



Shifting machine learning for healthcare from development to deployment and from models to data

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In the past decade, the application of machine learning (ML) to healthcare has helped drive the automation of physician tasks as well as enhancements in clinical capabilities and access to care. This progress has emphasized that, from model development to model deployment, data play central roles. In this Review, we provide a data-centric view of the innovations and challenges that are defining ML for healthcare. We discuss deep generative models and federated learning as strategies to augment datasets for improved model performance, as well as the use of the more recent transformer models for handling larger datasets and enhancing the modelling of clinical text. We also discuss data-focused problems in the deployment of ML, emphasizing the need to efficiently deliver data to ML models for timely clinical predictions and to account for natural data shifts that can deteriorate model performance.

In the past decade, machine learning (ML) for healthcare has been marked by particularly rapid progress. Initial groundwork has been laid for many healthcare needs that promise to improve patient care, reduce healthcare workload, streamline healthcare processes and empower the individual¹. In particular, ML for healthcare has been successful in the translation of computer vision through the development of image-based triage² and second readers³. There has also been rapid progress in the harnessing of electronic health records^{4,5} (EHRs) to predict the risk and progression of many diseases^{6,7}. A number of software platforms for ML are beginning to make their way into the clinic⁸. In 2018, iDX-DR, which detects diabetic retinopathy, was the first ML system for healthcare that the United States Food and Drug Administration approved for clinical use⁸. Babylon⁹, a chatbot triage system, has partnered with the United Kingdom's National Healthcare system. Furthermore, Viz.ai^{10,11} has rolled out their triage technology to more than 100 hospitals in the United States.

As ML systems begin to be deployed in clinical settings, the defining challenge of ML in healthcare has shifted from model development to model deployment. In bridging the gap between the two, another trend has emerged: the importance of data. We posit that large, well-designed, well-labelled, diverse and multi-institutional datasets drive performance in real-world settings far more than model optimization^{12–14}, and that these datasets are critical for mitigating racial and socioeconomic biases¹⁵. We realize that such rich datasets are difficult to obtain, owing to clinical limitations of data availability, patient privacy and the heterogeneity of institutional data frameworks. Similarly, as ML healthcare systems are deployed, the greatest challenges in implementation arise from problems with the data: how to efficiently deliver data to the model to facilitate workflow integration and make timely clinical predictions? Furthermore, once implemented, how can model robustness be maintained in the face of the inevitability of natural changes in

physician and patient behaviours? In fact, the shift from model development to deployment is also marked by a shift in focus: from models to data.

In this Review, we build on previous surveys^{1,16,17} and take a data-centric approach to reviewing recent innovations in ML for healthcare. We first discuss deep generative models and federated learning as strategies for creating larger and enhanced datasets. We also examine the more recent transformer models for handling larger datasets. We end by highlighting the challenges of deployment, in particular, how to process and deliver usable raw data to models, and how data shifts can affect the performance of deployed models.

Deep generative models

Generative adversarial networks (GANs) are among the most exciting innovations in deep learning in the past decade. They offer the capability to create large amounts of synthetic yet realistic data. In healthcare, GANs have been used to augment datasets¹⁸, alleviate the problems of privacy-restricted¹⁹ and unbalanced datasets²⁰, and perform image-modality-to-image-modality translation²¹ and image reconstruction²² (Fig. 1). GANs aim to model and sample from the implicit density function of the input data²³. They consist of two networks that are trained in an adversarial process under which one network, the 'generator', generates synthetic data while the other network, the 'discriminator', discriminates between real and synthetic data. The generative model aims to implicitly learn the data distribution from a set of samples to further generate new samples drawn from the learned distribution, while the discriminator pushes the generator network to sample from a distribution that more closely mirrors the true data distribution.

Over the years, a multitude of GANs have been developed to overcome the limitations of the original GAN (Table 1), and to optimize its performance and extend its functionalities. The original GAN²³ suffered from unstable training and low image diversity and

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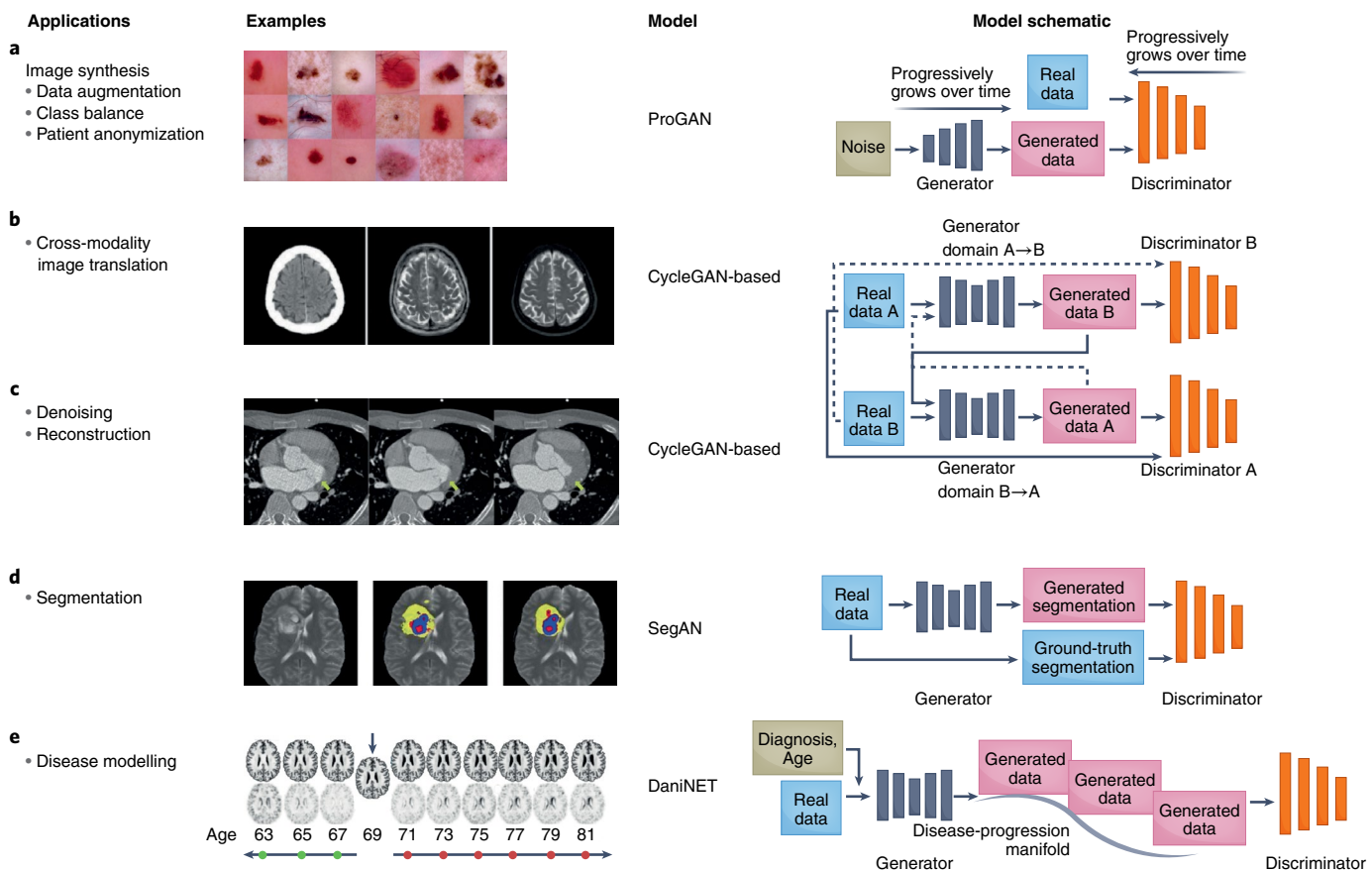


Fig. 1 | Roles of GANs in healthcare. **a**, GANs can be used to augment datasets to increase model performance and anonymize patient data. For example, they have been used to generate synthetic images of benign and malignant lesions from real images¹⁸³. **b**, GANs for translating images acquired with one imaging modality into another modality⁵¹. Left to right: input CT image, generated MR image and reference MR image. **c**, GANs for the denoising and reconstruction of medical images¹⁸⁴. Left, low-dose CT image of a patient with mitral valve prolapse, serving as the input into the GAN. Right, corresponding routine-dose CT image and the target of the GAN. Middle, GAN-generated denoised image resembling that obtained from routine-dose CT imaging. The yellow arrows indicate a region that is distinct between the input image (left) and the target denoised image (right). **d**, GANs for image classification, segmentation and detection³⁹. Left, input image of T2 MRI slice from the multimodal brain-tumour image-segmentation benchmark dataset. Middle, ground-truth segmentation of the brain tumour. Right, GAN-generated segmentation image. Yellow, segmented tumour; blue, tumour core; and red, Gd-enhanced tumour core. **e**, GANs can model a spectrum of clinical scenarios and predict disease progression⁶⁶. Top: given an input MR image (denoted by the arrow), DaniGAN can generate images that reflect neurodegeneration over time. Bottom, difference between the generated image and the input image. ProGAN, progressive growing of generative adversarial network; DaniNet, degenerative adversarial neuroimage net. Credit: Images ('Examples') reproduced with permission from: **a**, ref. ¹⁸³, Springer Nature Ltd; **b**, ref. ⁵¹, under a Creative Commons licence CC BY 4.0; **c**, ref. ¹⁸⁴, Wiley; **d**, ref. ³⁹, Springer Nature Ltd; **e**, ref. ⁶⁶, Springer Nature Ltd.

quality²⁴. In fact, training two adversarial models is, in practice, a delicate and often difficult task. The goal of training is to achieve a Nash equilibrium between the generator and the discriminator networks. However, simultaneously obtaining such an equilibrium for networks that are inherently adversarial is difficult and, if achieved, the equilibrium can be unstable (that is, it can be suddenly lost after model convergence). This has also led to sensitivity to hyperparameters (making the tuning of hyperparameters a precarious endeavour) and to mode collapse, which occurs when the generator produces a limited and repeated number of outputs. To remedy these limitations, changes have been made to GAN architectures and loss functions. In particular, the deep convolutional GAN (DCGAN²⁵), a popular GAN often used for medical-imaging tasks, aimed to combat instability by introducing key architecture-design decisions, including the replacement of fully connected layers with convolutional layers, and the introduction of batch normalization (to standardize the inputs to a layer when training deep neural networks) and ReLU (rectified linear unit) activation. The Laplacian

pyramid of adversarial networks GAN (LAPGAN²⁶) and the progressively growing GAN (ProGAN²⁷) build on DCGAN to improve training stability and image quality. Both LAPGAN and ProGAN start with a small image, which promotes training stability, and progressively grow the image into a higher-resolution image.

The conditional GAN (cGAN²⁸) and the auxiliary classifier GAN (AC-GAN²⁹) belong to a subtype of GANs that enable the model to be conditioned on external information to create synthetic data of a specific class or condition. This was found to improve the quality of the generated samples and increase the capability to handle the generation of multimodal data. The pix2pix GAN³⁰, which is conditioned on images, allows for image-to-image translation (also across imaging modalities) and has been popular in healthcare applications.

A recent major architectural change to GANs involve attention mechanisms. Attention was first introduced to facilitate language translation and has rapidly become a staple in deep-learning models, as it can efficiently capture longer-range global and spatial relations from input data. The incorporation of attention into GANs has

Table 1 | Popular GANs for medical imaging

Model	Description	Applications	Ref.
GAN	Original GAN; suffers from mode collapse; no guarantee of balance between the generator and the discriminator, which leads to the discriminator becoming too strong.	Multifarious	23
Changes to the loss function			
WGAN	Stabilizes training and prevents mode collapse by proposing the Wasserstein distance as loss function.	Unconditioned image synthesis Disease modelling ⁶⁵	34
WGAN-GP	Improves on WGAN to increase the stability of training and the quality of images.	Unconditioned image synthesis Disease modelling ⁶⁵	35
Conditional GANs			
cGAN	Original conditional GAN; auxiliary information is provided to the generator to produce synthetic data with a specific condition.	Image-to-image translation ⁵³ Lesion detection ^{53,59}	28
pix2pix	Conditional GAN in which the auxiliary information is an image.	Image reconstruction Image-to-image translation Data augmentation ¹⁹ Anonymization ¹⁹ Disease modelling ¹⁹	30
CycleGAN	Conditional GAN that can be used for image-to-image translation when paired training data are not available.	Image reconstruction Image-to-image translation ^{21,51,57,58,60} Segmentation ²¹ Data augmentation ⁴⁹ Anonymization ⁴⁹	60
Auxiliary GAN	Conditional GAN in which the discriminator is also asked to provide class probabilities.	Data augmentation ¹⁸	29
Changes to model architecture			
DCGAN	Replaced fully connected layers with convolutions.	Data augmentation ^{18,20,44} Class balance ²⁰	25
LAPGAN	Tackles image generation progressively instead of directly: proposed stack of GANs that add higher-frequency details to the generated image.	Unconditioned image synthesis	26
ProGAN	Tackles image generation progressively instead of directly: progressively grows the generator and discriminator with new layers, achieving higher-quality images.	Unconditioned image synthesis	27
Self-Attention GAN (SAGAN)	Introduces attention to obtain global and longer-range dependency modelling; uses conditioning; applies spectral normalization to improve training stability.	Conditioned image synthesis Image reconstruction ³²	31
BigGAN	Scales up SAGAN; applies orthogonal regularization to the generator to improve training stability.	Conditioned image synthesis	33

led to the development of self-attention GANs (SAGANs)^{31,32} and BigGAN;³³ the latter scales up SAGAN to achieve state-of-the-art performance.

Another primary strategy to mitigate the limitations of GANs involves improving the loss function. Early GANs used the Jensen-Shannon divergence and the Kullback-Leibler divergence as loss functions to minimize the difference in distribution between the synthetic generated dataset and the real-data dataset. However, the Jensen-Shannon divergence was found to fail in scenarios where there is no overlap (or little overlap) between distributions, while the minimization of the Kullback-Leibler divergence can lead to mode collapse. To address these problems, a number of GANs have used alternative loss functions. The most popular are arguably the Wasserstein GAN (WGAN)³⁴ and the Wasserstein GAN gradient penalty (WGAN-GP)³⁵. The Wasserstein distance measures the effort to minimize the distance between dataset distributions and has been shown to have a smoother gradient. Additional popular strategies that have been implemented to improve GAN performance and that do not involve modifying the model architecture include spectral normalization and varying how frequently the discriminator is updated (with respect to the update frequency of the generator).

The explosive progress of GANs has spawned many more offshoots of the original GAN, as documented by the diverse models that now populate the GAN Model Zoo³⁶.

Augmenting datasets. In the past decade, many deep-learning models for medical-image classification^{3,37}, segmentation^{38,39} and detection⁴⁰ have achieved physician-level performance. However, the success of these models is ultimately beholden to large, diverse, balanced and well-labelled datasets. This is a bottleneck that extends across domains, yet it is particularly restrictive in healthcare applications where collecting comprehensive datasets comes with unique obstacles. In particular, large amounts of standardized clinical data are difficult to obtain, and this is exacerbated by the reality that clinical data often reflects the patient population of one or few institutions (with the data sometimes overrepresenting common diseases or healthy populations and making the sampling of rarer conditions more difficult). Datasets with high class imbalance or insufficient variability can often lead to poor model performance, generalization failures, unintentional modelling of confounders⁴¹ and propagation of biases⁴². To mitigate these problems, clinical datasets can be augmented by using standard data-manipulation techniques, such as the flipping, rotation, scaling and translation of

images⁴³. However, these methods can lead to limited increases in performance and generate highly correlated training data.

GANs offer potent solutions to these problems. GANs can be used to augment training data to improve model performance. For example, a convolutional neural network (CNN) for the classification of liver lesions, trained on both synthetically and traditionally augmented data, boosted the performance of the model by 10% with respect to a CNN trained on only traditionally augmented datasets¹⁸. Moreover, when generating synthetic data across data classes, developing a generator for each class can result in higher model performance^{20,44}, as was shown via the comparison of two variants of GANs (a DCGAN that generated labelled examples for each of three lesion classes separately and an AC-GAN that incorporated class conditioning to generate labelled examples)¹⁸.

The aforementioned studies involved class-balanced datasets but did not address medical data with either simulated or real class imbalances. In an assessment of the capability of GANs to alleviate the shortcomings of unbalanced chest-X-ray datasets²⁰, it was found that training a classifier on real unbalanced datasets that had been augmented with DCGANs outperformed models that were trained with the unbalanced and balanced versions of the original dataset. Although there was an increase in classification accuracy across all classes, the greatest increase in performance was seen in the most imbalanced classes (pneumothorax and oedema), which had just one-fourth the number of training cases as the next class.

Protecting patient privacy. The protection of patient privacy is often a leading concern when developing clinical datasets⁴⁵. Sharing patient data when generating multi-institution clinical datasets can pose a risk to patient privacy⁴⁶. Even if privacy protocols are followed, patient characteristics can sometimes be inferred from the ML model and its outputs^{47,48}. In this regard, GANs may provide a solution. Data created by GANs cannot be attributed to a single patient, as they synthesize data that reflect the patient population in aggregate. GANs have thus been used as a patient-anonymization tool to generate synthetic data for model training^{9,49}. Although models trained on just synthetic data can perform poorly, models trained on synthetic data and fine-tuned with 10% real data resulted in similar performance to models trained on real datasets augmented with synthetic data¹⁹. Similarly, using synthetic data generated from GANs to train an image-segmentation model was sufficient to achieve 95% of the accuracy of the same model trained on real data⁴⁹. Hence, using synthetic data during model development can mitigate potential patient-privacy violations.

Image-to-image translation. One exciting use of GANs involves image-to-image translation. In healthcare, this capability has been used to translate between imaging modalities—between computed tomography (CT) and magnetic resonance (MR) images^{21,49–51}, between CT and positron emission tomography (PET)^{52–54}, between MR and PET^{55–57}, and between T1 and T2 MR images^{58,59}. Transfer between image modalities can reduce the need for additional costly and time-intensive image acquisitions, can be used in scenarios where imaging is not possible (as is the case for MR imaging in individuals with metal implants) and to expand the types of training data that can be created from image datasets. There are two predominant strategies for image-to-image translation: paired-image training (with pix2pix³⁰) and unpaired training (with CycleGAN⁶⁰). For example, pix2pix was used to generate synthetic CT images for accurate MR-based dose calculations for the pelvis⁶¹. Similarly, using paired magnetic resonance angiography and MR images, pix2pix was modified to generate a model for the translation of T1 and T2 MR images to retrospectively inspect vascular structures⁶².

Obtaining paired images can be difficult in scenarios involving moving organs or multimodal medical images that are in three dimensions and do not have cross-modality paired data. In such

cases, one can use CycleGAN⁶⁰, which handles image-to-image translation on unpaired images. A difficulty with unpaired images is the lack of ground-truth labels for evaluating the accuracy of the predictions (yet real cardiac MR images have been used to compare the performance of segmentation models trained on synthetic cardiac MR images translated from CT images⁴⁹). Another common problem is the need to avoid geometric distortions that destroy anatomical structures. Limitations with geometric distortions can be overcome by using two auxiliary mappings to constrain the geometric invariance of synthetic data²¹.

Opportunities. In the context of clinical datasets, GANs have primarily been used to augment or balance the datasets, and to preserve patient privacy. Yet a burgeoning application of GANs is their use to systematically explore the entire terrain of clinical scenarios and disease presentations. Indeed, GANs can be used to generate synthetic data to combat model deterioration in the face of domain shifts^{63,64}, for example, by creating synthetic data that simulate variable lighting or camera distortions, or that imitate data collected from devices from different vendors or from different imaging modalities. Additionally, GANs can be used to create data that simulate the full spectrum of clinical scenarios and disease presentations, from dangerous and rare clinical scenarios such as incorrect surgery techniques⁶³, to modelling the spectrum of brain-tumour presentation¹⁹, to exploring the disease progression of neurodegenerative diseases^{65,66}.

However, GANs can suffer from training instability and low image diversity and quality. These limitations could hamper the deployment of GANs in clinical practice. For example, one hope for image-to-image translation in healthcare involves the creation of multimodality clinical images (from CT and MR, for example) for scenarios in which only one imaging modality is possible. However, GANs are currently limited in the size and quality of the images that they can produce. This raises the question of whether these images can realistically be used clinically when medical images are typically generated at high resolution. Moreover, there may be regulatory hurdles involved in approving ML healthcare models that have been trained on synthetic data. This is further complicated by the current inability to robustly evaluate and control the quality of GANs and of the synthetic data that they generate⁶⁷. Still, in domains unrelated to healthcare, GANs have been used to make tangible improvements to deployed models⁶⁸. These successes may lay a foundation for the real-world application of GANs in healthcare.

Federated learning

When using multi-institutional datasets, model training is typically performed centrally: data siloed in individual institutions are aggregated into a single server. However, data used in such ‘centralized training’ represent a fraction of the vast amount of clinical data that could be harnessed for model development. Yet, openly sharing and exchanging patient data is restricted by many legal, ethical and administrative constraints; in fact, in many jurisdictions, patient data must remain local.

Federated learning is a paradigm for training ML models when decentralized data are used collaboratively under the orchestration of a central server^{69,70} (Fig. 2). In contrast to centralized training, where data from various locations are moved to a single server to train the model, federated learning allows for the data to remain in place. At the start of each round of training, the current copy of the model is sent to each location where the training data are stored. Each copy of the model is then trained and updated using the data at each location. The updated models are then sent from each location back to the central server, where they are aggregated into a global model. The subsequent round of training follows, the newly updated global model is distributed again, and the process is repeated until model convergence or training is stopped. At no point do the data

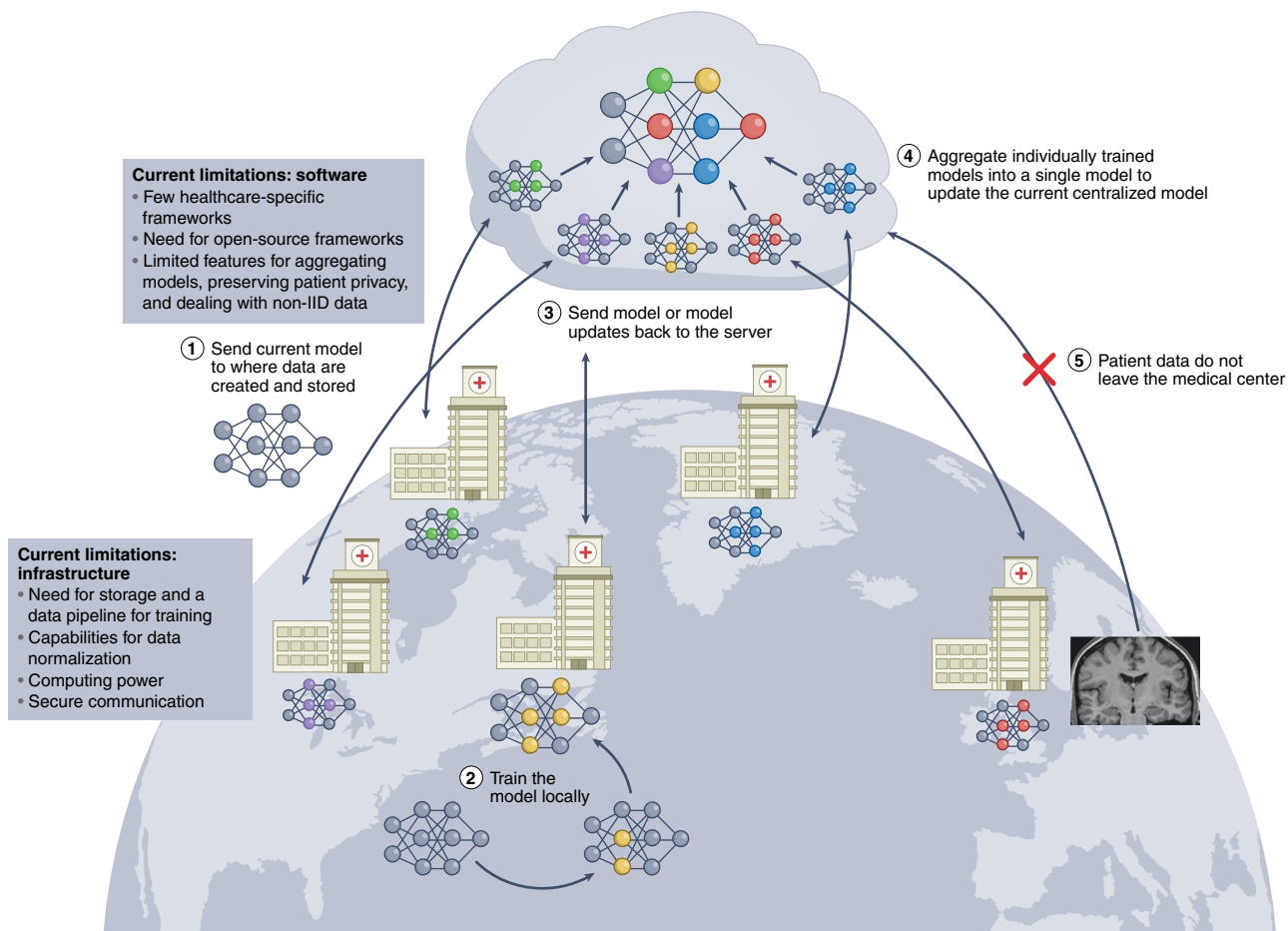


Fig. 2 | Cross-silo federated learning for healthcare. Multiple institutions collaboratively train an ML model. Federated learning begins when each institution notifies a central server of their intention to participate in the current round of training. Upon notification, approval and recognition of the institution, the central server sends the current version of the model to the institution (step 1). Then, the institution trains the model locally using the data available to it (step 2). Upon completion of local training, the institution sends the model back to the central server (step 3). The central server aggregates all of the models that have been trained locally by each of the individual institutions into a single updated model (step 4). This process is repeated in each round of training until model training concludes. At no point during any of the training rounds do patient data leave the institution (step 5). The successful implementation of federated learning requires healthcare-specific federated learning frameworks that facilitate training, as well as institutional infrastructure for communication with the central server and for locally training the model.

leave a particular location or institution, and only individuals associated with an institution have direct access to its data. This mitigates concerns about privacy breaches, minimizes costs associated with data aggregation, and allows training datasets to quickly scale in size and diversity. The successful implementation of federated learning could transform how deep-learning models for healthcare are trained. Here we focus on two applications: cross-silo federated learning and cross-device federated learning (Table 2).

Cross-silo federated learning. Cross-silo federated learning is an increasingly attractive solution to the shortcomings of centralized training⁷¹. It has been used to leverage EHRs to train models to predict hospitalization due to heart disease⁷², to promote the development of ‘digital twins’ or ‘Google for patients’⁷³, and to develop a Coronavirus disease 2019 (COVID-19) chest-CT lesion segmenter⁷⁴. Recent efforts have focused on empirically evaluating model-design parameters, and on logistical decisions to optimize model performance and overcome the unique implementation challenges of federated learning, such as bottlenecks in protecting privacy and in tackling the statistical heterogeneity of the data^{75,76}.

Compared with centralized training, one concern of federated learning is that models may encounter more severe domain shifts

or overfitting. However, models trained through federated learning were found to achieve 99% of the performance of traditional centralized training even with imbalanced datasets or with relatively few samples per institution, thus showing that federated learning can be realistically implemented without sacrificing performance or generalization^{77,78}.

Although federated learning offers greater privacy protection because patient data are no longer being transmitted, there are risks of privacy breaches⁷⁹. Communicating model updates during the training process can reveal sensitive information to third parties or to the central server. In certain instances, data leakage can occur, such as when ML models ‘memorize’ datasets^{80–82} and when access to model parameters and updates can be used to infer the original dataset⁸³. Differential privacy⁸⁴ can further reinforce privacy protection for federated learning^{70,85,86}. Selective parameter sharing⁸⁷ and the sparse vector technique⁸⁸ are two strategies for achieving greater privacy, but at the expense of model performance (this is consistent with differential-privacy findings in domains outside of medicine and healthcare^{80,89}).

Another active area of research for federated learning in healthcare involves the handling of data that are neither independent nor identically distributed (non-IID data). Healthcare data are

Table 2 | Federated learning

Characteristic	Centralized learning	Cross-silo federated learning	Cross-device federated learning
Setting			
Model location	Trained on a central server.	Model is sent to and trained at each institution.	Model is sent to and trained at each device.
Participants	1–1,000	2–100	$<10^{10}$
Participation frequency	Every participant participates in every round of training.		Not every participant participates in each round, owing to varied availability.
Data			
Location	Moved to central server.	Local and decentralized: data remains where they are generated.	
Privacy	Low; anyone with access can access patient data from any other institution.	Medium to high; participants cannot access data from other participants.	
Availability	Always available	Always available	Not always available; a fraction of devices are available, typically at night, when devices are idle.
Limitations			
Device dropout	Rare	Rare	$>5\%$ of devices expected to drop out owing to communication issues, battery depletion or the requirement of idleness.
Primary bottleneck	Computation; difficult to scale to large datasets.	Computation or communication	Communication, owing to issues of reliability or availability.

particularly susceptible to this problem, owing to a higher prevalence of certain diseases in certain institutions (which can cause label-distribution skew) or to institution-specific data-collection techniques (leading to ‘same label, different features’ or to ‘same features, different label’). Many federated learning strategies assume IID data, but non-IID data can pose a very real problem in federated learning; for example, it can cause the popular federated learning algorithm FedAvg⁷⁰ to fail to converge⁹⁰. The predominant strategies for addressing this issue have involved the reframing of the data to achieve a uniform distribution (consensus solutions) or the embracing of the heterogeneity of the data^{69,91,92} (pluralistic solutions). In healthcare, the focus has been on consensus solutions involving data sharing (a small subset of training data is shared among all institutions^{93,94}).

Cross-device federated learning to handle health data from individuals. ‘Smart’ devices can produce troves of continuous, passive and individualized health data that can be leveraged to train ML models and deliver personalized health insights for each user^{1,16,39,95,96}. As smart devices become increasingly widespread, and as computing and sensor technology become more advanced and cheaper to mass-produce, the amount of health data will grow exponentially. This will accentuate the challenges of aggregating large quantities of data into a single location for centralized training and exacerbate privacy concerns (such as any access to detailed individual health data by large corporations or governments).

Cross-device federated learning was developed to address the increasing amounts of data that are being generated ‘at the edge’ (that is, by decentralized smart devices), and has been deployed on millions of smart devices; for example, for voice recognition (by Apple, for the voice assistant Siri⁹⁷) and to improve query suggestions (by Google, for the Android operating system⁹⁸).

The application of cross-device federated learning to train healthcare models for smart devices is an emerging area of research. For example, using a human-activity-recognition dataset, a global model (FedHealth) was pre-trained using 80% of the data before deploying it to be locally trained and then aggregated⁹⁹. The aggregated model was then sent back to each user and fine-tuned on user-specific data to develop a personalized model for the user. Model personalization resolves issues arising from the highly

different probability distributions that may arise across users and the global model. This training strategy outperformed non-federated learning by nearly 5.3%.

Limitations and opportunities. In view of the initial promises and successes of federated learning, the next few years will be defined by progress towards the implementation of federated learning in healthcare. This will require a high degree of coordination across institutions at each step of the federated learning process. Before training, medical data will need to undergo data normalization and standardization. This can be challenging, owing to differences in how data are collected, stored, labelled and partitioned across institutions. Current data pre-processing pipelines could be adapted to create multi-institutional training datasets, yet in federated learning, the responsibility shifts from a central entity to each institution individually. Hence, methods to streamline and validate these processes across institutions will be essential for the successful implementation of federated learning.

Another problem concerns the inability of the developer of the model to directly inspect data during model development. Data inspection is critical for troubleshooting and for identifying any mislabelled data as well as general trends. Tools (such as Federated Analytics, developed by Google¹⁰⁰) that use GANs to create synthetic data that resemble the original training data¹⁰¹ and derive population-level summary statistics from the data, can be helpful. However, it is currently unclear whether tools that have been developed for cross-device settings can be applied to cross-silo healthcare settings while preserving institutional privacy.

Furthermore, federated learning will require robust frameworks for the implementation of federated networks. Many such software is proprietary, and many of the open-source frameworks are primarily intended for use in research. The primary concerns of federated learning can be addressed by frameworks designed to reinforce patient privacy, facilitate model aggregation and tackle the challenges of non-IID data.

One main hurdle is the need for each participating healthcare institution to acquire the necessary infrastructure. This implies ensuring that each institution has the same federated learning framework and version, that stable and encrypted network communication is available to send and receive model updates from

the central server, and that the computing capabilities (institutional graphics processing units or access to cloud computing) are sufficient to train the model. Although most large healthcare institutions may have the necessary infrastructure in place, it has typically been optimized to store and handle data centrally. The adaptation of infrastructure to handle the requirements of federated learning requires coordinated effort and time.

A number of ongoing federated learning initiatives in healthcare are underway. Specifically, the Federated Tumour Segmentation Initiative (a collaboration between Intel and the University of Pennsylvania) trains lesion-segmentation models collaboratively across 29 international healthcare institutions¹⁰². This initiative focuses on finding the optimal algorithm for model aggregation, as well as on ways to standardize training data from various institutions. In another initiative (a collaboration of NVIDIA and several institutions), federated learning was used to train mammography-classification models¹⁰³. These efforts may establish blueprints for coordinated federated networks applied to healthcare.

Natural language processing

Harnessing natural language processing (NLP)—the automated understanding of text—has been a long-standing goal for ML in healthcare^{1,16,17}. NLP has enabled the automated translation of doctor–patient interactions to notes^{5,104,105}, the summarization of clinical notes¹⁰⁶, the captioning of medical images^{107,108} and the prediction of disease progression^{6,7}. However, the inability to efficiently train models using the large datasets needed to achieve adept natural-language understanding has limited progress. In this section, we provide an overview of two recent innovations that have transformed NLP: transformers and transfer learning for NLP. We also discuss their applications in healthcare.

Transformers. When modelling sequential data, recurrent neural networks (RNNs) have been the predominant choice of neural network. In particular, long short-term memory networks¹⁰⁹ and gated units¹¹⁰ were staple RNNs in modelling EHR data, as these networks can model the sequential nature of clinical data^{111,112} and clinical text^{5,104,105,113}. However, RNNs harbour several limitations¹¹⁴. Namely, RNNs process data sequentially and not in parallel. This restricts the size of the input datasets and of the networks, which limits the complexity of the features and the range of relations that can be learned¹¹⁴. Hence, RNNs are difficult to train, deploy and scale, and are suboptimal for capturing long-range patterns and global patterns in data. However, learning global or long-range relationships are often needed when learning language representations. For example, sentences far removed from a word may be important for providing context for the word, and previous clinical events that have occurred can inform clinical decisions that are made years later. For a period, CNNs, which are adept at parallelization, were used to overcome some of the limitations of RNNs¹¹⁵, but were found to be inefficient when modelling longer global dependencies.

In 2017, a research team at Google (the Google Brain team) released the transformer, a landmark model that has revolutionized NLP¹¹⁶. Compared with RNN and CNN models, transformers are more parallelizable and less computationally complex at each layer, and thus can handle larger training data and learn longer-range and global relations. The use of only attention layers for the encoders and decoders while forgoing the use of RNNs or CNNs was critical to the success of transformers. Attention was introduced and refined^{117,118} to handle bottlenecks in sequence-to-sequence RNNs^{110,119}. Attention modules allow models to globally relate different positions of a sequence to compute a richer representation of the sequence¹¹⁶, and does so in parallel, allowing for increased computing efficiency and for the embedding of longer relations of the input sequence (Fig. 3).

Transfer learning for NLP. Simultaneous and subsequent work following the release of the transformer resolved another main problem in NLP: the formalization of the process of transfer learning. Transfer learning has been used most extensively in computer vision, owing to the success of the ImageNet challenge, which made pre-trained CNNs widely available¹²⁰. Transfer learning has enabled the broader application of deep learning in healthcare¹⁷, as researchers can fine-tune a pre-trained CNN adept at image classification on a smaller clinical dataset to accomplish a wide spectrum of healthcare tasks^{3,37,121,122}. Until recently, robust transfer learning for NLP models was not possible, which limited the use of NLP models in domain-specific applications. A series of recent milestones have enabled transfer learning for NLP. The identification of the ideal pre-training language task for deep-learning NLP models (for example, masked-language modelling, predicting missing words from surrounding context, next-sentence prediction or predicting whether two sentences follow one another) was solved by universal language model fine-tuning (ULM-FiT¹²³) and embeddings from language model (ELMo¹²⁴). The generative pre-trained transformer (GPT¹²⁵) from Open AI and the bidirectional encoder representations from transformers (BERT¹²⁶) from Google Brain then applied the methods formalized by ULM-FiT and ELMo to transformer models, delivering pre-trained models that achieved unprecedented capabilities on a series of NLP tasks.

Transformers for the understanding of clinical text. Following the success of transformers for NLP, their potential to handle domain-specific text, specifically clinical text, was quickly assessed. The performances of the transformer-based model BERT, the RNN-based model ELMo and traditional word-vector embeddings^{127,128} at clinical-concept extraction (the identification of the medical problems, tests and treatments) from EHR data were evaluated¹⁰⁶. BERT outperformed traditional word vectors by a substantial margin and was more computationally efficient than ELMo (it achieved higher performance with fewer training iterations)^{129–132}. Pre-training on a dataset of 2 million clinical notes (the dataset multiparameter intelligence monitoring in intensive care¹³²; MIMIC-III) increased the performance of all NLP models. This suggests that contextual embeddings encode valuable semantic information not accounted for in traditional word representations¹⁰⁶. However, the performance of MIMIC-III BERT began to decline after achieving its optimal model; this is perhaps indicative of the model losing information learned from the large open corpus and converging to a model similar to the one initialized from scratch¹⁰⁶. Hence, there may be a fine balance between learning from a large open-domain corpus and a domain-specific clinical corpus. This may be a critical consideration when applying pre-trained models to healthcare tasks.

To facilitate the further application of clinically pre-trained BERT¹²⁹ to downstream clinical tasks, a BERT pre-trained on large clinical datasets was publicly released. Because transformers and deep NLP models are resource-intensive to train (training the BERT model can cost US\$50,000–200,000¹³³, and pre-training BERT on clinical datasets required 18 d of continuous training, an endeavour that may be out of the reach of many institutions), openly releasing pre-trained clinical models can facilitate widespread advancements of NLP tasks in healthcare. Other large and publicly available clinically pre-trained models (Table 3) are ClinicalBERT¹³⁰, BioBERT¹³⁴ and SciBERT¹³⁵.

The release of clinically pre-trained models has spurred downstream clinical applications. ClinicalBERT, a BERT model pre-trained on MIMIC-III data using masked-language modelling and next-sentence prediction, was evaluated on the downstream task of predicting 30 d readmission¹³⁰. Compared with previous models^{136,137}, ClinicalBERT can dynamically predict readmission risk during a patient's stay and uses clinical text rather than

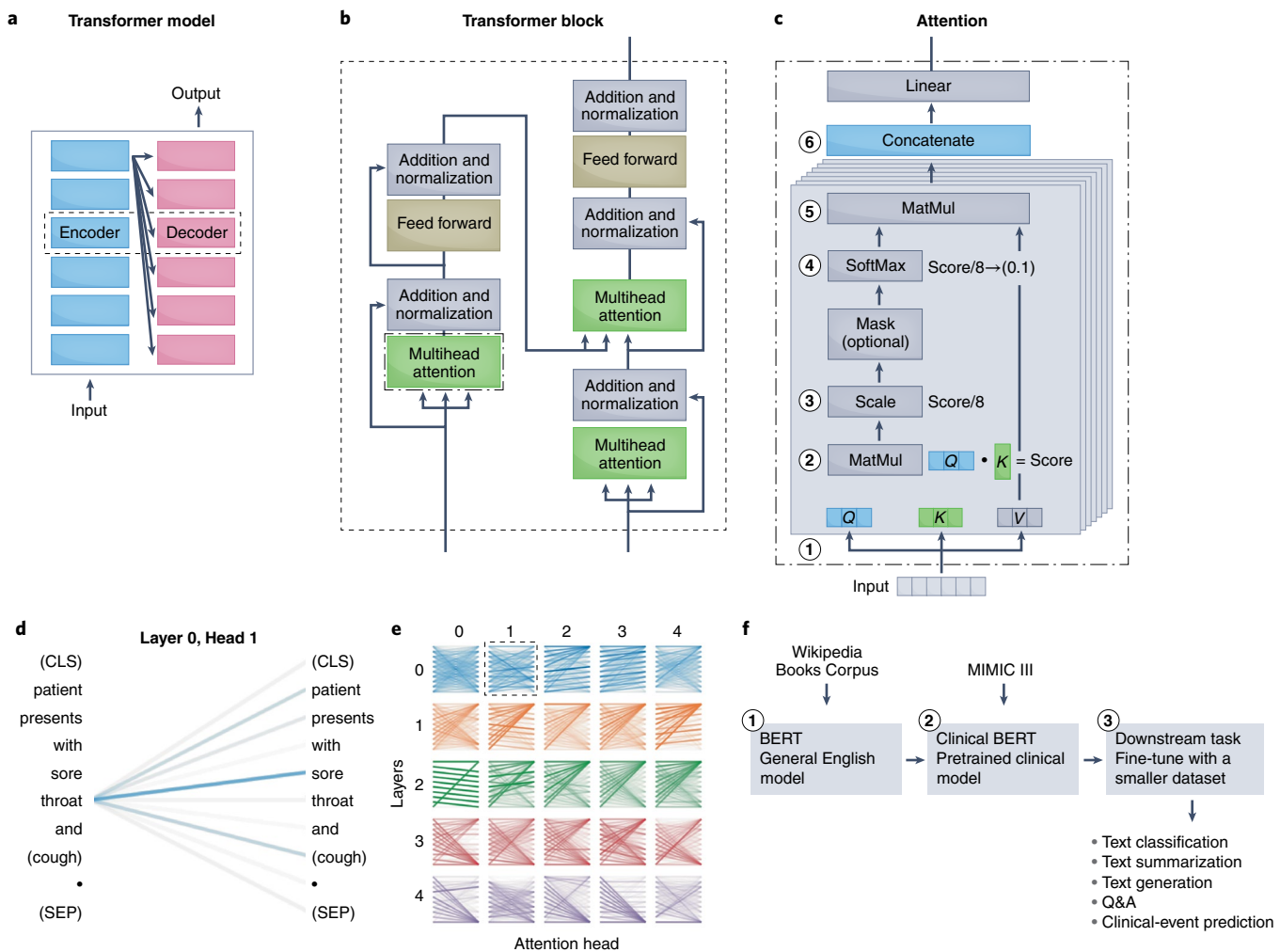


Fig. 3 | Transformers. **a**, The original transformer model performs language translation, and contains encoders that convert the input into an embedding and decoders that convert the embedding into the output. **b**, The transformer model uses attention mechanisms within its encoders and decoders. The attention module is used in three places: in the encoder (for the input sentence), in the decoder (for the output sentence) and in the encoder-decoder in the decoder (for embeddings passed from the encoder). **c**, The key component of the transformer block is the attention module. Briefly, attention is a mechanism to determine how much weight to place on input features when creating embeddings for downstream tasks. For NLP, this involves determining how much importance to place on surrounding text when creating a representation for a particular word. To learn the weights, the attention mechanism assigns a score to each pair of words from an input phrase to determine how strongly the words should influence the representation. To obtain the score, the transformer model first decomposes the input into three vectors: the query vector (Q ; the word of interest), the key vector (K ; surrounding words) and the value vector (V ; the contents of the input) (1). Next, the dot product is taken between the query and key vector (2) and then scaled to stabilize training (3). The SoftMax function is then applied to normalize the scores and ensure that they add to 1 (4). The output SoftMax score is then multiplied by the value vector to apply a weighted focus to the input (5). The transformer model has multiple attention mechanisms (termed attention heads); each learn a separate representation for the same word, which therefore increases the relations that can be learned. Each attention head is composed of stacked attention layers. The output of each attention mechanism is concatenated into a single matrix (6) that is fed into the downstream feed-forward layer. **d,e**, Visual representation of what is learned¹⁸⁵. Lines relate the query (left) to the words that are attended to the most (right). Line thickness denotes the magnitude of attention, and colours represent the attention head. **d**, The learned attention in one attention-mechanism layer of one head. **e**, Examples of what is learned by each layer of each attention head. Certain layers learn to attend to the next words (head 2, layer 0) or to the previous word (head 0, layer 0). **f**, Workflow for applying a transformer language model to a clinical task. Matmul, matrix multiplication; (CLS), classification token placed at the start of a sentence to store the sentence-level embedding; (SEP), separation token placed at the end of a sentence. BERT, bidirectional encoder representations from transformers; MIMIC, multiparameter intelligence monitoring in intensive care.

structured data (such as laboratory values, or codes from the international classification of diseases). This shows the power of transformers to unlock clinical text, a comparatively underused data source in EHRs. Similarly, clinical text from EHRs has been harnessed using SciBERT for the automated extraction of symptoms from COVID-19-positive and COVID-19-negative patients to identify the most discerning clinical presentation¹³⁸. ClinicalBERT has also been adapted to extract anginal symptoms from EHRs¹³⁹.

Others have used enhanced clinical-text understanding for the automatic labelling and summarization of clinical reports. BioBERT and ClinicalBERT have been harnessed to extract labels from radiology text reports, enabling an automatic clinical summarization tool and labeller¹⁴⁰. Transformers have also been used to improve clinical questioning and answering¹⁴¹, in clinical voice assistants^{142,143}, in chatbots for patient triage^{144,145}, and in medical-image-to-text translation and medical-image captioning¹⁴⁶.

Table 3 | Publicly available clinical BERT models

Model	Dataset	Evaluation task	Ref.
BERT base	BooksCorpus (800 million words)	GLUE	126
BERT large	English Wikipedia (2.5 billion words)	SQuAD v1.1 SQuAD v2.0 SWAG	
BioBERT	PubMed abstracts PubMed Central full articles	Named-entity recognition Relation classification Q&A (BioASQ)	134
SciBERT	1.14 million papers from Semantic Scholar (18% in computer science; 82% biomedical)	Named-entity recognition PICO extraction Text classification Relation classification Dependency parsing	135
Clinical BERT	MIMIC-III v1.4	MedNLI	129
Discharge summary BERT	MIMIC-III v1.4 discharge summaries only	Named-entity recognition (i2b2 2006, 2010, 2012, 2014)	
Bio+Clinical BERT	MIMIC-III v1.4		
Bio+Discharge summary BERT	MIMIC-III v1.4 discharge summaries only		
Clinical BERT	MIMIC-III	30 d hospital-readmission prediction	130
Med-BERT	Cerner HealthFacts	Disease prediction	150

GLUE, general language understanding evaluation; SQuAD, Stanford question-answering dataset; SWAG, situations with adversarial generations; BioASQ, a challenge on large-scale biomedical semantic indexing and question answering; PICO, medical questioning framework consisting of problem (patient problem), intervention, comparison with other interventions and outcomes; MedNLI, medical natural-language inference.

Transformers for the modelling of clinical events. In view of their adeptness to model the sequential nature of clinical text, transformers have also been harnessed to model the sequential nature of clinical events^{147–151}. A key challenge of modelling clinical events is properly capturing long-term dependencies—that is, previous clinical procedures that may preclude future downstream interventions. Transformers are particularly adept at exploring longer-range relationships and were recently used to develop BEHRT¹⁵², which leverages the parallels between sequences in natural language and clinical events in EHRs to portray diagnoses as words, visits as sentences and a patient’s medical history as a document¹⁵². When used to predict the likelihood of 301 conditions in future visits, BEHRT achieved an 8–13.2% improvement over the existing state-of-the-art EHR model¹⁵². BEHRT was also used to predict the incidence of heart failure from EHR data¹⁵³.

Data-limiting factors in the deployment of ML

The past decade of research in ML in healthcare has focused on model development, and the next decade will be defined by model deployment into clinical settings^{42,45,46,154,155}. In this section, we discuss two data-centric obstacles in model deployment: how to efficiently deliver raw clinical data (Table 4) to models, and how to monitor and correct for natural data shifts that deteriorate model performance.

Delivering data to models. A main obstacle to model deployment is associated with how to efficiently transform raw, unstructured and heterogeneous clinical data into structured data that can be inputted

into ML models. During model development, pre-processed structured data are directly inputted into the model. However, during deployment, minimizing the delay between the acquisition of raw data and the delivery of structured inputs requires an adept data pipeline for collecting data from their source, and for ingesting, preparing and transforming the data (Fig. 4). An ideal system would need to be high-throughput, have low latency and be scalable to a large number of data sources. A lack of optimization can result in major sources of inefficiency and delayed predictions from the model. In what follows, we detail the challenges of building a pipeline for clinical data and give an overview of the key components of such a pipeline.

The fundamental challenge of creating an adept data pipeline arises from the need to anticipate the heterogeneity of the data. ML models often require a set of specific clinical inputs (for example, blood pressure and heart rate), which are extracted from a suite of dynamically changing health data. However, it is difficult to extract the relevant data inputs. Clinical data vary in volume and velocity (the rate that data are generated), thus prompting the question of how frequently data should be collected. Furthermore, clinical data can vary in veracity (data quality), thus requiring different pre-processing steps. Moreover, the majority of clinical data exist in an unstructured format that is further complicated by the availability of hundreds of EHR products, each with its own clinical terminology, technical specifications and capabilities¹⁵⁶. Therefore, how to precisely extract data from a spectrum of unstructured EHR frameworks becomes critical.

Data heterogeneity must be carefully accounted for when designing the data pipeline, as it can influence throughput, latency and other performance factors. The data pipeline starts with the process of data ingestion (by which raw clinical data are moved from the data source and into the pipeline), a primary bottleneck in the throughput of the data through the pipeline. In particular, handling peaks of data generation may require the design and implementation of scalable ways to support a variable number of connected objects¹⁵⁷. Such data-elasticity issues can take advantage of software frameworks that scale up or down in real time to more effectively use computer resources in cloud data centres¹⁵⁸.

After the data enters the pipeline, the data-preparation stage involves the cleansing, denoising, standardization and shaping of the data into structured data that are ready for consumption by the ML system. In studies that developed data pipelines to handle healthcare data^{156,159,160}, the data-preparation stage was found to regulate the latency of the data pipeline, as latency depended on the efficiency of the data queue, the streaming of the data and the database for storing the computation results.

A final consideration is how data should move throughout the data pipeline; specifically, whether data should move in discrete batches or in continuous streams. Batch processing involves collecting and moving source data periodically, whereas stream processing involves sourcing, moving and processing data as soon as they are created. Batch processing has the advantages of being high-throughput, comprehensive and economical (and hence may be advantageous for scalability), whereas stream processing occurs in real time (and thus may be required for time-sensitive predictions). Many healthcare systems use a combination of batch processing and stream processing¹⁶⁰.

Established data pipelines are being harnessed to support real-time healthcare modelling. In particular, Columbia University Medical Center, in collaboration with IBM, is streaming physiological data from patients with brain injuries to predict adverse neurological complications up to 48 h before existing methods can¹⁶¹. Similarly, Yale School of Medicine has used a data pipeline to support real-time data acquisition for predicting the number of beds available, handling care for inpatients and patients in the intensive care unit (such as managing ventilator capacity) and tracking the

Table 4 | Commonly used clinical datasets

Dataset	Data types	Size of the dataset	Institutions	Applications
Multimodal brain-tumour image-segmentation benchmark dataset (BRATS)	Multiparametric MRI: T1, T1Gd, T2 and T2-FLAIR	~2,000 patients; ~8,000 scans	Multi-institution (13)	GANs: image-to-image translation Federated learning ⁷⁶
Alzheimer's disease neuroimaging initiative dataset (ADNI)	MRI, PET Genetics, cognitive tests and biomarkers	~2,000 patients	Multi-institution (63)	GANs: data augmentation, anonymization ¹⁹ , image-to-image translation ^{55,57} Federated learning ⁷⁵
Autism brain imaging data exchange	Functional MRI	~1,114 patients	Multi-institution (19)	Federated learning ⁷¹
NIH prostate, lung, colorectal and ovarian cancer dataset (NIH PLCO)	X-ray images (chest) Digital histopathology (prostate, lung, colorectal, ovarian, breast and bladder) Questionnaires and laboratory data	~155,000 patients	NCI	GANs
Medical segmentation decathlon	MRI images (brain, heart and prostate) CT images (lung, liver, spleen, pancreas, colon, hepatic vessels and prostate)	~2,633 images	Multi-institution	Federated learning
NIH DeepLesion	CT images	~4,400 patients; ~32,000 lesions	NIH	Federated learning
Cancer imaging archive	MRI, CT, PET and digital histopathology Multi-organ	~1,000–3,000 patients	Multi-institution	GANs Federated learning ⁷⁸
Medical information mart for intensive care (MIMIC)	Electronic medical records	~60,000 patients	Beth Israel Deaconess Medical Center	Clinical text and events modelling Federated learning ⁷³
IBM MarketScan research databases for life-science researchers	Electronic medical records and claims	~43.6 million	Multi-institution	Federated learning
EchoNet-Dynamic	Echocardiogram videos	~10,030 videos	Stanford Health Care	Video-based segmentation and classification; largest publicly available medical video dataset ³⁸

NCI, National Cancer Institute; NIH, National Institutes of Health; T1Gd, gadolinium-enhanced T1-weighted; T2-FLAIR, T2-weighted fluid-attenuated inversion recovery.

number of healthcare providers exposed to COVID-19¹⁶¹. However, optimizing the components of the data pipeline, particularly for numerous concurrent ML healthcare systems, remains a challenging task.

Deployment in the face of data shifts. A main obstacle in deploying ML systems for healthcare has been maintaining model robustness when faced with data shifts¹⁶². Data shifts occur when differences or changes in healthcare practices or in patient behaviour cause the deployment data to differ substantially from the training data, resulting in the distribution of the deployment data diverging from the distribution of the training data. This can lead to a decline in model performance. Also, failure to correct for data shifts can lead to the perpetuation of algorithmic biases, missing critical diagnoses¹⁶³ and unnecessary clinical interventions¹⁶⁴.

In healthcare, data shifts are common occurrences and exist primarily along the axes of institutional differences (such as local clinical practices, or different instruments and data-collection workflows), epidemiological shifts, temporal shifts (for example, changes in physician and patient behaviours over time) and differences in patient demographics (such as race, gender and age). A recent case study¹⁶⁵ characterizing data shifts caused by institutional differences reported that pneumothorax classifiers trained

on individual institutional datasets declined in performance when evaluated on data from external institutions. Similar phenomena have been observed in a number of studies^{41,163,166}. Institutional differences are among the most patent causes of data shifts because they frequently harbour underlying differences in patient demographics, disease incidence and data-collection workflows. For example, in an analysis of chest-X-ray classifiers and their potential to generalize to other institutions, it was found that one institution collected chest X-rays using portable radiographs, whereas another used stationary radiographs⁴¹. This led to differences in disease prevalence (33% vs 2% for pneumonia) and patient demographics (average age of 63 vs 45), as portable radiographs were primarily used for inpatients who were too sick to be transported, whereas stationary radiographs were used primarily in outpatient settings. Similarly, another study found that different image-acquisition and image-processing techniques caused the deterioration of the performance of breast-mammography classifiers to random performance (areas under the receiver operating characteristic curve of 0.4–0.6) when evaluated on datasets from four external institutions and countries¹⁶³. However, it is important to note that the models evaluated were trained on data collected during the 1990s and were externally tested on datasets created in 2014–2017. The decline in performance owing to temporal shifts is particularly relevant;

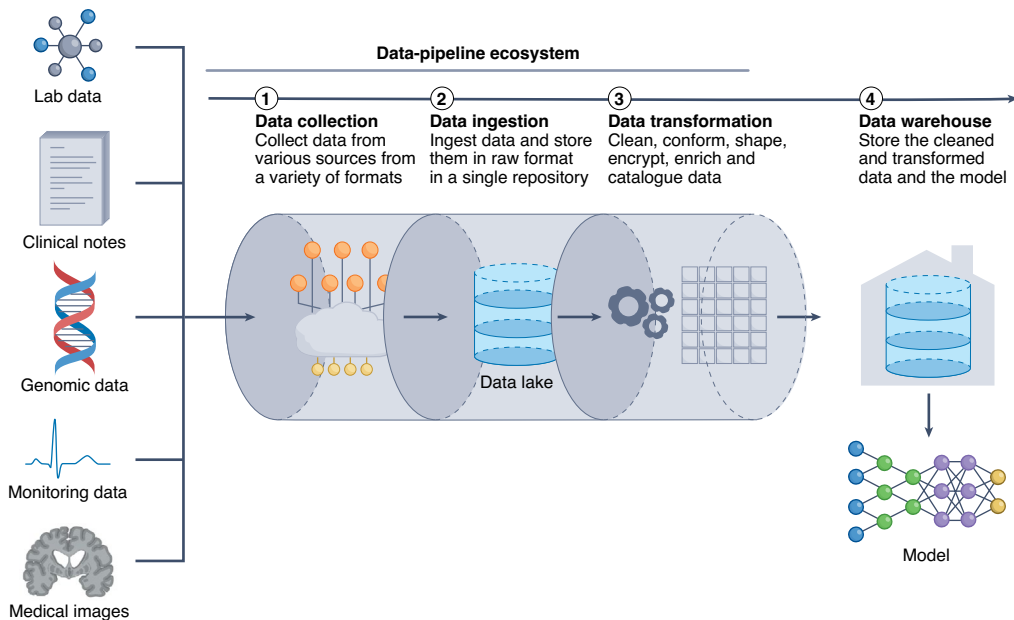


Fig. 4 | Data pipeline. Delivering data to a model is a key bottleneck in obtaining timely and efficient inferences. ML models require input data that are organized, standardized and normalized, often in tabular format. Therefore, it is critical to establish a pipeline for organizing and storing heterogeneous clinical data. The data pipeline involves collecting, ingesting and transforming clinical data from an assortment of data sources. Data can be housed in data lakes, in data warehouses or in both. Data lakes are central repositories to store all forms of data, raw and processed, without any predetermined organizational structure. Data in data lakes can exist as a mix of binary data (for example, images), structured data, semi-structured data (such as tabular data) and unstructured data (for example, documents). By contrast, data warehouses store cleaned, enriched, transformed and structured data with a predetermined organizational structure.

if deployed today, models that have been trained on older datasets would be making inferences on newly generated data.

Studies that have characterized temporal shifts have provided insights into the conditions under which deployed ML models should be re-evaluated. An evaluation of models that used data collected over a period of 9 years found that model performance deteriorated substantially, drifting towards overprediction as early as one year after model development¹⁶⁷. For the MIMIC-III dataset¹³² (commonly used for the development of models to predict clinical outcomes), an assessment of the effects of temporal shifts on model performance over time showed that, whereas all models experienced a moderate decline over time, the most significant drop in performance occurred owing to a shift in clinical practice, when EHRs transitioned systems¹⁶⁴ (from CareVue to MetaVision). A modern-day analogy would be how ML systems for COVID-19 (ref. ¹⁶⁸) that were trained on data¹⁶⁹ acquired during the early phase of the pandemic and before the availability of COVID-19 vaccines would perform when deployed in the face of shifts in disease incidence and presentation.

Data shifts and model deterioration can also occur when models are deployed on patients with gender, racial or socioeconomic backgrounds that are different from those of the patient population that the model was trained on. In fact, it has been shown that ML models can be biased against individuals of certain races¹⁷⁰ or genders⁴², or particular religious¹⁷¹ or socioeconomic¹⁵ backgrounds. For example, a large-scale algorithm used in many health institutions to identify patients for complex health needs underpredicted the health needs of African American patients and failed to triage them for necessary care¹⁷². Using non-representative or non-inclusive training datasets can constitute an additional source of gender, racial or socioeconomic biases. Popular chest-X-ray datasets used to train classifiers have been shown to be heavily unbalanced¹⁵: 67.6% of the patients in these datasets are Caucasian and only 8.98% are under Medicare insurance. Unsurprisingly, the performance of models

trained with these datasets deteriorates for non-Caucasian subgroups, and especially for Medicare patients¹⁵. Similarly, skin-lesion classifiers that were trained primarily on images of one skin tone decrease in performance when evaluated on images of different skin tones¹⁷³; in this case, the drop in performance could be attributed to variations in disease presentation that are not captured when certain patient populations are not adequately represented in the training dataset¹⁷⁴.

These findings exemplify two underlying limitations of ML models: the models can propagate existing healthcare biases on a large scale, and insufficient diversity in the training datasets can lead to an inadequate generalization of model outputs to different patient populations. Training models on multi-institutional datasets can be most effective at combating model deterioration¹⁵, and directly combating existing biases in the training data can also mitigate their impact¹⁷¹. There are also solutions for addressing data shifts that involve proactively addressing them during model development^{175–178} or retroactively by surveilling for data shifts during model deployment¹⁷⁹. A proactive attitude towards recognizing and addressing potential biases and data shifts will remain imperative.

Outlook

Substantial progress in the past decade has laid a foundation of knowledge for the application of ML to healthcare. In pursuing the deployment of ML models, it is clear that success is dictated by how data are collected, organized, protected, moved and audited. In this Review, we have highlighted methods that can address these challenges. The emphasis will eventually shift to how to build the tools, infrastructure and regulations needed to efficiently deploy innovations in ML in clinical settings. A central challenge will be the implementation and translation of these advances into healthcare in the face of their current limitations: for instance, GANs applied to medical images are currently limited by image resolution and image diversity, and can be challenging to train and scale; federated

learning promises to alleviate problems associated with small single-institution datasets, yet it requires robust frameworks and infrastructure; and large language models trained on large public datasets can subsume racial and ethnic biases¹⁷¹.

Another central consideration is how to handle the regulatory assessment of ML models for healthcare applications. Current regulation and approval processes are being adapted to meet the emerging needs; in particular, initiatives are attempting to address data shifts and patient representation in the training datasets^{165,180,181}. However, GANs, federated learning and transformer models add complexities to the regulatory process. Few healthcare-specific benchmarking datasets exist to evaluate the performance of these ML systems during clinical deployment. Moreover, the assessment of the performance of GANs is hampered by the lack of efficient and robust metrics to evaluate, compare and control the quality of synthetic data.

Notwithstanding the challenges, the fact that analogous ML technologies are being used daily by millions of individuals in other domains, most prominently in smartphones¹⁰⁰, search engines¹⁸² and self-driving vehicles⁶⁸, suggests that the challenges of deployment and regulation of ML for healthcare can also be addressed.

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References

1. Topol, E. J. High-performance medicine: the convergence of human and artificial intelligence. *Nat. Med.* **25**, 44–56 (2019).
2. Gulshan, V. et al. Development and validation of a deep learning algorithm for detection of diabetic retinopathy in retinal fundus photographs. *JAMA* **316**, 2402–2410 (2016).
3. Esteva, A. et al. Dermatologist-level classification of skin cancer with deep neural networks. *Nature* **542**, 115–118 (2017).
4. Rajkomar, A. et al. Scalable and accurate deep learning with electronic health records. *npj Digit. Med.* **1**, 18 (2018).
5. Rajkomar, A. et al. Automatically charting symptoms from patient-physician conversations using machine learning. *JAMA Intern. Med.* **179**, 836–838 (2019).
6. Henry, K. E., Hager, D. N., Pronovost, P. J. & Saria, S. A targeted real-time early warning score (TREWScore) for septic shock. *Sci. Transl. Med.* **7**, 299ra122 (2015).
7. Komorowski, M., Celi, L. A., Badawi, O., Gordon, A. C. & Faisal, A. A. The Artificial Intelligence Clinician learns optimal treatment strategies for sepsis in intensive care. *Nat. Med.* **24**, 1716–1720 (2018).
8. Abràmoff, M. D., Lavin, P. T., Birch, M., Shah, N. & Folk, J. C. Pivotal trial of an autonomous AI-based diagnostic system for detection of diabetic retinopathy in primary care offices. *npj Digit. Med.* **1**, 39 (2018).
9. Iacobucci, G. Babylon Health holds talks with 'significant' number of NHS trusts. *Brit. Med. J.* **368**, m266 (2020).
10. Hale, C. Medtronic to distribute Viz.ai's stroke-spotting AI imaging software. *Fierce Biotech* (23 July 2019); <https://www.fiercebiotech.com/medtech/medtronic-to-distribute-viz-ai-s-stroke-spotting-ai-imaging-software>
11. Hassan, A. E. et al. Early experience utilizing artificial intelligence shows significant reduction in transfer times and length of stay in a hub and spoke model. *Interv. Neuroradiol.* **26**, 615–622 (2020).
12. Ting, D. S. W. et al. Development and validation of a deep learning system for diabetic retinopathy and related eye diseases using retinal images from multiethnic populations with diabetes. *JAMA* **318**, 2211–2223 (2017).
13. McKinney, S. M. et al. International evaluation of an AI system for breast cancer screening. *Nature* **577**, 89–94 (2020).
14. Liu, Y. et al. A deep learning system for differential diagnosis of skin diseases. *Nat. Med.* **26**, 900–908 (2020).
15. Seyyed-Kalantari, L., Liu, G., McDermott, M., Chen, I. Y. & Ghassemi, M. CheXclusion: fairness gaps in deep chest X-ray classifiers. *Pac. Symp. Biocomput.* **26**, 232–243 (2021).
16. Yu, K.-H., Beam, A. L. & Kohane, I. S. Artificial intelligence in healthcare. *Nat. Biomed. Eng.* **2**, 719–731 (2018).
17. Esteva, A. et al. A guide to deep learning in healthcare. *Nat. Med.* **25**, 24–29 (2019).
18. Frid-Adar, M. et al. GAN-based synthetic medical image augmentation for increased CNN performance in liver lesion classification. *Neurocomputing* **321**, 321–331 (2018).
19. Shin, H.-C. et al. Medical image synthesis for data augmentation and anonymization using generative adversarial networks. In *Simulation and Synthesis in Medical Imaging SASHIMI 2018* (eds Gooya, A., Goksel, O., Oguz, I. & Burgos, N.) 1–11 (Springer Cham, 2018).
20. Salehinejad, H., Valaee, S., Dowdell, T., Colak, E. & Barfett, J. Generalization of deep neural networks for chest pathology classification in X-rays using generative adversarial networks. In *IEEE International Conference on Acoustics, Speech and Signal Processing (ICASSP)* 990–994 (ieeexplore.ieee.org, 2018).
21. Zhang, Z., Yang, L. & Zheng, Y. Translating and segmenting multimodal medical volumes with cycle-and shape-consistency generative adversarial network. In *Proceedings of IEEE/CVF Conference on Computer Vision and Pattern Recognition* 9242–9251 (IEEE, 2018).
22. Xu, F., Zhang, J., Shi, Y., Kang, K. & Yang, S. A fast low-rank matrix factorization method for dynamic magnetic resonance imaging restoration. In *5th International Conference on Big Data Computing and Communications (BIGCOM)* 38–42 (2019).
23. Goodfellow, I. J. et al. Generative adversarial networks. In *Advances in Neural Information Processing Systems 27* (eds Ghahramani, Z., Welling, M., Cortes, C., Lawrence, N. & Weinberger, K.Q.) Paper 1384 (Curran, 2014).
24. Wang, Z., She, Q. & Ward, T. E. Generative adversarial networks in computer vision: a survey and taxonomy. *ACM Comput. Surv.* **54**, 1–38 (2021).
25. Radford, A., Metz, L. & Chintala, S. Unsupervised representation learning with deep convolutional generative adversarial networks. Preprint at <https://arxiv.org/abs/1511.06434v2> (2016).
26. Denton, E. L., Chintala, S. & Fergus, R. Deep generative image models using a Laplacian pyramid of adversarial networks. In *Advances in Neural Information Processing Systems 28* (eds Cortes, C., Lawrence, N., Lee, D., Sugiyama, M. & Garnett, R.) Paper 903 (Curran, 2015).
27. Karras, T., Aila, T., Laine, S. & Lehtinen, J. Progressive growing of GANs for improved quality, stability, and variation. In *International Conference on Learning Representations 2018* Paper 447 (ICLR, 2018).
28. Mirza, M. & Osindero, S. Conditional generative adversarial nets. Preprint at <https://arxiv.org/abs/1411.1784v1> (2014).
29. Odena, A., Olah, C. & Shlens, J. Conditional image synthesis with auxiliary classifier GANs. In *Proceedings of the 34th International Conference on Machine Learning* (eds. Precup, D. & Teh, Y. W.) 2642–2651 (PMLR, 2017).
30. Isola, P., Zhu, J.-Y., Zhou, T. & Efros, A. A. Image-to-image translation with conditional adversarial networks. In *Proceedings of the IEEE Conference on Computer Vision and Pattern Recognition* 5967–5976 (2018).
31. Zhang, H., Goodfellow, I., Metaxas, D. & Odena, A. Self-attention generative adversarial networks. In *Proceedings of the 36th International Conference on Machine Learning* (eds. Chaudhuri, K. & Salakhutdinov, R.) 7354–7363 (PMLR, 2019).
32. Wu, Y., Ma, Y., Liu, J., Du, J. & Xing, L. Self-attention convolutional neural network for improved MR image reconstruction. *Inf. Sci.* **490**, 317–328 (2019).
33. Brock, A., Donahue, J. & Simonyan, K. Large scale GAN training for high fidelity natural image synthesis. In *International Conference on Learning Representations* Paper 564 (ICLR, 2019).
34. Arjovsky, M., Chintala, S. & Bottou, L. Wasserstein generative adversarial networks. In *Proceedings of the 34th International Conference on Machine Learning* (eds. Precup, D. & Teh, Y. W.) 214–223 (PMLR, 2017).
35. Gulrajani, I., Ahmed, F., Arjovsky, M., Dumoulin, V. & Courville, A. C. Improved training of Wasserstein GANs. In *Advances in Neural Information Processing Systems 30* (eds. Guyon, I. et al.) Paper 2945 (Curran, 2017).
36. Hindupur, A. The-gan-zoo. <https://github.com/hindupuravinash/the-gan-zoo> (2018).
37. Rajpurkar, P. et al. Deep learning for chest radiograph diagnosis: a retrospective comparison of the CheXNeXt algorithm to practicing radiologists. *PLoS Med.* **15**, e1002686 (2018).
38. Ouyang, D. et al. Video-based AI for beat-to-beat assessment of cardiac function. *Nature* **580**, 252–256 (2020).
39. Xue, Y., Xu, T., Zhang, H., Long, L. R. & Huang, X. SegAN: adversarial network with multi-scale L1 loss for medical image segmentation. *Neuroinformatics* **16**, 383–392 (2018).
40. Haque, A., Milstein, A. & Fei-Fei, L. Illuminating the dark spaces of healthcare with ambient intelligence. *Nature* **585**, 193–202 (2020).
41. Zech, J. R. et al. Variable generalization performance of a deep learning model to detect pneumonia in chest radiographs: a cross-sectional study. *PLoS Med.* **15**, e1002683 (2018).
42. Zou, J. & Schiebinger, L. AI can be sexist and racist — it's time to make it fair. *Nature* **559**, 324–326 (2018).
43. Perez, L. & Wang, J. The effectiveness of data augmentation in image classification using deep learning. Preprint at <https://arxiv.org/abs/1712.04621v1> (2017).
44. Madani, A., Moradi, M., Karargyris, A. & Syeda-Mahmood, T. Semi-supervised learning with generative adversarial networks for chest X-ray classification with ability of data domain adaptation. In *IEEE 15th International Symposium on Biomedical Imaging (ISBI)* 1038–1042 (IEEE, 2018).
45. He, J. et al. The practical implementation of artificial intelligence technologies in medicine. *Nat. Med.* **25**, 30–36 (2019).

46. Kelly, C. J., Karthikesalingam, A., Suleyman, M., Corrado, G. & King, D. Key challenges for delivering clinical impact with artificial intelligence. *BMC Med.* **17**, 195 (2019).

47. Rocher, L., Hendrickx, J. M. & de Montjoye, Y.-A. Estimating the success of re-identifications in incomplete datasets using generative models. *Nat. Commun.* **10**, 3069 (2019).

48. Schwarz, C. G. et al. Identification of anonymous MRI research participants with face-recognition software. *N. Engl. J. Med.* **381**, 1684–1686 (2019).

49. Chartsias, A., Joyce, T., Dharmakumar, R. & Tsafaris, S. A. Adversarial image synthesis for unpaired multi-modal cardiac data. In *Simulation and Synthesis in Medical Imaging* (eds. Tsafaris, S. A., Gooya, A., Frangi, A. F. & Prince, J. L.) 3–13 (Springer International Publishing, 2017).

50. Emami, H., Dong, M., Nejad-Davarani, S. P. & Glide-Hurst, C. K. Generating synthetic CTs from magnetic resonance images using generative adversarial networks. *Med. Phys.* <https://doi.org/10.1002/mp.13047> (2018).

51. Jin, C.-B. et al. Deep CT to MR synthesis using paired and unpaired data. *Sensors* **19**, 2361 (2019).

52. Bi, L., Kim, J., Kumar, A., Feng, D. & Fulham, M. In *Molecular Imaging, Reconstruction and Analysis of Moving Body Organs, and Stroke Imaging and Treatment* (eds. Cardoso, M. J. et al.) 43–51 (Springer International Publishing, 2017).

53. Ben-Cohen, A. et al. Cross-modality synthesis from CT to PET using FCN and GAN networks for improved automated lesion detection. *Eng. Appl. Artif. Intell.* **78**, 186–194 (2019).

54. Armanious, K. et al. MedGAN: medical image translation using GANs. *Comput. Med. Imaging Graph.* **79**, 101684 (2020).

55. Choi, H. & Lee, D. S. Alzheimer's Disease Neuroimaging Initiative. Generation of structural MR images from amyloid PET: application to MR-less quantification. *J. Nucl. Med.* **59**, 1111–1117 (2018).

56. Wei, W. et al. Learning myelin content in multiple sclerosis from multimodal MRI through adversarial training. In *Medical Image Computing and Computer Assisted Intervention — MICCAI 2018* (eds. Frangi, A. F., Schnabel, J. A., Davatzikos, C., Alberola-López, C. & Fichtinger, G.) 514–522 (Springer Cham, 2018).

57. Pan, Y. et al. Synthesizing missing PET from MRI with cycle-consistent generative adversarial networks for Alzheimer's disease diagnosis. In *Medical Image Computing and Computer Assisted Intervention — MICCAI 2018* (eds. Frangi, A. F., Schnabel, J. A., Davatzikos, C., Alberola-López, C. & Fichtinger, G.) 455–463 (Springer Cham, 2018).

58. Welander, P., Karlsson, S. & Eklund, A. Generative adversarial networks for image-to-image translation on multi-contrast MR images - a comparison of CycleGAN and UNIT. Preprint at <https://arxiv.org/abs/1806.07771v1> (2018).

59. Dar, S. U. H. et al. Image synthesis in multi-contrast MRI with conditional generative adversarial networks. *IEEE Trans. Med. Imaging* **38**, 2375–2388 (2019).

60. Zhu, J.-Y., Park, T., Isola, P. & Efros, A. A. Unpaired image-to-image translation using cycle-consistent adversarial networks. In *2017 IEEE International Conference on Computer Vision (ICCV)* (IEEE, 2017); <https://doi.org/10.1109/iccv.2017.244>

61. Maspero, M. et al. Dose evaluation of fast synthetic-CT generation using a generative adversarial network for general pelvis MR-only radiotherapy. *Phys. Med. Biol.* **63**, 185001 (2018).

62. Olut, S., Sahin, Y.H., Demir, U., Unal, G. Generative adversarial training for MRA image synthesis using multi-contrast MRI. In *Predictive Intelligence in MEDicine*. PRIME 2018. Lecture Notes in Computer Science (eds Rekik, I., Unal, G., Adeli, E. & Park, S.) (Springer Cham, 2018); https://doi.org/10.1007/978-3-030-00320-3_18

63. Chen, R. J., Lu, M. Y., Chen, T. Y., Williamson, D. F. K. & Mahmood, F. Synthetic data in machine learning for medicine and healthcare. *Nat. Biomed. Eng.* **5**, 493–497 (2021).

64. Kanakasabapathy, M. K. et al. Adaptive adversarial neural networks for the analysis of lossy and domain-shifted datasets of medical images. *Nat. Biomed. Eng.* **5**, 571–585 (2021).

65. Bowles, C., Gunn, R., Hammers, A. & Rueckert, D. Modelling the progression of Alzheimer's disease in MRI using generative adversarial networks. In *Medical Imaging 2018: Image Processing* (eds. Angelini, E. D. & Landman, B. A.) 397–407 (International Society for Optics and Photonics, 2018).

66. Ravi, D., Alexander, D.C., Oxtoby, N.P. & Alzheimer's Disease Neuroimaging Initiative. Degenerative adversarial neuroImage nets: generating images that mimic disease progression. In *Medical Image Computing and Computer Assisted Intervention — MICCAI 2019*. Lecture Notes in Computer Science. (eds Shen, D. et al.) 164–172 (Springer, 2019).

67. Borji, A. Pros and cons of GAN evaluation measures. *Comput. Vis. Image Underst.* **179**, 41–65 (2019).

68. Vincent, J. Nvidia uses AI to make it snow on streets that are always sunny. *The Verge* <https://www.theverge.com/2017/12/5/16737260/ai-image-translation-nvidia-data-self-driving-cars> (2017).

69. Kairouz, P. et al. Advances and open problems in federated learning. *Found. Trends Mach. Learn.* <https://doi.org/10.1561/2200000083> (2021).

70. McMahan, B., Moore, E., Ramage, D., Hampson, S. & Aguera y Arcas, B. Communication-efficient learning of deep networks from decentralized data. In *Proceedings of the 20th International Conference on Artificial Intelligence and Statistics* (eds. Singh, A. & Zhu, J.) 1273–1282 (ML Research Press, 2017).

71. Li, X. et al. Multi-site fMRI analysis using privacy-preserving federated learning and domain adaptation: ABIDE results. *Med. Image Anal.* **65**, 101765 (2020).

72. Brisimi, T. S. et al. Federated learning of predictive models from federated Electronic Health Records. *Int. J. Med. Inform.* **112**, 59–67 (2018).

73. Lee, J. et al. Privacy-preserving patient similarity learning in a federated environment: development and analysis. *JMIR Med. Inform.* **6**, e20 (2018).

74. Dou, Q. et al. Federated deep learning for detecting COVID-19 lung abnormalities in CT: a privacy-preserving multinational validation study. *npj Digit. Med.* **4**, 60 (2021).

75. Silva, S. et al. Federated learning in distributed medical databases: meta-analysis of large-scale subcortical brain data. In *2019 IEEE 16th International Symposium on Biomedical Imaging ISBI 2019* 18822077 (IEEE, 2019).

76. Sheller, M. J., Reina, G. A., Edwards, B., Martin, J. & Bakas, S. Multi-institutional deep learning modeling without sharing patient data: a feasibility study on brain tumor segmentation. *Brainlesion* **11383**, 92–104 (2019).

77. Sheller, M. J. et al. Federated learning in medicine: facilitating multi-institutional collaborations without sharing patient data. *Sci. Rep.* **10**, 12598 (2020).

78. Sarma, K. V. et al. Federated learning improves site performance in multicenter deep learning without data sharing. *J. Am. Med. Inform. Assoc.* **28**, 1259–1264 (2021).

79. Li, W. et al. Privacy-preserving federated brain tumour segmentation. In *Machine Learning in Medical Imaging* (eds. Suk, H.-I., Liu, M., Yan, P. & Lian, C.) 133–141 (Springer International Publishing, 2019).

80. Shokri, R., Stronati, M., Song, C. & Shmatikov, V. Membership inference attacks against machine learning models. In *IEEE Symposium on Security and Privacy SP 2017* 3–18 (IEEE, 2017).

81. Fredrikson, M., Jha, S. & Ristenpart, T. Model inversion attacks that exploit confidence information and basic countermeasures. In *Proceedings of the 22nd ACM SIGSAC Conference on Computer and Communications Security* 1322–1333 (Association for Computing Machinery, 2015).

82. Zhang, C., Bengio, S., Hardt, M., Recht, B. & Vinyals, O. Understanding deep learning (still) requires rethinking generalization. *Commun. ACM* **64**, 107–115 (2021).

83. Zhu, L., Liu, Z. & Han, S. Deep leakage from gradients. In *Advances in Neural Information Processing Systems 32* (eds Wallach, H. et al.) Paper 8389 (Curran, 2019).

84. Abadi, M. et al. Deep learning with differential privacy. In *Proceedings of the 2016 ACM SIGSAC Conference on Computer and Communications Security* 308–318 (Association for Computing Machinery, 2016).

85. Brendan McMahan, H. et al. A general approach to adding differential privacy to iterative training procedures. Preprint at <https://arxiv.org/abs/1812.06210v2> (2018).

86. McMahan, H. B., Ramage, D., Talwar, K. & Zhang, L. Learning differentially private recurrent language models. In *ICLR 2018 Sixth International Conference on Learning Representations* Paper 504 (ICLR, 2018).

87. Shokri, R. & Shmatikov, V. Privacy-preserving deep learning. In *Proceedings of the 22nd ACM SIGSAC Conference on Computer and Communications Security* 1310–1321 (Association for Computing Machinery, 2015).

88. Lyu, M., Su, D. & Li, N. Understanding the sparse vector technique for differential privacy. *Proc. VLDB Endow.* **10**, 637–648 (2017).

89. Hitaj, B., Ateniese, G. & Perez-Cruz, F. Deep models under the GAN: information leakage from collaborative deep learning. In *Proceedings of the 2017 ACM SIGSAC Conference on Computer and Communications Security* 603–618 (Association for Computing Machinery, 2017).

90. Li, X., Huang, K., Yang, W., Wang, S. & Zhang, Z. On the convergence of FedAvg on Non-IID Data. In *ICLR 2020 Eighth International Conference on Learning Representations* Paper 261 (2020).

91. Smith, V., Chiang, C.-K., Sanjabi, M. & Talwalkar, A. S. Federated multi-task learning. In *Advances in Neural Information Processing Systems 30* (eds Guyon, I. et al.) Paper 2307 (NeurIPS, 2017).

92. Xu, J. et al. Federated learning for healthcare informatics. *J. Healthc. Inform. Res.* **5**, 1–19 (2021).

93. Huang, L. et al. LoAdaBoost: loss-based AdaBoost federated machine learning with reduced computational complexity on IID and non-IID intensive care data. *PLoS ONE* **15**, e0230706 (2020).

94. Zhao, Y. et al. Federated learning with non-IID data. Preprint at <https://arxiv.org/abs/1806.00582v1> (2018).

95. Torres-Soto, J. & Ashley, E. A. Multi-task deep learning for cardiac rhythm detection in wearable devices. *npj Digit. Med.* **3**, 116 (2020).

96. Turakhia, M. P. et al. Rationale and design of a large-scale, app-based study to identify cardiac arrhythmias using a smartwatch: The Apple Heart Study. *Am. Heart J.* **207**, 66–75 (2019).

97. Synced. Apple reveals design of its on-device ML system for federated evaluation and tuning *SyncedReview* <https://syncedreview.com/2021/02/19/apple-reveals-design-of-its-on-device-ml-system-for-federated-evaluation-and-tuning> (2021).

98. McMahan, B. & Ramage, D. Federated learning: collaborative machine learning without centralized training data *Google AI Blog* <https://ai.googleblog.com/2017/04/federated-learning-collaborative.html> (2017).

99. Chen, Y., Qin, X., Wang, J., Yu, C. & Gao, W. FedHealth: a federated transfer learning framework for wearable healthcare. *IEEE Intell. Syst.* **35**, 83–93 (2020).

100. Ramage, D. & Mazzocchi, S. Federated analytics: collaborative data science without data collection *Google AI Blog* <https://ai.googleblog.com/2020/05/federated-analytics-collaborative-data.html> (2020).

101. Augenstein, S. et al. Generative models for effective ML on private, decentralized datasets. In *ICLR 2020 Eighth International Conference on Learning Representations* Paper 1448 (ICLR, 2020).

102. Pati, S. et al. The federated tumor segmentation (FeTS) challenge. Preprint at <https://arxiv.org/abs/2105.05874v2> (2021).

103. Flores, M. Medical institutions collaborate to improve mammogram assessment AI with Nvidia Clara federated learning *The AI Podcast* <https://blogs.nvidia.com/blog/2020/04/15/federated-learning-mammogram-assessment/> (2020).

104. Kannan, A., Chen, K., Jaunzeikare, D. & Rajkomar, A. Semi-supervised learning for information extraction from dialogue. In *Proc. Interspeech 2018* 2077–2081 (ISCA, 2018); <https://doi.org/10.21437/interspeech.2018-1318>

105. Chiu, C.-C. et al. Speech recognition for medical conversations. Preprint at <https://arxiv.org/abs/1711.07274v2>; <https://doi.org/10.1093/jamia/ocx073> (2017).

106. Si, Y., Wang, J., Xu, H. & Roberts, K. Enhancing clinical concept extraction with contextual embeddings. *J. Am. Med. Inform. Assoc.* **26**, 1297–1304 (2019).

107. Shin, H.-C. et al. Learning to read chest X-rays: recurrent neural cascade model for automated image annotation. In *2016 IEEE Conference on Computer Vision and Pattern Recognition (CVPR)* (IEEE, 2016); <https://doi.org/10.1109/cvpr.2016.274>

108. Wang, X., Peng, Y., Lu, L., Lu, Z. & Summers, R. M. TieNet: text-image embedding network for common thorax disease classification and reporting in chest X-rays. In *IEEE/CVF Conference on Computer Vision and Pattern Recognition 2018* (IEEE, 2018); <https://doi.org/10.1109/cvpr.2018.00943>

109. Hochreiter, S. & Schmidhuber, J. Long short-term memory. *Neural Comput.* **9**, 1735–1780 (1997).

110. Cho, K. et al. Learning phrase representations using RNN encoder-decoder for statistical machine translation. In *Proceedings of the 2014 Conference on Empirical Methods in Natural Language Processing (EMNLP)* (eds Moschitti, A., Pang, B. & Daelemans, W.) 1724–1734 (Association for Computational Linguistics, 2014).

111. Lipton, Z. C., Kale, D. C., Elkan, C. & Wetzel, R. Learning to diagnose with LSTM recurrent neural networks. Preprint at <https://arxiv.org/abs/1511.03677v7> (2015).

112. Choi, E., Bahadori, M. T., Schuetz, A., Stewart, W. F. & Sun, J. Doctor AI: predicting clinical events via recurrent neural networks. *JMLR Workshop Conf. Proc.* **56**, 301–318 (2016).

113. Zhu, Paschalidis & Tahmasebi. Clinical concept extraction with contextual word embedding. Preprint at <https://doi.org/10.48550/arXiv.1810.10566> (2018).

114. Cho, K., van Merriënboer, B., Bahdanau, D. & Bengio, Y. On the properties of neural machine translation: encoder–decoder approaches. In *Proceedings of SSST-8, Eighth Workshop on Syntax, Semantics and Structure in Statistical Translation* (eds Wu, D., Carpuat, M., Carreras, X. & Vecchi, E. M.) 103–111 (Association for Computational Linguistics, 2014).

115. Gehring, J., Auli, M., Grangier, D., Yarats, D. & Dauphin, Y. N. Convolutional sequence to sequence learning. In *Proceedings of the 34th International Conference on Machine Learning* (eds Precup, D. & Teh, Y. W.) 1243–1252 (PMLR, 2017).

116. Vaswani, A. et al. Attention is all you need. In *Advances in Neural Information Processing Systems 30* (eds Guyon, I. et al.) Paper 3058 (Curran, 2017).

117. Bahdanau, D., Cho, K. H. & Bengio, Y. Neural machine translation by jointly learning to align and translate. In *3rd International Conference on Learning Representations ICLR 2015* (ICLR, 2015).

118. Luong, T., Pham, H. & Manning, C. D. Effective approaches to attention-based neural machine translation. In *Proceedings of the 2015 Conference on Empirical Methods in Natural Language Processing* (eds Márquez, L., Callison-Burch, C. & Su, J.) 1412–1421 (Association for Computational Linguistics, 2015); <https://doi.org/10.18653/v1/d15-1166>

119. Sutskever, I., Vinyals, O. & Le, Q. V. Sequence to sequence learning with neural networks. In *Advances in Neural Information Processing Systems 27* (eds Ghahramani, Z., Welling, M., Cortes, C., Lawrence, N. & Weinberger, K. Q.) Paper 1610 (Curran, 2014).

120. Krizhevsky, A., Sutskever, I. & Hinton, G. E. in *Advances in Neural Information Processing Systems 25* (eds Bartlett, P. et al.) 1097–1105 (Curran, 2012).

121. Kiani, A. et al. Impact of a deep learning assistant on the histopathologic classification of liver cancer. *npj Digit. Med.* **3**, 23 (2020).

122. Park, S.-M. et al. A mountable toilet system for personalized health monitoring via the analysis of excreta. *Nat. Biomed. Eng.* **4**, 624–635 (2020).

123. Howard, J. & Ruder, S. Universal language model fine-tuning for text classification. In *Proceedings of the 56th Annual Meeting of the Association for Computational Linguistics* (eds Gurevych, I. & Miyao, Y.) 328–339 (Association for Computational Linguistics, 2018).

124. Peters, M. E. et al. Deep contextualized word representations. In *Proceedings of the 2018 Conference of the North American Chapter of the Association for Computational Linguistics: Human Language Technologies* (eds Walker, M., Ji, H. & Stent, A.) 2227–2237 (Association for Computational Linguistics, 2018).

125. Brown, T. et al. Language models are few-shot learners. In *Advances in Neural Information Processing Systems 33* (eds Larochelle, H., Ranzato, M., Hadsell, R., Balcan, M. F. & Lin, H.) 1877–1901 (Curran, 2020).

126. Kenton, J. D. M.-W. C. & Toutanova, L. K. BERT: pre-training of deep bidirectional transformers for language understanding. in *Proceedings of the 2019 Conference of the North American Chapter of the Association for Computational Linguistics: Human Language Technologies* (eds Burstein, J., Doran, C. & Solorio, T.) 4171–4186 (Association for Computational Linguistics, 2019).

127. Mikolov, T., Chen, K., Corrado, G. & Dean, J. Efficient estimation of word representations in vector space. Preprint at <https://arxiv.org/abs/1301.3781v3> (2013).

128. Pennington, J., Socher, R. & Manning, C. GloVe: global vectors for word representation. In *Proceedings of the 2014 Conference on Empirical Methods in Natural Language Processing* (eds Moschitti, A., Pang, B., Daelemans, W.) 1532–1543 (Association for Computational Linguistics, 2014).

129. Alsentzer, E. et al. Publicly available clinical BERT embeddings. In *Proceedings of the 2nd Clinical Natural Language Processing Workshop* (eds Rumshisky, A., Roberts, K., Bethard, S. & Naumann, T.) 72–78 (Association for Computational Linguistics, 2019).

130. Huang, K., Altosaar, J. & Ranganath, R. ClinicalBERT: modeling clinical notes and predicting hospital readmission. Preprint at <https://arxiv.org/abs/1904.05342v3> (2019).

131. Peng, Y., Yan, S. & Lu, Z. Transfer learning in biomedical natural language processing: an evaluation of BERT and ELMo on ten benchmarking datasets. In *Proceedings of the 18th BioNLP Workshop and Shared Task* (eds Demner-Fushman, D., Brettonn Cohen, K., Ananiadou, S. & Tsujii, J.) 58–65 (Association for Computational Linguistics, 2019).

132. Johnson, A. E. W. et al. MIMIC-III, a freely accessible critical care database. *Sci. Data* **3**, 160035 (2016).

133. Sharir, O., Peleg, B. & Shoham, Y. The cost of training NLP models: a concise overview. Preprint at <https://arxiv.org/abs/2004.08900v1> (2020).

134. Lee, J. et al. BioBERT: a pre-trained biomedical language representation model for biomedical text mining. *Bioinformatics* **36**, 1234–1240 (2020).

135. Beltagy, I., Lo, K. & Cohan, A. SciBERT: A pretrained language model for scientific text. In *Proceedings of the 2019 Conference on Empirical Methods in Natural Language Processing and the 9th International Joint Conference on Natural Language Processing* (eds Inui, K., Jiang, J., Ng, V. & Wan, X.) 3615–3620 (Association for Computational Linguistics, 2019).

136. Futoma, J., Morris, J. & Lucas, J. A comparison of models for predicting early hospital readmissions. *J. Biomed. Inform.* **56**, 229–238 (2015).

137. Caruana, R. et al. Intelligent models for healthcare: predicting pneumonia risk and hospital 30-day readmission. In *Proc. 21st ACM SIGKDD International Conference on Knowledge Discovery and Data Mining* 1721–1730 (Association for Computing Machinery, 2015).

138. Wagner, T. et al. Augmented curation of clinical notes from a massive EHR system reveals symptoms of impending COVID-19 diagnosis. *Elife* **9**, e58227 (2020).

139. Eismann, A. S. et al. Extracting angina symptoms from clinical notes using pre-trained transformer architectures. *AMIA Annu. Symp. Proc.* **2020**, 412–421 (American Medical Informatics Association, 2020).

140. Smit, A. et al. Combining automatic labelers and expert annotations for accurate radiology report labeling using BERT. In *Proceedings of the 2020 Conference on Empirical Methods in Natural Language Processing* (eds Webber, B., Cohn, T., He, Y. & Liu, Y.) 1500–1519 (Association for Computational Linguistics, 2020).

141. Soni, S. & Roberts, K. Evaluation of dataset selection for pre-training and fine-tuning transformer language models for clinical question answering. In *Proc. 12th Language Resources and Evaluation Conference* 5532–5538 (European Language Resources Association, 2020).

142. Sezgin, E., Huang, Y., Ramtekkar, U. & Lin, S. Readiness for voice assistants to support healthcare delivery during a health crisis and pandemic. *npj Digit. Med.* **3**, 122 (2020).

143. Sakthive, V., Kesaven, M. P. V., William, J. M. & Kumar, S. K. M. Integrated platform and response system for healthcare using Alexa. *Int. J. Commun. Computer Technol.* **7**, 14–22 (2019).

144. Comstock, J. Buoy Health, CVS MinuteClinic partner to send patients from chatbot to care. *mobihealthnews* <https://www.mobihealthnews.com/content/buoy-health-cvs-minuteclinic-partner-send-patients-chatbot-care> (2018).

145. Razzaki, S. et al. A comparative study of artificial intelligence and human doctors for the purpose of triage and diagnosis. Preprint at <https://doi.org/10.48550/arXiv.1806.10698> (2018).

146. Xiong, Y., Du, B. & Yan, P. Reinforced transformer for medical image captioning. In *Machine Learning in Medical Imaging* (eds. Suk, H.-I., Liu, M., Yan, P. & Lian, C.) 673–680 (Springer International Publishing, 2019).

147. Meng, Y., Speier, W., Ong, M. K. & Arnold, C. W. Bidirectional representation learning from transformers using multimodal electronic health record data to predict depression. *IEEE J. Biomed. Health Inform.* **25**, 3121–3129 (2021).

148. Choi, E. et al. Learning the graphical structure of electronic health records with graph convolutional transformer. *Proc. Conf. AAAI Artif. Intell.* **34**, 606–613 (2020).

149. Li, F. et al. Fine-tuning bidirectional encoder representations from transformers (BERT)-based models on large-scale electronic health record notes: an empirical study. *JMIR Med. Inform.* **7**, e14830 (2019).

150. Rasmy, L., Xiang, Y., Xie, Z., Tao, C. & Zhi, D. Med-BERT: pretrained contextualized embeddings on large-scale structured electronic health records for disease prediction. *npj Digital Medicine* **4**, 86 (2021).

151. Shang, J., Ma, T., Xiao, C. & Sun, J. Pre-training of graph augmented transformers for medication recommendation. In *Proceedings of the Twenty-Eighth International Joint Conference on Artificial Intelligence* (ed. Kraus, S.) 5953–5959 (International Joint Conferences on Artificial Intelligence Organization, 2019); <https://doi.org/10.24963/ijcai.2019/825>

152. Li, Y. et al. BEHRT: transformer for electronic health records. *Sci. Rep.* **10**, 7155 (2020).

153. Rao, S. et al. BEHRT-HF: an interpretable transformer-based, deep learning model for prediction of incident heart failure. *Eur. Heart J.* **41** (Suppl. 2), ehaa946.3553 (2020).

154. Qian, X. et al. Prospective assessment of breast cancer risk from multimodal multiview ultrasound images via clinically applicable deep learning. *Nat. Biomed. Eng.* **5**, 522–532 (2021).

155. Xing, L., Giger, M. L. & Min, J. K. *Artificial Intelligence in Medicine: Technical Basis and Clinical Applications* (Academic Press, 2020).

156. Reisman, M. EHRs: the challenge of making electronic data usable and interoperable. *P. T.* **42**, 572–575 (2017).

157. Cortés, R., Bonnaire, X., Marin, O. & Sens, P. Stream processing of healthcare sensor data: studying user traces to identify challenges from a big data perspective. *Procedia Comput. Sci.* **52**, 1004–1009 (2015).

158. Zhang, F., Cao, J., Khan, S. U., Li, K. & Hwang, K. A task-level adaptive MapReduce framework for real-time streaming data in healthcare applications. *Future Gener. Comput. Syst.* **43–44**, 149–160 (2015).

159. El Aboudi, N. & Benhlima, L. Big data management for healthcare systems: architecture, requirements, and implementation. *Adv. Bioinformatics* **2018**, 4059018 (2018).

160. Ta, V.-D., Liu, C.-M. & Nkabinde, G. W. Big data stream computing in healthcare real-time analytics. In *IEEE International Conference on Cloud Computing and Big Data Analysis* (ICCCBDA) 37–42 (ieeexplore.ieee.org, 2016).

161. *Data-Driven Healthcare Organizations Use Big Data Analytics for Big Gains* White Paper (IBM Software, 2017); <https://silo.tips/download/ibm-software-white-paper-data-driven-healthcare-organizations-use-big-data-analy>

162. Futoma, J., Simons, M., Panch, T., Doshi-Velez, F. & Celi, L. A. The myth of generalisability in clinical research and machine learning in health care. *Lancet Digit. Health* **2**, e489–e492 (2020).

163. Wang, X. et al. Inconsistent performance of deep learning models on mammogram classification. *J. Am. Coll. Radiol.* **17**, 796–803 (2020).

164. Nestor, B., McDermott, M. B. A. & Boag, W. Feature robustness in non-stationary health records: caveats to deployable model performance in common clinical machine learning tasks. Preprint at <https://doi.org/10.48550/arXiv.1908.00690> (2019).

165. Wu, E. et al. How medical AI devices are evaluated: limitations and recommendations from an analysis of FDA approvals. *Nat. Med.* <https://doi.org/10.1038/s41591-021-01312-x> (2021).

166. Barish, M., Bolourani, S., Lau, L. F., Shah, S. & Zanos, T. P. External validation demonstrates limited clinical utility of the interpretable mortality prediction model for patients with COVID-19. *Nat. Mach. Intell.* **3**, 25–27 (2020).

167. Davis, S. E., Lasko, T. A., Chen, G., Siew, E. D. & Matheny, M. E. Calibration drift in regression and machine learning models for acute kidney injury. *J. Am. Med. Inform. Assoc.* **24**, 1052–1061 (2017).

168. Wang, G. et al. A deep-learning pipeline for the diagnosis and discrimination of viral, non-viral and COVID-19 pneumonia from chest X-ray images. *Nat. Biomed. Eng.* **5**, 509–521 (2021).

169. Ning, W. et al. Open resource of clinical data from patients with pneumonia for the prediction of COVID-19 outcomes via deep learning. *Nat. Biomed. Eng.* **4**, 1197–1207 (2020).

170. Koenecke, A. et al. Racial disparities in automated speech recognition. *Proc. Natl Acad. Sci. USA* **117**, 7684–7689 (2020).

171. Abid, A., Farooqi, M. & Zou, J. Large language models associate muslims with violence. *Nat. Mach. Intell.* **3**, 461–463 (2021).

172. Obermeyer, Z., Powers, B., Vogeli, C. & Mullainathan, S. Dissecting racial bias in an algorithm used to manage the health of populations. *Science* **366**, 447–453 (2019).

173. Adamson, A. S. & Smith, A. Machine learning and health care disparities in dermatology. *JAMA Dermatol.* **154**, 1247–1248 (2018).

174. Han, S. S. et al. Classification of the clinical images for benign and malignant cutaneous tumors using a deep learning algorithm. *J. Invest. Dermatol.* **138**, 1529–1538 (2018).

175. Subbaswamy, A., Adams, R. & Saria, S. Evaluating model robustness and stability to dataset shift. In *Proceedings of The 24th International Conference on Artificial Intelligence and Statistics* (eds. Banerjee, A. & Fukumizu, K.) 2611–2619 (PMLR, 2021).

176. Izzo, Z., Ying, L. & Zou, J. How to learn when data reacts to your model: performative gradient descent. In *Proceedings of the 38th International Conference on Machine Learning* (eds. Meila, M. & Zhang, T.) 4641–4650 (PMLR, 2021).

177. Ghorbani, A., Kim, M. & Zou, J. A Distributional framework for data valuation. In *Proceedings of the 37th International Conference on Machine Learning* (eds. Ilii, H. D. & Singh, A.) 3535–3544 (PMLR, 2020).

178. Zhang, L., Deng, Z., Kawaguchi, K., Ghorbani, A. & Zou, J. How does mixup help with robustness and generalization? In *International Conference on Learning Representations 2021* Paper 2273 (ICLR, 2021).

179. Schulam, P. & Saria, S. Can you trust this prediction? Auditing pointwise reliability after learning. In *Proceedings of the Twenty-Second International Conference on Artificial Intelligence and Statistics* (eds. Chaudhuri, K. & Sugiyama, M.) 1022–1031 (PMLR, 2019).

180. Liu, X. et al. Reporting guidelines for clinical trial reports for interventions involving artificial intelligence: the CONSORT-AI extension. *Nat. Med.* **26**, 1364–1374 (2020).

181. Cruz Rivera, S. et al. Guidelines for clinical trial protocols for interventions involving artificial intelligence: the SPIRIT-AI extension. *Nat. Med.* **26**, 1351–1363 (2020).

182. Nayak, P. Understanding searches better than ever before. *Google The Keyword* <https://blog.google/products/search/search-language-understanding-bert/> (2019).

183. Baur, C., Albarqouni, S. & Navab, N. in *OR 2.0 Context-Aware Operating Theaters, Computer Assisted Robotic Endoscopy, Clinical Image-Based Procedures, and Skin Image Analysis* (eds Stoyanov, D. et al.) 260–267 (Springer International Publishing, 2018).

184. Kang, E., Koo, H. J., Yang, D. H., Seo, J. B. & Ye, J. C. Cycle-consistent adversarial denoising network for multiphase coronary CT angiography. *Med. Phys.* **46**, 550–562 (2019).

185. Vig, J. A multiscalar visualization of attention in the transformer model. In *Proceedings of the 57th Annual Meeting of the Association for Computational Linguistics: System Demonstrations* (Costa-jussà, M. R. & Alfonseca, E.) 37–42 (Association for Computational Linguistics, 2019).

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Competing interests

J.C.W. is a co-founder and scientific advisory board member of Greenstone Biosciences. The other authors declare no competing interests.

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