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Data Augmentation for medical imaging: a Systematic Literature Review

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Abstract

Recent advances in Deep Learning have largely benefited from larger and more diverse training sets. However, collecting large datasets for medical imaging is still a challenge due to privacy concerns and labeling costs. Data augmentation makes it possible to greatly expand the amount and variety of data available for training without actually collecting new data. Data augmentation techniques range from simple yet surprisingly effective transformations such as cropping, padding, and flipping, to complex generative models. Depending on the nature of the input and the visual task, different data augmentation strategies are likely to perform differently. For this reason, it is conceivable that medical imaging requires specific augmentation strategies that generate plausible data samples and enable effective regularization of deep neural networks. Data augmentation can also be used to augment specific classes that are underrepresented in the training set, e.g., to generate artificial lesions. The goal of this systematic literature review is to investigate which data augmentation strategies are used in the medical domain and how they affect the performance of clinical tasks such as classification, segmentation, and lesion detection. To this end, a comprehensive analysis of more than 300 articles published in recent years (2018-2022) was conducted. The results highlight the effectiveness of data augmentation across organs, modalities, tasks, and dataset sizes and suggest potential avenues for

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future research.

Keywords: data augmentation, deep learning, medical imaging, generative adversarial networks, MRI

1. Introduction

Deep learning (DL) has established state-of-the-art performance in many areas of computer vision and pattern recognition, including medical image analysis (1; 2). In order to successfully build well-generalizing deep neural networks (DNNs), we need, in most of the cases, a large dataset to avoid overfitting such large-capacity learners. Collecting a sufficient number of samples has become a significant obstacle that makes deep neural networks quite challenging to apply in medical image analysis, since the acquisition of high-quality reference standards requires a large amount of time, money, and human resources (3; 1). Moreover, most manually annotated datasets are imbalanced, with some specific classes that are often underrepresented. To mitigate the problem of limited medical training sets, several data augmentation techniques have been developed in the literature to generate synthetic training examples.

Data augmentation is a strategy allows to increase the diversity and size of data available for training models, without actually collecting new data (4). This is achieved by applying several image manipulation techniques on the original data or by creating new samples that employ other models. Data augmentation can be used to reduce class imbalance and dataset biases, avoid overfitting and, in general, increase the variability of the data. Another well-known property of data augmentation is that it promotes learning invariance with respect to transformations of the input data that should not affect the output. Thus, it provides a convenient way to learn invariance without designing the model architecture to be equivariant or invariant (5).

Different data augmentation techniques are likely to be more or less effective depending on the type of input, downstream visual task and application field. In the literature, there are many reviews and surveys that analyze different data augmentation techniques in different fields, such as (6), which presents existing methods for data augmentation, promising developments, and high-level guidelines for implementing data augmentation on RGB images. However, given the unique properties of medical images, it is likely that data augmentation should be tailored for this specific domain (7). To the best of our knowledge, existing surveys in the medical domain cover specific subfields: (8) as an example highlights the most promising research directions for synthesizing high-quality artificial brain tumor examples with the goal of improving the generalization capabilities of deep models. Considering the large number of data augmentation techniques applied in the most different subfields of medical image analysis, it is not trivial to identify the most appropriate one for a specific task.

Inspired by these premises, in this paper, we present a systematic literature review exploring the data augmentation techniques used in the context of medical image analysis. In particular, we identify, analyze, and summarize techniques categorizing them by the visual task to which they are applied and by other medical-specific factors such as pathology, organ, or data modality. As a result of our analysis, we provide a detailed comparison of the usage of different data augmentation techniques in the medical domain, as well as a discussion on their potential impacts on the performance of deep learning models. Moreover, we identify the current knowledge gaps, also by comparing with the corresponding literature in the non-medical domain. The objectives of our review can be summarized with the following research questions:

• RQ1: What are the most common study designs to present and assess data

augmentation methods in the medical domain? Are there studies comparing the effect of different data augmentation models on downstream tasks?

- RQ2: What types of data augmentation are used in the medical domain?
- **RQ3**: What are their effects on the performance of deep learning-based methods for medical image analysis?
- **RQ4**: Which data augmentation methods have not been explored in the medical domain?

2. Background

In this section, we first propose a taxonomy for data augmentation strategies used in the medical domain, illustrated in Fig. 1, and provide a brief introduction to each transformation family. Following previous reviews (6; 9), we divided data augmentation techniques into two broad categories: transformation of original data, detailed in Section 2.1 and generation of artificial data, detailed in Section 2.2.

The rest of this section introduces concepts that apply transversely to many augmentation techniques. In Section 2.3, specific challenges arising from the volumetric nature of many medical modalities are discussed. In Section 2.4, the concept of learnable data augmentation is introduced, which aims at learning to combine and adapt multiple transformations and their hyper-parameters to a specific dataset or task. Finally, we briefly mention how data augmentation can be exploited at test time to increase accuracy and reduce uncertainty (Section 2.5).

2.1. Transformation of original data

The first category of data augmentation strategies apply one or more image manipulation techniques to existing samples. They range from the simplest affine trans-

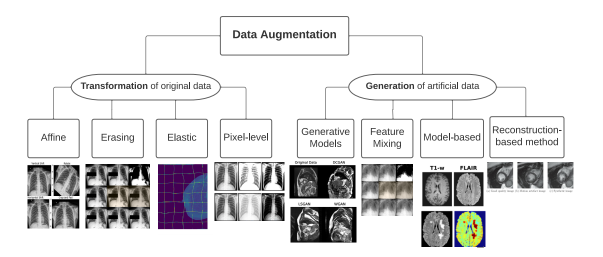


Figure 1: Data augmentation for medical imaging - a taxonomy.

formations (such as flipping or rotating an image) to pixel-level and elastic distortions. Most of the transformations included in this category are relatively easy to implement and are either built-in in deep learning frameworks or provided by easily integrated general purpose libraries (10). More recently, frameworks and libraries specifically targeting the medical domain, such as the Medical Open Network for AI (MONAI), have been proposed (11; 12). It must be observed that, being based on transformations of the original samples, these techniques cannot increase the generalization capabilities of the network beyond the initial training population, and tend to generate highly correlated samples.

2.1.1. Affine transformation

Affine transformations refer to a class of geometrical transformations that preserve lines and parallelism, but not necessarily distances and angles. This behavior is guaranteed by the constraint imposed during the transformation, which typically preserves the aspect ratio of the image along one or more axes of symmetry. Transformations in this group are translation, rotation, flipping, scaling, cropping, and shearing. All these geometrical transformations can be expressed in the form of matrices and thus can be combined to obtain composed affine transformations. Examples of affine transformations are reported in Fig. A.1.

2.1.2. Erasing transformation

Erasing transformations select a region of an image and replace the selected pixels with a fixed pixel intensity value or random noise. This transformation was originally proposed in the RGB domain to increase robustness to occlusions (which however are not present in the medical domain) and to force DNNs to avoid simplified detection patterns. It also reduces spurious correlations between the object of interest and other features, usually due to dataset biases. An example of an erasing transformation is reported in Fig. A.2.

2.1.3. Elastic transformation

Elastic transformation operate by applying a spatial deformation field to an image. Differently from affine transformations, elastic transformations do not impose any constraint on the preservation of col-linearity and aspect ratio, and thus can introduce local shape variation into an organ or lesion. These transformations can be used, for instance, to increase the robustness of segmentation algorithms (8). Elastic transformation may also simulate image deformations due to breathing and patient movements, thus making DNNs more robust to such common artifacts (13; 14). A critical aspect to consider is that unconstrained elastic transformation may yield highly unrealistic samples, although the impact on the performance of downstream tasks is still being debated (8).

An important class of elastic transformation are those based on *diffeomorphism*, i.e. a mapping between two differentiable and invertible mainfolds (8). Diffeomorphic

mapping can be used to transform a source image I to a target image J. By using an actual sample to guide the elastic transformation, it is more likely to obtain a plausible and realistic augmentation. An example of elastic transformation is shown in Fig. A.3.

2.1.4. Pixel-level transformation

Pixel-level transformations change the pixel values in order to modify image characteristics such as brightness, contrast, saturation and noise. Most medical imaging modalities are grayscale, and hence color-based transformations are not common. Pixel-level transformations are useful to increase DNNs robustness across different scanners and imaging protocols, which may affect the pixel distribution. Examples of pixel-level transformations are shown in Fig. A.4.

2.2. Generation of artificial data

Generating artificial or synthetic samples can yield more diverse and challenging samples, thus overcoming the limitations of transformation-based techniques. Generative networks, and specifically generative adversarial networks, are today the most common approach to medical image synthesis (15). Yet, artificial images can be generated also through feature mixing or leveraging ad-hoc modelling strategies tailored to a specific medical imaging task or modality. These techniques allow for a greater variety, at the expense of increased computational requirements and complexity. Artificial samples may also not reflect the visual characteristics or distribution of true samples.

2.2.1. Generative (Adversarial) Networks

Generative models have been used to generate realistic images (16; 17; 18), speech (19; 20), text (21; 22) and more. Generative models include variational autoen-

coders (23; 24; 25), generative adversarial networks (26; 23; 15), and, more recently, diffusion-based models (27; 28).

Today, the most common generative architecture is the Generative Adversarial Network (GAN), introduced by (26). GANs are composed by two components, namely a generator and a discriminator. While the generator tries to trick the discriminator into believing that the fake data is authentic, the discriminator attempts to discern if the image is real or synthetic. The two networks contest with each other in a zero-sum game, where the gain of one is the loss of the other. The success of GANs can be attributed to their exceptional visual fidelity, especially when compared to variational autoencoders.

A thorough analysis of GAN applications in medical imaging, including commonly used architectures and losses, is available in (15). We report some examples of synthetic images generated by GANs in Fig. A.5. It is worth mentioning that GANs allow for controlled image generation (29) that is particularly useful in the context of data augmentation. For instance, GANs conditioned on a label or a segmentation map can be used to generate synthetic lesions or, more generally, to balance a dataset by augmenting underrepresented groups (30). Image translation architectures, such as CycleGANs (31) have been used for cross-domain medical image synthesis, which allows transferring samples from modalities for which data is relatively abundant (e.g., CT) to more expensive or less widespread modalities (e.g., MRI) (32). It can also be used to hallucinate missing modalities for multi-parametric sequences, such as MRI (33).

However, leveraging GANs for data augmentation is tricky. Besides the complexity and computational cost of generating high resolution, realistic images, GANs are prone to mode collapse (34), in which the generator is prone to yield very similar examples and thus has limited impact on generalizability, and have been shown to

hallucinate image features that mimic or mask the presence of lesions (35).

In very recent years, following seminal work by Ho and colleagues (36), several diffusion-based generative models have been proposed achieving extraordinary results in photorealistic image synthesis, including unconstrained and text-to-image synthesis. These models generate samples by gradually removing noise from a signal and, compared to GANs, have more desirable properties in terms of distribution coverage, ease of training and scalability. Now that diffusion-models are now achieving competitive results compared to GANs (27), it is reasonable to expect that, in the future, this class of model will start to complement the use of GANs in the medical domain as well.

2.2.2. Feature mixing methods

Feature mixing methods combine two or more samples from the original dataset to create a new one. Mixing images together by averaging their pixel values or cutting and pasting some parts of an image into another may represent counter-intuitive approaches from a human perspective. In fact, generated images may not look meaningful to human observers (6). However, these techniques were shown to enhance generalization capabilities of DNNs, and increase robustness to adversarial examples (37). The mix-up technique, shown in Fig. A.7, is one of the most used feature mixing methods in the medical field.

2.2.3. Model-based methods

Under the umbrella of model-based techniques we include a variety of physically or biologically inspired models to generate new images, or modify existing ones. These transformations can be performed both using DNNs (not necessarily GANs) or traditional image processing techniques, such as shape modelling, shape deformation or image blending. Since medical datasets are often very skewed towards negative

cases, the most common model-based techniques involve the synthesis and injection of artificial lesions in otherwise healthy subjects. For instance, these techniques have been used to simulate multiple sclerosis lesions in brain MR images (38) or to add cancer signs to breast mammography images (39). An example of these model-based data augmentation is shown in Figure A.8.

Other techniques incorporate physiological time progression models, such as reproducing the effect of aging on organs or simulating disease progression and regression (40). Finally, some model-based techniques exploit prior knowledge on human anatomy and on images formation process to generate random variations representative of inter-subject variability, or to simulate common acquisition artifacts to increase the robustness of image analysis algorithms (41). Unlike GAN-based techniques, these techniques typically require minimal or no training, and directly incorporate anatomical constraints on the generated images. However, they can only be applied to specific organs for which mathematical models are available.

2.2.4. Reconstruction-based methods

Three-dimensional medical imaging modalities are reconstructed using a mathematical process that transforms the raw data collected by the image scanner into a three-dimensional volume that can be inspected by the radiologist. For example, image reconstruction in CT involves generating tomographic images from X-ray projections acquired at many different angles. Reconstruction algorithms and their settings influence the signal-to-noise ratio, resolution, presence of artifacts, and general quality of the reconstructed images. While the majority of data augmentation techniques operate directly on the reconstructed images, a few techniques have been proposed that operate directly on the raw data space and then reconstruct the distorted images. For example, in (42) realistic motion artifacts are simulated by corrupting the

raw data using Gaussian blurring, and then a segmentation network is trained on increasingly distorted data. These techniques may be relatively complex to apply, as they require access to the raw data acquired by the scanner (often not available in a clinical setting), and implementing the entire reconstruction algorithm. A subset of these techniques involves reconstructing 2D images (e.g., X-rays) from 3D volumes (e.g., CT scans), in order to simulate different acquisition angles and views (43). An example is shown in Figure A.6.

2.3. 2D vs. 3D data augmentation

One of the characteristics of modalities like CT and MRI is their volumetric nature. Nonetheless, not all DNNs employed in the medical domain employ 3D convolutions (44; 7). Many authors, especially toward the beginning of the deep learning era, have used 2D convolutions, by focusing on individual slices: 2D models have lower memory and computational footprint, and also allow the use of backbones pre-trained on ImageNet. 3D convolutions have the advantage of combining information from adjacent slices, and achieve higher performance if a suitable large dataset is available (45). An alternative is to use 2.5D convolutions, slices captured along three perpendicular planes (axial, coronal, and sagittal) mimic RGB images and are input to the three channels of ImageNet-pretrained models. Some image transformations (most notably, affine transformations) are widely available for 2D images, but may require extensions to 3D, which may not be straightforward or computationally efficient. Examples of 3D-aware data augmentations include 3D GANs (46), multiplanar image synthesis (47) and 3D affine transformations (48).

2.4. Learnable data augmentation

Learnable data augmentation is a recent subfield of deep learning research that studies approaches that can reduce the human effort required when selecting and validating a set of data augmentation techniques (49). The main idea is to discover automatically an optimal data augmentation strategy for a specific task. Aside from generative models, which we covered in Section 2.2.1, the most common approach is to learn an optimal data augmentation policy. Techniques based on this approach require the concurrent training of two networks, one for learning how to solve a task and the second to learn how to augment the data for the first one. The most common approach is augmentation policy learning or autoaugment, which learns the best policy to maximize a network performance by combining a list of known transformations (such as affine, pixel-level, and so forth) (50). The optimal policy can be determined by a neural network, trained using reinforcement learning (50), evolutionary algorithms (51), adversarial training (52) or parametric statistical models (53).

2.5. Train-Time & Test-Time Data Augmentation

Although most of the studies in this survey focus on improving data at training time, test time augmentation (TTA) has also been explored to increase DNN accuracy and robustness. In test-time augmentation, multiple transformations of a given input are generated, and then the DNN predictions are averaged to obtain a more robust estimate. TTA has been shown to increase the accuracy of a DNN (6), and can also be used to estimate the aleatoric uncertainty of the predictions (54; 55). Unlike epistemic uncertainty, which refers to the DNN parameters, aleatoric uncertainty reflects noise or randomness in the input image, and thus cannot be explained or reduced by increasing the training data (54).

3. Methodology

In this section we describe the methodology adopted to perform the systematic literature review. The analysis was performed following the Preferred Reporting Items for Systematic Reviews and Meta-Analyzes (PRISMA) guidelines.

3.1. Search strings and Study selection

The search was conducted on PubMed and on the search engine of the main publishers (ACM, IEEE, and Elsevier) in the fields of computer science, medical imaging and biomedical engineering. We searched for the keywords "data augmentation" and "medical imaging", as shown in Table B.7. We considered only studies published between January 2018 and July 2022.

We defined inclusion and exclusion criteria on parameters such as language, publication date, type of study, modality, context, and task. We selected only primary studies written in English from 2018 to 2022, including both proceedings and journal papers, but excluding preprints. We included papers related to the main medical imaging modalities: X-Ray (XR), Magnetic Resonance (MR), Computed Tomography (CT), Positron Emission Tomography (PET) and Ultrasound (US); we excluded papers in the dermatology, ophthalmology, endoscopy, and histopathology domains. We also excluded studies that do not use data augmentation or for which the full text was not available. The downstream tasks included in the review were classification, detection, and segmentation (in other words, we selected papers in which the efficacy of data augmentation was tested on classification, lesion detection or segmentation DNNs). Moreover, we specifically selected only papers that used DNNs to perform any of the downstream tasks selected, to increase the homogeneity of the selected studies. The selection was done in two steps: first, we selected papers based on their

titles and abstracts, and then performed a more in depth analysis on the full-texts. Inclusion and exclusion criteria are summarized in Table B.8.

3.2. Data extraction

To answer RQ1, we classified all retrieved studies in four categories: Type 1 (standard data augmentation techniques are used, impact on the performance of the downstream task not assessed); Type 2 (novel data augmentation techniques are proposed, impact on the performance of the downstream task not assessed); Type 3 (performances of the downstream task with and without data augmentation are compared) and Type 4 (performances of the downstream task with several data augmentation strategies are compared). Each paper was assigned to the highest number category (i.e., a paper that proposes a novel data augmentation technique, and compare its impact on a classification task against standard data augmentation, would be classified as Type 4, rather than Type 2). In addition, only papers in categories 3 and 4 were used to answer the question RQ3. The resulting categorization and an interpretation legend are reported in Table B.9.

3.3. Data analysis

The papers were reported grouping into five groups (brain, lung, heart, breast and others) according to the target anatomical district. The impact of data augmentation on the performance of downstream tasks was estimated by calculating the relative increase in performance with and without data augmentation, and the distribution of the relative increase (overall and by organ) was calculated using boxplots. The same technique was used to calculate the effect of different data augmentation techniques, focusing in particular on studies in which complex data augmentation techniques were compared against the simplest transformations (affine).

4. Results

The selection process was conducted in August 2022. The preliminary set initially contained 919 papers, but after the removal of 45 duplicates, the set was reduced to 874 papers. After applying the inclusion and exclusion criteria reported in Table B.8 to the titles and abstracts, the set of articles was reduced to 493 candidates. In particular, we discarded 53 studies that did not belong to the medical domain, 47 reviews or surveys, 74 studies that did not use any data augmentation, and 207 studies classified as out of scope (i.e., they did not meet the inclusion criteria related to the imaging modality or downstream task). Additional 10 studies were included through other sources (e.g., based on the authors' expertise, citations, or experts' suggestions). The full-texts of the remaining 503 papers were analyzed in detail; we further discarded 141 articles that did not meet inclusion criteria and selected the remaining 362 as candidates for the literature review. Abbreviations used in this section are reported in Table 1. The control flow diagram of the selection process is shown in Fig. 2.

4.1. RQ1 & RQ2: Which data augmentation techniques have been studied in the medical domain?

Each article was assigned to a category according to the methodology described in Section 3. As it can be noticed by analyzing the distribution shown in Fig. 3a, more than half of the papers, roughly 54% (196/362), quantitatively evaluate the effect of at least one data augmentation technique, while only 13% (47/362) propose a new technique without evaluating its impact on multiple downstream tasks.

The distribution of different data augmentation techniques is shown in Fig. 3b. The most used type of data augmentation is affine transformations, being used in

Abbreviation	Meaning	Abbreviation	Meaning
(*)	Test Time Augmentation	M-AF	Mix Affine
ACC	Accuracy	MD	Mean Distance (mm)
ACS	Acute Coronary Sindrome	MG	Mammography
AF	Affine	MOD	Model-based
ALZ	Alzheimer	MR	Magnetic Resonance
AUC	Area under the ROC Curve	MS	Multiple Sclerosis
BIF	Bifurcation	NMD	Neuromuscolar Diseases
\mathbf{C}	Classification	NOD	Nodule
CAN	Cancer	OS	Osteoporosis
CAS	Coronary Angioscopy	OSD	One Stage Detector
COV	COVID-19	PACC	Pixel Accuracy
CPM	Competition Performing Metric	PAR	Parkinson
CT	Computed Tomography	pGAN	custom GAN
CY	Cyst	PNE	Pneumothorax
D	Detection	POL	Polyp
DICE	DICE Score	PSNR	Peak Signal-to-Noise Ratio
EL	Elastic	PX	Pixel-level
ER	Erasing	REC	Reconstruction-based
F1	F1 Score	S	Segmentation
FM	Feature Mixing	S-AF	Single Affine
FPI	False Per Image	SNS	Sensitivity
FRC	Fractures	SP	Specificity
FROC	Free-Response Receiver Operating Characteristic	TPR	True Positive Rate
GAN	Generative Adversial Network	TUB	Tubercolosis
GN	Generative Network	URS	Urinary Stones
Н	Healthy	US	Ultrasound scan
HAND	Hand Injuries and Disorders	wGAN	Wasserstein GAN
iGAN	Info GAN	XR	X-Ray scan
L	Lesion	OTH	Other
TF	Transformer	LRN	Learnable data augmnetation

Table 1: List of the abbreviations used in Section 4

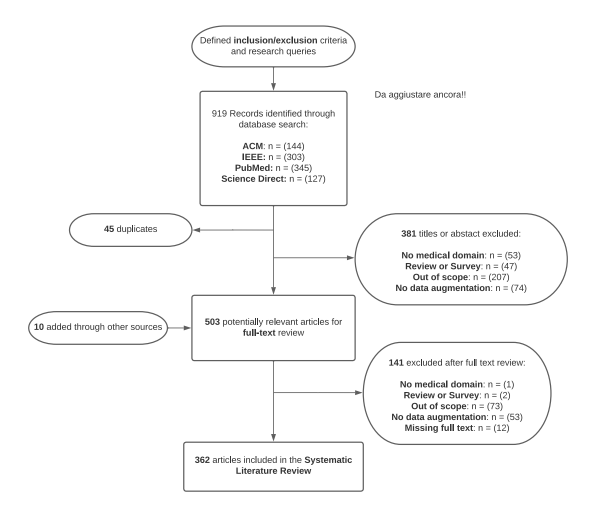


Figure 2: Systematic Literature Review Control Flow Diagram

64% (233/362) of the studies. This result was expected, as this type of data augmentation is fairly effective and extremely easy to implement. GAN-based and pixel-level transformations follow, appearing in 29% (104/362) and 22% (77/362) of the studies, respectively. Pixel-level transformations are popular, as they enhance generalization to different scanners and acquisition protocols (56). Model-based methods and elastic transformation are used in approximately 9% of the studies (33/362 and 34/362,

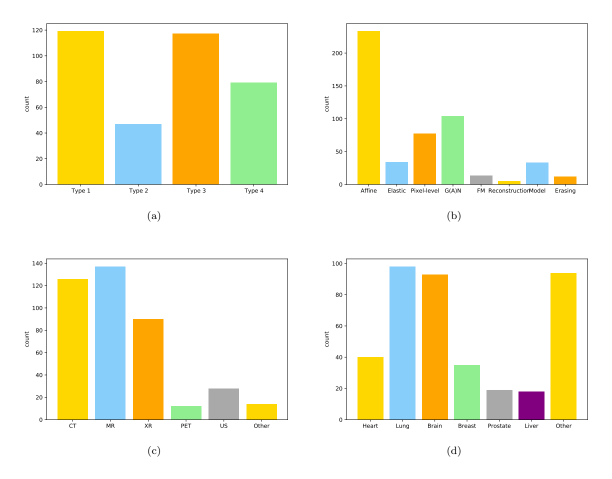


Figure 3: Distribution of studies by (a) study design, (b) data augmentation category, (c) modality and (d) organ.

respectively), and the remaining techniques only represent 8% (30/362) of our sample (42; 43; 57; 58; 59; 60).

As shown in Fig. 3c, CT, MR and XR (including mammography) are the most common modalities in our sample, reflecting their widespread adoption in clinical practice. Under other, we grouped less common modalities such as Coronary Angioscopy (CAS) (61), Myocardial perfusion imaging (MPI) (62) and Digital breast tomosynthesis (DBT) (63). We identified a few studies that leveraged cross-modality

image synthesis for data augmentation, e.g., setting MR as the source and CT as the target (64), CT as the source and MR as the target (65), and bidirectional style transfer between the two modalities (66).

Lastly, in Fig. 3d, we report the distribution of the studies with respect to different organs. As it can be noticed, the most studied organs are lungs and brain with 98 and 93 papers respectively. Under the category *other* we grouped organs included in less than 10 studies, such as the musculoskeletal system (arm, calcaneus, knee, hand, humerus, hip, maxillary, neck, pelvis, skull, shoulder, spine), specific organs in the abdomen (colon, kidney, lymph node, pancreas, stomach, rectum), the pituitary membrane, teeth, thyroid, and the urinary conduct.

Studies were further grouped by type of data augmentation and (i) organ, (ii) modality, (iii) type of paper, and (iv) downstream task (related bar plots are reported in the Appendix Appendix C). In general, there does not appear to be a qualitative correlation between organ, modality, and downstream task and the distribution of the related data enhancement strategies; affine transformations represent the most common technique in all cases. In most cases, type 1 papers include affine and pixel-level data augmentation techniques, whereas type 2, 3 and 4 papers tend to be biased towards more complex strategies, such as generative or model-based techniques. This particular result may be a side-effect of our selection process, which sought to highlight novel data augmentation strategies emerging in the literature. The results are consistent with a previous study by Nalepa et al., who compared the data augmentation strategies employed in the BraTS challenge, mostly affine or pixel-level transformations (9).

4.2. RQ3: What is the effect of data augmentation on the performance of the down-stream task?

To evaluate the effect of different data augmentation techniques, we only selected type 3 and type 4 papers that make a comparison with other data augmentation strategies or without data augmentation. Papers for which performance information could not be extracted (14 papers) were excluded from the quantitative analysis. We report the results separately for papers that include experiments on different organs or tasks, and hence the total number of records is 206 out of 196 papers.

The main characteristics of these studies are reported in the following sections, grouped by organ. For each study, we report the modality, the target disease, the type of DA used, the characteristics of the downstream task, and the performance metrics used. Whenever available, we report the percentage increase in performance with DA with respect to the experiments in which DA was not used (DA vs. no-DA). For type 4 papers, we selected the simplest form of data augmentation (typically affine transformations) as the baseline, the best performing data augmentation strategy as the proposed DA strategy, and report the relative difference in performance of the proposed vs. baseline DA. The use of relative differences allows us to compare studies based on different metrics.

4.2.1. Brain

Neuroimaging is used to assess (directly or indirectly) the structure and function of the nervous system, primarily with computed tomography (CT), magnetic resonance imaging (MR), and positron emission tomography (PET). The main pathologies analyzed in the different studies are Alzheimer's disease, Multiple Sclerosis, Parkinson's disease, and brain tumor.

The most common downstream tasks in brain imaging, as reported in Table

2, are classification (18/49) and segmentation (30/49). Classification is useful in determining the stage of progression in brain disease, such as in (67) for *glioma grading*. Segmentation, on the other hand, finds application both in healthy subjects and in the quantitative study of different brain diseases (68). We excluded from the analysis one paper (69) in which data augmentation was used to make the network robust to adversarial attacks, instead of improving the its generalization ability.

Every data augmentation technique used in these studies improves performance compared to those who do not use data augmentation. The highest relative increase is achieved by (30) with an increase of 26.56%, and the highest relative improvement with respect to baseline DA is achieved by (70) with an increase in performance of 10.36%.

Model-based techniques (9/39) and G(A)Ns (18/49) are used in more than half the selected studies. It is possible that the relatively stable structure of the brain, and the fact that accurate anatomical models have been developed over the course of many years, facilitate the application of these techniques. Many model-based techniques often involve the injection of synthetic lesions into otherwise healthy brains, such as the introduction of synthetic MS lesions (38), brain tumors (71) or microbleeds (72). Model-based techniques have also been used to simulate common image acquisition artifacts such as intensity inhomogeneity (73), with the aim of making DNNs more robust to artifacts. This is a unique feature of DNNs which can learn features that are invariant to the presence of specific artifacts, if exposed to them during training, shifting attention from removing such artifacts from the images (e.g., through normalization techniques) to selecting more realistic and varied training data.

In (74), the authors propose a framework based on GANs to create structural synthetic brain networks in multiple sclerosis. This paper shows that generative

models are competitive with respect to DA strategies that operate in the feature domain, such as Synthetic Minority Oversampling Technique (SMOTE) (75).

A few authors have proposed to use data augmentation techniques in order to train DNNs in few-shot learning scenarios by leveraging otherwise unlabelled data, thus pushing forward the boundaries of data augmentation. For instance, (76) and (70) used model-based and generative data augmentation techniques, respectively, to learn spatial and appearance transformations to match images of real healthy subjects to one or more reference atlases: since segmentation is known for the atlas, the images generated through the proposed methodology can be automatically labeled.

4.2.2. Heart

Cardiac imaging refers to non-invasive imaging of the heart using primarily ultrasound (US), magnetic resonance imaging (MR) and computed tomography (CT). Studies related to this organ are reported in Table 3; the most common downstream task is segmentation (15/21), followed by classification (4/21) and detection (2/21).

As detailed in Table 3, data augmentation techniques used in these studies usually improve performance with respect to the no-DA baseline and the other DA baselines. In interpreting the results, attention must be paid to the baseline performance, which in some cases is very low (114), likely reflecting poor hyper-parameter optimization and perhaps inflating comparison. Best results are achieved by Spectral augmentation (115) (DICE: 0.891 to 0.811) and BigAug (56) (DICE: 0.858 to 0.914). Some of the less performing studies, on the other hand, employ transformations that are not suited to this specific organ. For instance, horizontal flip was found to reduce performance, except in the very small data scenario (116), likely due to the fact that the heart has a specific orientation. Several GAN variants have been proposed for heart imaging, including SpeckleGAN (117), ScreenGAN (118) or CycleGAN (65).

Paper	Task	Network	Modality	Pathology	Proposed DA	Baseline DA	Metric	w/o DA	w/Baseline	w/Proposed	w/o DA -¿ Proposed	Baseline -; Proposed
(77)	С	CNN	MR	CAN	M-AF	S-AF	ACC	0.64	0.77	0.79	23.44%	2.60%
(78)	C	CNN	CT	H	AF	-	ACC	0.73	-	0.923	26.44%	-
(57)	C	CNN	PET	PAR	MOD	-	ACC	0.876	-	0.882	0.68%	-
(79)	C	CNN	MR	CAN	AF	-	ACC	0.976	-	1	2.46%	-
(80)	C	CNN	MR	CAN	AF	-	ACC	0.923	-	0.983	6.50%	-
(81)	C	CNN	PET-MR	ALZ	GAN	-	ACC	0.67	-	0.74	10.45%	-
(82)	C	CNN	MR	CAN	AF - PX	-	ACC	0.874	-	0.907	3.78%	-
(83)	C	CNN	MR	CAN	pGAN	GAN	SNS	0.689	0.74	0.789	14.51%	6.62%
(84)	C	CNN	MR	ALZ	AF - PX - ER	-	ACC	0.834	-	0.888	6.47%	-
(74)	C	AU + RF	MR	MS	GAN	-	F1	0.656	-	0.81	23.48%	-
(85)	D	Yolo-3	MR	CAN	pGAN	GAN	SNS	0.83	-	0.91	9.64%	-
(86)	S	U-Net	MR	MS	AF	-	DICE	0.803		0.866	7.85%	-
(87)	S	U-Net	MR	MS	MOD	AF	CPM	-	0.77	0.79	-	2.60%
(88)	S	AMRUNet	MR	CAN	FM	-	DICE	0.771	-	0.787	2.08%	-
(89)	S	U-Net	MR	CAN	GAN	AF - PX	DICE	-	0.841	0.851	-	1.19%
(90)	S	U-Net	MR	CAN	AF	-	DICE	0.636	-	0.701	10.22%	-
(71)	S	U-Net	MR	CAN	GAN - AF	AF	DICE	0.623	0.644	0.653	4.82%	1.40%
(91)	S	U-Net	MR	CAN	GAN	-	DICE	0.71		0.736	3.66%	-
(30)	S	U-Net	MR	CAN - ALZ	GAN	_	DICE	0.64		0.81	26.56%	-
(92)	S	U-Net	MR	L	GAN	_	DICE	0.438	-	0.55	25.57%	_
(90)	S	U-Net	MR	L	AF	_	DICE	0.427	_	0.46	7.73%	_
(73)	S	U-Net	MR	Н	MOD-AF	AF	DICE	-	0.876	0.892	-	1.83%
(93)	S	Custom	MR-CT	Н	pGAN	GAN	DICE	_	0.783	0.81	_	3.45%
94)	S	U-Net	MR	Н	EL	AF	DICE	_	0.884	0.906	_	2.49%
(68)	S	U-Net	PET	Н	GAN	-	DICE	0.872	-	0.889	1.95%	_
(95)	S	Custom	MR	Н	PX	_	DICE	0.805	_	0.82	1.86%	
(96)	s	U-Net	MR	Н	GN	-	DICE	0.83	-	0.88	6.02%	
(40)	S	U-Net	CT	CAN	MOD	AF	DICE	0.852	0.859	0.875	2.70%	1.86%
(76)	s	U-Net	MR	Н	MOD - AF - PX	AF - PX	DICE	0.76	0.775	0.815	7.24%	5.16%
97)	S	U-Net	MR	CAN	AF - EL	-	DICE	0.763	-	0.8	4.85%	0.1070
(38)	s	U-Net	MR	MS	MOD		DICE	0.57		0.63	10.53%	
(98)	S	U-Net	MR	CAN	GAN	-	DICE	0.72	-	0.82	13.89%	-
(99)	C	CNN	MR	ALZ	AF - PX - ER	_	ACC	0.786	-	0.92	17.05%	-
(100)	C	CNN	MR	OTH	AF - PX	-	AUC	0.780	-	0.92	2.07%	-
(100)	s	Custom	MR	CAN	AF - FA	-	DICE	0.967	-	0.987	3.33%	-
(101)	C	CUSTOM	MR	CAN	AF - PX	-	ACC	0.9	-	0.93	4.44%	-
(102)	s	U-Net	MR	H	FM	-	DICE	0.9	-	0.717	0.56%	-
` ′	C	CNN		ALZ		-	ACC		-			-
(104)			MR		GAN	-		0.806		0.863	7.07%	-
(105)	S	U-Net	MR	MS	PX	-	DICE	0.677	-	0.646	-4.58%	-
(106)	S	3D U-Net	DWI	OTH	AF-MOD-GAN	-	DICE	0.632	-	0.708	12.03%	
(72)	C	CNN	MR	ALZ	MOD	AF FI	AUC	0.984	0.993	0.995	1.12%	0.20%
(107)	S	U-Net	MR	CAN	GAN - MOD	AF - EL	DICE	0.784	0.801	0.79	0.77%	-1.37%
(70)	S	3D U-Net	MR	H	GAN	EL - MOD	DICE	-	0.743	0.82	-	10.36%
(108)	C	3D CNN	PET	ALZ	AF - PX	AF	ACC	-	0.395	0.352	-	-10.89%
(109)	S	3D U-Net	MR	CAN	FM	AF	DICE	-	0.853	0.864	-	1.29%
(110)	S	U-Net	MR	CAN	GAN	AF	ACC	0.9127	0.9366	0.9424	3.25%	0.62%
111)	C	CNN	MR	CAN	GAN	AF	ACC	-	0.888	0.92	-	3.60%
112)	C	CNN	MR	CAN	FM	AF	ACC	-	0.953	0.9676	-	1.53%
(113)	S	3D U-Net	MR	CAN	AF - PX - EL	FM	DICE	0.705	0.714	0.727	3.12%	1.82%

Table 2: Comparison on **49** studies on data augmentation techniques for brain imaging. See Table 1 for an explanation of the acronyms.

This section also includes one of the few studies that evaluated the performance of reconstruction-based methods (42). The authors presented an automatic spatio-temporal DNNs for cardiac motion artifact data which was trained by corrupting

data in the k-space and then reconstructing the images: this strategy was able to simulate different levels of realistic motion-related artifacts (e.g., breathing artifacts).

Finally, one of the studies investigated the role of data augmentation in a federated learning setting (119). Interestingly, the authors observed that in a federated training scheme, the impact of data augmentation was lower than in the standard centralized setting, in which the data is shared on a central server. Specifically, in the federated learning setting, AUC was 0.746 ± 0.001 without DA, 0.791 ± 0.001 for basic (affine) transformations, 0.766 ± 0.001 for shape transformations, and 0.731 ± 0.009 for shape and intensity transformations. In the centralized setting, the corresponding performance was generally lower and with higher variance, with AUCs equal to 0.732 ± 0.008 (w/o DA), 0.759 ± 0.016 (affine), 0.764 ± 0.022 (shape), and 0.776 ± 0.008 (shape and intensity), respectively.

Paper	Task	Network	Modality	Pathology	Proposed DA	Baseline DA	Metric	$\rm w/o~DA$	w/Baseline	w/Proposed	w/o DA -¿ Proposed	Baseline -¿ Proposed
(120)	S	U-Net	CT	Many	AF	-	DICE	0.911	-	0.927	1.76%	-
(121)	S	CNN	MR	H	PX	AF - EL	DICE	-	0.652	0.894	-	37.12%
(114)	S	CNN	MR - US	H	AF - PX - EL	GAN	DICE	0.188	0.575	0.85	352.13%	47.83%
(122)	S	U-Net	MR - CT	H	MOD	FM	DICE	0.332	0.652	0.747	125.00%	14.57%
(117)	S	U-Net	US	H	pGAN	GAN	DICE	0.824	0.833	0.846	2.67%	1.56%
(65)	S	U-Net	MR - CT	H	GAN	AF	DICE	0.613	0.676	0.712	16.15%	5.33%
(58)	D	CNN	CT	ACS	AF - FM	-	AUC	0.808	-	0.867	7.30%	-
(42)	D	3D CNN	MR	Many	AF - PX - REC	AF	AUC	0.581	0.674	0.735	26.51%	9.05%
(61)	C	CNN	CAS	Many	GAN	-	AUC	0.77	-	0.81	5.19%	-
(123)	C	CNN	CT	Many	AF - ER	-	ACC	0.74	-	0.903	22.03%	-
(118)	C	CNN	US	H	GAN	-	ACC	0.7	-	0.85	21.43%	-
(124)	S	3D nn-U-Net	US	H	GAN		DICE	0.822	-	0.823	0.12%	-
(125)	S	3D U-Net	CT	H	GAN	AF	DICE	0.876	0.878	0.893	1.94%	1.71%
(115)	S	Custom	CT	H	REC		DICE	0.811	-	0.891	9.86%	-
(126)	S	3D U-Net	MR - CT	H	pGAN	GAN	DICE	-	0.687	0.847	-	-
(127)	S	U-Net	MR - CT	H	EL	PX	DICE	0.703	0.721	0.731	3.99%	1.39%
(119)	C	3D CNN	MR	Many	AF - PX - EL	AF	DICE	0.746	0.791	0.731	-2.01%	-7.59%
(52)	S	U-Net	MR	H	AF - PX - EL (**)	FM	DICE	-	0.782	0.808	-	3.38%
(56)	S	3D U-Net	MR	H	AF - PX - EL	AF	DICE	0.919	0.913	0.914	-0.54%	0.11%
(56)	S	3D U-Net	US	H	AF - PX - EL	AF	DICE	0.858	0.910	0.921	7.34%	1.21%
(116)	S	U-Net	CT	H	AF		DICE	0.750	-	0.732	-2.38%	-

Table 3: Comparison of **21** studies on data augmentation techniques for heart imaging. See Table 1 for an explanation of the acronyms.

4.2.3. Lung

Lung imaging refers to non-invasive imaging of the lungs using primarily magnetic resonance imaging (MR), computed tomography (CT), and positron emission tomography (PET). Due to the fact that lung cancer is one of the leading causes of cancer-related deaths and due to the COVID-19 pandemic, the lung is probably the most studied and analyzed organ in this entire SLR. In lung imaging, the most common downstream task is classification (37/51), whereas detection (5/51) and segmentation (9/51) are less common, as shown in Table 4.

With few exceptions, data augmentation consistently improved performance with respect to the w/o DA baseline and the other DA baseline. In one study, data augmentation was associated with a a -14.45% decrease in COVID-19 classification performance (128). It is possible that the affine transformations employed in this study (flip, translation, and scaling) do not yield images consistent with the actual distribution at test time. However, this remains an isolated case.

Overall, 20/51 papers are based on transformations of existing data, including affine transformations, pixel-level transformations and, less frequently, random erasing and elastic deformations. The remainder of the studies are based on GANs or model-based techniques. Although most generative models are single-modality, some studies have also proposed cross-modality synthesis. An example of generative model is a tumor-aware unsupervised cross-domain generative network which translates images from CT to MRI while preserving tumor details using an ad-hoc loss (32). The synthetic MRI data were then used to train a U-Net segmentation model in a semi-supervised fashion leveraging segmented images already available in the CT domain.

As mentioned in the introduction, data augmentation techniques can be employed to build invariance with respect to the transformation used. Some authors have questioned whether the invariance could be encoded directly into the DNN architecture. A fitting example is invariance to rotation, which can be hard-coded into steerable convolution filters, bypassing the need and even surpassing the performance of rotational data augmentation (129; 130). However, it is hard to imagine that such strong inductive biases can be exploited for all relevant transformations.

Finally, a few studies have employed TTA (131; 132), observing an improvement in performance (DICE coefficient increased from 0.941 to 0.979 and 0.901 to 0.902, respectively). As introduced in Section 2.5, TTA augmentation can yield increases in performance.

4.2.4. Breast

Breast imaging is a sub-speciality of diagnostic radiology that involves imaging of the breast for cancer screening or diagnostic purposes. The most common modality is mammography (MG), complemented by breast US and DBT. In the reviewed studies (Table 5), 14/19 analyze whole-breast or patch-level classification, 3/19 lesion detection and 2/19 lesion segmentation. As in previous organs, all studies report an increase in performance when using DA with respect to non-DA.

Affine and pixel-level transformations are the most common data augmentations reported. Also elastic transformations are applicable to the breast. One study (172) compared elastic to affine transformations, decreasing the False Positives per Image (FPI) of about 25.66% in a lesion detection task.

Interestingly, several model-based techniques have been proposed in breast imaging. A first technique is based on synthetic images produced using a virtual three-dimensional anthropomorphic phantoms (39). The virtual phantom is generated using a procedural analytic model in which the main anatomical structures (including fat and glandular tissues, ductal tree, vasculature, and ligaments) are stochastically generated within a predefined breast volume. Artificial masses were inserted

Paper	Task	Network	Modality	Pathology	Proposed DA	Baseline DA	Metric	w/o DA	w/Baseline	w/Proposed	w/o DA -; Proposed	Baseline -; Proposed
128)	С	CNN	XR	COV	AF	-	ACC	0.868	-	0.745	-14.17%	-
133)	C	CNN	CT - XR	PNE	PX	-	ACC	0.928	-	0.988	6.47%	-
134)	C	CNN	CT	CAN	AF	-	ACC	0.9	-	1	11.11%	-
135)	C	CNN	XR	TUB	AF	-	ACC	0.78	-	0.8	2.56%	-
59)	C	CNN	XR	COV - PNE	AF - ER - FM	-	ACC	0.787	-	0.837	6.35%	-
136)	C	CNN	XR	PNE	GAN - PX - AF	-	ACC	0.795	-	0.91	14.47%	-
137)	C	CNN - RNN	CT - XR	COV	AF - GAN	-	AUC	0.91	-	0.99	8.79%	-
131)	C	CNN	XR	COV	AF (*)	-	ACC	0.941	-	0.979	4.04%	-
138)	C	CNN	XR	PNE	AF - PX	AF - PX	ACC	0.84	0.86	0.945	12.50%	9.88%
139)	C	CNN	CT	CAN	MOD	-	ACC	0.5	-	0.786	57.20%	-
46)	C	CNN	CT	CAN	AF - PX - GAN	AF - PX	ACC	0.5	0.761	0.825	65.00%	8.41%
129)	C	CNN	CT	CAN	AF	-	ACC	0.842	-	0.942	11.88%	-
47)	C	CNN	CT	CAN	AF - GAN	-	AUC	0.635	-	0.812	27.87%	-
140)	C	CNN	CT	NOD	MOD	-	CPM	0.77	-	0.79	2.60%	-
141)	C	CNN	CT	NOD	pGAN	wGAN	ACC	-	0.923	0.95	-	2.93%
142)	C	CNN	XR	PNE	AF	AF - PX	ACC	0.74	0.79	0.85	14.86%	7.59%
143)	C	PSSPNN	CT	COV	AF - PX	-	F1	0.939	-	0.958	2.02%	-
144)	C	CNN	CT	NOD	AF	AF (*)	ACC	-	0.848	0.88	-	3.77%
145)	C	CNN	CT	CAN	pGAN	wGAN	ACC	0.342	0.547	0.577	68.71%	5.48%
146)	C	CNN	CT	Н	MOD	AF	SNS	_	0.839	0.858	-	2.26%
60)	C	CNN	CT	NOD	AF - FM	_	ACC	0.694	_	0.74	6.63%	
147)	C	CNN	CT	CAN	AF	_	AUC	0.598	_	0.655	9.53%	_
148)	C	RNN	XR	TUB	AF - PX	_	ACC	0.471	_	0.928	97.03%	_
149)	C	CNN	CT	NOD	AF - EL	wGAN	ACC	_	0.57	0.65	_	14.04%
150)	C	CNN	XR	Many	GAN	-	ACC	0.662	-	0.957	44.56%	-
151)	C	CNN	CT	CAN	MOD	AF	ACC	-	0.83	0.92	-	10.84%
152)	D	3D CNN	CT	NOD	AF	-	SNS	0.75	-	0.9	20.00%	-
153)	D	-	CT	NOD	AF		ACC	0.697		0.788	13.06%	
154)	D	3D CNN	CT	NOD	AF		SNS	0.923		0.788	7.26%	-
155)	D	R-CNN	CT	NOD	GAN		CPM	0.518	-	0.55	6.18%	
132)	s	U-Net	XR	PNE	AF - PX - EL (*)	-	DICE	0.9011	-	0.9023	0.13%	
156)	S	U-Net	MR - CT	CAN	GAN	AF	DICE	0.9011	0.63	0.9023	0.1370	11.11%
32)	s	U-Net	CT CT	CAN	pGAN	GAN	DICE	0.55	0.63	0.8	45.45%	26.98%
40)	S	U-Net	XR	NOD	MOD	AF	DICE	0.725	0.794	0.809	11.59%	1.89%
	C	CNN	CT	COV		Ar	ACC	0.725	0.794			1.0970
157)	s		CT	COV	AF - PX AF - EL - PX		DICE		-	0.944	2.59%	-
158)	C	3D U-Net 3D CNN	CT	COPD			ACC	0.756 0.597	-	0.761	0.66%	-
159)					GAN					0.629	5.36%	-
160)	C	CNN	XR	PNE	GAN		ACC	0.795	-	0.903	13.52%	-
161)	S	U-Net	CT	Many	MOD		DICE	0.948	-	0.947	-0.07%	-
162)	C	Custom	CT	NOD	GAN		AUC	0.851	-	0.921	8.19%	-
163)	C	CNN	CT	COV	GAN		AUC	0.935	-	0.96	2.67%	-
164)	C	CNN	CT	NOD	GAN		ACC	0.851	-	0.879	3.19%	-
165)	C	CNN	CT	COV	AF - GAN	AF	ACC	-	0.929	0.9	-	-3.14%
166)	D	RetinaNet	XR	NOD	MOD - AF - PX	AF	SNS	-	0.494	0.52	-	5.26%
167)	S	U-Net	MRI	COPD	AF - GAN	AF	DICE	0.955	0.961	0.965	1.05%	0.42%
168)	C	CNN	XR	PNE	GAN - AF	AF	ACC	0.805	0.815	0.83	3.11%	1.84%
169)	C	CNN	XR	PNE	GAN	AF	AUC	0.941	0.951	0.962	2.23%	1.10%
170)	C	CNN	CT	CAN	GAN		ACC	0.342	-	0.577	68.71%	-
113)	S	3D U-Net	CT	CAN	AF - PX - EL	FM	DICE	0.564	0.581	0.639	13.30%	9.98%
116)	S	U-Net	CT	CAN	AF		DICE	0.983	-	0.985	0.26%	-
171)	C	CNN	XR	Many	GAN		ACC	0.93	-	0.95	2.15%	-

Table 4: Comparison of **51** studies on data augmentation techniques for lung imaging. See Table 1 for an explanation of the acronyms.

according to a previously defined and validated simulation model. Then, breast compression is simulated using a finite element model, and a Monte Carlo-based X-Ray transport simulation code was exploited to project the voxelized 3D phantoms into realistic-looking synthetic mammograms. Thanks to the relative simplicity of the breast organ, virtual phantoms are also increasingly used in image acquisition and reconstruction, denoising, and to conduct virtual trials (173). Likewise, their role in

training DNNs is likely to increase. Other authors have proposed a method called stochastic evolution (SE) to mimic the irregular deterioration and healing processes of the diseased tissue according to the direction of local distortion (40).

Paper	Task	Network	Modality	Pathology	Proposed DA	Baseline DA	Metric	$\mathrm{w/o}~\mathrm{DA}$	w/Baseline	w/Proposed	w/o DA - ¿ Proposed	Baseline - ¿ Proposed
(174)	С	CNN	MG	CAN	AF	-	ACC	0.77	-	0.82	6.49%	-
(175)	C	CNN	$_{ m MG}$	CAN	GAN	-	ACC	0.782	-	0.87	11.25%	-
(176)	C	CNN	US	CAN	AF	-	AUC	0.94	-	0.96	2.13%	-
(177)	D	OSD	$_{ m MG}$	CAN	AF - EL - PX	-	TPR	0.861	-	0.913	6.04%	-
(178)	S	U-Net	$_{ m MG}$	CAN	AF	-	DICE	0.922	-	0.951	3.15%	-
(179)	C	CNN	$_{ m MG}$	CAN	AF - PX	-	ACC	0.782	-	0.836	6.91%	-
(180)	C	CNN	$_{ m MG}$	CAN	AF	-	ACC	0.71	-	0.859	20.99%	-
(181)	C	CNN	$_{ m MG}$	CAN	AF - PX	-	ACC	0.721	-	0.938	29.00%	-
(172)	C	CNN	$_{ m MG}$	CAN	EL	AF	FPI	-	2.934	2.181	-	-25.66%
(182)	C	CNN	$_{ m MG}$	CAN	AF - GAN	AF	ACC	0.699	0.88	0.94	34.48%	6.82%
(183)	C	CNN	CT	CAN	GAN	AF	AUC	-	0.741	0.869	-	17.27%
(184)	D	R-CNN	$_{ m MG}$	CAN	GAN	AF	AUC	0.151	0.159	0.172	13.91%	8.18%
(185)	C	CNN	US	CAN	pGAN	iGAN	ACC	-	0.887	0.904	-	1.92%
(39)	D	R-CNN	CT	CAN	MOD	-	SNS	0.802	-	0.833	3.87%	-
(40)	S	U-Net	CT	CAN	MOD	AF	DICE	0.547	0.643	0.666	21.76%	3.58%
(186)	C	CAPSULE	$_{ m MG}$	CAN	AF		ACC	0.828	-	0.849	2.54%	-
(187)	C	CNN	US	CAN	MOD - ER		AUC	0.767	-	0.799	4.19%	-
(188)	C	CNN	$_{ m MG}$	CAN	GAN - AF		AUC	-	0.835	0.889	-	6.49%
(112)	C	CNN	$_{ m MG}$	CAN	$_{\mathrm{FM}}$		ACC	0.62	-	0.721	16.32%	-
(171)	C	CNN	US	CAN	GAN		ACC	0.863	-	0.891	3.20%	-

Table 5: Comparison of **19** studies on data augmentation techniques for breast imaging. See Table 1 for an explanation of the acronyms.

4.2.5. Other Organs

In Table 6 we collected studies conducted on less studied organs, including less frequent diseases such as detection of urinary stones in (189), classification of cyst and tumors of both jaws in (190) or classification of calcaneal fractures in (191). The most common organs *liver* and *prostate* with 18/66 and 16/66 studies, respectively. Despite a few exceptions (192; 193; 194; 195), data augmentation consistently improves performance in all organs.

One study uses train-time data augmentation to randomly increase the size of the training set and test-time data augmentation to estimate the uncertainty of the predictions (196). Both strategies contribute to increase the performance with respect w/o DA baseline. Model-based techniques were proposed to generate several types of images, such as spine (197) and urinary tract images (189).

In one study (198), a positive effect of augmentation was observed for the "severe" cases, whereas for the "moderate" cases there were no significant improvements, suggesting that the benefit of data augmentation may depend on the specific subpopulation.

It is worth highlighting the few studies in this SLR that investigate learnable data augmentation strategies (199; 200; 201). For instance, the optimal policy can be learned through reinforcement learning, choosing from a set of transformations that include flipping, rotation, cropping, elastic deformation, zooming, noise, and brightness adjustment (199). The authors observed an increase in performance with respect to a fixed data augmentation policy on kidney stone segmentation, but the results are rather preliminary and more studies are needed to assess the applicability to a wider range of organs and modalities.

Another interesting study investigates the application of three-dimensional transformations on various organs (48). The same model is trained across different views, which forces the network to use the same weights to capture structures at different viewing-angles. The results show that combining and augmenting three different views could further boost performance.

4.2.6. Effect of data augmentation on deep learning performance

In this section, we summarize the distribution of the relative increase in performance associated to the use of different data augmentation techniques. A total of 185 papers were included in this analysis, corresponding to different organs (Heart: 16, Brain: 39, Lung: 43, Breast: 16, Other: 71).

As shown in Figure 4, the median increase in performance associated to data

Paper (195)	Organ Abdomen	Task D	Network YOLO	Modality CT	Pathology H	Proposed DA GAN	Baseline DA	Metric MD	w/o DA 8.66	w/Baseline	w/Proposed 7.95	w/o DA -¿ Proposed -8.20%	Baseline -¿ Propose
(48)	Abdomen	S	U-Net	MR	Н	AF		DICE	0.897		0.9	0.33%	
202)	Abdomen	S	3D U-Net	CT	Н	EL	AF - EL - PX	DICE	0.84	0.85	0.86	2.38%	1.18%
202)	Abdomen	s	3D U-Net	CT	Н	EL	AF - EL - PX	DICE	0.84	0.85	0.86	2.38%	1.18%
194)	Abdomen	S	U-Net	CT	Н	AF	111 - 111 - 111	ACC	0.954	0.00	0.946	-0.84%	-
194)	Abdomen	S	U-Net	CT	Н	AF		ACC	0.954		0.946	-0.84%	
			U-Net U-Net										
203)	Arm	S		US	H	EL	AF	PACC	0.98	0.998	0.999	1.94%	0.10%
(204)	Arteries	S	U-Net		OTH	REC	AF	DICE	0.615	0.696	0.756	22.93%	8.62%
(204)	Arteries	S	U-Net	IVOCT	OTH	REC	AF	DICE	0.615	0.696	0.756	22.93%	8.62%
116)	Bladder	S	U-Net	CT	H	AF		DICE	0.904	-	0.914	1.11%	-
205)	Bones	C	TF	XR	FRC	ER	AF	ACC		0.786	0.805	-	2.35%
205)	Bones	C	TF	XR	FRC	ER	AF	ACC		0.786	0.805		2.35%
(206)	Colon	D	3D FCN	CT	POL	AF - EL	AF	FROC	-	0.9	0.97		7.78%
(207)	Colon	C	3D CNN	CT	POL	GN	GAN	AUC	0.811	0.846	0.881	8.63%	4.14%
200)	Eye	S	U-Net	OCT	H	MOD - FM - LRN	MOD - LRN	DICE	0.724	0.733	0.752	3.87%	2.59%
208)	Head	D	YoloX	CT	FRC	AF -FM	AF	AP	0.688	0.692	0.698	1.45%	0.87%
208)	Head	D	YoloX	CT	FRC	AF -FM	AF	AP	0.688	0.692	0.698	1.45%	0.87%
209)	Hip	C	Custom	CT	FRC	FM		ACC	0.519	-	0.977	88.19%	
209)	Hip	C	Custom	CT	FRC	FM		ACC	0.519		0.977	88.19%	
210)	Humerus	s	U-Net	US	Н	GAN - AF - PX	AF - PX	DICE	0.515	0.605	0.66	13.79%	9.09%
				CT									
199)	Kidney	S	U-Net		CAN	AF - ER - PX - LRN	AF	DICE	0.749	0.832	0.84	12.15%	0.96%
48)	Kidney	S	U-Net	MR	H	AF	-	DICE	0.935	-	0.954	2.03%	-
211)	Kidney	S	U-Net	CT	H	GAN - AF	AF	DICE	0.92	0.94	0.944	2.61%	0.43%
212)	Kidney	S	U-Net	US	H	MOD	-	DICE	0.93	-	0.942	1.29%	-
213)	Knee	S	U-Net	MR	H	GAN		DICE	0.836	-	0.847	1.34%	-
213)	Knee	S	U-Net	MR	H	GAN		DICE	0.836	-	0.847	1.34%	-
214)	Liver	C	CNN	CT	CAN	GAN	AF	SNS		0.797	0.865		8.53%
215)	Liver	s	U-Net	CT	CAN	AF - FM	-	DICE	0.915	-	0.945	3.28%	
41)	Liver	S	3D U-Net	CT	Н	MOD		ACC	0.861		0.883	2.56%	
216)	Liver	C	CNN	CT	CAN	GAN	GAN + AF	ACC	0.802		0.855	6.61%	
							GAN + AF						
48)	Liver	S	U-Net	MR	H	AF	-	DICE	0.961		0.962	0.10%	-
171)	Liver	S	CNN	CT	CAN	GAN		DICE	0.912	-	0.941	3.25%	-
217)	Liver	C	CNN	CT	L	GAN	AF	ACC	-	0.786	0.857	-	9.03%
218)	Liver	C	Custom	MR	CAN	AF		AUC	0.799	-	0.826	3.38%	-
218)	Liver	C	Custom	MR	CAN	AF		AUC	0.799	-	0.826	3.38%	-
219)	Liver	S	U-Net	CT	H	AF - PX	-	DICE	0.896	-	0.915	2.12%	
219)	Liver	S	U-Net	CT	CAN	AF - PX	-	DICE	0.659	-	0.702	6.53%	
201)	Liver	S	U-Net	MR	CAN	AF - EL - PX - LRN	AF	DICE		0.603	0.635		5.31%
201)	Liver	S	U-Net	MR	CAN	AF - EL - PX - LRN	AF	DICE		0.603	0.635		5.31%
2201)	Liver	C	CNN	CT	CAN	GAN - AF	AF	ACC		0.84	0.853		1.55%
												•	
220)	Liver	C	CNN	CT	CAN	GAN - AF	AF	ACC		0.84	0.853	•	1.55%
113)	Liver	S	3D U-Net	CT - MR	CAN	AF - PX - FM	FM	DICE	0.492	0.298	0.66	34.15%	121.48%
(113)	Liver	S	3D U-Net	CT - MR	CAN	AF - PX - FM	FM	DICE	0.492	0.298	0.66	34.15%	121.48%
(211)	Liver	S	U-Net	CT	H	GAN - AF	AF	DICE	0.944	0.941	0.947	0.32%	0.64%
221)	Muscle	S	U-Net	CT	H	AF - PX	-	DICE	0.833	-	0.915	9.84%	-
198)	Muscle	S	U-Net	MR	NMD	GAN - EL	EL	DICE	0.84	0.87	0.88	4.76%	1.15%
(48)	Pancreas	S	U-Net	MR	H	AF	-	DICE	0.86	-	0.864	0.47%	
13)	Pelvis	S	U-Net	MR	Many	EL		PACC	0.724	-	0.841	16.16%	
41)	Prostate	S	3D U-Net	MR	н	MOD		ACC	0.742		0.868	16.98%	
222)	Prostate	S	U-Net	MR	CAN	AF - GAN		DICE	0.678		0.738	8.85%	
52)	Prostate	S	U-Net	MR	Н	AF - EL - PX - LRN	MX	DICE	-	0.706	0.733		3.80%
52)	Prostate	S	U-Net	MR	Н	AF - EL - PX - LRN	MX	DICE		0.706	0.733		3.80%
							MA		-	0.706		-	3.80%
223)	Prostate	S	U-Net	MR - PET - CT	CAN	AF - FM	-	DICE	0.776	-	0.819	5.54%	-
224)	Prostate	C	CNN	MR	CAN	AF		ACC	0.802	-	0.85	6.00%	-
224)	Prostate	C	CNN	MR	CAN	AF		ACC	0.802	-	0.85	6.00%	-
225)	Prostate	S	U-Net	MR	H	MOD	-	DICE	0.813	-	0.863	6.15%	-
40)	Prostate	S	U-Net	XR	H	MOD	AF	DICE	0.785	0.81	0.848	8.03%	4.69%
226)	Prostate	S	AMS	MR	CAN	GAN	-	DICE	0.858	-	0.882	2.80%	-
116)	Prostate	S	U-Net	CT	CAN	AF		DICE	0.697	-	0.731	4.93%	-
227)	Prostate	S	U-Net	US	CAN	MOD		DICE	0.906	-	0.908	0.17%	
227)	Prostate	S	U-Net	US	CAN	MOD		DICE	0.906		0.908	0.17%	
228)	Prostate	C	Custom	MR	CAN	GAN	AF	ACC	-	0.815	0.892		9.45%
	Prostate	s	3D U-Net	MR	CAN	AF - EL - PX		DICE	0.896	0.908	0.913	1.90%	0.55%
56)			3D U-Net	MR	CAN	AF - EL - PX		DICE	0.896	0.908	0.913	1.90%	0.55%
										0.505			0.3376
56)	Prostate	S			H	AF - PX	-	DICE	0.938	-	0.943	0.53%	-
56) 229)	Prostate Rectal	S	MRSN	MR				DICE	0.732		0.742	1.37%	-
56) 229) 229)	Prostate Rectal Rectal	S S	MRSN MRSN	MR	CAN	AF - PX							
56) 229) 229) 116)	Prostate Rectal Rectal Rectum	s s	MRSN MRSN U-Net	MR CT	CAN CAN	AF		DICE	0.718	-	0.753	4.97%	-
(56) (56) (229) (229) (116) (197)	Prostate Rectal Rectal	S S	MRSN MRSN	MR	CAN CAN H	AF MOD			0.718 0.743	-		4.97% 27.59%	-
56) 229) 229) 116) 197)	Prostate Rectal Rectal Rectum	s s	MRSN MRSN U-Net	MR CT	CAN CAN H OTH	AF MOD AF - EL	-	DICE PACC DICE		:	0.753		
56) 229) 229) 116) 197)	Prostate Rectal Rectum Spine	S S S	MRSN MRSN U-Net U-Net	MR CT CT	CAN CAN H	AF MOD	-	DICE PACC	0.743		0.753 0.948	27.59%	
56) 229) 229) 116) 197) 193)	Prostate Rectal Rectum Spine Spine Spine	S S S S	MRSN MRSN U-Net U-Net 2D U-Net	MR CT CT MR	CAN CAN H OTH	AF MOD AF - EL	-	DICE PACC DICE	0.743 0.921		0.753 0.948 0.913	27.59% -0.87%	- - -
56) 229) 229) 116) 197) 193) 193)	Prostate Rectal Rectal Rectum Spine Spine Spine Spine Spine	s s s s	MRSN U-Net U-Net 2D U-Net 2D U-Net U-Net	MR CT CT MR MR	CAN CAN H OTH OTH	AF MOD AF - EL AF - EL AF	- - AF	DICE PACC DICE DICE DICE	0.743 0.921 0.921 0.944	:	0.753 0.948 0.913 0.913	27.59% -0.87% -0.87% 0.00%	- - - - - 3.26%
56) 229) 229) 116) 197) 193) 193) 48) 211)	Prostate Rectal Rectal Rectum Spine Spine Spine Spine Spleen Spleen	s s s s s	MRSN U-Net U-Net 2D U-Net 2D U-Net U-Net U-Net U-Net	MR CT CT MR MR CT	CAN CAN H OTH OTH H	AF MOD AF - EL AF - EL AF GAN - AF	- AF AF	DICE PACC DICE DICE DICE DICE	0.743 0.921 0.921	- - - 0.89	0.753 0.948 0.913 0.913 0.944 0.919	27.59% -0.87% -0.87%	- - - - - 3.26% 8.56%
56) 229) 229) 116) 197) 193) 193) 48) 211) 230)	Prostate Rectal Rectum Spine Spine Spine Spine Spieen Spleen Spleen	s s s s s s	MRSN U-Net U-Net 2D U-Net 2D U-Net U-Net U-Net U-Net U-Net U-Net	MR CT CT MR MR CT CT CT	CAN CAN H OTH OTH H OTH	AF MOD AF - EL AF - EL GAN - AF AF - GAN	AF	DICE PACC DICE DICE DICE DICE DICE	0.743 0.921 0.921 0.944 0.884	- - 0.89 0.488	0.753 0.948 0.913 0.913 0.944 0.919	27.59% -0.87% -0.87% 0.00%	8.56%
56) 229) 229) 116) 197) 193) 193) 48) 211) 230)	Prostate Rectal Rectum Spine Spine Spine Spine Spleen Spleen Spleen Spleen	s s s s s s s s s s	MRSN U-Net U-Net 2D U-Net 2D U-Net U-Net U-Net U-Net U-Net U-Net U-Net	MR CT CT MR MR CT CT CT CT	CAN CAN H OTH OTH H OTH OTH	AF MOD AF - EL AF - EL AF GAN - AF AF - GAN AF - GAN	AF AF	PACC DICE DICE DICE DICE DICE DICE	0.743 0.921 0.921 0.944 0.884	- - - 0.89	0.753 0.948 0.913 0.913 0.944 0.919 0.53	27.59% -0.87% -0.87% 0.00% 3.96%	
(56) (229) (229) (116) (197) (193) (193) (48) (211) (230) (230) (196)	Prostate Rectal Rectal Rectum Spine Spine Spine Spleen Spleen Spleen Spleen Spleen Spleen		MRSN U-Net U-Net 2D U-Net 2D U-Net U-Net U-Net U-Net U-Net U-Net U-Net U-Net	MR CT CT MR MR MR CT CT CT CT XR	CAN CAN H OTH OTH H OTH OTH H	AF MOD AF - EL AF - EL AF GAN - AF AF - GAN AF - GAN	AF	DICE PACC DICE DICE DICE DICE DICE DICE ACC	0.743 0.921 0.921 0.944 0.884	- - 0.89 0.488	0.753 0.948 0.913 0.913 0.944 0.919 0.53 0.53	27.59% -0.87% -0.87% 0.00% 3.96% - - 0.02%	8.56%
(56) (229) (229) (116) (197) (193) (193) (48) (211) (230) (230) (196) (190)	Prostate Rectal Rectal Rectum Spine Spine Spine Spleen Spleen Spleen Spleen Teeth	S S S S S S S S C	MRSN U-Net U-Net 2D U-Net 2D U-Net U-Net U-Net U-Net U-Net U-Net U-Net U-Net	MR CT CT MR MR MR CT CT CT XR	CAN CAN H OTH OTH H OTH OTH OTH COTH OTH COTH	AF MOD AF - EL AF - EL GAN - AF AF - GAN AF - GAN AF - GAN AF	AF AF	DICE PACC DICE DICE DICE DICE DICE DICE ACC AUC	0.743 0.921 0.921 0.944 0.884 - - 0.947	- - 0.89 0.488	0.753 0.948 0.913 0.913 0.944 0.919 0.53 0.53 0.948	27.59% -0.87% -0.87% -0.00% 3.96% - - - 0.02% 9.30%	8.56% 8.56% -
(56) (229) (229) (116) (197) (193) (193) (48) (211) (230) (230) (196) (190) (192)	Prostate Rectal Rectal Rectum Spine Spine Spine Spleen Spleen Spleen Spleen Spleen Spleen	S S S S S S C C C	MRSN U-Net U-Net 2D U-Net 2D U-Net U-Net U-Net U-Net U-Net U-Net U-Net U-Net U-Net CNN	MR CT CT MR MR CT CT CT XR XR	CAN CAN H OTH OTH H OTH OTH COTH H CAN - CY	AF MOD AF - EL AF - EL AF GAN - AF AF - GAN AF - GAN AF - PX AF - PX	AF AF	DICE PACC DICE DICE DICE DICE DICE ACC AUC AUC	0.743 0.921 0.921 0.944 0.884 - - 0.947 0.86 0.89	- - 0.89 0.488	0.753 0.948 0.913 0.913 0.944 0.919 0.53 0.53 0.948 0.94	27.59% -0.87% -0.87% 0.00% 3.96% - - 0.02% 9.30% -1.12%	8.56%
56) 229) 229) 116) 197) 193) 193) 48) 211) 230) 230) 196) 190) 192) 231)	Prostate Rectal Rectal Rectum Spine Spine Spine Spleen Spleen Spleen Spleen Teeth	S S S S S S C C C C	MRSN U-Net U-Net 2D U-Net 2D U-Net U-Net U-Net U-Net U-Net U-Net U-Net U-Net U-Net CONN CNN	MR CT CT MR MR CT CT CT XR XR XR	CAN CAN H OTH OTH H OTH OTH CAN - CY H OS	AF MOD AF - EL AF - EL AF GAN - AF AF - GAN AF - GAN AF - AF AF - PX AF - PX AF - PX AF	AF AF	DICE PACC DICE DICE DICE DICE DICE ACC AUC AUC ACC	0.743 0.921 0.921 0.944 0.884 - - 0.947 0.86 0.89	- - 0.89 0.488 0.488	0.753 0.948 0.913 0.913 0.914 0.919 0.53 0.53 0.948 0.94 0.98	27.59% -0.87% -0.87% -0.87% 3.96% 0.02% 9.30% -1.12% 5.95%	8.56% 8.56% -
(56) (229) (229) (116) (197) (193) (193) (48) (211) (230) (230) (196)	Prostate Rectal Rectal Rectum Spine Spine Spine Spleen Spleen Spleen Teeth Teeth	S S S S S S C C C	MRSN U-Net U-Net 2D U-Net 2D U-Net U-Net U-Net U-Net U-Net U-Net U-Net U-Net U-Net CNN	MR CT CT MR MR CT CT CT XR XR	CAN CAN H OTH OTH H OTH OTH COTH H CAN - CY	AF MOD AF - EL AF - EL AF GAN - AF AF - GAN AF - GAN AF - PX AF - PX	AF AF	DICE PACC DICE DICE DICE DICE DICE ACC AUC AUC	0.743 0.921 0.921 0.944 0.884 - - 0.947 0.86 0.89	- - 0.89 0.488 0.488	0.753 0.948 0.913 0.913 0.944 0.919 0.53 0.53 0.948 0.94	27.59% -0.87% -0.87% 0.00% 3.96% - - 0.02% 9.30% -1.12%	8.56% 8.56% - -
56) 229) 229) 116) 197) 193) 193) 48) 2211) 230) 196) 199) 192) 231)	Prostate Rectal Rectum Spine Spine Spine Spleen Spleen Spleen Spleen Teeth Teeth Teeth	S S S S S S C C C C	MRSN U-Net U-Net 2D U-Net 2D U-Net U-Net U-Net U-Net U-Net U-Net U-Net U-Net U-Net CONN CNN	MR CT CT MR MR CT CT CT XR XR XR	CAN CAN H OTH OTH H OTH OTH CAN - CY H OS	AF MOD AF - EL AF - EL AF GAN - AF AF - GAN AF - GAN AF - AF AF - PX AF - PX AF - PX AF	AF AF	DICE PACC DICE DICE DICE DICE DICE ACC AUC AUC ACC	0.743 0.921 0.921 0.944 0.884 - - 0.947 0.86 0.89	- - 0.89 0.488 0.488	0.753 0.948 0.913 0.913 0.914 0.919 0.53 0.53 0.948 0.94 0.98	27.59% -0.87% -0.87% -0.87% 3.96% 0.02% 9.30% -1.12% 5.95%	8.56% 8.56% - -
(56) (229) (229) (116) (197) (193) (193) (48) (211) (230) (230) (196) (190) (192) (231)	Prostate Rectal Rectum Spine Spine Spine Spleen Spleen Spleen Teeth Teeth Teeth Teeth	S S S S S S C C C C C	MRSN MRSN U-Net U-Net 2D U-Net 2D U-Net U-Net U-Net U-Net U-Net U-Net U-Net C-NN CNN CNN	MR CT CT MR MR MR CT CT CT CT XR XR XR XR XR	CAN CAN H OTH OTH OTH H H CAN - CY H OS CAN	AF MOD MOD AF - EL AF - EL AF GAN - AF AF - GAN AF - GAN AF - PX AF - PX AF AF	AF AF AF (*) - -	DICE PACC DICE DICE DICE DICE DICE ACC AUC AUC ACC ACC	0.743 0.921 0.921 0.944 0.884 - - 0.947 0.86 0.89 0.925 0.699	- - 0.89 0.488 0.488	0.753 0.948 0.913 0.913 0.944 0.919 0.53 0.948 0.94 0.88 0.994 0.98	27.59% -0.87% -0.87% 0.00% 3.99%	8.56% 8.56% - - -
56) 229) 229) 116) 197) 193) 48) 211) 230) 230) 196) 199) 192) 231) 232) 233)	Prostate Rectal Rectum Spine Spine Spine Spleen Spleen Spleen Spleen Teeth Teeth Teeth Teeth Treeth	S S S S S S C C C C C C	MRSN MRSN U-Net U-Net 2D U-Net 2D U-Net U-Net U-Net U-Net U-Net U-Net U-Net CNN CNN CNN CNN	MR CT CT MR MR MR CT CT CT CT XR XR XR XR XR XR XR US	CAN CAN H OTH OTH H COTH H COTH CON CAN CAN	AF MOD MOD AF - EL AF - EL AF GAN - AF AF - GAN AF - GAN AF - PX AF - PX AF GAN AF	AF AF AF (*) - - - - AF AF + PX	DICE PACC DICE DICE DICE DICE DICE ACC AUC ACC ACC ACC ACC	0.743 0.921 0.921 0.944 0.884 - - 0.947 0.86 0.89 0.925 0.699 0.701	0.89 0.488 0.488 - - - - 0.756	0.753 0.948 0.913 0.913 0.944 0.919 0.53 0.948 0.94 0.88 0.98 0.994 0.904	27.59% -0.87% -0.87% 0.00% 3.96% 0.02% 9.30% -1.12% 5.95% 29.33% 30.53%	8.56%
56) 229) 229) 116) 197) 193) 48) 211) 230) 230) 199) 192) 231) 232) 233) 234)	Prostate Rectal Rectal Rester Spine Spine Spine Spleen Spleen Spleen Spleen Spleen Teeth Teeth Teeth Teeth Thyroid Thyroid	s s s s s s s c c c c c c c c	MRSN MRSN U-Net CNN CNN CNN CNN CNN CNN CNN	MR CT CT MR MR MR CT CT CT CT XR XR XR XR XR XR US US	CAN CAN H OTH OTH H OTH OTH H CAN - CY H CAN CAN CAN	AF MOD AF - EL AF - EL AF AF GAN - AF AF - GAN AF - GAN AF AF - PX AF - PX AF GAN AF GAN AF	AF AF AF (*) - - - - AF	DICE PACC DICE DICE DICE DICE DICE ACC AUC ACC ACC ACC ACC ACC	0.743 0.921 0.921 0.944 0.884 - - 0.947 0.86 0.89 0.925 0.699 0.701 0.845		0.753 0.948 0.913 0.913 0.914 0.919 0.53 0.53 0.948 0.94 0.98 0.904 0.915 0.953	27.59% -0.87% -0.87% 0.00% 3.96% -0.22% 9.30% -1.12% 5.95% 29.33% 30.53% 12.75%	8.56% - - - - - 21.03% 7.46%
(56) (229) (229) (116) (197) (193) (193) (48) (211) (230) (196) (190) (192) (231) (232) (233)	Prostate Rectal Rectum Spine Spine Spine Spleen Spleen Spleen Spleen Teeth Teeth Teeth Teeth Treeth	s s s s s s s c c c c c c c	MRSN MRSN U-Net U-Net 2D U-Net 2D U-Net U-Net U-Net U-Net U-Net U-Net U-Net U-Net U-Net CNN CNN CNN CNN CNN CNN CNN	MR CT CT MR MR MR CT CT CT CT CT XR XR XR XR XR XR US US	CAN CAN H OTH OTH OTH OTH OTH OTH OTH OTH CAN - CY H OS CAN CAN CAN	AF MOD MOD AF - EL AF - EL AF GAN - AF AF - GAN AF - GAN AF - PX AF - PX AF GAN AF	AF AF AF (*) - - - - AF AF + PX	DICE PACC DICE DICE DICE DICE DICE ACC AUC ACC ACC ACC ACC	0.743 0.921 0.921 0.944 0.884 - - 0.947 0.86 0.89 0.925 0.699 0.701	0.89 0.488 0.488 - - - - 0.756	0.753 0.948 0.913 0.913 0.944 0.919 0.53 0.53 0.53 0.944 0.88 0.98 0.904 0.915 0.953	27.59% -0.87% -0.87% 0.00% 3.96%	8.56%

Table 6: Comparison of **66** studies on data augmentation techniques for other organs, not covered by previous tables. See Table 1 for an explanation of the acronyms.

augmentation is around 10% across all organs. Studies related to the heart, lung and breast report a higher benefit associated to data augmentation than brain and other organs. This difference could be explained by various factors: better data augmentation techniques are available for organs that are more heavily researched; datasets are typically smaller; some organs/modalities are susceptible to higher intersubject or inter-scanner variability, thus benefitting more from data augmentation.

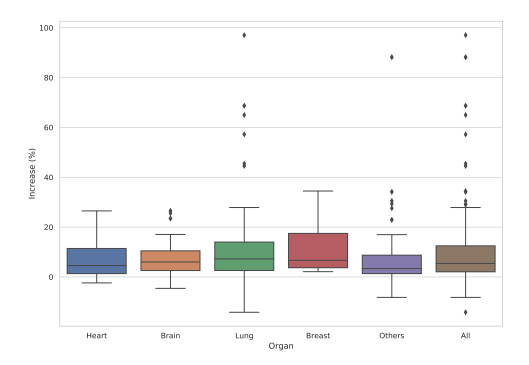


Figure 4: Box plot of the relative performance increase associated to data augmentation (DA vs. no-DA) for each organ.

We further evaluated the relative performance benefit that can be ripped by exploiting more complex data augmentation techniques. For this analysis, we split data augmentation techniques in two categories: affine and non-affine (other). We choose affine transformations as the baseline given their simplicity, ease of use and widespread adoption in the medical domain. Methods that combine multiple data augmentation techniques were tagged with the most complex transformation (in reverse order: GAN, model-based, reconstruction-based, feature mixing, erasing, pixel-level, affine).

Figure 5 reports the distribution of the relative performance increase for three groups of papers: affine DA vs. no-DA (41 studies), non-affine DA vs. no-DA (144 studies) and non-affine DA vs. affine DA (56 studies). Results show that simple affine transformations offer a substantial performance advantage with respect to not using DA at all. More complex techniques yield higher performance, at the expense of greater complexity. While these distributions must be interpreted with great caution, since we are pooling very heterogeneous studies without adjustments, they overall suggest that "simple" data augmentation techniques should always be tried before moving to more complex solutions.

Finally, in Figure 6 we report the average relative increase in performance associated with the use of DA for classification (78), segmentation (93), and detection (14) tasks. Overall, classification tasks seem to benefit more from data augmentation than segmentation tasks. However, these results should be interpreted with caution as tasks are not evenly distributed across organs and modalities, and segmentation tasks are often associated with volumetric (3D) modalities, which are more complex to augment than 2D images.

We concluded this analysis by plotting the relative increase in performance with respect to the size of the training set, on a logarithmic scale (reported in Figure 7). Only studies comparing the results with and without DA (182 studies) were included in this graph. More than half of the datasets are small dataset with less than 1000 samples. However, correlation between relative performance increase and training

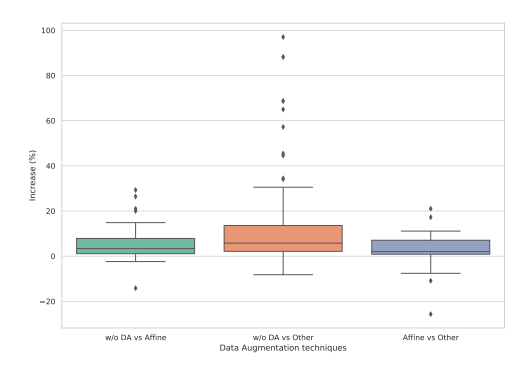


Figure 5: Box plot of the relative performance increase associated to different types of data augmentation (affine and non-affine).

set size is poor (-5.11e-05), suggesting that DA can be beneficial regardless of the size of the training set. We attribute this to two phenomena: i) data augmentation promotes learned invariances that, in turn, enable more effective generalization, and ii) data augmentation is often used to balance the distribution of the training set, typically by generating additional lesion cases. In the case of severe data imbalance, the number of examples from minority classes is likely insufficient even in medium-to large-scale datasets.

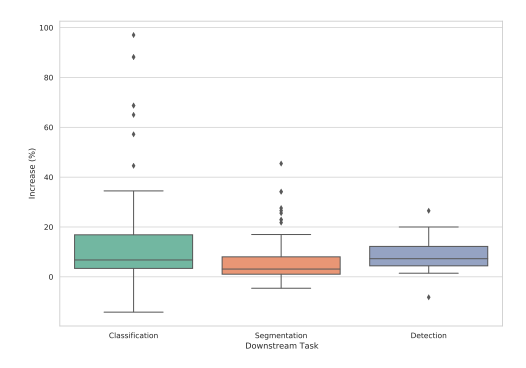


Figure 6: Box plot of the relative performance increase associated to data augmentation (DA vs. no-DA) for each task.

4.3. RQ4: Which data augmentation methods have not been explored in the medical domain?

To answer RQ4, we compared the results of our SLR with previous literature review tackling the more general computer vision field (6). Most of the techniques adopted in general computer vision have found widespread adoption in the medical domain, with a few notable exceptions: random erasing, feature mixing, and learnable data augmentation. These differences could be attributed to publication bias, since authors are more likely to publish techniques that yield positive rather than negative results. For instance, random erasing was introduced to make DNNs

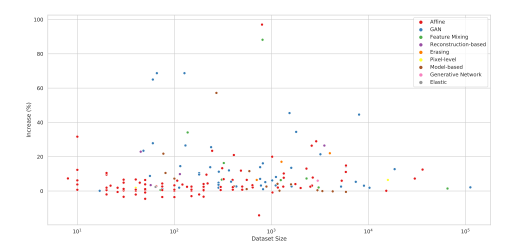


Figure 7: Scatter plot of percentage increase associated to data augmentation with respect to the training set size (in logarithmic scale). (Best viewed in color)

robust to occlusions, which however do not occur in medical imaging. Hence, it is conceivable that their use is not particularly effective in this domain.

On the other hand, a more widespread adoption of learnable DA techniques could in principle discover policies that are uniquely tailored to the medical domain. However, many techniques that have been proposed, such as those based on reinforcement learning, are computationally intensive, as they require training multiple networks to determine which transformations are more challenging at a given stage of training and should be selected with higher probabilities. This fact, coupled with the fact that medical images are typically bigger than most RGB datasets, may have made these techniques less appealing for the medical imaging community.

5. Comparison with previous research

We have compared our findings with those of Nalepa et al., who reviewed the data augmentation techniques used in 20 submissions to the Multimodal Brain Tumor Segmentation Challenge (BraTS 2018) challenge (9). Participants in the BraTS challenge received multimodal MRI data of brain tumor patients, all co-registered to a common anatomical reference. The task is to build a supervised segmentation model, which is then tested on unseen data released during the testing phase. Although most proposed augmentations are single-modal, they can be easily applied to each co-registered series, thus yielding multi-modal samples.

Briefly summarized, Nalepa and colleagues found that 75% of the submissions included affine transformations (mostly horizontal flip); all but one paper employed 2D, rather than 3D transformations. Pixel-level transformations were less common, but appeared in the three top-performing solutions (236; 237; 238). Interestingly, one of the submissions experimentally observed that simpler networks combined with carefully designed data augmentation outperformed more complex solutions (236). Only one submission used generative models (a combination of synthetic images and cycle-GANs). In the small number of submissions that reported performance with and without data augmentation, the relative DICE score increased between -5% to +7%, and the relative Hausdorff distance decreased between -86% to +26%. For comparison, in the 49 studies related to brain image analysis (analyzed in detail in Section 4.2.1), one out of two papers described generative models or model-based synthetic image generation techniques, sometimes in combination with affine and pixel-level transformations. Studies focusing on segmentation techniques reported a relative increase in DICE score between 2% and 26%.

The comparison between Nalepa's findings and ours highlights the increasing role

that GANs are playing in the data augmentation landscape. Indeed, the present SLR covers articles published from 2018 (when the BraTS challenge analyzed in (9) took place) to 2022. On the other hand, articles that evaluate the impact of data augmentation are likely to analyze more complex techniques (and thus more interesting, from an editorial perspective). It is possible, however, that practical adoption of generative models is lagging behind, given the significant gap in implementation time and computational resources with respect to transformations that are available out of the box from most deep learning frameworks.

Another study that provides key insights to integrate and complement of our findings is that of Zhang et al. (239), who investigated the performance of BigAug on different modalities and organs. BigAug is a sequence of n stacked transformations, selected from a pool of options that the authors divide in three categories: image quality, appearance and spatial configuration. Image quality and appearance include pixel-level transformations: noise insertion, blurriness and sharpness simulate the effect of varying image quality, whereas brightness, contrast and intensity perturbation emulate the properties of different scanners and imaging protocols. Spatial configuration transformations, on the other hand, include both affine and elastic transformations that simulate inter- and intra-patient differences due to, e.g., positioning in the scanner.

The main conclusion of this study, as reported also in Tables 3 and 6, is that BigAug outperforms any individual transformation, which is line with the findings of this review. Additionally, the impact of each individual transformation is thoroughly investigated on datasets from two modalities (MR and US). The results suggest an important correlation between the modality and the optimal transformations. In the case of MRI, transformations such as sharpening, contrast enhancement, brightness, and intensity perturbation appear to be the most beneficial, whereas in the case of

US, the most effective transformations are scaling, brightness, blurring, and contrast. Thus, in the case of MRI, generalizing to different intensity distributions appears to be more important than accounting for shape variations, which on the other hand are most prominent in US.

In addition, Zhang et al. prove that the benefits of data augmentation persist, and are even amplified, when the trained model is transferred to a different target dataset, such as one acquired in a different center (239). As an example, in the case of prostate segmentation in MR images, the model trained without data augmentation has a good performance on the source dataset (DICE 0.896), which drops significantly on the three target datasets (DICE 0.604, 0.580 and 0.768, respectively). On the other hand, the model trained with BigAug, despite having similar performance on the source dataset (DICE 0.913), generalizes significantly better to different target datasets (DICE 0.802, 0.850 and 0.865, respectively). When averaged across all experiments, BigAug slightly outperformed the baseline on the source dataset (mean DICE 0.916 vs. 0.891), but showed much stronger generalization (mean DICE 0.80 vs. 0.498). In conclusion, even when data augmentation shows minor improvements on the source dataset (which is the setting analyzed in this SLR), further benefits could be obtained in terms of cross-domain generalization. These properties should and could be further investigated in modalities and organs not covered by Zhang et al. (239).

6. Discussion, limitations, and key findings

In this SLR, we have analyzed more than 300 articles that proposed and / or evaluated data augmentation techniques for modalities such as CT, MR, and XR imaging. Our key findings can be summarized as follows:

- Although affine and pixel-level transformations remain the most adopted transformations, papers published between 2018 and 2022 have substantially shifted their attention towards the generation of synthetic data for training, either based on generative models or, for selected organs for which computational models are available, through model-based computer-simulated imagery.
- Traditional data augmentation techniques, based on transformation of existing data, and generative techniques serve two different purposes. Thus, practitioners exploiting generative models should consider including also conventional transformations in their data pipelines. In particular, traditional data augmentation plays a fundamental role in making DNNs robust to variations introduced by the acquisition system, as well as to inter- and intra-patient variability. Among traditional data augmentation, pixel-level transformations appear to be under-utilized with respect to their potential.
- Generative techniques, on the other hand, are more flexible and can be used to balance training data sets by generating cases with specific characteristics (149; 30), inserting artificial lesions (155; 72; 30), transferring data across domains (156) and alleviating the need for expensive data labeling, as labels can be co-generated with the corresponding image (76).
- Learnable data augmentation and feature mixing, despite showing promising results in other domains, have been scarcely investigated in medical imaging. Recent advances in generative models, that have led to photographic images of unprecedented visual fidelity (27), have also untapped potential in the medical domain.
- Data augmentation can be particularly effective in more complex training set-

ting, such as one-shot/few-shot learning (70; 76; 32; 52), cross-domain transfer learning (239), and federated learning (119). Authors proposing new techniques for data augmentation should consider including experiments on cross-domain generalization, as done in (239).

- The uptake of more complex data augmentation strategies could be facilitated by the availability of open-source tools, including code and pre-trained models. Many authors tend to adopt transformations available in standard deep learning libraries, which however do not fully support the needs of medical imagery. For instance, adapting even standard affine transformations for on-the-fly data augmentation of volumetric 3D images is not trivial (239). Recently, dedicated medical imaging platforms, such as MONAI (12), provide tailored data augmentation pipelines, including several 3D transformations.
- In the future, available resources could and should extend to generative models and synthetic imagery, to facilitate their adoption. As an example, open-source tools are already available to support in silico virtual clinical trials for regulatory purposes (240), and may in the future extend to other uses as well. Synthetic datasets will likely play a prominent role in future medical imaging deep learning research. For instance, NVIDIA and King's College London have recently announced the release of a large-scale dataset containing more than 100,000 synthetic MRI images, generated by a combination of variational autoencoder and transformers (241).

Despite the large number of papers included in the analysis, our study has two potential limitations:

• Like all SLRs, our results may reflect publication bias. Since we primarily

included papers that specifically reported performance with and without data augmentation, we may risk overestimating the practical relevance of generative and model-based techniques that, while undoubtedly beneficial in terms of performance, are much more complex and computationally expensive to implement than more traditional data transformations. Papers that mentioned the use of data augmentation without investigating its impact on downstream performance predominantly made use of standard affine and pixel-level transformations. As mentioned before, the gap between research and practice could be alleviated by the development of open-source AI platforms and tools.

• We attempted to quantitatively combine results from heterogeneous studies, which may not be directly comparable due to different datasets used or experimental conditions. In this respect, it is encouraging to see how our results align with previous studies which compared a wide range of data augmentation strategies on the same dataset, as detailed in Section 5. In the future, more controlled experiments should compare different data augmentation strategies on the same datasets, especially considering how the results may vary across modalities.

7. Conclusions

In this SLR, we have analyzed more than 300 articles that proposed and / or evaluated data augmentation techniques for different imaging modalities (e.g., CT, MR, XR, MG and functional imaging), organs (e.g., brain, heart, lung, breast, liver, prostate and many others), and tasks (classification, segmentation and detection). Overall, we found consistent benefits, across all organs, modalities, and tasks, with the use of data augmentation, from the simplest affine transformations to the most

complex generative models. Through extensive qualitative and quantitative analysis, as well as comparison with previous surveys and experimental studies, we draw a comprehensive and up-to-date picture of the state of the art of this lively field of research, as well as highlight current research gaps and directions of improvement.

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Appendix A. Examples of data augmentation techniques

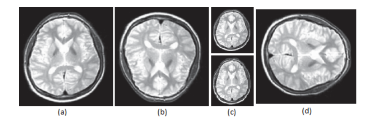


Figure A.1: Example of affine transformations applied to a brain magnetic resonance image. Respectively: (a) original image (b) vertical flipping (c) scaling (d) horizontal flipping

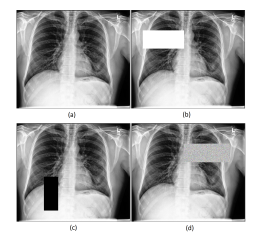


Figure A.2: Example of random erasing transformation applied to chest x-ray image from CheXpert (242). Respectively: (a) original image (b) white erasing value (c) black erasing value (d) random erasing value

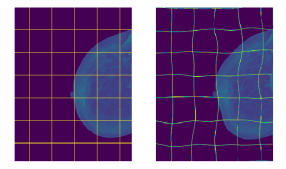


Figure A.3: Example of elastic transformation applied to a mammography. A Gaussian filter is applied after the distortion. (Image credit: (172))

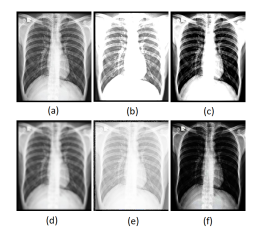


Figure A.4: Example of pixel-level transformations applied to a chest x-ray image. Respectively: (a) original image (b) brightness (c) contrast (d) blur (e) light gamma correction (f) dark gamma correction

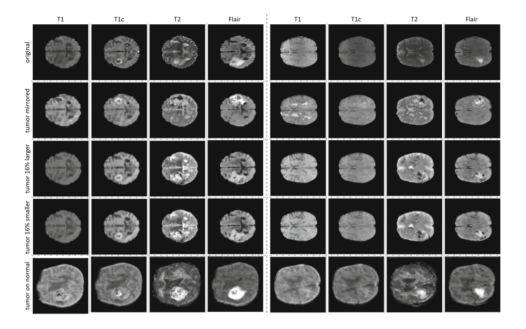


Figure A.5: Example of GAN generated images. The first row depicts the real images on which the synthetic tumors were based. Generated images without adjustment of the segmentation label are shown in the second row. Examples of generated images with various adjustments to the tumor segmentation label are shown in the third through fifth rows. The last row depicts examples of synthetic images where a tumor label is placed on a tumor-free brain label from the ADNI data set. (Image credit: (30))

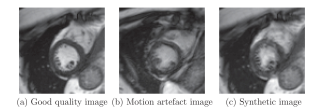


Figure A.6: Example of reconstruction-based method: a good quality cine CMR image (a), an image with blurring motion artifacts (b), and a k-space corrupted image (c). The k-space corruption process is able to simulate realistic motion-related artifacts (Image credit: (42))

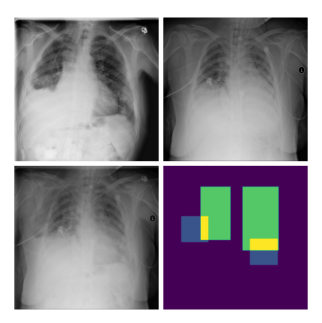


Figure A.7: Example of mix-up generated images. In the first row there original chest x-rays, in the bottom left there is the result of mix-up augmentation and in the bottom right the corresponding mask.(Image credit: (243))

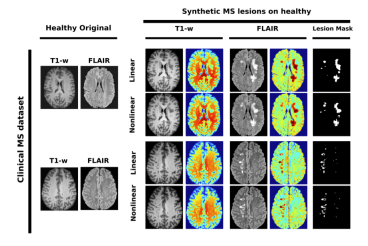


Figure A.8: Example of model-based method: synthetic multiple sclerosis lesions generated on a healthy subject. Slices are also displayed using jet color maps to visually enhance the intensities (Image credit: (38))

Appendix B. Supplementary material: methods

Digital Library	Search string
ACM Digital Library	[All: "data augmentation"] AND [All: "medical imaging"]
IEL-IEEE	("Abstract":Data Augmentation) AND ("Abstract":Medical Imaging)
PubMed	("data augmentation") AND ("imaging")
Science Direct	Title, abstract, keywords: ("Data Augmentation") AND ("Medical")

Table B.7: Digital libraries with respective URL and search string.

	Inclusion criteria	Exclusion criteria
Language	English	Any other languages
Time window	January 2018 to July	Published before 2018
	2022	
Study	Original research	Review, Survey, Commentaries,
		Opinion paper, Abstract
Data Augmentation	Used	Not used
Full text	Available	Unavailable
Modality	CT, XR, MR, PET,	Dermatology, Ophthalmology,
	US	Histopathology
Context	Deep Learning	Any other type of machine learning
Downstream task	Classification, Detec-	Any other tasks
	tion, Segmentation	

Table B.8: Inclusion and exclusion criteria.

Column Name	Description	Possible Values
Source(*)	Digital library which the study came from	ACM, IEEE, PubMed, ScienceDirect
Pubblication Type(*)	Type of publication	Conference paper, Journal article
Year(*)	Year of publication	-
$\operatorname{Author}(^*)$	List of authors	-
$\mathrm{Title}(*)$	Title of the study	-
Abstract(*)	Abstract of the study	-
$\operatorname{Language}(^*)$	Language of the study	English
Exclusion Reason (1)	Reason of exclusion in selection process	No D.A., Out of scope, No Medical domain
Deep Learning architecture	Deep Learning architecture used in the study	-
Task	Task object of experiment	Classification, Detection, Segmentation
Data-set size	Number of images in original dataset	-
Modality	Imaging modalities	CT, MR, XR, US, PET
Organ	Organs involved in the study	-
Pathology	Pathologies involved in the study	-
D.A. Objective	Purpose of Data Augmentation implementation	Balance Dataset, Random
Train/Test	D.A. applied to train or test phase	Train, Test
D.A. Type	Data augmentation techniques used in the study	-
D.A. Details	Brief description of the D.A. used	-
Paper Type	Type of paper according to our division in the next chapter	1, 2, 3, 4
Results	All the results obtained in relative task.	-
Exclusion Reason (2)	Reason of exclusion in extraction process	No D.A., Out of scope, No Medical domain, Missing tex

Table B.9: Description and possible values of data extracted from the papers

Appendix C. Supplementary material: results

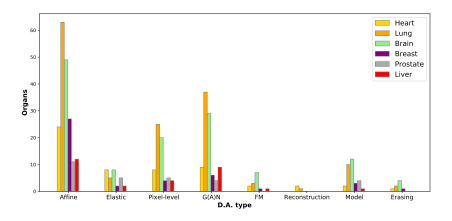


Figure C.9: Number of different data augmentation techniques for each organ

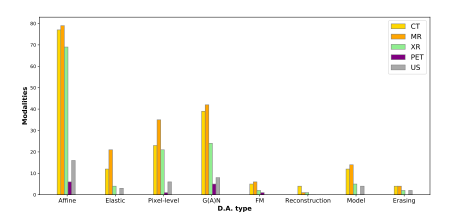


Figure C.10: Number of different modality for each data augmentation technique

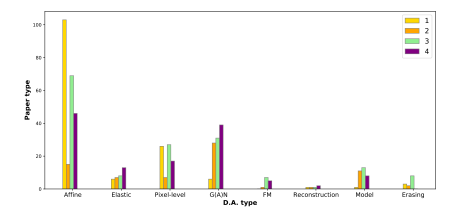


Figure C.11: Number of different data augmentation techniques for each type of paper

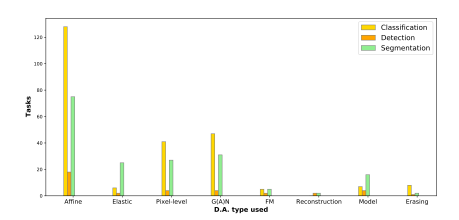


Figure C.12: Number of different data augmentation techniques for each task