and resurfaces as the tagline of annual conferences on MRI methods development. Meanwhile, applied research studies leveraging MRI measurements to test biomedical hypotheses continue to proliferate, spawning new subfields and career trajectories. However, the widening gap between the ever-expanding body of rejected null hypotheses and the scarcity of QIBs that have led to changes in patient management has raised concerns among clinicians about the validity of research priorities and their practical impact on patient care.

"Stop da music... stop da music" & "You cannot be serious"

Opening quotes from two articles by MDs Weinberger and Radulescu [123, 124].

Following these staggering opening quotes in their articles on structural MRI, the authors warn against overinterpreting data-driven QIBs and emphasize that such findings should be framed strictly as "differences in MRI measurements" rather than definitive neurobiological conclusions

such as "cortical thinning". They further argue that the ongoing overinterpretation of confounded MRI measurements is not only problematic but also a disservice to patients and their families, potentially by misdirecting valuable tax-payer dollars—funds that could be used more effectively to improve patient care, such as by better staffing hospitals and enhancing clinical services.

Given the unresolved translational gap, it remains too soon to determine whether numerical metrics derived from MRI offer clear clinical advantages over visual assessment alone. However, significant efforts are being made to improve the utility of QIBs, and we briefly discuss their potential benefits and challenges below.

To standardize or to harmonize?

MRI does not directly measure anatomy; apparent tissue boundaries are influenced by systematic biases and biophysical confounders. Most data-driven QIBSs rely on these boundaries, which may reflect secondary physiological effects rather than true morphology [124]. Similarly, the "scanner (or batch) effects" linked to *Acq* are neither linear nor separable from biology [56]. Therefore, both simple transformations of QIBs into a common coordinate system—such as z-scores [125] or histogram matching [126]—and more sophisticated methods that attempt to harmonize raw scanner signals [127] remain fundamentally limited. Other limitations associated with such methods include over-harmonization, i.e., bleach-washing true biological variability, and the challenge of choosing the right technique from a plethora of harmonization options [45, 128].

Can we dispense with the differences in k-space encoding within the latent space of deep learning models? For example, CALAMITI claims to disentangle anatomy and contrast into distinct latent representations [129], achieving state-of-the-art harmonization performance. Even though these methods show promising results for improving downstream processes such as segmentation, their value in establishing reliable QIBs remains uncertain.

Simulation-based bias correction strategies targeting the opaque zone (Fig. 1a) can reduce vendor-specific differences to some extent [63]. However, some sources of variation, such as prescan calibrations, remain unaddressed. From a system theory perspective [130], it is important to note that simulation-driven corrections rely on idealized models of both the acquisition and spin systems. As a result, their effectiveness in correcting known differences between sequence implementations is constrained.

An interesting avenue for further exploration is examining inter-site effects starting from the k-space data acquired using vendor-native sequences. In the context of data-driven QIBs, variability in the reconstruction of images from multichannel receive coils is a notable source of inconsistency



[131]. For physics-driven QIBs, differences in the reconstruction of undersampled data or phase images can lead to variations in measurements [102]. A key challenge in this area is the lack of reference k-space datasets. Moreover, using synthetic k-space data (Fourier-transformed from images) as a reference has been shown to artificially improve reconstruction results [132]. It remains to be seen whether post-hoc workflow unification (Fig. 2), aligned with the frosted-box benchmark (Fig. 1b), can provide the same benefits as glass-box workflows in improving the clinical value of QIBs.

It is crucial to recognize that even the simplest QIB in MRI involves far more degrees of freedom in its calculation than the Hounsfield Unit in CT. The final form of an MR image is influenced by a complex interplay of factors, including spin dynamics, coil sensitivities, gradient nonlinearities, B0 inhomogeneities, eddy currents, susceptibility effects, RF field distortions, and physiological noise, among others. This complexity underscores the importance of returning to the first principles of MRI measurements for conscious reduction of scanner effects. We argue that taking a step back and striving for standardization in a metrological framework is a leap toward unlocking the true clinical potential of QIBs, and adopting glass-box workflows can show the way forward.

Our pipelines can now span vendor boundaries, but can they bridge the gap between research labs and PACS workstations?

The consideration of user-friendliness is still in its early stages for deploying vendor-neutral acquisition platforms coupled with glass-box workflows. Nevertheless, exciting developments are underway both by open-source developers and vendors.

Currently, only the RTHawk proprietary platform [57] offers the flexibility to design and standardize the same user interface across vendors. It also enables the scaling of RF and gradient waveforms, as well as modifying timing parameters through the user interface. In contrast, open-source alternatives must integrate separately with each vendor's interface. While earlier versions of Pulseq did not support changing sequence parameters, the latest release (v1.5.0) has introduced support for this feature [58]. Also working towards a new major release, Gammastar provides support for changing parameters through the interface and even for AI-driven protocol optimizations [98]. Meanwhile, some major manufacturers are putting in the effort to allow easy integration of open-source *Recon* [133, 134] pipelines and even native support for interpreting Pulseq descriptions of a sequence.

The modular and portable approach to creating a comprehensive measurement pipeline presented in this review has great potential to complement these efforts, facilitate collaboration, and lower the barriers to standardizing QIB implementations. Once we standardize MRI measurement procedures through efficient use of reference objects [135, 136] and metrology principles [2], precision medicine using MRI will become more feasible and reliable [137].

Conclusion

Unlike standardized biochemical assays which have universally accepted calibration standards across laboratories, MRI measurements remain highly variable across scanners and sites. This variability stems from the complex interplay of factors in MRI signal generation and site-specific differences in the measurement chain. Rather than constructing a Rube Goldberg machine of harmonization methods and normative modeling, we argue for returning to the first principles of MRI measurements, using standardization within a metrological framework. By embracing workflows that strive for standardization, we can bridge the gap between research innovations and clinical applications [138]. The modular and portable approach to creating OIB pipelines presented in this review offers a practical path forward, facilitating collaboration and lowering barriers to standardizing QIB implementations. This standardization, coupled with efficient use of reference objects and rigorous application of metrology principles [139], will ultimately enable more reliable and clinically meaningful QIBs, towards redefining MRI as a measurement device.

Acknowledgements NYUAD Center for Brain and Health, funded by Tamkeen under NYU Abu Dhabi Research Institute grant CG012.

Funding Open Access funding provided by the IReL Consortium.

Declarations

Conflict of interest Agah Karakuzu, Nadia Blostein, Alex Valcourt, Arnaud Boré, François Rheault, Nikola Stikov declares that he has no conflict of interest. Maxime Descoteaux declares that he is a shareholder at Imeka Solutions. He has no conflict of interest with respect to the content of this work.

Ethical approval This article does not contain any studies with human participants or animals performed by any of the authors.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in



the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

- DIN 1319–2 2005–10 DIN Media. [cited 29 Nov 2024]. Available: https://www.dinmedia.de/de/norm/din-1319-2/81716151
- Cashmore MT, McCann AJ, Wastling SJ, McGrath C, Thornton J, Hall MG (2021) Clinical quantitative MRI and the need for metrology. Br J Radiol 94:20201215
- 3. Tofts P (ed) (2005) Quantitative MRI of the brain, 1st edn. John Wiley & Sons, Chichester
- Kessler LG, Barnhart HX, Buckler AJ, Choudhury KR, Kondratovich MV, Toledano A et al (2015) The emerging science of quantitative imaging biomarkers terminology and definitions for scientific studies and regulatory submissions. Stat Methods Med Res 24:9–26
- Novikov DS, Kiselev VG, Jespersen SN (2018) On modeling. Magn Reson Med 79:3172–3193
- Zhang H, Schneider T, Wheeler-Kingshott CA, Alexander DC (2012) NODDI: practical in vivo neurite orientation dispersion and density imaging of the human brain. Neuroimage 61:1000-1016
- Stikov N, Campbell JSW, Stroh T, Lavelée M, Frey S, Novek J et al (2015) In vivo histology of the myelin g-ratio with magnetic resonance imaging. Neuroimage 118:397–405
- Gillies RJ, Kinahan PE, Hricak H (2016) Radiomics: Images are more than pictures, they are data. Radiology 278:563–577
- Zwanenburg A, Vallières M, Abdalah MA, Aerts HJWL, Andrearczyk V, Apte A et al (2020) The image biomarker standardization initiative: standardized quantitative radiomics for high-throughput image-based phenotyping. Radiology 295:328–338
- Avanzo M, Wei L, Stancanello J, Vallières M, Rao A, Morin O et al (2020) Machine and deep learning methods for radiomics. Med Phys 47:e185–e202
- 11. Klontzas ME (2024) Radiomics feature reproducibility: the elephant in the room. Eur J Radiol 175:111430
- Kushol R, Parnianpour P, Wilman AH, Kalra S, Yang Y-H (2023) Effects of MRI scanner manufacturers in classification tasks with deep learning models. Sci Rep 13:16791
- Warrington S, Ntata A, Mougin O, Campbell J, Torchi A, Craig M et al (2023) A resource for development and comparison of multimodal brain 3T MRI harmonisation approaches. Imaging Neurosci. https://doi.org/10.1162/imag_a_00042
- Rutherford S, Barkema P, Tso IF, Sripada C, Beckmann CF, Ruhe HG et al (2023) Evidence for embracing normative modeling. Elife 12:e85082
- Sled JG, Pike GB (2000) Correction for B(1) and B(0) variations in quantitative T(2) measurements using MRI. Magn Reson Med 43:589–593
- Bojorquez JZ, Bricq S, Acquitter C, Brunotte F, Walker PM, Lalande A (2017) What are normal relaxation times of tissues at 3 T? Magn Reson Imaging 35:69–80
- Karakuzu A, Boudreau M, Stikov N (2024) Reproducible research practices in magnetic resonance neuroimaging: a review informed by advanced language models. Magn Reson Med Sci 23:252–267
- Stikov N, Trzasko JD, Bernstein MA (2019) Reproducibility and the future of MRI research. Magn Reson Med 82:1981–1983

- Soher BJ, Clarke WT, Wilson M, Near J, Oeltzschner G (2022) Community-organized resources for reproducible MRS data analysis. Magn Reson Med 88:1959–1961
- Guimaraes AR (2021) Quantitative Imaging Biomarker Alliance (QIBA): protocols and profiles. Quantitative imaging in medicine. AIP Publishing, New York, pp 1–22
- QIBA Profile Stages. In: Quantitative Imaging Biomarker Alliance [Internet]. 2023 [cited 23 Feb 2025]. Available: https://gibawiki.rsna.org/index.php/OIBA Profile Stages
- Pepin K (2023) MR elastography of the liver, clinically feasible Profile. Radiological Society of North America (RSNA)/Quantitative Imaging Biomarkers Alliance (QIBA). https://doi.org/10. 1148/qiba/20231107
- MR (diffusion-Weighted Imaging (DWI) of the Apparent Diffusion Coefficient (ADC), clinically feasible Profile (2022) Radiological Society of North America (RSNA)/Quantitative Imaging Biomarkers Alliance (QIBA). https://doi.org/10.1148/qiba/20221
- DSC-MRI Consensus QIBA Profile (2020) Radiological Society of North America (RSNA)/Quantitative Imaging Biomarkers Alliance (QIBA). https://doi.org/10.1148/qiba/20201022
- MR MSK Cartilage for Joint Disease, Consensus Profile (2021) Radiological Society of North America (RSNA)/Quantitative Imaging Biomarkers Alliance (QIBA). https://doi.org/10.1148/ qiba/20210925
- DCE-MRI V.2, Consensus QIBA Profile (2023) Radiological Society of North America (RSNA)/Quantitative Imaging Biomarkers Alliance (QIBA). https://doi.org/10.1148/qiba/20231206
- Morton W (2023) RSNA changes tack on quantitative imaging efforts. In: Aunt Minnie [Internet]. 2023 [cited 27 Feb 2025]. Available: https://www.auntminnie.com/imaging-informatics/ advanced-visualization/image-processing/article/15634192/ rsna-changes-tack-on-quantitative-imaging-efforts
- Webb AG (2024) Accessible MRI: No Surrender. ISMRM & ISMRT Annual Meeting and Exhibition; 2024; Singapore. Available: https://ismrm2024.blazestreaming.com/sessions/ ismrm-2024-p-os
- 29. Khan ME, Khan F (2012) A comparative study of white box, black box and grey box testing techniques. Int J Adv Comput Sci Appl. Available: https://citeseerx.ist.psu.edu/document?repid=rep1&type=pdf&doi=ab0ed151c010e03e2d34284ae5f89756372e7690#page=22. Accessed 24 Feb 2025
- Di Tommaso P, Chatzou M, Floden EW, Barja PP, Palumbo E, Notredame C (2017) Nextflow enables reproducible computational workflows. Nat Biotechnol 35:316–319
- Hafyane T, Karakuzu A, Duquette C, Mongeon F-P, Cohen-Adad J, Jerosch-Herold M, et al. (2018) Let's talk about cardiac T1 mapping. bioRxivorg. 2018; 343079.
- Schelbert EB, Messroghli DR (2016) State of the art: clinical applications of cardiac T1 mapping. Radiology 278:658–676
- Sinha U, Sinha S (2024) Magnetic resonance imaging biomarkers of muscle. Tomography 10:1411–1438
- Henninger B, Alustiza J, Garbowski M, Gandon Y (2020) Practical guide to quantification of hepatic iron with MRI. Eur Radiol 30:383–393
- Bauer CM, Jara H, Killiany R (2010) Alzheimer's disease neuroimaging initiative. Whole brain quantitative T2 MRI across multiple scanners with dual echo FSE: applications to AD, MCI, and normal aging. Neuroimage. 52:508–514
- Jack CR Jr, Bernstein MA, Fox NC, Thompson P, Alexander G, Harvey D et al (2008) The Alzheimer's Disease Neuroimaging Initiative (ADNI): MRI methods. J Magn Reson Imaging 27:685–691
- Boudreau M, Karakuzu A, Cohen-Adad J, Bozkurt E, Carr M, Castellaro M et al (2024) Repeat it without me: crowdsourcing



- the T1 mapping common ground via the ISMRM reproducibility challenge. Magn Reson Med. 92:1115
- Stikov N, Karakuzu A (2023) The relaxometry hype cycle. Front Physiol 14:1281147
- Pierpaoli C, Jezzard P, Basser PJ, Barnett A, Di Chiro G (1996)
 Diffusion tensor MR imaging of the human brain. Radiology. https://doi.org/10.1148/radiology.201.3.8939209
- Campbell JSW, Leppert IR, Narayanan S, Boudreau M, Duval T, Cohen-Adad J et al (2018) Promise and pitfalls of g-ratio estimation with MRI. Neuroimage 182:80–96
- Niso G, Botvinik-Nezer R, Appelhoff S, De La Vega A, Esteban O, Etzel JA et al (2022) Open and reproducible neuroimaging: from study inception to publication. Neuroimage 263:119623
- 42. Hernando D, Sharma SD, Aliyari Ghasabeh M, Alvis BD, Arora SS, Hamilton G et al (2017) Multisite, multivendor validation of the accuracy and reproducibility of proton-density fat-fraction quantification at 1.5T and 3T using a fat-water phantom. Magn Reson Med. 77:1516–1524
- Dupuis A, Chen Y, Hansen M, Chow K, Sun JEP, Badve C et al (2024) Quantifying 3D MR fingerprinting (3D-MRF) reproducibility across subjects, sessions, and scanners automatically using MNI atlases. Magn Reson Med 91:2074–2088
- Keenan KE, Gimbutas Z, Dienstfrey A, Stupic KF (2019) Assessing effects of scanner upgrades for clinical studies. J Magn Reson Imaging 50:1948–1954
- Gebre RK, Senjem ML, Raghavan S, Schwarz CG, Gunter JL, Hofrenning EI et al (2023) Cross-scanner harmonization methods for structural MRI may need further work: a comparison study. Neuroimage 269:119912
- Karakuzu A, Samson P, Stikov N (2023) Vendor-neutrality and upgrade immunity: Post-upgrade assessment of vendor-neutral qMRI from two perspectives.
- International Vocabulary of Metrology—Basic and General Concepts and Associated Terms (VIM) Third Edition. 2017 [cited 23 Feb 2025]. Available: https://jcgm.bipm.org/vim/en/index.html
- Tay JSW, Parker RH (1990) Measuring international harmonization and standardization. Abacus 26:71–88
- Greenberg N (2014) Update on current concepts and meanings in laboratory medicine—standardization, traceability and harmonization. Clin Chim Acta 432:49–54
- 50. Miller WG, Greenberg N (2021) Harmonization and standardization: where are we now? J Appl Lab Med 6:510–521
- Fuertes I (2008) Towards harmonization or standardization in governmental accounting? The international public sector accounting standards board experience. J Comp Pol Anal: Res Pract 10:327–345
- Gaspar AS, Silva NA, Ferreira AM, Nunes RG (2024) Repeatability of Open-MOLLI: an open-source inversion recovery myocardial T1 mapping sequence for fast prototyping. Magn Reson Med 92:741–750
- 53. Keenan KE, Tasdelen B, Javed A, Ramasawmy R, Rizzo R, Martin MN et al (2025) T1 and T2 measurements across multiple 0.55T MRI systems using open-source vendor-neutral sequences. Magn Reson Med. 93:289–300
- Veldmann M, Ehses P, Chow K, Nielsen J-F, Zaitsev M, Stöcker T (2022) Open-source MR imaging and reconstruction workflow. Magn Reson Med 88:2395–2407
- 55. Helms G, Dathe H, Kallenberg K, Dechent P (2008) High-resolution maps of magnetization transfer with inherent correction for RF inhomogeneity and T1 relaxation obtained from 3D FLASH MRI: saturation and relaxation in MT FLASH. Magn Reson Med 60:1396–1407
- Karakuzu A, Biswas L, Cohen-Adad J, Stikov N (2022) Vendor-neutral sequences and fully transparent workflows improve

- inter-vendor reproducibility of quantitative MRI. Magn Reson Med 88:1212–1228
- Santos JM, Wright GA, Pauly JM (2004) Flexible real-time magnetic resonance imaging framework. Conf Proc IEEE Eng Med Biol Soc 2004:1048–1051
- 58. Layton KJ, Kroboth S, Jia F, Littin S, Yu H, Leupold J et al (2017) Pulseq: a rapid and hardware-independent pulse sequence prototyping framework. Magn Reson Med 77:1544–1552
- Liu Q, Ning L, Shaik IA, Liao C, Gagoski B, Bilgic B et al (2024) Reduced cross-scanner variability using vendor-agnostic sequences for single-shell diffusion MRI. Magn Reson Med 92:246–256
- 60. Nunes RG, Ravi KS, Geethanath S, Vaughan JT Jr (2020) Implementation of a Diffusion-Weighted Echo Planar Imaging sequence using the Open Source Hardware-Independent PyPulseq Tool. ISMRM 28th annual meeting and exhibition.
- Liu Q, Gagoski B, Shaik IA, Westin C-F, Wilde EA, Schneider W et al (2024) Time-division multiplexing (TDM) sequence removes bias in T2 estimation and relaxation-diffusion measurements. bioRxivorg. https://doi.org/10.1101/2024.06.03.597138
- Hannum AJ, Loecher M, Chen Q, Arbes E, Littin S, Setsompop K et al (2025) Towards open-source spin-echo cardiac diffusion tensor imaging. J Cardiovasc Magn Reson 27:101389
- 63. Chhetri G, McPhee KC, Wilman AH (2021) Alzheimer's disease neuroimaging initiative. Bloch modelling enables robust T2 mapping using retrospective proton density and T2-weighted images from different vendors and sites. Neuroimage. 237:118116
- 64. Rowley CD, Campbell JSW, Wu Z, Leppert IR, Rudko DA, Pike GB et al (2021) A model-based framework for correcting B 1 + inhomogeneity effects in magnetization transfer saturation and inhomogeneous magnetization transfer saturation maps. Magn Reson Med 86:2192–2207
- Rui Pedro Teixeira AG, Malik SJ, Hajnal JV (2019) Fast quantitative MRI using controlled saturation magnetization transfer. Magn Reson Med. 81:907–920
- Crooijmans HJA, Scheffler K, Bieri O (2011) Finite RF pulse correction on DESPOT2. Magn Reson Med 65:858–862
- Zaiss M, Schmitt B, Bachert P (2011) Quantitative separation of CEST effect from magnetization transfer and spillover effects by Lorentzian-line-fit analysis of z-spectra. J Magn Reson 211:149–155
- 68. Mossahebi P, Alexander AL, Field AS, Samsonov AA (2015) Removal of cerebrospinal fluid partial volume effects in quantitative magnetization transfer imaging using a three-pool model with nonexchanging water component: removal of CSF partial volume effects in qMT imaging. Magn Reson Med 74:1317–1326
- Boudreau M, Tardif CL, Stikov N, Sled JG, Lee W, Pike GB (2017) B1 mapping for bias-correction in quantitative T1 imaging of the brain at 3T using standard pulse sequences. J Magn Reson Imaging 46:1673–1682
- Lee Y, Callaghan MF, Acosta-Cabronero J, Lutti A, Nagy Z (2019) Establishing intra- and inter-vendor reproducibility of T1 relaxation time measurements with 3T MRI. Magn Reson Med 81:454–465
- Fortin J-P, Parker D, Tunç B, Watanabe T, Elliott MA, Ruparel K et al (2017) Harmonization of multi-site diffusion tensor imaging data. Neuroimage 161:149–170
- Chen AA, Beer JC, Tustison NJ, Cook PA, Shinohara RT, Shou H et al (2022) Mitigating site effects in covariance for machine learning in neuroimaging data. Hum Brain Mapp 43:1179–1195
- Radua J, Vieta E, Shinohara R, Kochunov P, Quidé Y, Green MJ et al (2020) Increased power by harmonizing structural MRI site differences with the ComBat batch adjustment method in ENIGMA. Neuroimage 218:116956



- Bell TK, Godfrey KJ, Ware AL, Yeates KO, Harris AD (2022) Harmonization of multi-site MRS data with ComBat. Neuroimage 257:119330
- 75. Bayer JMM, Thompson PM, Ching CRK, Liu M, Chen A, Panzenhagen AC et al (2022) Site effects how-to and when: an overview of retrospective techniques to accommodate site effects in multi-site neuroimaging analyses. Front Neurol 13:923988
- 76. Zhou C, Zhou J, Lv Y, Batuer M, Huang J, Zhong J et al (2025) The impact of the novel CovBat harmonization method on enhancing radiomics feature stability and machine learning model performance: a multi-center, multi-device study. Eur J Radiol 184:111956
- Zhang R, Oliver LD, Voineskos AN, Park JY (2023) RELIEF: a structured multivariate approach for removal of latent interscanner effects. Imaging Neurosci (Camb) 1:1–16
- 78. De Luca A, Karayumak SC, Leemans A, Rathi Y, Swinnen S, Gooijers J et al (2022) Cross-site harmonization of multi-shell diffusion MRI measures based on rotational invariant spherical harmonics (RISH). Neuroimage 259:119439
- Dewey BE, Zhao C, Reinhold JC, Carass A, Fitzgerald KC, Sotirchos ES et al (2019) DeepHarmony: a deep learning approach to contrast harmonization across scanner changes. Magn Reson Imaging 64:160–170
- Torbati ME, Minhas DS, Laymon CM, Maillard P, Wilson JD, Chen C-L et al (2023) MISPEL: a supervised deep learning harmonization method for multi-scanner neuroimaging data. Med Image Anal 89:102926
- Lee G, Ye DH, Oh S-H (2025) A preliminary attempt to harmonize using physics-constrained deep neural networks for multisite and multiscanner MRI datasets (PhyCHarm). medRxiv. https://doi.org/10.1101/2025.02.07.25321867
- Dennis EL, Veer IM, Descoteaux M, Tate DF, Wilde EA (2023) Editorial: harmonization strategies and considerations in neuroimaging studies. Front Neurol 14:1165263
- Stamoulou E, Spanakis C, Manikis GC, Karanasiou G, Grigoriadis G, Foukakis T et al (2022) Harmonization strategies in multicenter MRI-based radiomics. J Imaging 8:303
- 84. Abbasi S, Lan H, Choupan J, Sheikh-Bahaei N, Pandey G, Varghese B (2024) Deep learning for the harmonization of structural MRI scans: a survey. Biomed Eng Online 23:90
- Zhu AH, Moyer DC, Nir TM, Thompson PM, Jahanshad N (2019) Challenges and opportunities in dMRI data harmonization. Computational diffusion MRI. Springer International Publishing, Cham, pp 157–172
- Gulban OF, Nielson D, Lee J, Poldrack R, Gorgolewski C, Vanessasaurus, et al. poldracklab/pydeface: PyDeface v2.0.2. Zenodo; 2022. 10.5281/ZENODO.6856482
- 87. Mölder F, Jablonski KP, Letcher B, Hall MB, Tomkins-Tinch CH, Sochat V et al (2021) Sustainable data analysis with Snakemake. F1000Res. 10:33
- 88. Gorgolewski K, Burns CD, Madison C, Clark D, Halchenko YO, Waskom ML et al (2011) Nipype: a flexible, lightweight and extensible neuroimaging data processing framework in python. Front Neuroinform 5:13
- Harenslak BP, de Ruiter JR (2021) Data pipelines with Apache airflow. Manning Publications, New York
- Ferreira da Silva R, Filgueira R, Pietri I, Jiang M, Sakellariou R, Deelman E (2017) A characterization of workflow management systems for extreme-scale applications. Future Gener Comput Syst. 75:228–238
- Inati SJ, Naegele JD, Zwart NR, Roopchansingh V, Lizak MJ, Hansen DC et al (2017) ISMRM Raw data format: a proposed standard for MRI raw datasets. Magn Reson Med 77:411–421
- Gorgolewski KJ, Auer T, Calhoun VD, Craddock RC, Das S, Duff EP et al (2016) The brain imaging data structure, a format

- for organizing and describing outputs of neuroimaging experiments. Scientific Data 3:1-9
- 93. Karakuzu A, Appelhoff S, Auer T, Boudreau M, Feingold F, Khan AR et al (2022) qMRI-BIDS: an extension to the brain imaging data structure for quantitative magnetic resonance imaging data. Scientific Data 9:1–9
- Gorgolewski KJ, Alfaro-Almagro F, Auer T, Bellec P, Capotă M, Chakravarty MM et al (2017) BIDS apps: Improving ease of use, accessibility, and reproducibility of neuroimaging data analysis methods. PLoS Comput Biol 13:e1005209
- Hayashi S, Caron BA, Heinsfeld AS, Vinci-Booher S, McPherson B, Bullock DN et al (2024) Brainlife. Io: a decentralized and open-source cloud platform to support neuroscience research. Nat Methods. 21:809–813
- Sherif T, Rioux P, Rousseau M-E, Kassis N, Beck N, Adalat R et al (2014) CBRAIN: a web-based, distributed computing platform for collaborative neuroimaging research. Front Neuroinform 8:54
- Markiewicz CJ, Gorgolewski KJ, Feingold F, Blair R, Halchenko YO, Miller E et al (2021) The OpenNeuro resource for sharing of neuroscience data. Elife. https://doi.org/10.7554/eLife.71774
- Hoinkiss DC, Huber J, Plump C, Lüth C, Drechsler R, Günther M (2023) AI-driven and automated MRI sequence optimization in scanner-independent MRI sequences formulated by a domainspecific language. Front Neuroimaging 2:1090054
- Karakuzu A, Boudreau M, Duval T, Boshkovski T, Leppert IR, Cabana J-F et al (2020) qMRLab: quantitative MRI analysis, under one umbrella. J Open Sourc Softw 5:2343
- 100. Harding RJ, Bermudez P, Bernier A, Beauvais M, Bellec P, Hill S et al (2023) The Canadian open neuroscience platform-an open science framework for the neuroscience community. PLoS Comput Biol 19:e1011230
- 101. Marques JP, Kober T, Krueger G, van der Zwaag W, de Moortele PFV, Gruetter R (2010) MP2RAGE, a self bias-field corrected sequence for improved segmentation and T1-mapping at high field. NeuroImage. https://doi.org/10.1016/j.neuroimage.2009. 10.002
- 102. Karakuzu A, Hagiwara A, Uchida W, Kamagata K, Fujita S, Tasdelen B, et al. MP2RAGE against the machine: Scanning in the name of reproducibility with Pulseq in VENUS. ESMRMB 40th Annual Meeting. 2024. Available: https://www.esmrmb2024.org/abstracts-form/posters-e/abstract-data/69b7381e52950bd1a549918c156faf0a. Accessed 24 Feb 2025
- Van Essen DC, Ugurbil K, Auerbach E, Barch D, Behrens TEJ, Bucholz R et al (2012) The human connectome project: a data acquisition perspective. Neuroimage 62:2222–2231
- 104. Fan Q, Witzel T, Nummenmaa A, Van Dijk KRA, Van Horn JD, Drews MK et al (2016) MGH–USC human connectome project datasets with ultra-high b-value diffusion MRI. Neuroimage 124:1108–1114
- 105. Sudlow C, Gallacher J, Allen N, Beral V, Burton P, Danesh J et al (2015) UK biobank: an open access resource for identifying the causes of a wide range of complex diseases of middle and old age. PLoS Med 12:e1001779
- 106. Marek K, Jennings D, Lasch S, Siderowf A, Tanner C, Simuni T et al (2011) The Parkinson progression marker initiative (PPMI). Prog Neurobiol 95:629–635
- Casey BJ, Cannonier T, Conley MI, Cohen AO, Barch DM, Heitzeg MM et al (2018) The Adolescent Brain Cognitive Development (ABCD) study: imaging acquisition across 21 sites. Dev Cogn Neurosci 32:43–54
- Theaud G, Houde J-C, Boré A, Rheault F, Morency F, Descoteaux M (2020) TractoFlow: a robust, efficient and reproducible diffusion MRI pipeline leveraging Nextflow & Singularity. Neuroimage 218:116889



- Scilpy: library supporting research and development at the Sherbrooke Connectivity Imaging Lab. In: Scilpy [Internet]. [cited 24 Feb 2025]. Available: https://github.com/scilus/scilpy
- 110. Garyfallidis E, Brett M, Amirbekian B, Rokem A, van der Walt S, Descoteaux M et al (2014) Dipy, a library for the analysis of diffusion MRI data. Front Neuroinform 8:8
- 111. Tournier J-D, Smith R, Raffelt D, Tabbara R, Dhollander T, Pietsch M et al (2019) MRtrix3: a fast, flexible and open software framework for medical image processing and visualisation. Neuroimage 202:116137
- 112. Avants B, Tustison NJ, Song G (2009) Advanced normalization tools: V1.0. Insight J. https://doi.org/10.54294/uvnhin
- Jenkinson M, Beckmann CF, Behrens TEJ, Woolrich MW, Smith SM (2012) FSL. Neuroimage 62:782–790
- 114. Fischl B (2012) FreeSurfer. Neuroimage 62:774-781
- St-Onge E, Schilling KG, Rheault F (2023) BundleSeg: a versatile, reliable and reproducible approach to white matter bundle segmentation. International Workshop on Computational Diffusion MRI. Springer, Cham, pp 47–57
- Daducci A, Canales-Rodríguez EJ, Zhang H, Dyrby TB, Alexander DC, Thiran J-P (2015) Accelerated Microstructure Imaging via Convex Optimization (AMICO) from diffusion MRI data. Neuroimage 105:32–44
- Pasternak O, Sochen N, Gur Y, Intrator N, Assaf Y (2009) Free water elimination and mapping from diffusion MRI. Magn Reson Med 62:717–730
- 118. Cousineau M, Jodoin P-M, Morency FC, Rozanski V, Grand'Maison M, Bedell BJ et al (2017) A test-retest study on Parkinson's PPMI dataset yields statistically significant white matter fascicles. NeuroImage Clin. 16:222–233
- Connectoflow. In: GitHub [Internet]. [cited 24 Feb 2025]. Available: https://github.com/scilus/connectoflow
- 120. Valcourt Caron A, Shmuel A, Hao Z, Descoteaux M (2023) versaFlow: a versatile pipeline for resolution adapted diffusion MRI processing and its application to studying the variability of the PRIME-DE database. Front Neuroinform 17:1191200
- Neff EP (2018) PRIME-DE is primed with primate data. Lab Anim 48:26–26
- 122. Keenan KE, Biller JR, Delfino JG, Boss MA, Does MD, Evelhoch JL et al (2019) Recommendations towards standards for quantitative MRI (qMRI) and outstanding needs: Recommendations for qMRI Standards. J Magn Reson Imaging 49:e26–e39
- Weinberger DR, Radulescu E (2021) Structural magnetic resonance imaging all over again. JAMA Psychiat 78:11–12
- Weinberger DR, Radulescu E (2016) Finding the elusive psychiatric "lesion" with 21st-century neuroanatomy: a note of caution. Am J Psychiatry 173:27–33
- 125. Kochunov P, Jahanshad N, Sprooten E, Nichols TE, Mandl RC, Almasy L et al (2014) Multi-site study of additive genetic effects on fractional anisotropy of cerebral white matter: comparing meta and megaanalytical approaches for data pooling. Neuroimage 95:136–150
- 126. Um H, Tixier F, Bermudez D, Deasy JO, Young RJ, Veeraraghavan H (2019) Impact of image preprocessing on the scanner dependence of multi-parametric MRI radiomic features and

- covariate shift in multi-institutional glioblastoma datasets. Phys Med Biol 64:165011
- Mirzaalian H, Ning L, Savadjiev P, Pasternak O, Bouix S, Michailovich O et al (2016) Inter-site and inter-scanner diffusion MRI data harmonization. Neuroimage 135:311–323
- Eshaghzadeh Torbati M, Minhas DS, Ahmad G, O'Connor EE, Muschelli J, Laymon CM et al (2021) A multi-scanner neuroimaging data harmonization using RAVEL and ComBat. Neuroimage 245:118703
- Zuo L, Dewey BE, Liu Y, He Y, Newsome SD, Mowry EM et al (2021) Unsupervised MR harmonization by learning disentangled representations using information bottleneck theory. Neuroimage 243:118569
- Kamen EW, Heck BS (2007) Fundamentals of signals and systems using the web and MATLAB, 3rd edn. Pearson, Upper Saddle River
- 131. Beauferris Y, Teuwen J, Karkalousos D, Moriakov N, Caan M, Yiasemis G et al (2022) Multi-coil MRI reconstruction challenge-assessing brain MRI reconstruction models and their generalizability to varying coil configurations. Front Neurosci 16:919186
- Shimron E, Tamir JI, Wang K, Lustig M (2022) Implicit data crimes: machine learning bias arising from misuse of public data. Proc Natl Acad Sci USA 119:e2117203119
- Hansen MS, Sørensen TS (2013) Gadgetron: an open source framework for medical image reconstruction. Magn Reson Med 69:1768–1776
- Uecker M, Ong F, Tamir JI, Bahri D, Virtue P, Cheng JY, et al. (2015) Berkeley advanced reconstruction toolbox. Proc Intl Soc Mag Reson Med. p. 9.
- 135. Stupic KF, Ainslie M, Boss MA, Charles C, Dienstfrey AM, Evelhoch JL et al (2021) A standard system phantom for magnetic resonance imaging. Magn Reson Med 86:1194–1211
- 136. Lavdas I, Behan KC, Papadaki A, McRobbie DW, Aboagye EO (2013) A phantom for diffusion-weighted MRI (DW-MRI): a phantom for DW-MRI. J Magn Reson Imaging 38:173–179
- 137. Hollander W, Weiant C, Chung C, Hylton N, Rosen M. A Quantitative Imaging Infrastructure to Revolutionize AI-Enabled Precision Medicine. In: Federation of American Scientists [Internet]. 2024 [cited 27 Feb 2025]. Available: https://fas.org/publication/ai-enabled-precision-medicine/
- Tamir JI, Blumenthal M, Wang J, Oved T, Shimron E, Zaiss M (2025) MRI acquisition and reconstruction cookbook: recipes for reproducibility, served with real-world flavour. Magn Reson Mater Phy 1–9
- 139. Hall MG, Cashmore M, Cho HM, Ittermann B, Keenan KE, Kolbitsch C, Lee C, Li C, Ntata A, Obee K, Pu Z (2025) Metrology for MRI: the field you've never heard of. Magn Reson Mater Phy 1–26

Publisher's Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

